

ARCA biopharma, Inc.  
Form 424B4  
November 13, 2013

**Prospects Supplement No. 10**

**Filed pursuant to Rule 424(b)(4)**

**(to Prospectus dated May 30, 2013)**

**Registration No. 333-187508**

**125,000 Shares of Series A Convertible Preferred Stock**

**12,500,000 Shares of Common Stock Underlying the Preferred Stock**

**Warrants to Purchase up to 6,250,000 Shares of Common Stock and**

**6,250,000 Shares of Common Stock Underlying the Warrants**

**ARCA biopharma, Inc.**

This prospectus supplement supplements the prospectus dated May 30, 2013 (the "Prospectus"), as supplemented by that certain Prospectus Supplement No. 1 dated July 17, 2013 ("Supplement No. 1"), by that certain Prospectus Supplement No. 2 dated July 19, 2013 ("Supplement No. 2"), by that certain Prospectus Supplement No. 3 dated July 24, 2013 ("Supplement No. 3"), by that certain Prospectus Supplement No. 4 dated July 30, 2013 ("Supplement No. 4"), by that certain Prospectus Supplement No. 5 dated August 6, 2013 ("Supplement No. 5"), by that certain Prospectus Supplement No. 6 dated September 4, 2013 ("Supplement No. 6"), by that certain Prospectus Supplement No. 7 dated September 23, 2013 ("Supplement No. 7"), by that certain Prospectus Supplement No. 8 dated October 29, 2013 ("Supplement No. 8"), and by that certain Prospectus Supplement No. 9 dated November 6, 2013 ("Supplement No. 9"), and together with Supplement No. 1, Supplement No. 2, Supplement No. 3, Supplement No. 4, Supplement No. 5, Supplement No. 6, Supplement No. 7, and Supplement No. 8, the "Supplements"), which form a part of our Registration Statement on Form S-1 (Registration No. 333-187508). This prospectus supplement is being filed to update and supplement the information in the Prospectus and the Supplements with the information contained in our quarterly report on Form 10-Q, filed with the Securities and Exchange Commission (the "Commission") on November 13, 2013 (the "Quarterly Report"). Accordingly, we have attached the Quarterly Report to this prospectus supplement.

The Prospectus, the Supplements and this prospectus supplement relate to the offer and sale of up to 125,000 shares of Series A Convertible Preferred Stock ("Preferred Stock") which are convertible into 12,500,000 shares of Common Stock, warrants to purchase up to 6,250,000 shares of our Common Stock and 6,250,000 shares of Common Stock underlying the warrants.

This prospectus supplement should be read in conjunction with the Prospectus and the Supplements. This prospectus supplement updates and supplements the information in the Prospectus and the Supplements. If there is any inconsistency between the information in the Prospectus, the Supplements and this prospectus supplement, you should rely on the information in this prospectus supplement.

Our common stock is traded on the Nasdaq Global Market under the trading symbol "ABIO". On November 13, 2013, the last reported sale price of our common stock was \$1.47 per share.

**Investing in our securities involves a high degree of risk. You should review carefully the risks and uncertainties described under the heading "Risk Factors" beginning on page 5 of the Prospectus and beginning on page 23 of our quarterly report on Form 10-Q for the quarterly period ended September 30, 2013 before you decide whether to invest in shares of our common stock.**

**Neither the Securities and Exchange Commission nor any state securities commission has approved or disapproved of these securities or determined if the Prospectus or this prospectus supplement is truthful or complete. Any representation to the contrary is a criminal offense.**

**The date of this prospectus supplement is November 13, 2013**

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

FORM 10-Q

(Mark One)

☒ QUARTERLY REPORT UNDER SECTION 13 OR 15(d) OF THE SECURITIES EXCHANGE ACT OF 1934  
FOR THE QUARTERLY PERIOD ENDED SEPTEMBER 30, 2013

OR

☐ TRANSITION REPORT UNDER SECTION 13 OR 15(d) OF THE SECURITIES EXCHANGE ACT OF 1934  
FOR THE TRANSITION PERIOD FROM                      TO

Commission File Number 000-22873

ARCA BIOPHARMA, INC.

(Exact Name of Registrant as Specified in Its Charter)

Delaware  
(State or Other Jurisdiction of

36-3855489  
(I.R.S. Employer

Incorporation or Organization)  
11080 CirclePoint Road, Suite 140, Westminster, CO

Identification Number)  
80020

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(Address of Principal Executive Offices)  
(720) 940-2200

(Zip Code)

(Registrant's Telephone Number, including Area Code)

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

Indicate by check mark whether the registrant 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 (§232.405 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. See the definitions of large accelerated filer, accelerated filer and smaller reporting company in Rule 12b-2 of the Exchange Act.

Large accelerated filer ☐

Accelerated filer ☐

Non-accelerated filer ☐ (Do not check if smaller reporting company) 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 ☒

Indicate the number of shares outstanding of each of the issuer's classes of common stock, as of the latest practicable date.

| Class                          | Number of<br>Shares Outstanding |
|--------------------------------|---------------------------------|
| Common Stock \$0.001 par value | On November 8, 2013: 15,675,562 |



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## PART I. FINANCIAL INFORMATION

## ITEM 1. CONSOLIDATED FINANCIAL STATEMENTS

## ARCA BIOPHARMA, INC.

(a development stage enterprise)

## CONSOLIDATED BALANCE SHEETS

(Unaudited)

|                                                                                                                                                                                                                   | September 30,<br>2013                                    | December 31,<br>2012 |
|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------|----------------------|
|                                                                                                                                                                                                                   | (in thousands, except<br>share and per share<br>amounts) |                      |
| ASSETS                                                                                                                                                                                                            |                                                          |                      |
| Current assets:                                                                                                                                                                                                   |                                                          |                      |
| Cash and cash equivalents                                                                                                                                                                                         | \$ 18,632                                                | \$ 2,920             |
| Other current assets                                                                                                                                                                                              | 379                                                      | 125                  |
| Total current assets                                                                                                                                                                                              | 19,011                                                   | 3,045                |
| Property and equipment, net                                                                                                                                                                                       | 27                                                       | 23                   |
| Other assets                                                                                                                                                                                                      | 100                                                      | 144                  |
| Total assets                                                                                                                                                                                                      | \$ 19,138                                                | \$ 3,212             |
| LIABILITIES AND STOCKHOLDERS' EQUITY                                                                                                                                                                              |                                                          |                      |
| Current liabilities:                                                                                                                                                                                              |                                                          |                      |
| Accounts payable                                                                                                                                                                                                  | \$ 447                                                   | \$ 65                |
| Accrued compensation and employee benefits                                                                                                                                                                        | 75                                                       | 103                  |
| Accrued expenses and other liabilities                                                                                                                                                                            | 247                                                      | 121                  |
| Deferred rent, current portion                                                                                                                                                                                    |                                                          | 16                   |
| Total current liabilities                                                                                                                                                                                         | 769                                                      | 305                  |
| Total liabilities                                                                                                                                                                                                 | 769                                                      | 305                  |
| Commitments and contingencies                                                                                                                                                                                     |                                                          |                      |
| Stockholders' equity:                                                                                                                                                                                             |                                                          |                      |
| Series A convertible preferred stock, \$0.001 par value; 135,000 shares authorized; 24,902 shares issued and outstanding at September 30, 2013; no shares authorized, issued and outstanding at December 31, 2012 |                                                          |                      |
| Common stock, \$0.001 par value; 100 million shares authorized; 13,195,362 shares issued and outstanding at September 30, 2013; 2,660,315 shares issued and outstanding at December 31, 2012.                     | 13                                                       | 3                    |
| Additional paid-in capital                                                                                                                                                                                        | 90,343                                                   | 70,898               |

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|                                                  |           |          |
|--------------------------------------------------|-----------|----------|
| Deficit accumulated during the development stage | (71,987)  | (67,994) |
| Total stockholders' equity                       | 18,369    | 2,907    |
| Total liabilities and stockholders' equity       | \$ 19,138 | \$ 3,212 |

See accompanying notes to consolidated financial statements

## ARCA BIOPHARMA, INC.

(a development stage enterprise)

## CONSOLIDATED STATEMENTS OF OPERATIONS AND COMPREHENSIVE LOSS

(unaudited)

|                                                              | Three Months Ended<br>September 30,                |          | Nine Months Ended<br>September 30, |            | Period from<br>December 17,<br>2001 (date of<br>inception) to<br>September 30,<br>2013 |
|--------------------------------------------------------------|----------------------------------------------------|----------|------------------------------------|------------|----------------------------------------------------------------------------------------|
|                                                              | 2013                                               | 2012     | 2013                               | 2012       |                                                                                        |
|                                                              | (in thousands, except share and per share amounts) |          |                                    |            |                                                                                        |
| Costs and expenses:                                          |                                                    |          |                                    |            |                                                                                        |
| Research and development                                     | \$ 714                                             | \$ 156   | \$ 1,141                           | \$ 901     | \$ 43,818                                                                              |
| Selling, general and<br>administrative                       | 1,011                                              | 772      | 2,852                              | 2,579      | 45,435                                                                                 |
| Merger transaction costs                                     |                                                    |          |                                    |            | 5,470                                                                                  |
| Restructuring expense, net                                   |                                                    |          |                                    |            | 2,413                                                                                  |
| Loss on impairment of in-process<br>research and development |                                                    |          |                                    |            | 6,000                                                                                  |
| Total costs and expenses                                     | 1,725                                              | 928      | 3,993                              | 3,480      | 103,136                                                                                |
| Loss from operations                                         | (1,725)                                            | (928)    | (3,993)                            | (3,480)    | (103,136)                                                                              |
| Gain on assignment of patent<br>rights                       |                                                    |          |                                    |            | 2,000                                                                                  |
| Gain on bargain purchase                                     |                                                    |          |                                    |            | 25,282                                                                                 |
| Interest and other income                                    | 2                                                  | 2        | 3                                  | 3          | 2,031                                                                                  |
| Interest and other expense                                   |                                                    | (1)      | (3)                                | (3)        | (445)                                                                                  |
| Loss before income taxes                                     | (1,723)                                            | (927)    | (3,993)                            | (3,480)    | (74,268)                                                                               |
| Benefit from income taxes                                    |                                                    |          |                                    |            | 2,281                                                                                  |
| Net loss and comprehensive loss                              | \$ (1,723)                                         | \$ (927) | \$ (3,993)                         | \$ (3,480) | \$ (71,987)                                                                            |
| Less: Accretion of redeemable<br>convertible preferred stock |                                                    |          |                                    |            | (245)                                                                                  |
| Less: Deemed preferred stock<br>dividend                     |                                                    |          | (2,026)                            |            | (2,807)                                                                                |
| Net loss attributable to common<br>stockholders              | \$ (1,723)                                         | \$ (927) | \$ (6,019)                         | \$ (3,480) | \$ (75,039)                                                                            |

Net loss attributable to common  
stockholders per share:

|                   |    |        |    |        |    |        |    |        |
|-------------------|----|--------|----|--------|----|--------|----|--------|
| Basic and diluted | \$ | (0.16) | \$ | (0.41) | \$ | (0.96) | \$ | (1.65) |
|-------------------|----|--------|----|--------|----|--------|----|--------|

Weighted average shares  
outstanding:

|                   |            |           |           |           |
|-------------------|------------|-----------|-----------|-----------|
| Basic and diluted | 10,832,516 | 2,283,736 | 6,299,828 | 2,113,679 |
|-------------------|------------|-----------|-----------|-----------|

See accompanying notes to consolidated financial statements

## ARCA BIOPHARMA, INC.

(a development stage enterprise)

## CONSOLIDATED STATEMENTS OF PREFERRED STOCK AND STOCKHOLDERS' EQUITY (DEFICIT)

(unaudited)

|                                                              |  | Stockholders                                    |        |                                                 |        | Equity (Deficit) |        |                            |                                                  |       |
|--------------------------------------------------------------|--|-------------------------------------------------|--------|-------------------------------------------------|--------|------------------|--------|----------------------------|--------------------------------------------------|-------|
|                                                              |  | Series B Redeemable Convertible Preferred Stock |        | Series A Redeemable Convertible Preferred Stock |        | Common Stock     |        | Additional Paid In Capital | Deficit Accumulated During the Development Stage | Total |
|                                                              |  | Shares                                          | Amount | Shares                                          | Amount | Shares           | Amount |                            |                                                  |       |
| (in thousands, except share and per share amounts)           |  |                                                 |        |                                                 |        |                  |        |                            |                                                  |       |
| e,<br>ber 17,<br>late of<br>(on)                             |  |                                                 | \$     |                                                 | \$     |                  | \$     | \$                         | \$                                               | \$    |
| ce of<br>on stock<br>ders on<br>ber 31,<br>or cash,<br>6 per |  |                                                 |        |                                                 |        | 2,588            |        | 1                          | (116)                                            |       |
| s<br>e,<br>ber 31,<br>ce of<br>on stock                      |  |                                                 |        |                                                 |        | 2,588            |        | 1                          | (116)                                            |       |
| ber 30,<br>or cash,<br>6 per                                 |  |                                                 |        |                                                 |        | 19,720           |        | 7                          | (511)                                            |       |
| s<br>e,<br>ber 31,<br>ce of<br>on stock                      |  |                                                 |        |                                                 |        | 22,308           |        | 8                          | (627)                                            |       |
| uary 3,                                                      |  |                                                 |        |                                                 |        | 2,922            |        | 1                          |                                                  |       |

|                                                                                                    |           |       |        |       |         |
|----------------------------------------------------------------------------------------------------|-----------|-------|--------|-------|---------|
| or cash,<br>6 per                                                                                  |           |       |        |       |         |
| ce of<br>on stock<br>uary 3,<br>upon<br>sion of<br>ayable<br>ated<br>d<br>at<br>er                 |           |       | 2,978  | 1     |         |
| ce of<br>on stock<br>bber 14,<br>or<br>tual<br>y<br>rights,<br>84 per                              |           |       | 903    | 44    |         |
| ce of<br>on stock<br>bber 14,<br>upon<br>sion of<br>ayable<br>ated<br>d<br>t<br>s<br>e,<br>ber 31, |           |       | 31,095 | 1,354 | (1,459) |
| ce of<br>on stock<br>ry 21,<br>or<br>tual<br>y<br>rights,<br>2 per                                 |           |       | 60,206 | 1,408 | (2,086) |
| ce of<br>A on<br>ry 22,<br>or cash,<br>265 per                                                     | 5,727,354 | 9,316 | 17,372 | 75    |         |

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420,817                  684

8,019                                  3

See accompanying notes to consolidated financial statements

|                                                                                              | Stockholders                                    |        |                                                 |        |                                      |        | Equity (Deficit)           |                                                  | Total |
|----------------------------------------------------------------------------------------------|-------------------------------------------------|--------|-------------------------------------------------|--------|--------------------------------------|--------|----------------------------|--------------------------------------------------|-------|
|                                                                                              | Series A Redeemable Convertible Preferred Stock |        | Series B Redeemable Convertible Preferred Stock |        | Series A Convertible Preferred Stock |        | Additional Paid In Capital | Deficit Accumulated During the Development Stage |       |
|                                                                                              | Shares                                          | Amount | Shares                                          | Amount | Shares                               | Amount |                            |                                                  |       |
|                                                                                              |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| (in thousands, except share and per share amounts)                                           |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at January 22, 2019                                                                  |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for actual net proceeds, net of offering costs, of \$2.00 per share |                                                 |        |                                                 |        |                                      | 13,907 |                            | 60                                               |       |
| Balance at January 23, 2019                                                                  |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for actual net proceeds, net of offering costs, of \$2.00 per share |                                                 |        |                                                 |        |                                      | 2,505  |                            | 15                                               |       |
| Balance at January 24, 2019                                                                  |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for actual net proceeds, net of offering costs, of \$2.00 per share |                                                 |        |                                                 |        |                                      | 38     |                            |                                                  |       |
| Balance at January 25, 2019                                                                  |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                | 3,074,086                                       | 5,000  |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 | (98)   |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at January 26, 2019                                                                  |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at January 27, 2019                                                                  |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at January 28, 2019                                                                  |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at January 29, 2019                                                                  |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at January 30, 2019                                                                  |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at January 31, 2019                                                                  |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 1, 2019                                                                  |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 2, 2019                                                                  |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 3, 2019                                                                  |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 4, 2019                                                                  |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 5, 2019                                                                  |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 6, 2019                                                                  |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 7, 2019                                                                  |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 8, 2019                                                                  |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 9, 2019                                                                  |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 10, 2019                                                                 |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 11, 2019                                                                 |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 12, 2019                                                                 |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 13, 2019                                                                 |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 14, 2019                                                                 |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 15, 2019                                                                 |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 16, 2019                                                                 |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 17, 2019                                                                 |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 18, 2019                                                                 |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 19, 2019                                                                 |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 20, 2019                                                                 |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 21, 2019                                                                 |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 22, 2019                                                                 |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 23, 2019                                                                 |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 24, 2019                                                                 |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 25, 2019                                                                 |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 26, 2019                                                                 |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 27, 2019                                                                 |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 28, 2019                                                                 |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at February 29, 2019                                                                 |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 1, 2019                                                                     |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 2, 2019                                                                     |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 3, 2019                                                                     |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 4, 2019                                                                     |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 5, 2019                                                                     |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 6, 2019                                                                     |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 7, 2019                                                                     |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 8, 2019                                                                     |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 9, 2019                                                                     |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 10, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 11, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 12, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 13, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 14, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 15, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 16, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 17, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 18, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 19, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 20, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 21, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 22, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 23, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 24, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 25, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 26, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 27, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 28, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 29, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 30, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at March 31, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 1, 2019                                                                     |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 2, 2019                                                                     |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 3, 2019                                                                     |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 4, 2019                                                                     |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 5, 2019                                                                     |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 6, 2019                                                                     |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 7, 2019                                                                     |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 8, 2019                                                                     |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 9, 2019                                                                     |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 10, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 11, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 12, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 13, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 14, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 15, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 16, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 17, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 18, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 19, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 20, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 21, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 22, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 23, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 24, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 25, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 26, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 27, 2019                                                                    |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Issuance of common stock for cash, net of offering costs, of \$2.00 per share                |                                                 |        |                                                 |        |                                      |        |                            |                                                  |       |
| Balance at April 28, 2019                                                                    |                                                 |        |                                                 |        |                                      | </     |                            |                                                  |       |

|                                                                                                                                             |           |           |       |         |         |            |
|---------------------------------------------------------------------------------------------------------------------------------------------|-----------|-----------|-------|---------|---------|------------|
| based<br>nsation<br>ion of<br>g costs<br>emable<br>tible<br>ed                                                                              | 17        |           |       | (17)    | (5,241) | (5         |
| ss<br>e,<br>ber 31,                                                                                                                         | 9,222,257 | 14,919    |       | 102,047 | 1,583   | (7,327) (5 |
| ce of<br>B<br>tible<br>nable<br>ed<br>on<br>1, 2007<br>439 per                                                                              |           | 3,688,902 | 9,000 |         |         |            |
| ce of<br>B<br>tible<br>nable<br>ed<br>on<br>ber 28,<br>or<br>per                                                                            |           | 2,766,677 | 9,000 |         |         |            |
| B<br>g Costs<br>ion of<br>A<br>g costs<br>ion of<br>B<br>g costs<br>ce of<br>on stock<br>ellectual<br>ty<br>rights,<br>uary 18,<br>t<br>per | 19        |           | (147) |         | (19)    |            |
| ce of<br>on stock<br>ellectual                                                                                                              |           |           | 18    |         | (18)    |            |
|                                                                                                                                             |           |           |       | 1,303   | 13      |            |
|                                                                                                                                             |           |           |       | 642     | 7       |            |

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See accompanying notes to consolidated financial statements

