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

#### FORM 8-K

## CURRENT REPORT Pursuant to Section 13 OR 15(d) of The Securities Exchange Act of 1934

Date of Report (Date of earliest event reported): May 6, 2014 (May 6, 2014)

#### **NewLink Genetics Corporation**

(Exact name of registrant as specified in its charter)

Delaware001-3534242-1491350(State or other jurisdiction<br/>of incorporation)(Commission<br/>File Number)(IRS Employer<br/>Identification No.)

#### 2503 South Loop Drive Ames, IA

50010

(Address of principal executive offices)

(Zip Code)

Registrant's telephone number, including area code: (515) 296-5555

#### Not applicable

(Former name or former address, if changed since last report.)  $% \label{eq:continuous} % \label{eq:c$ 

Check the appropriate box below if the Form 8-