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|                 |             |              |          |         |   |        |  |        |
|-----------------|-------------|--------------|----------|---------|---|--------|--|--------|
| Common          |             |              |          |         |   |        |  |        |
| Conversion      |             |              |          |         |   |        |  |        |
| Preferred       |             |              |          |         |   |        |  |        |
| Stock dividend  |             |              |          |         |   |        |  |        |
| Additional      |             |              |          |         |   |        |  |        |
| Common          |             |              |          |         |   |        |  |        |
| Shares          |             |              |          |         |   |        |  |        |
| Available under |             |              |          |         |   |        |  |        |
| Anti-dilution   |             |              |          |         |   |        |  |        |
| Provision       |             |              | 781      |         |   | (781)  |  | (781)  |
| Retention of    |             |              |          |         |   |        |  |        |
| Series A        |             |              |          |         |   |        |  |        |
| Issuance costs  | 42          |              |          |         |   | (42)   |  | (42)   |
| Retention of    |             |              |          |         |   |        |  |        |
| Series B        |             |              |          |         |   |        |  |        |
| Issuance costs  |             |              | 93       |         |   | (93)   |  | (93)   |
| Conversion of   |             |              |          |         |   |        |  |        |
| Preferred       |             |              |          |         |   |        |  |        |
| Stock           | (9,222,257) | (16,466,579) | (18,781) | 507,123 | 1 | 33,780 |  | 33,780 |
| Restricted      |             |              |          |         |   |        |  |        |
| Stock release   |             |              |          |         |   |        |  |        |
| Penalty         |             |              |          |         |   |        |  |        |
| Restriction     |             |              |          |         |   | 75     |  | 75     |
| Conversion of   |             |              |          |         |   |        |  |        |
| Convertible     |             |              |          |         |   |        |  |        |
| Shares and      |             |              |          |         |   |        |  |        |
| Warrants        |             |              |          |         |   |        |  |        |
| Issued          |             |              |          |         |   |        |  |        |
| Interest        |             |              |          | 145,465 |   | 8,501  |  | 8,501  |
| Conversion of   |             |              |          |         |   |        |  |        |
| Warrants for    |             |              |          |         |   |        |  |        |
| Preferred       |             |              |          |         |   | 36     |  | 36     |
| Stock           |             |              |          |         |   |        |  |        |
| Agreement with  |             |              |          | 447,826 |   | 11,913 |  | 11,913 |
| Novelo, Inc.    |             |              |          |         |   |        |  |        |
| Adjustment      |             |              |          |         |   |        |  |        |
| Fractional      |             |              |          |         |   |        |  |        |
| Shares          |             |              |          | (102)   |   |        |  |        |
| Share-based     |             |              |          |         |   |        |  |        |
| Compensation    |             |              |          |         |   | 845    |  | 845    |
| Assurance of    |             |              |          |         |   |        |  |        |
| Common stock    |             |              |          |         |   |        |  |        |
| On exercise     |             |              |          |         |   |        |  |        |
| Stock           |             |              |          |         |   |        |  |        |
| Options for     |             |              |          |         |   |        |  |        |
| Which           |             |              |          | 10,521  |   | 114    |  | 114    |
| Assurance of    |             |              |          | 177     |   | 2      |  |        |
| Common stock    |             |              |          |         |   |        |  |        |
| Over            |             |              |          |         |   |        |  |        |
| Employee        |             |              |          |         |   |        |  |        |

|                                                                                                                                                                                                                   |           |   |        |                            |                       |
|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------|---|--------|----------------------------|-----------------------|
| stock purchase<br>and upon<br>the<br>of<br>restricted<br>stock units<br>estimated fair<br>value of<br>warrants<br>vested in<br>connection<br>with lease<br>termination<br>loss<br>income,<br>December 31,<br>2019 | 1,270,074 | 1 | 57,301 | 377<br>(9,138)<br>(49,890) | 37<br>(9,138)<br>7,41 |
|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------|---|--------|----------------------------|-----------------------|

See accompanying notes to consolidated financial statements

|                                                                  | Stockholders' Equity (Deficit) |                             |                             |                             |                             |                             |                             |                             |                             |        |                                                  |         |
|------------------------------------------------------------------|--------------------------------|-----------------------------|-----------------------------|-----------------------------|-----------------------------|-----------------------------|-----------------------------|-----------------------------|-----------------------------|--------|--------------------------------------------------|---------|
|                                                                  | Series A                       |                             | Series B                    |                             | Series A                    |                             | Common Stock                |                             | Additional Paid In Capital  |        | Deficit Accumulated During the Development Stage | Total   |
|                                                                  | Convertible Preferred Stock    | Convertible Preferred Stock | Convertible Preferred Stock | Convertible Preferred Stock | Convertible Preferred Stock | Convertible Preferred Stock | Convertible Preferred Stock | Convertible Preferred Stock | Convertible Preferred Stock |        |                                                  |         |
|                                                                  | Shares                         | Amount                      | Shares                      | Amount                      | Shares                      | Amount                      | Shares                      | Amount                      | Shares                      | Amount |                                                  |         |
| (in thousands, except share and per share amounts)               |                                |                             |                             |                             |                             |                             |                             |                             |                             |        |                                                  |         |
| Issuance of common stock for cash, net of offering costs         |                                |                             |                             |                             |                             |                             | 194,100                     |                             |                             | 7,182  |                                                  | 7,182   |
| Issuance of common stock upon exercise of stock options for cash |                                |                             |                             |                             |                             |                             | 8,248                       |                             |                             | 139    |                                                  | 139     |
| Share-based compensation                                         |                                |                             |                             |                             |                             |                             |                             |                             |                             | 458    |                                                  | 458     |
| Net loss                                                         |                                |                             |                             |                             |                             |                             |                             |                             |                             |        | (8,420)                                          | (8,420) |
| Balance, December 31, 2010                                       |                                |                             |                             |                             |                             |                             | 1,472,422                   | 1                           | 65,080                      |        | (58,310)                                         | 6,771   |
| Issuance of common stock for cash, net of offering costs         |                                |                             |                             |                             |                             |                             | 557,890                     | 1                           | 4,016                       |        |                                                  | 4,017   |
| Issuance of common stock upon exercise of stock options for cash |                                |                             |                             |                             |                             |                             | 188                         |                             |                             |        |                                                  |         |
| Share-based compensation                                         |                                |                             |                             |                             |                             |                             |                             |                             |                             | 308    |                                                  | 308     |
| Net loss                                                         |                                |                             |                             |                             |                             |                             |                             |                             |                             |        | (5,364)                                          | (5,364) |
| Balance, December 31, 2011                                       |                                |                             |                             |                             |                             |                             | 2,030,500                   | 2                           | 69,404                      |        | (63,674)                                         | 5,732   |
| Issuance of common stock for cash, net of offering costs         |                                |                             |                             |                             |                             |                             | 629,815                     | 1                           | 1,188                       |        |                                                  | 1,189   |
| Share-based compensation                                         |                                |                             |                             |                             |                             |                             |                             |                             |                             | 306    |                                                  | 306     |

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|                                                                                                                            |           |            |    |         |          |
|----------------------------------------------------------------------------------------------------------------------------|-----------|------------|----|---------|----------|
| Net loss                                                                                                                   |           |            |    | (4,320) | (4,320)  |
| Balance,<br>December 31,<br>2012                                                                                           |           | 2,660,315  | 3  | 70,898  | (67,994) |
| Issuance of<br>common stock<br>for cash, net of<br>offering costs                                                          |           | 521,066    |    | 1,421   |          |
| Adjustment for<br>fractional<br>shares                                                                                     |           | (64)       |    |         |          |
| Issuance of<br>common stock<br>upon exercise<br>of warrants for<br>cash                                                    |           | 4,245      |    | 12      |          |
| Issuance of<br>Series A<br>convertible<br>preferred<br>stock, net of<br>offering costs                                     | 125,000   |            |    | 17,917  |          |
| Deemed<br>preferred stock<br>dividend for<br>beneficial<br>conversion<br>feature                                           |           |            |    | 2,026   |          |
| Impact of<br>deemed<br>preferred stock<br>dividend for<br>beneficial<br>conversion<br>feature on<br>common<br>stockholders |           |            |    | (2,026) |          |
| Conversion of<br>preferred stock<br>to common<br>stock                                                                     | (100,098) | 10,009,800 | 10 | (10)    |          |
| Costs of<br>issuance of<br>common stock<br>and conversion<br>of preferred<br>stock to<br>common stock                      |           |            |    | (20)    |          |
| Share-based<br>compensation                                                                                                |           |            |    | 125     |          |
| Net loss                                                                                                                   |           |            |    | (3,993) | (3,993)  |

Balance,  
September 30,  
2013

|    |    |        |    |            |    |    |    |        |    |          |    |        |
|----|----|--------|----|------------|----|----|----|--------|----|----------|----|--------|
| \$ | \$ | 24,902 | \$ | 13,195,362 | \$ | 13 | \$ | 90,343 | \$ | (71,987) | \$ | 18,369 |
|----|----|--------|----|------------|----|----|----|--------|----|----------|----|--------|

See accompanying notes to consolidated financial statements

## ARCA BIOPHARMA, INC.

(a development stage enterprise)

## CONSOLIDATED STATEMENTS OF CASH FLOWS

(unaudited)

|                                                                             | Nine Months Ended<br>September 30, |            | Period from<br>December 17, 2001<br>(date of inception)<br>to September 30,<br>2013 |
|-----------------------------------------------------------------------------|------------------------------------|------------|-------------------------------------------------------------------------------------|
|                                                                             | 2013                               | 2012       |                                                                                     |
|                                                                             |                                    |            | (in thousands)                                                                      |
| Cash flows used in operating activities:                                    |                                    |            |                                                                                     |
| Net loss                                                                    | \$ (3,993)                         | \$ (3,480) | \$ (71,987)                                                                         |
| Adjustments to reconcile net loss to net cash used in operating activities: |                                    |            |                                                                                     |
| Gain on patent rights assignment                                            |                                    |            | (2,000)                                                                             |
| Gain on bargain purchase                                                    |                                    |            | (25,282)                                                                            |
| Depreciation and amortization                                               | 24                                 | 34         | 1,804                                                                               |
| Non-cash interest expense                                                   |                                    |            | 211                                                                                 |
| Share-based compensation                                                    | 125                                | 243        | 2,713                                                                               |
| Issuance of warrants for lease termination                                  |                                    |            | 377                                                                                 |
| Accretion of liabilities                                                    |                                    |            | 152                                                                                 |
| Impairment of property and equipment                                        |                                    |            | 125                                                                                 |
| Impairment of in-process research and development                           |                                    |            | 6,000                                                                               |
| Write-off of deferred tax liability                                         |                                    |            | (2,281)                                                                             |
| Gain on marketable securities available for sale                            |                                    |            | (263)                                                                               |
| (Gain) loss from disposal of property and equipment                         |                                    |            | 83                                                                                  |
| Other, net                                                                  |                                    |            | 267                                                                                 |
| Change in operating assets and liabilities (net of amounts acquired):       |                                    |            |                                                                                     |
| Other current assets                                                        | (80)                               | 219        | 2,740                                                                               |
| Other assets                                                                | 44                                 | 60         | 7,370                                                                               |
| Accounts payable                                                            | 382                                | (141)      | (1,743)                                                                             |
| Accrued expenses and other liabilities                                      | 98                                 | (318)      | (19,123)                                                                            |
| Deferred rent                                                               | (16)                               | (25)       |                                                                                     |
| Net cash used in operating activities                                       | (3,416)                            | (3,408)    | (100,837)                                                                           |
| Cash flows (used in) provided by investing activities:                      |                                    |            |                                                                                     |
| Cash received from Merger                                                   |                                    |            | 30,392                                                                              |
| Payment of deferred transaction costs                                       |                                    |            | (1,186)                                                                             |
| Purchase of property and equipment                                          | (28)                               | (1)        | (1,907)                                                                             |
| Proceeds from sale of marketable securities                                 |                                    |            | 15,369                                                                              |
| Proceeds from sale of property and equipment                                |                                    |            | 358                                                                                 |

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|                                                                                           |           |          |           |
|-------------------------------------------------------------------------------------------|-----------|----------|-----------|
| Proceeds from patent rights assignment                                                    |           |          | 2,000     |
| Net cash (used in) provided by investing activities                                       | (28)      | (1)      | 45,026    |
| Cash flows provided by (used in) financing activities:                                    |           |          |           |
| Proceeds from issuance of convertible notes payable and related warrants for common stock |           |          | 10,841    |
| Proceeds from issuance of bank note payable                                               |           |          | 4,000     |
| Proceeds from stock subject to repurchase                                                 |           |          | 38        |
| Proceeds from the issuance of preferred stock                                             | 20,000    |          | 52,316    |
| Payment of preferred stock offering costs                                                 | (2,083)   |          | (2,329)   |
| Proceeds from the issuance of common stock                                                | 1,741     | 953      | 15,850    |
| Payment of common stock offering costs                                                    | (328)     | (272)    | (1,714)   |
| Repayment of principal on bank note payable                                               |           |          | (4,000)   |
| Repayment of principal on convertible notes payables                                      |           |          | (105)     |
| Repayment of principal on vendor finance agreement                                        | (174)     | (134)    | (454)     |
| Net cash provided by (used in) financing activities                                       | 19,156    | 547      | 74,443    |
| Net increase (decrease) in cash and cash equivalents                                      | 15,712    | (2,862)  | 18,632    |
| Cash and cash equivalents, beginning of period                                            | 2,920     | 5,943    |           |
| Cash and cash equivalents, end of period                                                  | \$ 18,632 | \$ 3,081 | \$ 18,632 |
| Supplemental cash flow information:                                                       |           |          |           |
| Interest paid                                                                             | \$ 3      | \$ 3     | \$ 118    |
| Supplemental disclosure of noncash investing and financing transactions:                  |           |          |           |
| Accrued interest on notes payable converted to equity                                     | \$        | \$       | \$ 163    |
| Warrant issued in connection with credit facility                                         | \$        | \$       | \$ 111    |
| Accrued deferred transaction costs                                                        | \$        | \$       | \$ 482    |
| Vendor finance agreement                                                                  | \$        | \$       | \$        |

See accompanying notes to consolidated financial statements

ARCA BIOPHARMA, INC.

(a development stage enterprise)

NOTES TO CONSOLIDATED FINANCIAL STATEMENTS

(unaudited)

(1) The Company and Summary of Significant Accounting Policies

Description of Business

ARCA biopharma, Inc., or the Company or ARCA, a Delaware corporation, is headquartered in Westminster, Colorado and is a biopharmaceutical company principally focused on developing genetically-targeted therapies for cardiovascular diseases. The Company's lead product candidate, Gencaro (bucindolol hydrochloride), a pharmacologically unique beta-blocker and mild vasodilator that ARCA plans to evaluate in a new clinical trial for the treatment of atrial fibrillation, or AF, in patients with heart failure and left ventricular dysfunction, or HFREF. The Company has identified common genetic variations in receptors in the cardiovascular system that it believes interact with Gencaro's pharmacology and may predict patient response to the drug.

The Company plans to test this hypothesis in a Phase 2B/3 clinical trial of Gencaro, known as GENETIC-AF. The AF indication for Gencaro was chosen based on prior clinical data from the previously conducted Phase 3 heart failure (HF) trial of Gencaro in 2,708 HF patients, or the BEST trial, suggest that Gencaro may be successful in reducing or preventing AF. GENETIC-AF is planned as a multi-center, randomized, double-blind clinical trial designed to compare the safety and efficacy of Gencaro to an active comparator in HFREF patients recently diagnosed with persistent AF and having beta-1 389 arginine homozygous genotype, the genotype the Company believes responds most favorably to Gencaro. The primary endpoint of GENETIC-AF is time to recurrent symptomatic AF or all-cause mortality.

ARCA has created an adaptive design for GENETIC-AF which it plans to initiate with a Phase 2B study in approximately 200 HFREF patients. The GENETIC-AF Data Safety Monitoring Board (DSMB) will analyze selected data from the Phase 2B portion of the trial and recommend whether the trial should proceed to Phase 3 and enroll an additional 420 patients. The DSMB will make their recommendation based on analysis of selected trial data after 200 patients have been enrolled and have completed 24 weeks of follow-up, the period for measuring the trial's primary end-point. The interim analysis will focus on available data regarding AF event rates, AF burden, and safety. Should the DSMB interim analysis conclude the data is consistent with the pre-trial statistical assumptions and that the data indicates potential for achieving statistical significance for the Phase 3 endpoint, then the DSMB may recommend the study proceed to Phase 3. The DSMB may also recommend changes to the study design before potentially proceeding to Phase 3, or it may recommend that the study not proceed to Phase 3. The Company, in consultation with the trial's clinical steering committee and the DSMB, will make the final determination on the trial's development steps. The Company believes the Phase 2B portion of the study would take approximately two and one-half years to complete from the time the first patient is enrolled until the planned DSMB interim analysis of data from the initial 200 patients.

The Company has been granted patents in the U.S., Europe, and other jurisdictions for methods of treating AF and HF patients with Gencaro based on genetic testing, which the Company believes may provide market exclusivity for these uses of Gencaro into at least 2026 in the U.S. and into 2025 in Europe. In addition, the Company believes that if Gencaro is approved, a Gencaro patent will be eligible for patent term extension based on our current clinical trial plans which, if granted, may provide market exclusivity for Gencaro into 2029 or 2030 in the U.S. and Europe.

To complete both phases of the GENETIC-AF clinical trial and submit for FDA approval, the Company will need to raise additional funding through public or private equity transactions or a strategic combination or partnership. If the Company is unable to obtain additional funding or is unable to complete a strategic transaction, it may have to discontinue development activities on Gencaro or discontinue its operations.

#### Development Stage Risks, Liquidity and Going Concern

The Company is in the development stage and devotes substantially all of its efforts towards obtaining regulatory approval for Gencaro and raising capital necessary to fund its operations. The Company has not generated revenue to date and is subject to a number of risks similar to those of other development stage companies, including dependence on key individuals, the development of and regulatory approval of commercially viable products, the need to raise adequate additional financing necessary to fund the development and commercialization of its products, and competition from larger companies. The Company has historically funded its operations through issuances of common and preferred stock, as well as through the business combination with Nuvelo, Inc, or Nuvelo.

Since ARCA was founded on December 17, 2001, or Inception, the Company has incurred substantial losses and negative cash flows from operations. Since Inception, the Company incurred a loss from operations of \$103.1 million and had negative cash flows from operations of \$100.8 million.

To support the continued development of Gencaro, the Company completed a public equity offering in June 2013 to initiate the Phase 2B/3 GENETIC-AF trial and fund ongoing operations. In light of the substantial additional time and costs associated with the development of Gencaro, the Company will need to raise a significant amount of capital on acceptable terms to finance the completion of GENETIC-AF and the Company's ongoing operations. The Company currently believes its cash and cash equivalents balance as of September 30, 2013 will be sufficient to fund its operations through at least 2014. However, changing circumstances may cause the Company to consume capital significantly faster or slower than is currently anticipated. These estimates are based upon assumptions that may prove to be wrong, and ARCA could exhaust its available financial resources sooner than currently projected.

The Company's liquidity, and its ability to raise additional capital or complete any strategic transaction, depends on a number of factors, including, but not limited to, the following:

- the costs and timing for the planned GENETIC-AF clinical trial ;
  - the market price of the Company's stock and the availability and cost of additional equity or debt capital;
  - the Company's ability to retain the listing of its common stock on the Nasdaq Capital Market;
  - general economic and industry conditions affecting the availability and cost of capital;
  - the Company's ability to control costs associated with its operations;
  - the costs of filing, prosecuting, defending and enforcing any patent claims and other intellectual property rights; and
  - the terms and conditions of the Company's existing collaborative and licensing agreements.
- The sale of additional equity or convertible debt securities would likely result in additional dilution to the Company's existing stockholders. If the Company raises additional funds through the incurrence of indebtedness, the obligations related to such indebtedness would be senior to rights of holders of the Company's capital stock and could contain covenants that would restrict the Company's operations. The Company also cannot predict what consideration might be available, if any, to the Company or its stockholders, in connection with any strategic transaction. Should strategic alternatives or additional capital not be available to the Company, or not be available on acceptable terms, the Company may be unable to realize value from its assets and discharge its liabilities in the normal course of business which may, among other alternatives, cause the Company to further delay, substantially reduce or discontinue operational activities to conserve its cash resources.

#### Reverse Stock Split

On March 4, 2013, the Company completed a 1-for-6 reverse split of its common stock. All common shares and per common share amounts in the financial statements and footnotes have been adjusted retroactively to reflect the effects of this action.

#### Basis of Presentation

The accompanying unaudited consolidated financial statements of the Company were prepared in accordance with generally accepted accounting principles for interim financial information and instructions to Form 10-Q and Rule 10-01 of Regulation S-X. Accordingly, these financial statements do not include all of the information and footnotes required by accounting principles generally accepted in the United States of America for complete financial statements. In the opinion of management, these financial statements include all normal and recurring adjustments considered necessary for a fair presentation of these interim consolidated financial statements. The results of operations for the nine months ended September 30, 2013 are not necessarily indicative of results expected for the full year ending December 31, 2013. The Company has generated no revenue to date and its activities have consisted of seeking regulatory approval, research and development, exploring strategic alternatives for further developing and commercializing Gencaro, and raising capital. Accordingly, the Company continues to be considered in the development stage at September 30, 2013. These unaudited consolidated financial statements should be read in conjunction with the audited consolidated financial statements and footnotes thereto for the year ended December 31, 2012 included in the Company's Annual Report on Form 10-K filed with the Securities and Exchange Commission, as amended. Amounts presented are rounded to the nearest thousand, where indicated, except per share data and par values.

#### Concentrations of Credit Risk

Financial instruments that potentially subject the Company to significant concentrations of credit risk consist primarily of cash and cash equivalents. The Company has no off-balance-sheet concentrations of credit risk, such as foreign exchange contracts, option contracts, or foreign currency hedging arrangements. The Company maintains cash and cash equivalent balances in the form of bank demand deposits, money market fund accounts and debt securities with financial institutions that management believes are creditworthy. Such balances may at times exceed the insured amount.

## Accrued Expenses

As part of the process of preparing its financial statements, the Company is required to estimate accrued expenses. This process involves identifying services that third parties have performed on the Company's behalf and estimating the level of service performed and the associated cost incurred for these services as of the balance sheet date. Examples of estimated accrued expenses include contract service fees, such as fees payable to contract manufacturers in connection with the production of materials related to the Company's drug product, and professional service fees, such as attorneys, consultants, and clinical research organizations. The Company develops estimates of liabilities using its judgment based upon the facts and circumstances known at the time.

## (2) Loss Per Share

The Company calculates basic loss per share by dividing loss attributable to common stockholders by the weighted average common shares outstanding during the period, excluding common stock subject to vesting provisions. In the three and nine month periods ended September 30, 2013, the calculation of net loss attributable to common stockholders includes the effect of the deemed dividend to Series A Preferred Stock purchasers (see Note 7).

Diluted loss per share is computed by dividing loss attributable to common stockholders by the weighted average number of common shares outstanding during the period increased to include, if dilutive, the number of additional common shares that would have been outstanding if the potential common shares had been issued. The Company's potentially dilutive shares include Series A Convertible Preferred Stock, stock options and warrants for common stock.

A reconciliation of the numerator and denominator used in the calculation of basic and diluted loss per share follows:

| (In thousands, except shares and per share data)                                                                                                                                                                                                                                                                                                 | Three Months<br>Ended September 30, |           | Nine Months<br>Ended September 30, |            |
|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------|-----------|------------------------------------|------------|
|                                                                                                                                                                                                                                                                                                                                                  | 2013                                | 2012      | 2013                               | 2012       |
| Net loss                                                                                                                                                                                                                                                                                                                                         | \$ (1,723)                          | \$ (927)  | \$ (3,993)                         | \$ (3,480) |
| Less: Series A Preferred Stock deemed dividend                                                                                                                                                                                                                                                                                                   |                                     |           | (2,026)                            |            |
| Net loss available to common shareholders                                                                                                                                                                                                                                                                                                        | \$ (1,723)                          | \$ (927)  | \$ (6,019)                         | \$ (3,480) |
| Weighted average shares of common stock outstanding                                                                                                                                                                                                                                                                                              | 10,835,299                          | 2,286,519 | 6,302,611                          | 2,116,462  |
| Less: Weighted-average shares of unvested common stock                                                                                                                                                                                                                                                                                           | (2,783)                             | (2,783)   | (2,783)                            | (2,783)    |
| Total weighted-average shares used in computing net loss per share attributed to common stockholders                                                                                                                                                                                                                                             | 10,832,516                          | 2,283,736 | 6,299,828                          | 2,113,679  |
| Basic and diluted loss per share                                                                                                                                                                                                                                                                                                                 | \$ (0.16)                           | \$ (.41)  | \$ (0.96)                          | \$ (1.65)  |
| Potentially dilutive securities representing 13.1 million and 0.8 million weighted average shares of common stock were excluded for the three months ended September 30, 2013 and 2012, respectively, and potentially dilutive securities representing 6.7 million and 0.7 million weighted average shares of common stock were excluded for the |                                     |           |                                    |            |

nine months ended September 30, 2013 and 2012, respectively, because including them would have an anti-dilutive effect on net loss per share.

(3) Merger with Nuvelo, Inc. on January 27, 2009

On January 27, 2009, ARCA Colorado, Inc. (ARCA Colorado) completed the Merger with Nuvelo in accordance with the terms of the Merger Agreement, in which a wholly-owned subsidiary of Nuvelo merged with and into ARCA Colorado, with ARCA Colorado continuing after the Merger as the surviving corporation and a wholly-owned subsidiary of Nuvelo. Immediately following the Merger, the Company changed its name from Nuvelo, Inc. to ARCA biopharma, Inc., and its common stock began trading on the Nasdaq Global Market under the symbol ABIO on January 28, 2009. On March 7, 2011, the listing of the Company's common stock was transferred from the Nasdaq Global Market to the Nasdaq Capital Market.

The Merger was treated as a reverse merger and accounted for as a business combination using the acquisition method of accounting in accordance with ASC 805. For accounting purposes, ARCA Colorado was considered to have acquired Nuvelo in the Merger, as the stockholders of ARCA Colorado prior to the Merger had a controlling interest in the combined company and the Company's management is the former management of ARCA Colorado. The results of operations and cash flows include the activities of Nuvelo since the date of the Merger. Pursuant to the rules and regulations of the United States Securities and Exchange Commission, or the SEC, the historical financial statements of ARCA Colorado replaced the historical financial statements of Nuvelo, and the disclosures in this report relating to the pre-Merger business of the Company, unless noted as being the business of Nuvelo prior to the Merger, pertain to the business of ARCA Colorado prior to the Merger.

The estimated total acquisition consideration of \$11.9 million to acquire Nuvelo was based on the market capitalization of Nuvelo as of January 27, 2009 and the estimated fair values of its vested stock options and warrants outstanding on that date, as this was deemed the most reliable measure of the consideration effectively transferred to acquire Nuvelo on that date. The Company estimated the net assets acquired in the Merger to be \$37.2 million, including \$45.5 million of cash, cash equivalents and marketable securities. In accordance with ASC 805, any excess of fair value of net assets acquired in a business combination over the acquisition consideration results in a gain on bargain purchase, and as a result, the Company recorded a gain on bargain purchase of \$25.3 million.

#### (4) Fair Value Disclosures

As of September 30, 2013, the Company had \$18.6 million of cash equivalents consisting of money market funds with maturities of 90 days or less. The Company has the ability to liquidate these investments without restriction. The Company determines fair value for these money market funds and equity securities with Level 1 inputs through quoted market prices. There were no transfers of assets between fair value hierarchy levels during the three or nine month period ended September 30, 2013.

Fair value is defined as the price that would be received to sell an asset or paid to transfer a liability in an orderly transaction between market participants at the measurement date (exit price). Inputs used to measure fair value are classified into the following hierarchy:

·Level 1 Unadjusted quoted prices in active markets for identical assets or liabilities

·Level 2 Unadjusted quoted prices in active markets for similar assets or liabilities; unadjusted quoted prices for identical or similar assets or liabilities in markets that are not active; or inputs other than quoted prices that are observable for the asset or liability

·Level 3 Unobservable inputs for the asset or liability

#### Fair Value of Other Financial Instruments

The carrying amount of other financial instruments, including cash, accounts payable, and short-term notes payable approximated fair value due to their short maturities.

#### (5) Property and Equipment

Property and equipment consist of the following (in thousands):

|                        | Estimated Life | September 30,<br>2013 | December 31,<br>2012 |
|------------------------|----------------|-----------------------|----------------------|
| Computer equipment     | 3 years        | \$ 97                 | \$ 104               |
| Lab equipment          | 5 years        | 142                   | 142                  |
| Furniture and fixtures | 5 years        | 86                    | 93                   |
| Computer software      | 3 years        | 176                   | 176                  |
| Leasehold improvements |                | 8                     | 18                   |

Lesser of useful life or life of the  
lease

|                                                |       |       |
|------------------------------------------------|-------|-------|
|                                                | 509   | 533   |
| Less accumulated depreciation and amortization | (482) | (510) |
|                                                | \$ 27 | \$ 23 |

For the nine months ended September 30, 2013 and 2012, and for the period from Inception through September 30, 2013, depreciation and amortization expense was \$24,000, \$34,000, and \$1.8 million respectively.

#### (6) Commitments and Contingencies

The Company has or is subject to the following commitments and contingencies:

##### Employment Agreements

The Company maintains employment agreements with several executive employees. Most of these agreements provide for payments to be made under certain conditions related to a change in control of the Company and entitle the employee to wages and certain benefits payments not exceeding one calendar year from the date of termination without cause or by the employee for good reason. The agreements may be terminated at any time by the Company with or without cause upon written notice to the employee.

## Operating Lease

On February 8, 2008, the Company entered into a lease agreement for approximately 15,000 square feet of newly constructed office facilities in Broomfield, Colorado. The office location served as the Company's primary business office. In June 2011, the Company and its landlord amended the lease mutually agreeing for the Company to relocate to another suite, comprising approximately 4,500 square feet, within the same building. The original five year term of the Lease remained unchanged. The Company's facility lease which would have originally expired in June 2013 was extended by a letter of extension through September 30 2013.

Rent expense under this lease for the nine months ended September 30, 2013 and 2012 was \$44,000 and \$36,000, respectively, and was \$559,000 from Inception through September 30, 2013.

Effective August, 1, 2013 the Company entered into a lease agreement for approximately 5,300 square feet of office facilities in Westminster, Colorado that has served as the Company's primary business office since October 1, 2013. Below is a summary of the future minimum lease payments committed for the Company's facility in Westminster, Colorado as of September 30, 2013:

|                                     |            |
|-------------------------------------|------------|
| 2013                                | \$ 19,000  |
| 2014                                | 78,000     |
| 2015                                | 81,000     |
| 2016                                | 62,000     |
| Total future minimum lease payments | \$ 240,000 |

## Cardiovascular Pharmacology and Engineering Consultants, LLC, or CPEC

Under the terms of its strategic license agreement with CPEC, a licensing subsidiary of Indevus Pharmaceuticals Inc. (a wholly owned subsidiary of Endo Pharmaceuticals), holding ownership rights to certain clinical trial data of Gencaro, the Company will incur milestone and royalty obligations upon the occurrence of certain events. In August 2008, the Company paid CPEC a milestone payment of \$500,000 based on the July 31, 2008 submission of its NDA to the FDA. If the FDA grants marketing approval for Gencaro, the agreement provides that the Company will owe CPEC another milestone payment of \$8.0 million within six months after FDA approval. The agreement also states that the Company has the obligation to make milestone payments of up to \$5.0 million in the aggregate upon regulatory marketing approval in Europe and Japan. The agreement states that the Company's royalty obligation ranges from 12.5% to 25% of revenue from the related product based on achievement of specified product sales levels, including a 5% royalty that CPEC is obligated to pay under its original license agreement for Gencaro. The Company has the right to buy down the royalties to a range of 12.5% to 17% by making a payment to CPEC within six months of regulatory approval.

## (7) Equity Financings and Warrants

### Private Investment in Public Equity (PIPE) Transaction

On January 22, 2013, the Company entered into a Subscription Agreement (the "Purchase Agreement") with various accredited investors and its Chief Executive Officer in connection with a private placement of its common stock and

warrants. Pursuant to the Purchase Agreement, the Company sold an aggregate of 356,430 shares of its common stock and warrants to purchase up to 249,501 additional shares of its common stock for aggregate gross proceeds of approximately \$1 million, before deducting estimated offering expenses payable by the Company. The net proceeds to the Company were approximately \$805,000, and the private placement closed on January 25, 2013.

The common stock and warrants were sold in units consisting of one share of common stock and a warrant to purchase 0.70 shares of common stock. The purchase price for each unit was \$2.81. The warrants were exercisable upon issuance, expire seven years from the date of issuance, and have an exercise price of \$2.28 per share, equal to 100% of the closing bid price of ARCA's common stock on the Nasdaq Capital Market on January 22, 2013.

The Company filed a registration statement for the resale of the shares underlying the units sold in these private placements. That registration statement was declared effective by the Securities and Exchange Commission on February 14, 2013.

In connection with this transaction, the Company agreed that, subject to certain exceptions, it would not, while the warrants are outstanding, effect or enter into an agreement to effect any issuance of common stock or securities convertible into, exercisable for or exchangeable for common stock in a variable rate transaction, which means a transaction in which the Company issues or sells any convertible securities either (A) at a conversion price, exercise price or exchange rate or other price that is based upon and/or varies with the trading prices of, or quotations for, the shares of common stock at any time after the initial issuance of such convertible securities, or (B) with a conversion, exercise or exchange price that is subject to being reset at some future date after the initial issuance of the convertible securities or upon the occurrence of the specified or contingent events directly or indirectly related to our business or the market for our common stock. In addition, the Company agreed that, subject to certain exceptions, if it issues securities within one year following the closing of the offering, each investor would have the right to purchase its pro rata share of a specified portion of the securities in the future offering on the same terms, conditions and price provided for in the proposed issuance of securities.

#### Registered Direct Offering

On January 31, 2013, the Company entered into a subscription agreement with certain institutional investors (the Investors ) in connection with its Registered Direct public offering (the Offering ), pursuant to which the Company sold an aggregate of 164,636 shares of its common stock and warrants to purchase up to 65,855 additional shares of its common stock to the Investors for aggregate gross proceeds of approximately \$730,000, before deducting placement agent fee and other estimated offering expenses payable by the Company. The net proceeds to the Company were approximately \$616,000, and the Offering closed on February 4, 2013.

The common stock and warrants were sold in units consisting of one share of common stock and a warrant to purchase 0.40 shares of common stock. The purchase price for each unit was \$4.43. The warrants were exercisable upon issuance, expire five years from the date of issuance, and have an exercise price of \$4.13 per share, equal to the closing bid price of ARCA's common stock on the Nasdaq Capital Market on January 31, 2013. The Offering was effected as a takedown off the Company's S-3 Registration Statement, which became effective on April 4, 2011, pursuant to a prospectus supplement filed with the Securities and Exchange Commission on February 1, 2013. The warrant agreements provide for settlement of the warrants in unregistered shares should an effective registration statement or current prospectus not be in place at the time a warrant is exercised.

#### Public Offering

On June 4, 2013, the Company sold shares of its Series A Convertible Preferred Stock (Preferred Stock) and warrants to purchase common stock in a public offering for aggregate gross proceeds of \$20.0 million. The Company issued 125,000 shares of Preferred Stock and warrants to purchase up to 6,250,000 shares of common stock at a purchase price of \$160 per share of Preferred Stock. The net proceeds, after deducting placement agent fees and other offering expenses payable by the Company, were approximately \$17.9 million. ARCA's Director and Chief Executive Officer participated in the offering, purchasing 781 shares of Preferred Stock and warrants to purchase 39,050 shares of common stock.

Each share of Preferred Stock is initially convertible into 100 shares of the Company's common stock at any time at the option of the holder; provided, that the holder will be prohibited from converting to the extent that, as a result of such conversion, the holder, together with its affiliates, would beneficially own more than 9.99% of the total number of shares of the Company's common stock then issued and outstanding. Each share of Preferred Stock has a liquidation preference of \$.001 per share. The shares of Preferred Stock have no preferential dividends or redemption rights, and no voting rights except as required by law. As of September 30, 2013, 100,098 shares of the Preferred Stock had been converted into 10,009,800 shares of ARCA common stock. An additional 24,802 shares of Preferred Stock were converted into 2,480,200 shares of common stock between October 1, 2013 and October 31, 2013.

Each purchaser in the offering was issued a warrant to purchase 50 shares of the Company's common stock for each share of Preferred Stock purchased. The warrants have an exercise price of \$1.60 per share, will expire on the five year anniversary of the date of issuance, and were exercisable immediately upon issuance, provided that the holder will be prohibited from exercising the warrants if, as a result of such exercise, the holder, together with its affiliates, would beneficially own more than 9.99% of the total number of shares of common stock then issued and outstanding.

The securities were sold pursuant to a placement agreement and have been registered under the Securities Act of 1933 pursuant to the Company's Registration Statement on Form S-1, as amended (No.333-187508), which was declared effective by the Securities and Exchange Commission on May 29, 2013, and the Preferred Stock and Warrants were offered and sold pursuant to a prospectus dated May 30, 2013.

In connection with the Preferred Stock financing, the Company recorded a non-cash dividend of approximately \$2.0 million to recognize the intrinsic value of the embedded beneficial conversion feature. Typically, such a deemed dividend would be represented as a reduction in a company's retained earnings and an increase in additional paid in capital in recognition of the reapportionment of common shareholder value to the preferred stock purchasers. However, since ARCA has an accumulated deficit, the deemed dividend

is recognized by a reapportionment of additional paid in capital from common shareholders to additional paid in capital of preferred stock purchasers, which are combined in the Company's statement of stockholders' equity.

#### Warrants

As of September 30, 2013, warrants to purchase approximately 8.2 million shares of common stock were outstanding at exercise prices ranging from \$1.60 to \$116.89, with a weighted average exercise price per share of \$2.53. These warrants, which were granted as part of various financing and business agreements, expire at various times between October 2013 and January 2020. Warrants were recorded in additional paid-in capital at their estimated fair market value at the date of grant using a Black-Scholes option-pricing model.

#### (8) Share-based Compensation

For the three and nine month periods ended September 30, 2013 and 2012 and for the period from Inception through September 30, 2013, the Company recognized the following non-cash, share-based compensation expense in the consolidated statement of operations (in thousands):

|                                     | Three Months<br>Ended<br>September 30, |       | Nine Months<br>Ended September<br>30, |        | Period from<br>December 17,<br>2001 (date of<br>inception) to<br>September 30, 2013 |
|-------------------------------------|----------------------------------------|-------|---------------------------------------|--------|-------------------------------------------------------------------------------------|
|                                     | 2013                                   | 2012  | 2013                                  | 2012   |                                                                                     |
| Research and Development            | \$ 7                                   | \$ 25 | \$ 36                                 | \$ 76  | \$ 626                                                                              |
| Selling, General and Administrative | 42                                     | 49    | 89                                    | 167    | 1,700                                                                               |
| Restructuring Expense               |                                        |       |                                       |        | 387                                                                                 |
| Total                               | \$ 49                                  | \$ 74 | \$ 125                                | \$ 243 | \$ 2,713                                                                            |

Stock option transactions for the nine month period ended September 30, 2013 under all plans are as follows:

|                                           | # of<br>Options | Weighted<br>Average<br>Exercise Price | Weighted<br>Average<br>Remaining<br>Contractual<br>Term<br>(in years) |
|-------------------------------------------|-----------------|---------------------------------------|-----------------------------------------------------------------------|
| Options outstanding at December 31, 2012  | 144,019         | \$ 18.28                              | 4.91                                                                  |
| Changes during the period:                |                 |                                       |                                                                       |
| Granted                                   | 731,535         | 1.38                                  |                                                                       |
| Exercised                                 |                 |                                       |                                                                       |
| Forfeited, cancelled or expired           | (39,113)        | 11.07                                 |                                                                       |
| Options outstanding at September 30, 2013 | 836,441         | \$ 3.84                               | 9.39                                                                  |
| Options exercisable at September 30, 2013 | 115,435         | \$ 18.08                              | 6.12                                                                  |
| Options vested and expected to vest       | 721,216         | \$ 4.20                               | 9.31                                                                  |

(9) Income Taxes

In accordance with United States Generally Accepted Accounting Principles, a valuation allowance should be provided if it is more likely than not that some or all of the Company's deferred tax assets will not be realized. The Company's ability to realize the benefit of its deferred tax assets will depend on the generation of future taxable income. Due to the uncertainty of future profitable operations and taxable income, the Company has recorded a full valuation allowance against its net deferred tax assets. The Company believes its tax filing positions and deductions related to tax periods subject to examination will be sustained upon audit and, therefore, has no reserve for uncertain tax positions.

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

This Management's Discussion and Analysis of Financial Condition and Results of Operations contains

forward-looking statements within the meaning of Section 27A of the Securities Act of 1933, as amended, Section 21E of the Securities Exchange Act of 1934, as amended and the Private Securities Litigation Reform Act of 1995. These statements include, but are not limited to, statements regarding the Company's anticipated timing for initiation or completion of its clinical trials for any of its product candidates; the potential for Gencaro to be an effective potential treatment for atrial fibrillation and, the Company's ability to fund future operations. Such statements are based on management's current expectations and involve risks and uncertainties. Actual results and performance could differ materially from those projected in the forward-looking statements as a result of many factors, including, without limitation, the risks and uncertainties associated with: the Company's financial resources and whether they will be sufficient to meet the Company's business objectives and operational requirements; the Company's ability to complete a strategic transaction to support the continued development of Gencaro, and/or obtain additional financing; the Company's anticipated timing for initiation or completion of its clinical trials for any of its product candidates; the Company's ability to identify, develop and achieve commercial success for products and technologies; drug discovery and the regulatory approval process; estimated timelines for regulatory filings and the implications of interim or final results of the Company's clinical trials; the extent to which the Company's issued and pending patents may protect its products and technology; the potential of the Company's clinical development program to lead to the approval of the Company's New Drug Application for Gencaro; and, the impact of competitive products and technological changes. Actual results and performance could differ materially from those projected in the forward-looking statements as a result of many factors discussed herein and elsewhere. These and other factors are identified and described in more detail in ARCA's filings with the SEC, including without limitation the Company's annual report on Form 10-K for the year ended December 31, 2012, as amended, the Company's Registration Statement on Form S-1 (Registration No. 333-187508), and subsequent filings. Forward-looking statements may be identified by words including will, plan, anticipate, believe, intend, estimates, expect, should, may, potential and similar expressions. The Company disclaims any intent or obligation to update these forward-looking statements.

The terms ARCA, we, us, our and similar terms refer to ARCA biopharma, Inc.

### Overview

We are a biopharmaceutical company whose principal focus is developing genetically-targeted therapies for cardiovascular diseases. Our lead product candidate is Gencaro (bucindolol hydrochloride), a pharmacologically unique beta-blocker and mild vasodilator that we plan to evaluate in a new clinical trial for the treatment of atrial fibrillation, or AF, in patients with heart failure and left ventricular dysfunction, or HFREF. We have identified common genetic variations in receptors in the cardiovascular system that we believe interact with Gencaro's pharmacology and may predict patient response to the drug.

AF is a disorder in which the normally regular and coordinated contraction pattern of the heart's two small upper chambers (the atria) becomes irregular and uncoordinated. The irregular contraction pattern associated with AF causes blood to pool in the atria, predisposing the formation of clots potentially resulting in stroke. AF is considered an epidemic cardiovascular disease with an estimated prevalence of at least 2.7 million Americans in 2010. The approved therapies for the treatment or prevention AF have certain disadvantages in HFREF patients, such as toxic or cardiovascular adverse effects, and most of the approved drugs for AF are contra indicated or have warnings in their prescribing information for such patients. We believe there is an unmet medical need for new AF treatments that have fewer side effects than currently available therapies and are more effective, particularly in HFREF patients.

GENETIC-AF is planned as a multi-center, randomized, double-blind clinical trial designed to compare the safety and efficacy of Gencaro to an active comparator in HFREF patients recently diagnosed with persistent AF and having beta-1 389 arginine homozygous genotype, the genotype we believe responds most favorably to Gencaro. The

primary endpoint of GENETIC-AF, time to recurrent symptomatic AF or all-cause mortality, will be measured over a twenty-four week period after the patient's AF has been electrically cardioverted through the administration of a direct current shock to restore normal heart rhythm.

We have created an adaptive design for GENETIC-AF which we plan to initiate with a Phase 2B study in approximately 200 HFREF patients with recent onset, persistent AF who have a genetic variant of the beta-1 adrenergic receptor that we believe responds most favorably to Gencaro. In addition to measuring the primary endpoint of recurrent symptomatic AF or all-cause mortality, an additional efficacy measure in the Phase 2B portion of GENETIC-AF will be AF burden, defined as a patient's percentage of time in AF per day, regardless of symptoms. All 200 patients in the Phase 2B portion of the trial will have AF burden measured by continuous monitoring, either by previously implanted cardiac resynchronization or defibrillation devices, or newly or previously inserted loop recorders. The GENETIC-AF Data Safety Monitoring Board (DSMB) will analyze selected data from the Phase 2B portion of the trial and recommend whether the trial should proceed to Phase 3 and enroll an additional 420 patients. The DSMB will make their recommendation based on analysis of selected trial data after 200 patients have been enrolled and have completed 24 weeks of follow-up, the period for measuring the trial's primary end-point. The interim analysis will focus on available data regarding AF event rates,

AF burden, and safety. Should the DSMB interim analysis conclude the data is consistent with the pre-trial statistical assumptions and that the data indicates potential for achieving statistical significance for the Phase 3 endpoint, then the DSMB may recommend the study proceed to Phase 3. The DSMB may also recommend changes to the study design before potentially proceeding to Phase 3, or it may recommend that the study not proceed to Phase 3. The Company, in consultation with the trial's clinical steering committee and the DSMB, will make the final determination on the trial's development steps. The Company believes the Phase 2B portion of the study would take approximately two and one-half years to complete from the time the first patient is enrolled until the planned DSMB interim analysis of data from the initial 200 patients. The trial is designed to compare Gencaro to the beta-blocker Toprol XL in patients with the beta-1 389 arginine homozygous genotype, which we believe responds most favorably to Gencaro. We believe data from the BEST trial indicate that Gencaro may have a genetically regulated effect in reducing or preventing AF, whereas we believe the therapeutic benefit of Toprol XL does not appear to be enhanced in patients with this genotype. A retrospective analysis of data from the BEST trial shows that the entire cohort of patients in the BEST trial treated with Gencaro had a 41% reduction in the risk of new onset AF (time-to-event) compared to placebo ( $p = 0.0004$ ). In the BEST DNA substudy, patients with the beta-1 389 arginine homozygous genotype experienced a 74% ( $p = 0.0003$ ) reduction in risk of AF when receiving Gencaro, based on the same analysis. The beta-1 389 arginine homozygous genotype was present in about 47% of the patients in the BEST pharmacogenetic substudy, and we estimate it is present in about 50% of the US general population.

Medtronic, Inc., a leader in medical technologies to improve the treatment of chronic diseases including cardiac rhythm disorders, has signed an agreement with us to collaborate on the GENETIC-AF trial. Under the collaboration with Medtronic, ARCA plans to conduct a substudy that will include continuous monitoring of the cardiac rhythms of all 200 patients enrolled during the Phase 2B portion, and an additional 100 patients in the Phase 3 portion, of GENETIC-AF. The collaboration will be administered by a joint ARCA-Medtronic committee. Medtronic will use its proprietary CareLink System to collect and analyze the cardiac rhythm data from the implanted Medtronic devices and provide the data to us at the close of the Phase 2B portion of the trial. Medtronic will support the reimbursement process for patients enrolled in the Phase 2B portion, and will provide financial support of unreimbursed costs for a certain number of patients in the Phase 2B portion up to a certain maximum amount per patient. If GENETIC-AF proceeds to Phase 3, we will seek to enroll an additional 100 patients in the substudy, and Medtronic will provide the agreed-on CareLink System cardiac rhythm data collection and analysis for the Phase 3 portion of the substudy, and support the reimbursement process.

We have been granted patents in the U.S., Europe, and other jurisdictions for methods of treating AF and HF patients with Gencaro based on genetic testing, which we believe may provide market exclusivity for these uses of Gencaro into at least 2026 in the U.S. and into 2025 in Europe. In addition, we believe that if Gencaro is approved, a Gencaro patent will be eligible for patent term extension based on our current clinical trial plans which, if granted, may provide market exclusivity for Gencaro into 2029 or 2030 in the U.S. and Europe.

To support the continued development of Gencaro, we completed a public equity offering in June 2013 to initiate the Phase 2B/3 GENETIC-AF trial and fund ongoing operations. In light of the substantial additional time and costs associated with the development of Gencaro, we will need to raise a significant amount of capital on acceptable terms to finance the completion of GENETIC-AF and our ongoing operations. We anticipate that our current cash and cash equivalents will be sufficient to fund our operations through at least the end of 2014. However, changing circumstances may cause us to consume capital significantly faster or slower than we currently anticipate. We have based these estimates on assumptions that may prove to be wrong, and we could exhaust our available financial resources sooner than we currently anticipate.

## Results of Operations

### Research and Development Expenses

Research and development, or R&D, expense is comprised of clinical, regulatory, and manufacturing process development activities and costs. Our R&D expense continues to be almost entirely generated by our activities relating to the development of Gencaro. Research and Development expense for the three months ended September 30, 2013 was \$714,000 compared to \$156,000 for the corresponding period of 2012, an increase of approximately \$558,000. R&D expense was \$1,141,000 for the nine months ended September 30, 2013 as compared to \$901,000 for the corresponding period of 2012, an increase of \$240,000.

Clinical expense increased approximately \$134,000 for the three months ended September 30, 2013 and decreased approximately \$37,000 for the nine months ended September 30, 2013. The increase in the three month period is primarily due to increase personnel costs as we have added staff to initiate and oversee our GENETIC-AF clinical trial. The decrease in the nine-month period was primarily due to reduced personnel costs from staff furloughs implemented in the third quarter of 2012.

Regulatory and manufacturing process costs increased \$422,000 for the three months and \$277,000 for the nine months ended September 30, 2013 compared to the corresponding periods of 2012. The increases in both the three and nine month periods ended September 30, 2013 compared to the corresponding period of 2012 is primarily due to costs of initiating production of clinical trial drug materials as well as regulatory audits of suppliers to be used in our GENETIC-AF clinical trial. A portion of the increase is also attributable to increased personnel costs as we have increased staff during the quarter to prepare and initiate our clinical trial.

R&D expenses for the remainder of 2013 are expected to increase from those of the first nine months of 2013 as we initiate our GENETIC-AF clinical trial.

#### Selling, General and Administrative Expenses

Selling, general and administrative expenses, or SG&A, primarily consist of personnel costs, consulting and professional fees, insurance, facilities and depreciation expenses, and various other administrative costs.

SG&A expense was \$1.0 million for the three months ended September 30, 2013 as compared to \$772,000 for the corresponding period in 2012, an increase of \$239,000. The increase in the three months ended September 30, 2013 as compared to the corresponding period of 2012 is comprised of increased legal, accounting and other professional services of approximately \$133,000 primarily related to our proxy report, annual shareholder meeting, and quarterly financial reporting activities, along with increased personnel costs and consulting fees of approximately \$98,000 as we returned personnel from furlough to support initiating our GENETIC-AF clinical trial.

For the nine month period ended September 30, 2013 SG&A expense was approximately \$2.9 million as compared to approximately \$2.6 million for the corresponding period of 2012, an increase of \$273,000. The change included increased costs for legal, consulting, and other professional services of approximately \$307,000 primarily attributable to the special shareholder meeting held in the first quarter of 2013 and in support of our financing efforts completed in the first half of the year. These increases were offset by reduced board of director advisory costs of approximately \$32,000 attributable to cost reduction efforts implemented in the third quarter of 2012.

SG&A expenses for the remainder of 2013 are expected to increase from those of the first nine months of 2013 as we increase our activities to support initiation of our GENETIC-AF clinical trial.

#### Interest and Other Income

Interest and other income was \$2,000 in the three months and \$3,000 in the nine months ended September 30, 2013.

Interest and other income for the comparative three month and nine month periods ended September 30, 2012 were the same. We expect interest income to continue to be nominal for the remainder of 2013 due to low investment yields and utilizing our cash and cash equivalents to fund our operations

#### Interest and Other Expense

Interest and other expense was less than \$1,000 in the three months ended September 30, 2013, compared to \$1,000 for the three months ended September 30, 2012. Interest and other expense was \$3,000 for the nine month periods ended September 30, 2013 and 2012. Based on our current capital structure, interest expense for the remainder of 2013 is expected to be minimal.

#### Liquidity and Capital Resources

##### Cash and Cash Equivalents

|                              | September 30,<br>2013 | December 31,<br>2012 |
|------------------------------|-----------------------|----------------------|
| Cash and cash equivalents \$ | 18,632                | \$ 2,920             |

As of September 30, 2013, we had total cash and cash equivalents of approximately \$18.6 million, as compared to \$2.9 million as of December 31, 2012. The net increase of \$15.7 million in the nine month period reflects the \$19.3 million of net proceeds from our stock offerings completed, less approximately \$3.4 million of cash used to fund operating activities and approximately \$174,000 in payments on a vendor financing arrangement during the nine months ended September 30, 2013.

## Cash Flows from Operating, Investing and Financing Activities

|                                                      | Nine Months Ended     |            |
|------------------------------------------------------|-----------------------|------------|
|                                                      | September 30,<br>2013 | 2012       |
| Net cash (used in) provided by:                      |                       |            |
| Operating activities                                 | \$ (3,416)            | \$ (3,408) |
| Investing activities                                 | (28)                  | (1)        |
| Financing activities                                 | 19,156                | 547        |
| Net increase (decrease) in cash and cash equivalents | \$ 15,712             | \$ (2,862) |

Net cash used in operating activities for the nine months ended September 30, 2013 increased approximately \$8,000 compared with the same period in 2012 primarily due to increased expenses discussed above.

Net cash used in investing activities for the nine months ended September 30, 2013 was approximately \$28,000 representing investment in leasehold improvements associated with our office relocation compared to \$1,000 used in investing activities in the nine months ended September 30, 2012.

Net cash provided by financing activities was \$19.2 million for the nine months ended September 30, 2013 representing \$19.3 million of net proceeds from three equity financings completed during the period, less \$174,000 in payments on a vendor finance agreement. Net cash provided by financing activities of \$547,000 for the nine months ended September 30, 2012 was comprised of \$741,000 of net proceeds from our registered direct offering completed in August 2012, less approximately \$134,000 for payments on a vendor financing arrangement and approximately \$60,000 of costs incurred in 2012 for preparing and filing a registration statement for an equity financing transaction completed in December 2011.

## Sources and Uses of Capital

Our primary sources of liquidity to date have been capital raised from issuances of shares of our preferred and common stock and funds provided by the merger with Nuvelo. The primary uses of our capital resources to date have been to fund operating activities, including research, clinical development and drug manufacturing expenses, license payments, and spending on capital items.

We have completed three equity-financing transactions in 2013 and raised approximately \$19.3 million, net of offering costs. On January 22, 2013, we sold approximately \$1 million of our common stock and warrants for common stock in a private placement transaction with accredited investors including our Chief Executive Officer. We issued 356,430 shares of common stock together with warrants to purchase 249,501 shares of common stock. The net proceeds, after deducting placement agent fees and other offering expenses, were approximately \$805,000. Each unit, consisting of a share of common stock and a warrant to purchase 0.70 shares of common stock, was sold at a purchase price of \$2.81 per unit. The warrants were exercisable upon issuance, expire seven years from the date of issuance, and have an exercise price of \$2.28 per share. Pursuant to the terms of the Registration Rights Agreements (the Rights Agreements) entered into as part of this and prior Private Placement transactions, we filed a registration statement for the resale of the shares underlying the units sold in these private placements. That registration statement was declared effective by the Securities and Exchange Commission on February 14, 2013.

On January 31, 2013, we sold approximately \$730,000 of ARCA's common stock and warrants for common stock in a Registered Direct Offering in which we issued 164,636 shares of common stock and warrants to purchase 65,855

shares of common stock. The net proceeds, after deducting placement agent fees and other offering expenses payable by us, was approximately \$616,000. Each unit, consisting of a share of common stock and a warrant to purchase 0.40 shares of common stock, was sold at a purchase price of \$4.43 per unit. The warrants were exercisable upon issuance, expire five years from the date of issuance, and have an exercise price of \$4.13 per share. The Registered Direct Offering was effected pursuant to a prospectus supplement filed with the Securities and Exchange Commission on February 1, 2013. The warrant agreements provide for settlement of the warrants in unregistered shares should an effective registration statement or current prospectus not be in place at the time a warrant is exercised.

On June 4, 2013, we sold shares of our Series A Convertible Preferred Stock (Preferred Stock) and warrants to purchase common stock in a public offering for aggregate gross proceeds of \$20 million. We issued 125,000 shares of Preferred Stock and warrants to up to purchase 6,250,000 shares of common stock at a purchase price of \$160 per share of Preferred Stock. The net proceeds, after deducting placement agent fees and other offering expenses payable by us, were approximately \$17.9 million. Each share of Preferred Stock is convertible into 100 shares of the Company's Common Stock at any time at the option of the holder. As of November 8, 2013, 100 shares of the Preferred Stock remained outstanding as 124,900 shares of the Preferred Stock had been converted into 12,490,000 shares of common stock since June 4, 2013. The Warrants have an exercise price of \$1.60 per share, will expire on the five year anniversary of the date of issuance, and were exercisable immediately upon issuance. Our Chief Executive

Officer participated in the offering, purchasing 781 shares of Preferred Stock and warrants to purchase 39,050 shares of common stock.

We believe these financings have positioned us to initiate our GENETIC-AF Phase 2B/3 clinical trial for which we currently anticipate initiating patient enrollment in the first quarter of 2014. Our ability to execute our GENETIC-AF Phase 2B trial in accordance with our projected time line depends on a number of factors, including, but not limited to, the following:

- recruitment and formation of key oversight committees;
- selection and successfully entering into agreements with clinical research organizations for managing the clinical trial;
- recruitment of sufficient clinical trial sites and enrollment of patients;
- our ability to control costs associated with the clinical trial and our operations;
- our ability to retain the listing of our common stock on the Nasdaq Capital Market;
- the market price of our stock and the availability and cost of additional equity capital from existing and potential new investors; general economic and industry conditions affecting the availability and cost of capital;
- the costs of filing, prosecuting, defending and enforcing any patent claims and other intellectual property rights; and
- the terms and conditions of our existing collaborative and licensing agreements.

The sale of additional equity or convertible debt securities will be necessary for us to complete both Phase 2B and Phase 3 of the GENETIC-AF clinical trial and submit for FDA approval of Gencaro. Such financing would likely result in additional dilution to our existing stockholders. If we raise additional funds through the incurrence of indebtedness, the obligations related to such indebtedness would be senior to rights of holders of our capital stock and could contain covenants that would restrict our operations. We anticipate that our current cash and cash equivalents will be sufficient to fund our operations through at least the end of 2014. However, our forecast of the period of time through which our financial resources will be adequate to support our current and forecasted operations could vary materially.

#### Critical Accounting Policies and Estimates

A critical accounting policy is one that is both important to the portrayal of our financial condition and results of operation and requires our 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. Our significant accounting policies are described in Note 1 of Notes to Consolidated Financial Statements included within our 2012 Annual Report on Form 10-K filed with the Securities and Exchange Commission. Following is a discussion of the accounting policies that we believe involve the most difficult, subjective or complex judgments and estimates.

## Accrued Expenses

As part of the process of preparing our financial statements, we are required to estimate accrued expenses. This process involves identifying services that third parties have performed on our the Company's behalf and estimating the level of service performed and the associated cost incurred for these services as of the balance sheet date. Examples of estimated accrued expenses include contract service fees, such as fees payable to contract manufacturers in connection with the production of materials related to our drug product, and professional service fees, such as attorneys, consultants, and clinical research organizations. We develop estimates of liabilities using our judgment based upon the facts and circumstances known at the time.

## Share-based Compensation

Our share-based compensation cost recognized includes: (a) compensation costs for current period vesting of all share-based awards granted prior to January 1, 2006, based on the intrinsic value method, and (b) compensation cost for current period vesting of all share-based awards granted or modified subsequent to January 1, 2006, based on the estimated grant date fair value. We recognize compensation costs for our share-based awards on a straight-line basis over the requisite service period for the entire award, as adjusted for expected forfeitures.

From Inception through December 31, 2005, we accounted for issuances of share-based compensation under the intrinsic-value-based method of accounting. Under this method, compensation expense is generally recorded on the date of grant only if the estimated fair value of the underlying stock exceeds the exercise price.

#### Off-Balance Sheet Arrangements

We have not participated in any transactions with unconsolidated entities, such as special purpose entities, which would have been established for the purpose of facilitating off-balance sheet arrangements.

#### Indemnifications

In the ordinary course of business, we enter into contractual arrangements under which we may agree to indemnify certain parties from any losses incurred relating to the services they perform on our behalf or for losses arising from certain events as defined within the particular contract. Such indemnification obligations may not be subject to maximum loss clauses. We have entered into indemnity agreements with each of our directors, officers and certain employees. Such indemnity agreements contain provisions, which are in some respects broader than the specific indemnification provisions contained in Delaware law. We also maintain an insurance policy for our directors and executive officers insuring against certain liabilities arising in their capacities as such.

#### ITEM 3. QUANTITATIVE AND QUALITATIVE DISCLOSURES ABOUT MARKET RISK

Not applicable.

#### ITEM 4. CONTROLS AND PROCEDURES

##### Evaluation of Disclosure Controls and Procedures

We maintain disclosure controls and procedures that are designed to ensure that information required to be disclosed in the reports that we file under the Securities Exchange Act of 1934 is recorded, processed, summarized and reported within the time periods specified in the rules and forms of the SEC, and that such information is accumulated and communicated to our management, including our Chief Executive Officer and Chief Financial Officer, as appropriate, to allow timely decisions regarding required disclosures. In designing and evaluating the disclosure controls and procedures, management recognized that any controls and procedures, no matter how well designed and operated, can provide only reasonable assurance of achieving the desired control objectives, and management necessarily was required to apply its judgment in evaluating the cost-benefit relationship of possible controls and procedures.

As required by Rule 13a-15(b) of the Securities Exchange Act of 1934, an evaluation was carried out under the supervision and with the participation of management, including our Chief Executive Officer and Chief Financial Officer, of the effectiveness of our disclosure controls and procedures as of the end of the quarter covered by this report. Based on the foregoing, our Chief Executive Officer and Chief Financial Officer concluded that our disclosure controls and procedures were effective at a reasonable level of assurance.

##### Changes in Internal Control over Financial Reporting

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

PART II. OTHER INFORMATION

ITEM 1. LEGAL PROCEEDINGS

None

Item 1A. Risk Factors

An investment in ARCA's securities involves certain risks, including those set forth below and elsewhere in this report. In addition to the risks set forth below and elsewhere in this report, other risks and uncertainties not known to ARCA, that are beyond its control or that ARCA deems to be immaterial may also materially adversely affect ARCA's business operations. You should carefully consider the risks described below as well as other information and data included in this report.

Risks Related to Our Business and Financial Condition

Our management and our independent registered public accountant, in their report on our financial statements as of and for the year ended December 31, 2012, have concluded that due to our need for additional capital, and the uncertainties surrounding our ability to raise such funding, substantial doubt exists as to our ability to continue as a going concern.

Our audited consolidated financial statements for the fiscal year ended December 31, 2012 and our unaudited condensed consolidated financial statements for the nine months ended September 30, 2013, were prepared assuming that we will continue as a going concern. The going concern basis of presentation assumes that we will continue in operation for the foreseeable future and will be able to realize our assets and discharge our liabilities and commitments in the normal course of business and do not include any adjustments to reflect the possible future effects on the recoverability and classification of assets or the amounts and classification of liabilities that may result from our inability to continue as a going concern. Our management and our independent registered public accountants concluded as of December 31, 2012 that due to our need for additional capital, and the uncertainties surrounding our ability to raise such funding, substantial doubt exists as to our ability to continue as a going concern. In June 2013 the Company raised aggregate net proceeds of \$17.9 million positioning ARCA to initiate the phase 2B portion of GENETIC-AF and we now anticipate that our current cash and cash equivalents will be sufficient to fund our operations through at least 2014. However, changing circumstances may cause us to consume capital significantly faster or slower than we currently anticipate.

We have based these estimates on assumptions that may prove to be wrong, and we could exhaust our available financial resources sooner than we currently anticipate. We may be forced to reduce our operating expenses and raise additional funds to meet our working capital needs, principally through the additional sales of our securities or debt financings. However, we cannot guarantee that we will be able to obtain sufficient additional funds when needed or that such funds, if available, will be obtainable on terms satisfactory to us. If we are unable to raise sufficient additional capital or complete a future strategic transaction, we may be unable to continue to fund our operations, develop Gencaro or our other product candidates, or realize value from our assets and discharge our liabilities in the normal course of business. If we cannot raise sufficient funds, we may have to liquidate our assets, and might realize significantly less than the values at which they are carried on our financial statements, and stockholders may lose all or part of their investment in our common stock or Preferred Stock.

We will need to raise substantial additional funds through public or private equity transactions and/or complete one or more strategic transactions, to continue development of Gencaro. If we are unable to raise such financing or complete such a transaction, we may not be able to continue operations.

In light of the expected development timeline to potentially obtain FDA approval for Gencaro, if at all, the substantial additional costs associated with the development of Gencaro, including the costs associated with the planned GENETIC-AF clinical trial, and the substantial cost of commercializing Gencaro, if it is approved, we will need to raise substantial additional funding through public or private equity transactions or a strategic combination or partnership. If we are delayed in obtaining funding or are unable to complete a strategic transaction, we may discontinue our development activities on Gencaro or discontinue our operations. Even if we are able to fund continued development and Gencaro is approved, we expect that we will need to complete a strategic transaction or raise substantial additional funding through public or private debt or equity securities to successfully commercialize Gencaro.

We believe our cash and cash equivalents balance as of September 30, 2013 will be sufficient to fund our operations, at our projected cost structure, through at least 2014. Changing circumstances may cause us to consume capital significantly faster or slower than we currently anticipate. We have based these estimates on assumptions that may prove to be wrong, and we could exhaust our available financial resources sooner than we currently anticipate.

Our liquidity, and our ability to raise additional capital or complete any strategic transaction, depends on a number of factors, including, but not limited to, the following:

- the costs and timing for additional clinical trials in order to gain possible FDA approval for Gencaro;
- the market price of our stock and the availability and cost of additional equity capital from existing and potential new investors;
- our ability to retain the listing of our common stock on the Nasdaq Capital Market;
- general economic and industry conditions affecting the availability and cost of capital;
- our ability to control costs associated with our operations;
- the costs of filing, prosecuting, defending and enforcing any patent claims and other intellectual property rights; and
- the terms and conditions of our existing collaborative and licensing agreements.

The sale of additional equity or convertible debt securities would likely result in substantial dilution to our stockholders. If we raise additional funds through the incurrence of indebtedness, the obligations related to such indebtedness would be senior to rights of holders of our capital stock and could contain covenants that would restrict our operations. We also cannot predict what consideration might be available, if any, to us or our stockholders, in connection with any strategic transaction. Should strategic alternatives or additional capital not be available to us in the near term, or not be available on acceptable terms, we may be unable to realize value from our assets and discharge our liabilities in the normal course of business which may, among other alternatives, cause us to further delay, substantially reduce or discontinue operational activities to conserve our cash resources.

If we are not able to successfully develop, obtain FDA approval for and provide for the commercialization of Gencaro in a timely manner, we may not be able to continue our business operations.

We currently have no products that have received regulatory approval for commercial sale. The process to develop, obtain regulatory approval for and commercialize potential product candidates is long, complex and costly. We plan to conduct a Phase 2B/Phase 3 clinical study of Gencaro in 200 hundred HFREF patients with AF initially, and it could expand to approximately 620 HFREF patients with AF. Clinical trials are typically lengthy, complex and expensive and we do not currently have the resources to fully fund such a trial.

Failure to demonstrate that a product candidate, particularly Gencaro, is safe and effective, or significant delays in demonstrating such safety and efficacy, would adversely affect our business. Failure to obtain marketing approval of Gencaro from appropriate regulatory authorities, or significant delays in obtaining such approval, would also adversely affect our business and could, among other things, preclude us from completing a strategic transaction or obtaining additional financing necessary to continue as a going concern.

Even if approved for sale, a product candidate must be successfully commercialized to generate value. We do not currently have the capital resources or management expertise to commercialize Gencaro and, as a result, will need to complete a strategic transaction, or, alternatively, raise substantial additional funds to enable commercialization of Gencaro, if it is approved. Failure to successfully provide for the commercialization of Gencaro, if it is approved, would damage our business.

Our clinical trials for our product candidates may not yield results that will enable us to further develop our products and obtain the regulatory approvals necessary to sell them.

We will receive regulatory approval for our product candidates only if we can demonstrate in carefully designed and conducted clinical trials that the product candidate is safe and effective. We do not know whether any future clinical trials, including the planned GENETIC-AF clinical trial for Gencaro, will demonstrate sufficient safety and efficacy to

obtain the requisite regulatory approvals or will result in marketable products. For example, GENETIC-AF is designed to be an adaptive trial. If we do not see sufficient efficacy and safety in the Phase 2B portion of the trial, we will not complete the Phase 3 portion of the trial. Clinical trials are lengthy, complex and expensive processes with uncertain results. We have spent, and expect to continue to spend, significant amounts of time and money in the clinical development of our product candidates. We have never conducted a Phase 2 or Phase 3 clinical trial and do not currently have sufficient staff with the requisite experience to do so, and we therefore expect that we will have to rely on contract research organizations to conduct certain of our clinical trials. While certain of our employees have experience in designing and administering clinical trials, these employees have no such experience since being with us.

The results we obtain in preclinical testing and early clinical trials may not be predictive of results that are obtained in later studies. We may suffer significant setbacks in advanced clinical trials, even after seeing promising results in earlier studies. Based on results at any stage of clinical trials, we may decide to repeat or redesign a trial or discontinue development of one or more of our product candidates. If we fail to adequately demonstrate the safety and efficacy of our products under development, we will not be

able to obtain the required regulatory approvals to commercialize our product candidates, and our business, results of operations and financial condition would be materially adversely affected.

Administering our product candidates to humans may produce undesirable side effects. These side effects could interrupt, delay or halt clinical trials of our product candidates and could result in the FDA or other regulatory authorities denying approval of our product candidates for any or all targeted indications.

If clinical trials for a product candidate are unsuccessful, we will be unable to commercialize the product candidate. If one or more of our clinical trials are delayed, we will be unable to meet our anticipated development timelines. Either circumstance could cause the market price of our common stock to decline.

We are relying on contract research organizations (CROs) to conduct substantial portions of our GENETIC AF clinical trial, and as a result, we will be unable to directly control the timing, conduct and expense of the clinical trial.

We do not currently have sufficient staff with the requisite experience to conduct our clinical trial and are therefore relying primarily on third parties to conduct our clinical trial. For example, we have contracted with Duke Clinical Research Institute (DCRI), as our CRO to conduct the clinical component of our GENETIC-AF trial. As a result of this contract, we will have less control over many details and steps of the trial, the timing and completion of the trial, the required reporting of adverse events and the management of data developed through the trial than would be the case if we were relying entirely upon our own staff. Communicating with outside parties can also be challenging, potentially leading to mistakes as well as difficulties in coordinating activities. Outside parties, such as CROs, may have staffing difficulties, may undergo changes in priorities or may become financially distressed, adversely affecting their willingness or ability to conduct our trial. We may experience unexpected cost increases that are beyond our control. Problems with the timeliness or quality of the work of a CRO may lead us or any strategic partner to seek to terminate the relationship and use an alternative service provider. However, making this change may be costly and may delay ongoing trials, and contractual restrictions may make such a change difficult or impossible. Additionally, it may be impossible to find a replacement organization that can conduct clinical trials in an acceptable manner and at an acceptable cost.

Even though we are using a CRO to conduct our clinical trial, we have to devote substantial resources and rely on the expertise of our employees to manage the work being done by the CRO. We have never conducted a clinical trial and the inability of our current staff to adequately manage any CRO that we engage may exacerbate the risks associated with relying on a CRO.

If we encounter difficulties enrolling patients in our clinical trials, our trials could be delayed or otherwise adversely affected.

The GENETIC-AF clinical trial for Gencaro requires that we identify and enroll a large number of patients with the condition under investigation and the trial will enroll only those patients having a specific genotype. Because of the rigorous enrollment criteria, we may not be able to enroll a sufficient number of patients to complete our clinical trial in a timely manner.

Patient enrollment is affected by factors including:

- design of the protocol;
- the size of the patient population;
- eligibility criteria for the study in question;

perceived risks and benefits of the drug under study;  
availability of competing therapies, including the off-label use of therapies approved for related indications;  
efforts to facilitate timely enrollment in clinical trials;  
the success of our personnel in making the arrangements with potential clinical trial sites necessary for those sites to begin enrolling patients;  
patient referral practices of physicians;  
availability of clinical trial sites; and  
other clinical trials seeking to enroll subjects with similar profiles.

If we have difficulty enrolling a sufficient number of patients to conduct our clinical trials as planned, we may need to delay or terminate ongoing or planned clinical trials, either of which would have a negative effect on our business. Delays in enrolling patients in our clinical trials would also adversely affect our ability to generate any product, milestone and royalty revenues under collaboration agreements, if any, and could impose significant additional costs on us or on any future collaborators.

We may not achieve our projected development goals in the time frames we announce and expect.

We set goals for, and make public statements regarding, the timing of certain accomplishments, such as, the commencement and completion of clinical trials, particularly with respect to steps for commencing and continuing GENETIC-AF, the disclosure of trial results, the obtainment of regulatory approval and the sale of drug product, which we sometimes refer to as milestones. These milestones may not be achieved, and the actual timing of these events can vary dramatically due to a number of factors such as delays or failures in our clinical trials, disagreements with current or future collaborative partners, the uncertainties inherent in the regulatory approval process and manufacturing scale-up and delays in achieving manufacturing or marketing arrangements sufficient to commercialize our products. FDA approval of Gencaro, if it occurs, is expected to require years of additional clinical development, including the completion of a new multi-year clinical study of Gencaro in approximately 200 patients and, depending on the outcome of the Phase 2B portion, may be expanded to a Phase 3 study with up to an estimated additional 420 patients. There can be no assurance that our clinical trials will be completed, or that we will make regulatory submissions or receive regulatory approvals as planned. If we fail to achieve one or more of these milestones as planned, our business will be materially adversely affected.

If we are not able to maintain the requirements for listing on the Nasdaq Capital Market, we could be delisted, which could have a materially adverse effect on our ability to raise additional funds as well as the price and liquidity of our common stock.

Our common stock is currently listed on the Nasdaq Capital Market. To maintain the listing of our common stock on the Nasdaq Capital Market we are required to meet certain listing requirements, including, among others, either: (i) a minimum closing bid price of \$1.00 per share, a market value of publicly held shares (excluding shares held by our executive officers, directors and 10% or more stockholders) of at least \$1 million and stockholders' equity of at least \$2.5 million; or (ii) a minimum closing bid price of \$1.00 per share, a market value of publicly held shares (excluding shares held by our executive officers, directors and 10% or more stockholders) of at least \$1 million and a total market value of listed securities of at least \$35 million.

During 2012 our stock price fell below the Nasdaq Capital Market's minimum bid price requirements and we became subject to delisting from the exchange. On March 4, 2013 we executed a 1 for 6 reverse split of our common stock and have subsequently regained compliance with the minimum bid price requirements. In future periods, if we do not meet the minimum stockholders' equity, minimum closing bid price requirements, or any other listing requirements, we would be subject to delisting from the Nasdaq Capital Market.

As of November 8, 2013, the closing price of our common stock was \$1.53 per share, and the total market value of our listed securities was approximately \$24 million. As of September 30, 2013, we had stockholders' equity of \$19.1 million.

We expect to depend on existing and future collaborations with third parties for the development of some of our product candidates. If those collaborations are not successful, we may not be able to complete the development of these product candidates.

We currently have a collaboration agreement with Medtronic, Inc. or Medtronic for the support of our GENETIC-AF trial. Medtronic can terminate its collaboration with us for various reasons including uncured material breach, an ARCA bankruptcy, if, after FDA communication, it is reasonably concluded that the FDA will not allow GENETIC-AF to enroll or proceed, if the trial has not begun by December 1, 2014, or if Medtronic's obligations are unilaterally expanded. We may seek additional third party collaborators for the development of Gencaro or other product candidates.

Under our current arrangement with Medtronic, we have limited control over the amount and timing of resources that they dedicate to the development of Gencaro. This is also likely to be true in any future collaborations with third

parties. Our ability to generate revenues from these arrangements will depend on our collaborators' abilities to successfully perform the functions assigned to them in these arrangements.

Collaborations involving our product candidates pose the following risks to us:

- collaborators have significant discretion in determining the efforts and resources that they will apply to these collaborations;

- collaborators may not pursue development and commercialization of our product candidates or may elect not to continue or renew development or commercialization programs based on clinical trial results, changes in the collaborator's strategic focus or available funding, or external factors such as an acquisition that diverts resources or creates competing priorities;

- collaborators may delay clinical trials, provide insufficient funding for a clinical trial program, stop a clinical trial or abandon a product candidate, repeat or conduct new clinical trials or require a new formulation of a product candidate for clinical testing;

- collaborators could independently develop, or develop with third parties, products that compete directly or indirectly with our product candidates if the collaborators believe that competitive products are more likely to be successfully developed or can be commercialized under terms that are more economically attractive than ours;

collaborators may not properly maintain or defend our intellectual property rights or may use our proprietary information in such a way as to invite litigation that could jeopardize or invalidate our proprietary information or expose us to potential litigation;

disputes may arise between the collaborators and us that result in the delay or termination of the research, development or commercialization of our product candidates or that result in costly litigation or arbitration that diverts management attention and resources;

collaborations may be terminated and, if terminated, may result in a need for additional capital to pursue further development or commercialization of the applicable product candidates;

collaborators may elect to take over manufacturing rather than retain us as manufacturers and may encounter problems in starting up or gaining approval for their manufacturing facility and so be unable to continue development of product candidates;

we may be required to undertake the expenditure of substantial operational, financial and management resources in connection with any collaboration;

we may be required to issue equity securities to collaborators that would dilute our existing stockholders' percentage ownership;

we may be required to assume substantial actual or contingent liabilities;

collaborators may not commit adequate resources to the marketing and distribution of our product candidates, limiting our potential revenues from these products; and

collaborators may experience financial difficulties.

We face a number of challenges in seeking additional collaborations. Collaborations are complex and any potential discussions may not result in a definitive agreement for many reasons. For example, whether we reach a definitive agreement for a collaboration will depend, among other things, upon our assessment of the collaborator's resources and expertise, the terms and conditions of the proposed collaboration, and the proposed collaborator's evaluation of a number of factors, such as the design or results of our clinical trials, the potential market for our product candidates, the costs and complexities of manufacturing and delivering our product candidates to patients, the potential of competing products, the existence of uncertainty with respect to ownership or the coverage of our intellectual property, and industry and market conditions generally. If we were to determine that additional collaborations for our Gencaro development is necessary and were unable to enter into such collaborations on acceptable terms, we might elect to delay or scale back the development or commercialization of Gencaro in order to preserve our financial resources or to allow us adequate time to develop the required physical resources and systems and expertise ourselves.

Collaboration agreements may not lead to development or commercialization of product candidates in the most efficient manner, or at all. In addition, there have been a significant number of recent business combinations among large pharmaceutical companies that have resulted in a reduced number of potential future collaborators. If a present or future collaborator of ours were to be involved in a business combination, the continued pursuit and emphasis on our product development or commercialization program could be delayed, diminished or terminated.

Our GENETIC-AF clinical trial will require the use of a third-party diagnostic services provider to administer the genetic test needed to identify the patient receptor genotypes of clinical trial participants, and as a result, we will be unable to directly control the timing, conduct and expense of the genetic test.

The GENETIC-AF clinical trial we intend to conduct with Gencaro requires a companion diagnostic test that identifies the patient's receptor genotype and the trial will only enroll those patients with the receptor that has the potential for enhanced efficacy, the beta-1 389 Arg receptor as detected by a beta-1 389 Arg/Arg genotype. Accordingly, the GENETIC-AF trial will require use of a third-party diagnostic service to perform the genetic test. There has been limited experience in our industry in prospective development of companion diagnostics required to perform the required molecular profiling. We recently entered into an agreement with Laboratory Corporation of

America, LabCorp, to provide the diagnostic services of the genetic test needed to support our GENETIC-AF trial. The FDA and similar regulatory authorities outside the United States regulate companion diagnostics. Companion diagnostics require separate or coordinated regulatory approval prior to commercialization. The regulatory pathway for co-development of therapeutics and companion diagnostics is uncertain. Changes to regulatory advice could delay our development programs or delay or prevent eventual marketing approval for our product candidates that may otherwise be approvable. In July 2011, the FDA issued draft guidance that stated that if safe and effective use of a therapeutic depends on an in vitro diagnostic, then the FDA generally will not approve the therapeutic unless the FDA approves or clears this in vitro companion diagnostic device at the same time that the FDA approves the therapeutic. The approval or clearance of the companion diagnostic would occur through the FDA's Center for Devices and Radiological Health Office of In Vitro Diagnostic Device Evaluation and Safety. It is unclear whether the FDA will finalize this guidance in its current form, or when it will do so. Even if the FDA does finalize the guidance, it is difficult to predict how it will implement the guidance. For example, the draft guidance allows for flexibility by the FDA in the case of disease indications with serious unmet medical needs, but it is unclear how this discretion will be applied by the agency. Therefore, even with

the issuance of the draft guidance, the FDA's expectations for companion diagnostics remain unclear. The FDA's evolving position on the topic of companion diagnostics could affect our clinical development programs that utilize companion diagnostics. In particular, the FDA may limit our ability to use retrospective data, otherwise disagree with our approaches to trial design, biomarker qualification, clinical and analytical validity, and clinical utility, or make us repeat aspects of a trial or initiate new trials.

Assays that can be used as companion diagnostics for detecting beta-1 389 Arg receptor are commercially available, but in most cases they do not yet have regulatory approval for use as companion diagnostics. In GENETIC-AF, we plan to use commercially available companion diagnostics, and the diagnostic will be required to obtain regulatory approval for use in our GENETIC-AF trial. Any delay in such approval would delay our clinical trial.

Given our limited experience in developing diagnostics, we expect to rely primarily on third parties for their design and manufacture. If we, or any third parties that we engage to assist us, are unable to successfully develop companion diagnostics for our product candidates that require such diagnostics, or experience delays in doing so, the development of our product candidates may be adversely affected, our product candidates may not receive marketing approval and we may not realize the full commercial potential of any products that receive marketing approval. As a result, our business could be materially harmed.

We will need to establish a collaborative arrangement with a third-party diagnostics services provider to obtain marketing clearance or approval of the companion genetic test. There is no guarantee that the FDA will grant timely clearance or approval of the genetic test, if at all, and failure to obtain such timely clearance or approval would adversely affect our ability to market Gencaro.

The drug label we intend to seek for Gencaro would identify the patient receptor genotype for which the drug is approved. Accordingly, we believe developing a genetic test that is simple to administer and widely available will be critical to the successful commercialization of Gencaro and also to the ability to conduct our planned GENETIC-AF clinical trial. The genetic test will be subject to regulation by the FDA and by comparable agencies in various foreign countries. The process of complying with the requirements of the FDA and comparable agencies is costly, time consuming and burdensome.

Despite the time and expense expended, regulatory clearance or approval is never guaranteed. If regulatory clearance or approval is delayed, or if one or more third-party diagnostic services providers are unable to obtain FDA approval of the genetic test at all or in parallel with the approval of Gencaro, or are unable to commercialize the test successfully and in a manner that effectively supports the commercial efforts for Gencaro, or if the information concerning the differential response to Gencaro resulting from certain genetic variation is not included in the approval label for Gencaro, the commercial launch of Gencaro may be significantly and adversely affected.

Regulatory approval will be required for the genetic test to be used in the GENETIC-AF trial and to support the commercialization of the test, if approved. Delays or failures in obtaining such regulatory approval, including any required validation analyses may prevent a third-party diagnostics provider from commercializing such genetic test and will adversely affect our business, operating results and prospects.

The GENETIC-AF trial requires the submission to, and approval from, the FDA of an investigational device exemption, or IDE, before the GENETIC-AF trial can begin. LabCorp has not yet submitted the IDE for the test we intend to use in GENETIC-AF and they may not submit the IDE for approval when we anticipate, or at all. There is no guarantee that the FDA will approve LabCorp's IDE submission if it is made or the potential timing of such approval. A delay in submission of the IDE or delay or denial by the FDA of the IDE would impair the timing of, and possibly our ability to perform, our GENETIC-AF clinical trial.

Before a genetic test can be used commercially, including in conjunction with Gencaro, if it is approved for marketing, the third-party diagnostics provider must obtain FDA Premarket Approval, or PMA, for such test. The

FDA may require additional validation of the genetic test we plan to use in GENETIC-AF prior to any approval of Gencaro or the genetic test. We anticipate the genetic test will be required as a condition to prescribing Gencaro.

There is no guarantee the FDA will approve the anticipated PMA submission for the genetic test. Even if the genetic test is eventually approved, performing additional validation work necessary to support the PMA, if required, for current or future genetic test products, including one associated with Gencaro, would require additional time and expense and the outcome would be uncertain. Moreover, such delays or increased costs or failures could adversely affect our business, operating results and prospects for commercializing the genetic test.

If a third-party diagnostics provider responsible for the genetic test or certain of its third-party suppliers fails to comply with ongoing FDA or other foreign regulatory authority requirements, or if there are unanticipated problems with the genetic test, these products could be subject to restrictions or withdrawal from use in trial or from the market.

Any diagnostic for which a third-party diagnostics provider obtains clearance or approval, and the manufacturing processes, reporting requirements, post-approval clinical data and promotional activities for such product, will be subject to continued regulatory review, oversight and periodic inspections by the FDA and other domestic and foreign regulatory bodies. With respect to the genetic

test, to the extent applicable, any third-party diagnostics provider and certain of its suppliers will be required to comply with the FDA's Quality System Regulation, or QSR, and International Standards Organization, or ISO, requirements which cover the methods and documentation of the design, testing, production, control, quality assurance, labeling, packaging, storage and shipping of any product for which clearance or approval is obtained. Regulatory bodies, such as the FDA, enforce the QSR and other regulations through periodic inspections. The failure by a third-party diagnostics provider, or certain of its third-party manufacturers or suppliers, as the case may be, to comply with applicable statutes and regulations administered by the FDA and other regulatory bodies, or the failure to timely and adequately respond to any adverse inspectional observations or product safety issues, could result in, among other things, enforcement actions. If any of these actions were to occur, it could harm our reputation and cause product sales and profitability of Gencaro to suffer and may prevent us from generating revenue or utilizing the genetic test further in any clinical trial. Even if regulatory clearance or approval is granted, such clearance or approval may be subject to limitations on the intended uses for which the product may be marketed and reduce our potential to successfully commercialize the product and generate revenue from the product.

Future sales of Gencaro may suffer if its marketplace acceptance is negatively affected by the genetic test.

The genetic test is an important component of the commercial strategy for Gencaro in addition to being required to proceed with our planned AF trial. We believe that the genetic test helps predict patient response to Gencaro, and that this aspect of the drug is important to its ability to compete effectively with current therapies. The genetic test adds an additional step in the prescribing process, an additional cost for the patient and payors, the risk that the test results may not be rapidly available and the possibility that it may not be available at all to hospitals and medical centers. Although we anticipate that Gencaro, if approved in a timely manner, would be the first genetically-targeted cardiovascular drug, Gencaro will be one of a number of successful drugs in the beta-blocker class currently on the market. Prescribers may be more familiar with these other beta-blockers, and may be resistant to prescribing Gencaro as an AF therapy in patients with HF. Any one of these factors could affect prescriber behavior, which in turn may substantially impede market acceptance of the genetic test, which could cause significant harm to Gencaro's ability to compete, and in turn harm our business.

Our failure to raise substantial additional funding or enter into a strategic transaction may materially and adversely affect our business.

Unless we are able to raise substantial additional funding through other means, we will need to complete a strategic transaction to continue the development of Gencaro or our other operations. The strategic transactions that we may consider include a potential combination or partnership. Our board of directors and management team has and will continue to devote substantial time and resources to obtaining additional capital or the consideration and implementation of any such strategic transaction. In addition, conditions in the financial markets may lead to an increased number of biotechnology companies that are also seeking to enter into strategic transactions, which may limit our ability to negotiate favorable terms for any such transaction. Further, our current employees do not have experience in the strategic transaction process, and our previous efforts to enter into a strategic transaction have not been successful. As a result of these and other factors, there is substantial risk that we may not be able to complete a strategic transaction on favorable terms, or at all. The failure to complete a strategic transaction may materially and adversely affect our business.

We may be limited in our ability to access sufficient funding through a private equity or convertible debt offering.

Nasdaq rules impose restrictions on our ability to raise funds through a private offering of our common stock, convertible debt or similar instruments without obtaining stockholder approval. Under Nasdaq rules, an offering of more than 20% of our total shares outstanding for less than the greater of book or market value requires stockholder approval unless the offering qualifies as a public offering for purposes of the Nasdaq rules. As of November 8, 2013, we had approximately 15.6 million shares of common stock outstanding, 20% of which is approximately 3.1 million shares. To the extent we seek to raise funds through a private offering of stock, convertible debt or similar

instruments, we are limited in how much funding we could raise privately without requiring a stockholder vote.

In addition, we are currently subject to certain contractual rights of investors arising from our public and private equity financing transactions that limit the nature and price of future public and private financing transactions that we may effect. For example, in January 2013, we entered into separate subscription agreements with certain institutional investors in connection with a private investment in public equity, pursuant to which we sold shares of our common stock and warrants to purchase shares of our common stock to the investors. In connection with this transaction, we agreed that, subject to certain exceptions, we would not, while the warrants issued in such financing are outstanding, effect or enter into an agreement to effect any issuance of common stock or securities convertible into, exercisable for or exchangeable for common stock in a variable rate transaction, which means a transaction in which we issue or sell any convertible securities either (A) at a conversion price, exercise price or exchange rate or other price that is based upon and/or varies with the trading prices of, or quotations for, the shares of common stock at any time after the initial issuance of such convertible securities, or (B) with a conversion, exercise or exchange price that is subject to being reset at some future date after the initial issuance of the convertible securities or upon the occurrence of the specified or contingent events

directly or indirectly related to our business or the market for our common stock. The restrictions imposed by the terms of our previous offerings, and that could be imposed in future offerings, may limit our access to capital on agreeable terms and delay or make impossible certain otherwise available equity financing opportunities and could severely restrict our access to the capital necessary to conduct our business.

Unless we are able to generate sufficient product revenue, we will continue to incur losses from operations and may not achieve or maintain profitability. We are years away from commercializing a product and generating product revenue.

Our historical losses have had and will continue to have an adverse effect on our stockholders' equity and working capital, among other things. We are years away from commercializing a product and generating any product revenue. As a result, we expect to continue to incur significant operating losses for the foreseeable future. Even if we ultimately receive regulatory approval for Gencaro or our other product candidates, sales of such products may not generate sufficient revenue for it to achieve or maintain profitability. Because of the numerous risks and uncertainties associated with developing therapeutic drugs, we may experience larger than expected future losses and may never reach profitability.

Our product candidates are subject to extensive regulation, which can be costly and time-consuming, and unsuccessful or delayed regulatory approvals could increase our future development costs or impair our future revenue.

The preclinical and clinical development, testing, manufacture, safety, efficacy, labeling, storage, recordkeeping, and subsequent advertising, promotion, sale, marketing, and distribution, if approved, of our product candidates are subject to extensive regulation by the FDA and other regulatory authorities in the United States and elsewhere. These regulations also vary in important, meaningful ways from country to country. We are not permitted to market a potential drug in the United States until we receive approval of an NDA from the FDA. We have not received an NDA approval from the FDA for Gencaro or any of our other product candidates. There can be no guarantees with respect to our product candidates that clinical studies will adequately support an NDA, that the products will receive necessary regulatory approvals, or that they will prove to be commercially successful.

To receive regulatory approval for the commercial sale of any product candidates, we must demonstrate safety and efficacy in humans to the satisfaction of regulatory authorities through preclinical studies and adequate and well-controlled clinical trials of the product candidates. This process is expensive and can take many years, and failure can occur at any stage of the testing. Our failure to adequately demonstrate the safety and efficacy of our product candidates will prevent regulatory approval and commercialization of such products. In 2008, we submitted and the FDA accepted our NDA filing for Gencaro for the treatment of chronic HF. In 2009, the FDA issued a Complete Response Letter (CRL) in which the FDA stated that it could not approve the Gencaro NDA in its current form and specified actions required for approval of the NDA, including conducting an additional Phase 3 clinical trial of Gencaro in patients with HF. We plan to conduct a clinical study of Gencaro in HFREF patients to assess its efficacy in reducing or preventing AF. We currently anticipate that GENETIC-AF will begin patient enrollment in the first quarter of 2014. This trial is planned to begin as a Phase 2B study in approximately 200 patients and, depending on the outcome of the Phase 2B portion, may be expanded to a Phase 3 study with up to an estimated additional 420 patients. We believe the Phase 2B study would take approximately two and a half years to complete. This product candidate will require years of clinical development. Even if we conduct additional studies in accordance with further FDA guidance and submit or file a new or amended NDA, the FDA may ultimately decide that the NDA does not satisfy the criteria for approval.

In the event that we or our collaborators conduct preclinical studies that do not comply with Good Laboratory Practices or GLP or incorrectly design or carry out human clinical trials in accordance with Good Clinical Practices or GCP or those clinical trials fail to demonstrate clinical significance, it is unlikely that we will be able to obtain FDA approval for product development candidates. Our inability to successfully and effectively complete clinical trials for any product candidate on schedule, or at all, will severely harm our business. Significant delays in clinical

development could materially increase product development costs or allow our competitors to bring products to market before we do, impairing our ability to effectively commercialize any future product candidate. We do not know whether planned clinical trials will begin on time, will need to be redesigned or will be completed on schedule, if at all. Clinical trials can be delayed for a variety of reasons, including:

delays or failures in obtaining regulatory authorization to commence a trial because of safety concerns of regulators relating to our product candidates or similar product candidates of our competitors or failure to follow regulatory guidelines;

delays or failures in obtaining clinical materials and manufacturing sufficient quantities of the product candidates for use in trials;

delays or failures in reaching agreement on acceptable terms with prospective study sites;

delays or failures in obtaining approval of our clinical trial protocol from an institutional review board, or IRB, to conduct a clinical trial at a prospective study site;

delays in recruiting patients to participate in a clinical trial, which may be due to the size of the patient population, eligibility criteria, protocol design, perceived risks and benefits of the drug, availability of other approved and standard of care therapies, availability of clinical trial sites;  
other clinical trials seeking to enroll subjects with similar profile;  
failure of our clinical trials and clinical investigators to be in compliance with the FDA's Good Clinical Practices;  
unforeseen safety issues, including negative results from ongoing preclinical studies;  
inability to monitor patients adequately during or after treatment;  
difficulty monitoring multiple study sites; and  
failure of our third-party contract research organizations, clinical site organizations and other clinical trial managers, to satisfy their contractual duties, comply with regulations or meet expected deadlines.

In addition, any approvals we may obtain may not cover all of the clinical indications for which we seek approval or permit us to make claims of superiority over currently marketed competitive products. Also, an approval might contain significant limitations in the form of narrow indications, warnings, precautions or contraindications with respect to conditions of use. If the FDA determines that a risk evaluation and mitigation strategy, or REMS, is necessary to ensure that the benefits of the drug outweigh the risks, we may be required to include as part of the NDA a proposed REMS that may include a package insert directed to patients, a plan for communication with healthcare providers, restrictions on a drug's distribution, or a Medication Guide, to provide better information to consumers about the drug's risks and benefits. Finally, an approval could be conditioned on our commitment to conduct further clinical trials, which we may not have the resources to conduct or which may negatively impact our financial situation.

The manufacture and tableting of Gencaro is done by third party suppliers, who must also meet current Good Manufacturing Practices, or cGMP, requirements and pass a pre-approval inspection of their facilities before we can obtain marketing approval.

All of our product candidates are prone to the risks of failure inherent in drug development. The results from preclinical animal testing and early human clinical trials may not be predictive of results obtained in later human clinical trials. Further, although a new product may show promising results in preclinical or early human clinical trials, it may subsequently prove unfeasible or impossible to generate sufficient safety and efficacy data to obtain necessary regulatory approvals. The data obtained from preclinical and clinical studies are susceptible to varying interpretations that may delay, limit or prevent regulatory approval, and the FDA and other regulatory authorities in the United States and elsewhere exercise substantial discretion in the drug approval process. The numbers, size and design of preclinical studies and clinical trials that will be required for FDA or other regulatory approval will vary depending on the product candidate, the disease or condition for which the product candidate is intended to be used and the regulations and guidance documents applicable to any particular product candidate. The FDA or other regulators can delay, limit or deny approval of any product candidate for many reasons, including, but not limited to:

Side effects;

Safety and efficacy;

Defects in the design of clinical trials;

The fact that the FDA or other regulatory officials may not approve our or our third party manufacturer's processes or facilities; or

The fact that new regulations may be enacted by the FDA or other regulators may change their approval policies or adopt new regulations requiring new or different evidence of safety and efficacy for the intended use of a product candidate.

In light of widely publicized events concerning the safety of certain drug products, regulatory authorities, members of Congress, the Government Accountability Office, medical professionals and the general public have raised concerns about potential drug safety issues. These events have resulted in the withdrawal of certain drug products, revisions to certain drug labeling that further limit use of the drug products and establishment of risk management programs that may, for instance, restrict distribution of drug products. The increased attention to drug safety issues may result in a more cautious approach by the FDA to clinical trials and approval. Data from clinical trials may receive greater scrutiny with respect to safety and the product's risk/benefit profile, which may make the FDA or other regulatory authorities more likely to terminate clinical trials before completion, or require longer or additional clinical trials that may result in substantial additional expense, and a delay or failure in obtaining approval or approval for a more limited indication than originally sought. Aside from issues concerning the quality and sufficiency of submitted preclinical and clinical data, the FDA may be constrained by limited resources from reviewing and determining the approvability of the Gencaro NDA in a timely manner.

In pursuing clinical development of Gencaro for an AF indication, we will be required to amend the Gencaro HF NDA or prepare a new NDA. The FDA could approve Gencaro, but without including some or all of the prescribing information that we have requested. For instance, the FDA could approve Gencaro for AF in a more limited patient population or included additional warnings in the drug's label. This, in turn, could substantially and detrimentally impact our ability to successfully commercialize Gencaro and effectively protect our intellectual property rights in Gencaro.

If our product candidates receive regulatory approval, we would be subject to ongoing regulatory obligations and restrictions, which may result in significant expenses and limit our ability to develop and commercialize other potential products.

If a product candidate of ours is approved by the FDA or by another regulatory authority, we would be held to extensive regulatory requirements over product manufacturing, testing, distribution, labeling, packaging, adverse event reporting and other reporting to regulatory authorities, storage, advertising, marketing, promotion, distribution, and record keeping. Regulatory approvals may also be subject to significant limitations on the indicated uses or marketing of the product candidates. Potentially costly follow-up or post-marketing clinical studies may be required as a condition of approval to further substantiate safety or efficacy, or to investigate specific issues of interest to the regulatory authority. Previously unknown problems with the product candidate, including adverse events of unanticipated severity or frequency, may result in additional regulatory controls or restrictions on the marketing or use of the product or the need for post marketing studies, and could include suspension or withdrawal of the products from the market.

Furthermore, our third-party manufacturers and the manufacturing facilities that they use to make our product candidates are regulated by the FDA. Quality control and manufacturing procedures must continue to conform to cGMP after approval. Drug manufacturers and their subcontractors are required to register their facilities and products manufactured annually with the FDA and certain state agencies and are subject to periodic unannounced inspections by the FDA, state and/or other foreign authorities. Any subsequent discovery of problems with a product, or a manufacturing or laboratory facility used by us or our collaborators, may result in restrictions on the product, or on the manufacturing or laboratory facility, including a withdrawal of the drug from the market or suspension of manufacturing. Any changes to an approved product, including the way it is manufactured or promoted, often require FDA approval before the product, as modified, can be marketed. We and our third-party manufacturers will also be subject to ongoing FDA requirements for submission of safety and other post-market information.

The marketing and advertising of our drug products by our collaborators or us will be regulated by the FDA, certain state agencies or foreign regulatory authorities. Violations of these laws and regulations, including promotion of our products for unapproved uses or failing to disclose risk information, are punishable by criminal and civil sanctions and may result in the issuance of enforcement letters or other enforcement action by the FDA, U.S. Department of Justice, state agencies, or foreign regulatory authorities that could jeopardize our ability to market the product.

In addition to the FDA, state or foreign regulations, the marketing of our drug products by us or our collaborators will be regulated by federal, state or foreign laws pertaining to health care fraud and abuse, such as the federal anti-kickback law prohibiting bribes, kickbacks or other remuneration for the order or recommendation of items or services reimbursed by federal health care programs. Many states have similar laws applicable to items or services reimbursed by commercial insurers. Violations of these laws are punishable by criminal and civil sanctions, including, in some instances, imprisonment and exclusion from participation in federal and state health care programs, including the Medicare, Medicaid and Veterans Affairs healthcare programs. Because of the far-reaching nature of these laws, we may be required to discontinue one or more of our practices to be in compliance with these laws. Health care fraud and abuse regulations are complex, and even minor irregularities can potentially give rise to claims that a statute or prohibition has been violated. Any violations of these laws, or any action against us for violations of these laws, even if we successfully defend against it, could have a material adverse effect on our business, financial condition and results of operations.

We could also become subject to false claims litigation under federal statutes, which can lead to civil money penalties, restitution, criminal fines and imprisonment, and exclusion from participation in Medicare, Medicaid and other federal and state health care programs. These false claims statutes include the False Claims Act, which allows any person to bring a suit on behalf of the federal government alleging submission of false or fraudulent claims, or causing to present such false or fraudulent claims, under federal programs or contracts claims or other violations of the statute and to share in any amounts paid by the entity to the government in fines or settlement. These suits against pharmaceutical companies have increased significantly in volume and breadth in recent years. Some of these suits have been brought on the basis of certain sales practices promoting drug products for unapproved uses. This new growth in litigation has increased the risk that a pharmaceutical company will have to defend a false claim action, pay fines or restitution, or be excluded from the Medicare, Medicaid, Veterans Affairs and other federal and state healthcare programs as a result of an investigation arising out of such action. We may become subject to such litigation and, if we are not successful in defending against such actions, those actions may have a material adverse effect on our business, financial condition and results of operations. We could also become subject to false claims litigation and consumer protection claims under state statutes, which also could lead to civil monetary penalties, restitution, criminal fines and imprisonment, and exclusion from participation in state health care programs.

Of note, over the past few years there has been an increased focus on the sales and marketing practices of the pharmaceutical industry at both the federal and state level. Additionally, the law or regulatory policies governing pharmaceuticals may change. New

statutory requirements may be enacted or additional regulations may be adopted that could prevent or delay regulatory approval of our product candidates or limit our ability to commercialize our products. We cannot predict the likelihood, nature or extent of adverse government regulation that may arise from future legislation or administrative action, either in the U.S. or elsewhere.

If we, our collaborators or our third-party manufacturers fail to comply with applicable continuing regulatory requirements, our business could be seriously harmed because a regulatory agency may:

- issue untitled or warning letters;
- suspend or withdraw our regulatory approval for approved products;
- seize or detain products or recommend a product recall of a drug or medical device, or issue a mandatory recall of a medical device;
- refuse to approve pending applications or supplements to approved applications filed by us;
- suspend our ongoing clinical trials;
- restrict our operations, including costly new manufacturing requirements, or restrict the sale, marketing and/or distribution of our products;
- seek an injunction;
- pursue criminal prosecutions;
- close the facilities of our contract manufacturers; or
- impose civil or criminal penalties.

Reliance on third parties to commercialize Gencaro could negatively impact our business. If we are required to establish a direct sales force in the U.S. and are unable to do so, our business may be harmed.

Commercialization of Gencaro, particularly the establishment of a sales organization, will require substantial additional capital resources. We currently intend to pursue a strategic partnership alternative for the commercialization of Gencaro, if it is approved, and we have suspended our efforts to build internal sales, marketing and distribution capabilities. If we elect to rely on third parties to sell Gencaro and any other products, then we may receive less revenue than if we sold such products directly. In addition, we may have little or no control over the sales efforts of those third parties.

If we are unable to complete a strategic transaction, we would be unable to commercialize Gencaro or any other product candidate without substantial additional capital. Even if such capital were secured, we would be required to build internal sales, marketing and distribution capabilities to market Gencaro in the U.S. None of our current employees have experience in establishing and managing a sales force.

In the event we are unable to sell Gencaro and other selected product candidates, either directly or through third parties via a strategic transaction, the commercialization of Gencaro, if it is approved, may be delayed indefinitely.

We are dependent on our key personnel.

The success of our business is highly dependent on the principal members of our board of directors and executive management, including our President and Chief Executive Officer, Michael R. Bristow. The loss of the services of any such individual might seriously harm our product development, partnering and financing efforts. Recruiting and training personnel with the requisite skills is challenging and we compete for talent with companies that are larger and have more financial resources.

We have no manufacturing capacity which puts us at risk of lengthy and costly delays of bringing our products to market.

We do not currently operate manufacturing facilities for clinical or commercial production of our product candidates, including their active pharmaceutical ingredients, or API. We have no experience in drug formulation or manufacturing, and we lack the resources and the capabilities to manufacture any of our product candidates on a clinical or commercial scale. We do not intend to develop facilities for the manufacture of product candidates for clinical trials or commercial purposes in the foreseeable future. We have contracted with Groupe Novasep to manufacture commercial quantities of the API for Gencaro. For drug production, we have contracted with Patheon, Inc. to manufacture the Gencaro tablets. These contract manufacturers may not perform as agreed or may not remain in the contract manufacturing business for the time required to successfully produce, store and distribute our products. In addition, these manufacturers may have staffing difficulties, may not be able to manufacture our products on a timely basis or may become financially distressed. In the event of errors in forecasting production quantities required to meet demand, natural disaster, equipment malfunctions or failures, technology malfunctions, strikes, lock-outs or work stoppages, regional power outages, product tampering, war or terrorist activities, actions of regulatory authorities, business failure, strike or other difficulty, we may be unable to

find an alternative third-party manufacturer in a timely manner and the production of our product candidates would be interrupted, resulting in delays and additional costs, which could impact our ability to commercialize and sell our product candidates. We or our contract manufacturers may also fail to achieve and maintain required manufacturing standards, which could result in patient injury or death, product recalls or withdrawals, an order by governmental authorities to halt production, delays or failures in product testing or delivery, cost overruns or other problems that could seriously hurt our business. Contract manufacturers also often encounter difficulties involving production yields, quality control and quality assurance, as well as shortages of qualified personnel. In addition, our contract manufacturers are subject to ongoing inspections and regulation by the FDA, the U.S. Drug Enforcement Agency and corresponding foreign and state agencies and they may fail to meet these agencies' acceptable standards of compliance. If our contract manufacturers fail to comply with applicable governmental regulations, such as quality control, quality assurance and the maintenance of records and documentation, we may not be able to continue production of the API or finished product. If the safety of any API or product supplied is compromised due to failure to adhere to applicable laws or for other reasons, this may jeopardize our regulatory approval for Gencaro and other product candidates, and we may be held liable for any injuries sustained as a result. Upon the occurrence of one of the aforementioned events, the ability to switch manufacturers may be difficult for a number of reasons, including:

- the number of potential manufacturers is limited and we may not be able to negotiate agreements with alternative manufacturers on commercially reasonable terms, if at all;
- long lead times are often needed to manufacture drugs;
- the manufacturing process is complex and may require a significant learning curve; and
- the FDA must approve any replacement prior to manufacturing, which requires new testing and compliance inspections.

Transitioning from a developmental stage company will require successful completion of a number of steps, many of which are outside of our control and, consequently, we can provide no assurance of our successful and timely transition from a developmental stage company.

We are a development stage biopharmaceutical company with a limited operating history. To date we have not generated any product revenue and have historically funded our operations through investment capital. Our future growth depends on our ability to emerge from the developmental stage and successfully commercialize or provide for the commercialization of Gencaro and our other product candidates which in turn, will depend, among other things, on our ability to:

- conduct an additional clinical trial and develop and obtain regulatory approval for Gencaro or other product candidates;
- successfully partner a companion genetic test with the commercial launch of Gencaro;
- enter into a strategic transaction enabling the continued development and commercialization of Gencaro, or alternatively, raise significant additional capital to enable these activities;
- pursue additional indications for Gencaro and develop other product candidates, including other cardiovascular therapies; and
- obtain commercial quantities of Gencaro or other product candidates at acceptable cost levels.

Any one of these factors or other factors discussed in this report could affect our ability to successfully commercialize Gencaro and other product candidates, which could impact our ability to earn sufficient revenues to transition from a developmental stage company and continue our business.

If approved by the FDA, Gencaro will be entering a competitive marketplace and may not succeed.

Gencaro is a new type of beta-blocker and vasodilator being developed for AF. While we anticipate that this drug, if approved, would be the first genetically-targeted cardiovascular drug, and potentially the only beta-blocker approved for AF, Gencaro will be one of a number of accepted treatments for AF. In addition, our proposed prescribing information for Gencaro is expected to include a requirement for genetic testing of the patient to ascertain if they have the genotype that we believe responds most favorably to Gencaro. This additional step will add incremental cost and procedures to prescribing Gencaro, which could make it more difficult to compete against existing therapies.

Our commercial opportunity may be reduced or eliminated if competitors develop and commercialize products that are safer, more effective, have fewer side effects, are more convenient or are less expensive than Gencaro. If products with any of these properties are developed, or any of the existing products are better marketed, then prescriptions of Gencaro by physicians and patient use of Gencaro could be significantly reduced or rendered obsolete and noncompetitive. Further, public announcements regarding the development of any such competing drugs could adversely affect the market price of our common stock and the value of our assets.

Future sales of our products may suffer if they are not accepted in the marketplace by physicians, patients and the medical community.

Gencaro or our other product candidates may not gain market acceptance among physicians, patients and the medical community. The degree of market acceptance of Gencaro or our other product candidates will depend on a number of factors, such as its effectiveness and tolerability, as compared with competitive drugs. Also, prevalence and severity of side-effects could negatively affect market acceptance of Gencaro or our other product candidates. Failure to achieve market acceptance of Gencaro would significantly harm our business.

If we are unable to obtain acceptable prices or adequate reimbursement from third-party payors for Gencaro, or any other product candidates that we may seek to commercialize, then our revenues and prospects for profitability will suffer.

Our or any strategic partner's ability to commercialize Gencaro, or any other product candidates that we may seek to commercialize, is highly dependent on the extent to which coverage and reimbursement for these product candidates will be available from:

- governmental payors, such as Medicare and Medicaid;
- private health insurers, including managed-care organizations; and
- other third-party payors.

Many patients will not be capable of paying for our potential products themselves and will rely on third-party payors to pay for their medical needs. A primary current trend in the U.S. health care industry is toward cost containment. Large private payors, managed-care organizations, group purchasing organizations and similar organizations are exerting increasing influence on decisions regarding the use of, and reimbursement levels for, particular treatments. Such third-party payors, including Medicare, are challenging the prices charged for medical products and services, and many third-party payors limit reimbursement for newly approved health care products.

Cost-control initiatives could decrease the price we might establish for products, which could result in product revenues lower than anticipated. If the prices for our product candidates decrease, or if governmental and other third-party payors do not provide adequate coverage and reimbursement levels, then our revenue and prospects for profitability will suffer.

Health care reform measures could materially and adversely affect our business.

The business and financial condition of pharmaceutical and biotechnology companies are affected by the efforts of governmental and third-party payors to contain or reduce the costs of health care. The U.S. Congress has enacted legislation to reform the health care system. While we anticipate that this legislation may, over time, increase the number of patients who have insurance coverage for pharmaceutical products, it also imposes cost containment measures that may adversely affect the amount of reimbursement for pharmaceutical products. These measures include increasing the minimum rebates for products covered by Medicaid programs and extending such rebates to drugs dispensed to Medicaid beneficiaries enrolled in Medicaid managed care organizations as well as expansion of the 340(B) Public Health Services drug discount program. In addition, such legislation contains a number of provisions designed to generate the revenues necessary to fund the coverage expansion, including new fees or taxes on certain health-related industries, including medical device manufacturers. Beginning in 2013, each medical device manufacturer will have to pay an excise tax (or sales tax) in an amount equal to 2.3% of the price for which such manufacturer sells its medical devices. Such excise taxes may impact any potential sales of the genetic test if it is approved for marketing. In foreign jurisdictions there have been, and we expect that there will continue to be, a

number of legislative and regulatory proposals aimed at changing the health care system. For example, in some countries other than the United States, pricing of prescription drugs is subject to government control and we expect to see continued efforts to reduce healthcare costs in international markets.

Some states are also considering legislation that would control the prices of drugs, and state Medicaid programs are increasingly requesting manufacturers to pay supplemental rebates and requiring prior authorization by the state program for use of any drug for which supplemental rebates are not being paid. Managed care organizations continue to seek price discounts and, in some cases, to impose restrictions on the coverage of particular drugs. Government efforts to reduce Medicaid expenses may lead to increased use of managed care organizations by Medicaid programs. This may result in managed care organizations influencing prescription decisions for a larger segment of the population and a corresponding constraint on prices and reimbursement for drugs. It is likely that federal and state legislatures and health agencies will continue to focus on additional health care reform in the future although we are unable to predict what additional legislation or regulation, if any, relating to the health care industry or third-party coverage and reimbursement may be enacted in the future or what effect such legislation or regulation would have on our business. We or any strategic partner's ability to commercialize Gencaro, or any other product candidates that we may seek to commercialize, is highly dependent on the extent to which coverage and reimbursement for these product candidates will be available from government payors, such as Medicare and Medicaid, private health insurers, including managed care organizations, and other third-party payors, and any change in reimbursement levels could materially and adversely affect our business. Further, the pendency or approval of future

proposals or reforms could result in a decrease in our stock price or limit our ability to raise capital or to obtain strategic partnerships or licenses.

Our competitors may be better positioned in the marketplace and thereby may be more successful than us at developing, manufacturing and marketing approved products.

Many of our competitors currently have significantly greater financial resources and expertise in conducting clinical trials, obtaining regulatory approvals, managing manufacturing and marketing approved products than us. Other early-stage companies may also prove to be significant competitors, particularly through collaborative arrangements with large and established companies. In addition, these third parties compete with us in recruiting and retaining qualified scientific and management personnel, establishing clinical trial sites and patient registration for clinical trials, as well as in acquiring therapies and therapy licenses complementary to our programs or advantageous to our business. We expect that our ability to compete effectively will depend upon our ability to:

- successfully and rapidly complete clinical trials for any product candidates and obtain all requisite regulatory approvals in a cost-effective manner;
- build an adequate sales and marketing infrastructure, raise additional funding, or enter into strategic transactions enabling the commercialization of our products;
- develop competitive formulations of our product candidates;
- attract and retain key personnel; and
- identify and obtain other product candidates on commercially reasonable terms.

If we fail to identify and license or acquire other products or product candidates, then we may be unable to expand our business, and the acquisition or licensing of other products or product candidates may put a strain on our operations and will likely require us to seek additional financing.

One of our strategies is to license or acquire clinical-stage products or product candidates and further develop them for commercialization. The market for licensing and acquiring products and product candidates is intensely competitive and many of our competitors may have greater resources than us. If we undertake any additional acquisitions, whether of product candidates or other biopharmaceutical companies, the process of integrating an acquired product candidate or complementary company into our business may put a strain on our operations, divert personnel, financial resources and management's attention. In 2013, we expect our research and development activities will be dedicated to Gencaro. If we are not able to substantially expand our research and development efforts, or identify, or license or acquire other products or product candidates or complete future acquisitions, then we will likely be unable to expand our pipeline of product candidates. In addition, any future acquisition would give rise to additional operating costs and will likely require us to seek additional financing. Future acquisitions could result in additional issuances of equity securities that would dilute the ownership of existing stockholders. Future acquisitions could also result in the incurrence of debt, contingent liabilities or the amortization of expenses related to other intangible assets, any of which could adversely affect our operating results.

We would be subject to applicable regulatory approval requirements of the foreign countries in which we market our products, which are costly and may prevent or delay us from marketing our products in those countries.

In addition to regulatory requirements in the United States, we would be subject to the regulatory approval requirements in each foreign country where we market our products. In addition, we might be required to identify one or more collaborators in these foreign countries to develop, seek approval for and manufacture our products and any companion genetic test for Gencaro. If we decide to pursue regulatory approvals and commercialization of our product candidates internationally, we may not be able to obtain the required foreign regulatory approvals on a timely basis, if

at all, and any failure to do so may cause us to incur additional costs or prevent us from marketing our products in foreign countries, which may have a material adverse effect on our business, financial condition and results of operations.

If our internal control over financial reporting is not considered effective, our business and stock price could be adversely affected.

Section 404 of the Sarbanes-Oxley Act of 2002 requires us to evaluate the effectiveness of our internal control over financial reporting as of the end of each fiscal year, and to include a management report assessing the effectiveness of our internal control over financial reporting in our annual report on Form 10-K for that fiscal year. Our management, including our chief executive officer and chief financial officer, does not expect that our internal control over financial reporting will prevent all error and all fraud. During the first quarter of 2011 there was a reduction in our workforce which included personnel involved in financial reporting and our internal control processes. Since that time we have continued to operate with a reduced staff for financial reporting. Though the process and design of our internal controls over financial reporting have not been altered, the small number of staff involved in financial reporting may limit our ability to properly segregate internal control procedures which could result in deficiencies or material weaknesses in our

internal controls in the future. A control system, no matter how well designed and operated, can provide only reasonable, not absolute, assurance that the control system's objectives will be met. Further, the design of a control system must reflect the fact that there are resource constraints, and the benefits of controls must be considered relative to their costs. Because of the inherent limitations in all control systems, no evaluation of controls can provide absolute assurance that all control issues and instances of fraud involving a company have been, or will be, detected. The design of any system of controls is based in part on certain assumptions about the likelihood of future events, and we cannot assure you that any design will succeed in achieving its stated goals under all potential future conditions. Over time, controls may become ineffective because of changes in conditions or deterioration in the degree of compliance with policies or procedures. Because of the inherent limitations in a cost-effective control system, misstatements due to error or fraud may occur and not be detected. We cannot assure you that we or our independent registered public accounting firm will not identify a material weakness in our internal control over financial reporting in the future. A material weakness in our internal control over financial reporting would require management to consider our internal control over financial reporting as ineffective. If our internal control over financial reporting is not considered effective, we may experience a loss of public confidence, which could have an adverse effect on our business and on the market price of our common stock.

#### Risks Related to Intellectual Property and Other Legal Matters

If product liability lawsuits are successfully brought against us, then we will incur substantial liabilities and may be required to limit commercialization of Gencaro or other product candidates.

We face product liability exposure related to the testing of our product candidates in human clinical trials, and may face exposure to claims by an even greater number of persons once we begin marketing and distributing our products commercially. If we cannot successfully defend against product liability claims, then we will incur substantial liabilities.

Regardless of merit or eventual outcome, liability claims may result in:

- decreased demand for our products and product candidates;
- injury to our reputation;
- withdrawal of clinical trial participants;
- costs of related litigation;
- substantial monetary awards to patients and others;
- loss of revenues; and
- the inability to commercialize our products and product candidates.

We have obtained limited product liability insurance coverage. Such coverage, however, may not be adequate or may not continue to be available to us in sufficient amounts or at an acceptable cost, or at all. We may not be able to obtain commercially reasonable product liability insurance for any product candidate.

Defending against claims relating to improper handling, storage or disposal of hazardous chemicals, radioactive or biological materials could be time consuming and expensive.

Our research and development of product candidates may involve the controlled use of hazardous materials, including chemicals, radioactive and biological materials. We cannot eliminate the risk of accidental contamination or discharge and any resultant injury from the materials. Various laws and regulations govern the use, manufacture, storage, handling and disposal of hazardous materials. We may be sued or be required to pay fines for any injury or contamination that results from our use or the use by third parties of these materials. Compliance with environmental

laws and regulations may be expensive, and current or future environmental regulations may impair our research, development and production efforts.

The loss of any rights to market key products would significantly impair our operating results.

We have licensed from CPEC, who has licensed rights in Gencaro from Bristol Meyers Squibb (BMS), the exclusive rights to Gencaro for all therapeutic and diagnostic uses in any country until the later of (i) 10 years from the first commercial sale of Gencaro in such country, or (ii) the termination of our commercial exclusivity in such country. This license includes a sublicense to us from BMS. We are obligated to use commercially reasonable efforts to develop and commercialize Gencaro, including obtaining regulatory approvals. Our ability to develop and commercialize Gencaro is dependent on numerous factors, including some factors that are outside of our control. CPEC has the right to terminate our license if we materially breach our obligations under the license agreement and fail to cure any such breach within the terms of the license.

If our license agreement with CPEC is terminated for reasons related to non-payment of fees, or for any other breach, then we would have no further rights to develop and commercialize Gencaro for any indication. The termination of this license, or of any other agreement which enables us to market a key product or product candidate, could significantly and adversely affect our business.

Certain intellectual property licensed by us is the subject of additional licensing arrangements to which the party that has licensed rights to us is subject. If such parties were to breach the terms of such licenses or such licenses were otherwise to terminate, our and our partners' rights to use such technology and develop and commercialize their products such as the genetic test may terminate and our business would be materially harmed.

Third parties may own or control patents or patent applications that we may be required to license to commercialize our product candidates or that could result in litigation that would be costly and time consuming.

Our or any strategic partner's ability to commercialize Gencaro and other product candidates depends upon our ability to develop, manufacture, market and sell these drugs without infringing the proprietary rights of third parties. A number of pharmaceutical and biotechnology companies, universities and research institutions have or may be granted patents that cover technologies similar to the technologies owned by or licensed to us. We may choose to seek, or be required to seek, licenses under third party patents, which would likely require the payment of license fees or royalties or both. We may also be unaware of existing patents that may be infringed by Gencaro, the genetic testing we intend to use in connection with Gencaro or our other product candidates. Because patent applications can take many years to issue, there may be other currently pending applications that may later result in issued patents that are infringed by Gencaro or our other product candidates. Moreover, a license may not be available to us on commercially reasonable terms, or at all.

There is a substantial amount of litigation involving patent and other intellectual property rights in the biotechnology and biopharmaceutical industries generally. If a third party claims that we are infringing on its technology, then our business and results of operations could be harmed by a number of factors, including:

- infringement and other intellectual property claims, even if without merit, are expensive and time-consuming to litigate and can divert management's attention from our core business;
- monetary damage awards for past infringement can be substantial;
- a court may prohibit us from selling or licensing product candidates unless the patent holder chooses to license the patent to us; and
- if a license is available from a patent holder, we may have to pay substantial royalties.

We may also be forced to bring an infringement action if we believe that a competitor is infringing our protected intellectual property. Any such litigation will be costly, time-consuming and divert management's attention, and the outcome of any such litigation may not be favorable to us.

Our intellectual property rights may not preclude competitors from developing competing products and our business may suffer.

Our competitive success will depend, in part, on our ability to obtain and maintain patent protection for our inventions, technologies and discoveries, including intellectual property that we license. The patent positions of biotechnology companies involve complex legal and factual questions, and we cannot be certain that our patents and licenses will successfully preclude others from using our technology. Consequently, we cannot be certain that any of our patents will provide significant market protection or will not be circumvented or challenged and found to be unenforceable or invalid. In some cases, patent applications in the U.S. and certain other jurisdictions are maintained

in secrecy until patents issue, and since publication of discoveries in the scientific or patent literature often lags behind actual discoveries, we cannot be certain of the priority of inventions covered by pending patent applications. Moreover, we may have to participate in interference proceedings declared by the U.S. Patent and Trademark Office to determine priority of invention or in opposition proceedings in a foreign patent office, any of which could result in substantial cost to us, even if the eventual outcome is favorable. There can be no assurance that a court of competent jurisdiction would hold any claims in any issued patent to be valid. An adverse outcome could subject us to significant liabilities to third parties, require disputed rights to be licensed from third parties or require us to cease using such technology. Regardless of merit, the listing of patents in the FDA Orange Book for Gencaro may be challenged as being improperly listed. We may have to defend against such claims and possible associated antitrust issues. We could also incur substantial costs in seeking to enforce our proprietary rights against infringement.

While the composition of matter patents on the compound that comprises Gencaro have expired, we hold the intellectual property concerning the interaction of Gencaro with the polymorphisms of the  $\beta 1$  and 2C receptors. We have obtained patents that claim methods involving Gencaro after a patient's receptor genotype has been determined. Our NDA requested a label that will include a claim that efficacy varies based on receptor genotype and a recommendation in the prescribing information that prospective patients be tested for their receptor genotype. We believe that under applicable law, a generic bucindolol label would likely be

required to include this recommendation as it pertains directly to the safe or efficacious use of the drug. Such a label may be considered as inducing infringement, carrying the same liability as direct infringement. If the label with the genotype information for Gencaro is not approved, or if generic labels are not required to copy the approved label, competitors could have an easier path to introduce competing products and our business may suffer. The approved label may not contain language covered by the patents, or we may be unsuccessful in enforcing them.

We may not be able to effectively protect our intellectual property rights in some foreign countries, as our patents are limited by jurisdiction and many countries do not offer the same level of legal protection for intellectual property as the U.S.

We require our employees, consultants, business partners and members of our scientific advisory board to execute confidentiality agreements upon the commencement of employment, consulting or business relationships with us. These agreements provide that all confidential information developed or made known during the course of the relationship with us be kept confidential and not disclosed to third parties except in specific circumstances. In the case of employees, the agreements provide that all inventions resulting from work performed for us, utilizing the property or relating to our business and conceived or completed by the individual during employment shall be our exclusive property to the extent permitted by applicable law.

Third parties may breach these and other agreements with us regarding our intellectual property and we may not have adequate remedies for the breach. Third parties could also fail to take necessary steps to protect our licensed intellectual property, which could seriously harm our intellectual property position.

If we are not able to protect our proprietary technology, trade secrets and know-how, then our competitors may develop competing products. Any issued patent may not be sufficient to prevent others from competing with us. Further, we have trade secrets relating to Gencaro, and such trade secrets may become known or independently discovered. Our issued patents and those that may issue in the future, or those licensed to us, may be challenged, opposed, invalidated or circumvented, which could allow competitors to market similar products or limit the patent protection term of our product candidates. All of these factors may affect our competitive position.

If the manufacture, use or sale of our products infringe on the intellectual property rights of others, we could face costly litigation, which could cause us to pay substantial damages or licensing fees and limit our ability to sell some or all of our products.

Extensive litigation regarding patents and other intellectual property rights has been common in the biopharmaceutical industry. Litigation may be necessary to assert infringement claims, enforce patent rights, protect trade secrets or know-how and determine the enforceability, scope and validity of certain proprietary rights. Litigation may even be necessary to defend disputes of inventorship or ownership of proprietary rights. The defense and prosecution of intellectual property lawsuits, U.S. Patent and Trademark Office interference proceedings, and related legal and administrative proceedings (e.g., a reexamination) in the U.S. and internationally involve complex legal and factual questions. As a result, such proceedings are costly and time-consuming to pursue, and their outcome is uncertain.

Regardless of merit or outcome, our involvement in any litigation, interference or other administrative proceedings could cause us to incur substantial expense and could significantly divert the efforts of our technical and management personnel. Any public announcements related to litigation or interference proceedings initiated or threatened against us could cause our stock price to decline. Adverse outcomes in patent litigation may potentially subject us to antitrust litigation which, regardless of the outcome, would adversely affect our business. An adverse determination may subject us to the loss of our proprietary position or to significant liabilities, or require us to seek licenses that may include substantial cost and ongoing royalties. Licenses may not be available from third parties, or may not be obtainable on satisfactory terms. An adverse determination or a failure to obtain necessary licenses may restrict or prevent us from manufacturing and selling our products, if any. These outcomes could materially harm our business, financial condition and results of operations.

### Risks Related to Stock Price Volatility

Ownership of our common stock is highly concentrated, and it may prevent you and other stockholders from influencing significant corporate decisions and may result in conflicts of interest that could cause our stock price to decline.

Our executive officers, directors and their affiliates beneficially owned approximately 7% of our outstanding common stock as of September 30, 2013. Accordingly, these executive officers, directors and their affiliates, acting individually or as a group, have substantial influence over the outcome of a corporate action of ours requiring stockholder approval, including the election of directors, any merger, consolidation or sale of all or substantially all of our assets or any other significant corporate transaction. These stockholders may also delay or prevent a change in control of us, even if such change in control would benefit our other stockholders. The significant concentration of stock ownership may adversely affect the value of our common stock due to investors' perception that conflicts of interest may exist or arise.

Our stock price is expected to be volatile.

Our common stock could be subject to significant fluctuations. Market prices for securities of early-stage pharmaceutical, biotechnology and other life sciences companies have historically been particularly volatile. Some of the factors that may cause the market price of our common stock to fluctuate include:

- the regulatory status of Gencaro and the genetic test, and whether and when they are approved for sale, if at all, and the labeling or other conditions of use imposed by the FDA;
- our ability to secure substantial additional funding or complete a strategic transaction or to complete development of and commercialize Gencaro;
- potential receipt of government or third party funding to further develop Gencaro;
- the results of our future clinical trials and any future NDAs of our current and future product candidates;
- the entry into, or termination of, key agreements, including key strategic alliance agreements;
- the results and timing of regulatory reviews relating to our product candidates;
- failure of any of our product candidates, if approved, to achieve commercial success;
- general and industry-specific economic conditions that may affect our research and development expenditures;
- the results of clinical trials conducted by others on drugs that would compete with our product candidates;
- issues in manufacturing our product candidates or any approved products;
- the initiation of or material developments in or the conclusion of litigation to enforce or defend any of our intellectual property rights;
- the loss of key employees;
- the introduction of technological innovations or new commercial products by our competitors;
- changes in estimates or recommendations by securities analysts, if any, who cover our common stock;
- future sales of our common stock;
- changes in the structure of health care payment systems;
- period-to-period fluctuations in our financial results; and
- our ability to retain the listing of our common stock on the Nasdaq Capital Market.

Moreover, the stock markets in general have experienced substantial volatility that has often been unrelated to the operating performance of individual companies. These broad market fluctuations may also adversely affect the trading price of our common stock. In the past, following periods of volatility in the market price of a company's securities, stockholders have often instituted class action securities litigation against those companies. Such litigation, if instituted, could result in substantial costs and diversion of management attention and resources, which could significantly harm our profitability and reputation.

Future sales or the possibility of future sales of our common stock may depress the market price of our common stock.

Sales in the public market of substantial amounts of our common stock could depress prevailing market prices of our common stock. As of September 30, 2013, approximately 13.2 million shares of common stock were outstanding. All of these shares are freely transferable without restriction or further registration under the Securities Act, except for shares held by our directors, officers and other affiliates and unregistered shares held by non-affiliates. The sale of these additional shares, or the perception that such sales may occur, could depress the market price of our common stock.

As of September 30, 2013 approximately 8.2 million shares of our common stock were issuable upon the exercise of outstanding warrants. Once a warrant is exercised, if the shares of our common stock issued upon the exercise of any such warrant are not available for sale in the open market without further registration under the Securities Act, then the holder can arrange for the resale of shares either by invoking any applicable registration rights, causing the shares

to be registered under the Securities Act and thus freely transferable, or by relying on an exemption to the Securities Act. If these registration rights, or similar registration rights that may apply to securities we may issue in the future, are exercised, it could result in additional sales of our common stock in the market, which may have an adverse effect on our stock price.

As of September 30, 2013, there were approximately 836,000 shares of our common stock which may be issued upon exercise of outstanding stock options. If and when these options are exercised, such shares will be available for sale in the open market without further registration under the Securities Act. The existence of these outstanding options may negatively affect our ability to complete future equity financings at acceptable prices and on acceptable terms. The exercise of those options, and the prompt resale of shares of our common stock received, may also result in downward pressure on the price of our common stock.

In the absence of a significant strategic transaction, we will need to raise significant additional capital to finance the research, development and commercialization of our drug products. If future securities offerings occur, they would dilute our current stockholders' equity interests and could reduce the market price of our common stock.

We do not expect to pay cash dividends, and accordingly, stockholders must rely on stock appreciation for any return on their investment.

We anticipate that we will retain our earnings, if any, for future growth and therefore do not anticipate paying cash dividends in the future. As a result, only appreciation of the price of our common stock will provide a return to stockholders. Investors seeking cash dividends should not invest in our common stock or Preferred Stock.

We have implemented anti-takeover provisions that could discourage, prevent or delay a takeover, even if the acquisition would be beneficial to our stockholders.

Provisions of our certificate of incorporation and bylaws, as well as provisions of Delaware law, could make it more difficult for a third party to acquire us, even if doing so would benefit our stockholders. These provisions:

- establish a classified board of directors so that not all members of our board may be elected at one time;
- authorize the issuance of up to approximately 4.9 million additional shares of preferred stock that could be issued by our board of directors to increase the number of outstanding shares and hinder a takeover attempt;
- limit who may call a special meeting of stockholders;
- prohibit stockholder action by written consent, thereby requiring all stockholder actions to be taken at a meeting of our stockholders; and
- establish advance notice requirements for nominations for election to our board of directors or for proposing matters that can be acted upon at a stockholder meeting.

Specifically, our certificate of incorporation provides that all stockholder action must be effected at a duly called meeting and not by a written consent. The bylaws provide, however, that our stockholders may call a special meeting of stockholders only upon a request of stockholders owning at least 50% of our outstanding common stock. These provisions of our certificate of incorporation and bylaws could discourage potential acquisition proposals and could delay or prevent a change in control. We designed these provisions to reduce our vulnerability to unsolicited acquisition proposals and to discourage certain tactics that may be used in proxy fights. These provisions, however, could also have the effect of discouraging others from making tender offers for our shares. As a consequence, they also may inhibit fluctuations in the market price of our shares that could result from actual or rumored takeover attempts. Such provisions also may have the effect of preventing changes in our management.

We are permitted to issue shares of our preferred stock without stockholder approval upon such terms as our board of directors determines. Therefore, the rights of the holders of our common stock are subject to, and may be adversely affected by, the rights of the holders of our preferred stock that may be issued in the future. In addition, the issuance of preferred stock could have a dilutive effect on the holdings of our current stockholders.

We are subject to the Delaware anti-takeover laws regulating corporate takeovers. These anti-takeover laws prevent a Delaware corporation from engaging in a merger or sale of more than 10% of its assets with any stockholder, including all affiliates and associates of the stockholder, who owns 15% or more of the corporation's outstanding voting stock, for three years following the date that the stockholder acquired 15% or more of the corporation's stock unless:

the board of directors approved the transaction where the stockholder acquired 15% or more of the corporation's stock;

after the transaction in which the stockholder acquired 15% or more of the corporation's stock, the stockholder owned at least 85% of the corporation's outstanding voting stock, excluding shares owned by directors, officers and employee stock plans in which employee participants do not have the right to determine confidentially whether shares held under the plan will be tendered in a tender or exchange offer; or

on or after this date, the merger or sale is approved by the board of directors and the holders of at least two-thirds of the outstanding voting stock that is not owned by the stockholder.

The provisions of our governing documents and current Delaware law may, collectively:

lengthen the time required for a person or entity to acquire control of us through a proxy contest for the election of a majority of our board of directors;

discourage bids for our common stock at a premium over market price; and

generally deter efforts to obtain control of us.

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

Not applicable.

ITEM 3. DEFAULTS UPON SENIOR SECURITIES

Not applicable.

ITEM 4. MINE SAFETY DISCLOSURES

Not applicable.

ITEM 5. OTHER INFORMATION

Not applicable.

ITEM 6. EXHIBITS

The following documents are filed as part of this quarterly report on Form 10-Q. The Company will furnish a copy of any exhibit listed to requesting stockholders upon payment of the Company's reasonable expenses in furnishing those materials.

Exhibit

| Number  | Description                                                                                                                                                                                       |
|---------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| 3.1     | Amended and Restated Certificate of Incorporation of the Registrant, as amended.(1)                                                                                                               |
| 3.1(a)  | Certificate of Amendment to Restated Certificate of Incorporation.(2)                                                                                                                             |
| 3.1(b)  | Form of Certificate of Designation of Preferences, Rights and Limitations of Series A Convertible Preferred Stock of the Registrant.(5)                                                           |
| 3.2     | Second Amended and Restated Bylaws of the Registrant.(3)                                                                                                                                          |
| 10.1    | Letter Agreement between ARCA biopharma, Inc. and Medtronic, Inc. dated as of July 26, 2013. (4) .                                                                                                |
| 10.2    | Office Lease Agreement by and between ARCA biopharma, Inc. and Circle Point Properties, LLC, effective August 1, 2013. (5)                                                                        |
| 10.3    | ARCA biopharma, Inc. 2013 Equity Incentive Plan. (6)                                                                                                                                              |
| 10.4    | Form of Stock Option Agreement and Option Grant Notice under 2013 Equity Incentive Plan (Standard). (6)                                                                                           |
| 10.5    | Form of Stock Option Agreement and Option Grant Notice under 2013 Equity Incentive Plan (Officer). (6)                                                                                            |
| 10.6    | Form of Stock Option Agreement and Option Grant Notice under 2013 Equity Incentive Plan (Director). (6)                                                                                           |
| 10.7    | Form of Restricted Stock Unit Award Agreement and Notice of Grant Award under 2013 Equity Incentive Plan (Standard). (6)                                                                          |
| 10.8    | Form of Restricted Stock Unit Award Agreement and Notice of Grant Award under 2013 Equity Incentive Plan (Officer). (6)                                                                           |
| 31.1*   | Certification of Chief Executive Officer pursuant to Rule 13a-14(a) or 15d-14(a) under the Securities Exchange Act of 1934, as adopted pursuant to Section 302 of the Sarbanes-Oxley Act of 2002. |
| 31.2*   | Certification of Chief Financial Officer pursuant to Rule 13a-14(a) or 15d-14(a) under the Securities Exchange Act of 1934, as adopted pursuant to Section 302 of the Sarbanes-Oxley Act of 2002. |
| 32.1*   | Certification of Chief Executive Officer and Chief Financial Officer pursuant to 18 U.S.C. sec. 1350, as adopted pursuant to Section 906 of the Sarbanes-Oxley Act of 2002.                       |
| 101.INS | XBRL Instance Document (filed electronically herewith)                                                                                                                                            |
| 101.SCH | XBRL Taxonomy Extension Schema Document (filed electronically herewith)                                                                                                                           |
| 101.CAL | XBRL Taxonomy Extension Calculation Linkbase Document (filed electronically herewith)                                                                                                             |
| 101.LAB | XBRL Taxonomy Extension Label Linkbase Document (filed electronically herewith)                                                                                                                   |
| 101.PRE | XBRL Taxonomy Extension Presentation Linkbase Document (filed electronically herewith)                                                                                                            |

101.DEF XBRL Taxonomy Extension Definition Linkbase Document (filed electronically herewith)

- (1) Previously filed with the SEC as an Exhibit to and incorporated herein by reference from ARCA biopharma, Inc. s Form 10-K, filed March 27, 2009, File No. 000-22873.
- (2) Previously filed with the SEC as an Exhibit to and incorporated herein by reference from ARCA biopharma, Inc. s Form 8-K, filed on March 5, 2013, File No. 000-22873.
- (3) Previously filed with the SEC as an Exhibit to and incorporated herein by reference from ARCA biopharma, Inc. s Form 10-Q, filed November 16, 2009, File No. 000-22873.
- (4) Previously filed with the SEC as an Exhibit to and incorporated herein by reference from ARCA biopharma, Inc. s Form 10-Q, filed August 13, 2013, File No. 000-22873.
- (5) Previously filed with the SEC as an Exhibit and incorporated herein by reference from ARCA biopharma, Inc. s Form 8-K, filed on August 6, 2013.

Previously filed with the SEC as an Exhibit and incorporated herein by reference from ARCA biopharma, Inc. s  
(6) Form 8-K, filed on September 23, 2013.

\* Filed herewith.

SIGNATURE

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

ARCA biopharma, Inc.

(Registrant)

By: /s/ Patrick M. Wheeler

Patrick M. Wheeler

Chief Financial Officer

Dated: November 13 , 2013

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\* Filed herewith.



CERTIFICATION

I, Michael R. Bristow, certify that:

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

Date: November 13, 2013

/s/ Michael R. Bristow

Michael R. Bristow  
President and Chief Executive Officer  
(Principal Executive Officer)

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CERTIFICATION

I, Patrick M. Wheeler, certify that:

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

Date: November 13, 2013

/s/ Patrick M. Wheeler

Patrick M. Wheeler  
Chief Financial Officer

(Principal Financial and Accounting Officer)

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Exhibit 32.1

ARCA BIOPHARMA, INC.

CERTIFICATION PURSUANT TO

18 U.S.C. SEC. 1350

AS ADOPTED PURSUANT TO

SECTION 906 OF THE SARBANES-OXLEY ACT OF 2002

Pursuant to the requirement set forth in Rule 13a-14(b) of the Securities Exchange Act of 1934, as amended (the "Exchange Act") and Section 1350 of Chapter 63 of Title 18 of the United States Code (18 U.S.C. §1350), Michael R. Bristow, Chief Executive Officer of ARCA biopharma, Inc. (the "Company"), and Patrick M. Wheeler, Chief Financial Officer of the Company, each hereby certifies that, to the best of his/her knowledge:

(1) The Company's Quarterly Report on Form 10-Q for the period ended September, 30, 2013, to which this Certification is attached as Exhibit 32.1 (the "Periodic Report") fully complies with the requirements of section 13(a) or 15(d) of the Exchange Act; and

(2) The information contained in the Periodic Report fairly presents, in all material respects, the financial condition and results of operations of the Company.

In Witness Whereof, the undersigned have set their hands hereto as of the 13<sup>th</sup> day of November 2013.

/s/ Michael R. Bristow

/s/ Patrick M. Wheeler

Michael R. Bristow  
President and Chief Executive Officer

Patrick M. Wheeler  
Chief Financial Officer

(Principal Executive Officer)

(Principal Financial and Accounting Officer)

A signed original of this written statement required by Section 906 of the Public Company Accounting Reform and Investor Protection Act of 2002 (18 U.S.C. § 1350, as adopted) has been provided to the Company and will be retained by the Company and furnished to the Securities and Exchange Commission or its staff upon request. This certification accompanies the Form 10-Q to which it relates, is not deemed filed with the Securities and Exchange Commission and is not to be incorporated by reference into any filing of ARCA biopharma, Inc. under the Securities Act of 1933, as amended, or the Securities Exchange Act of 1934, as amended (whether made before or after the date of the Form 10-Q), irrespective of any general incorporation language contained in such filing.

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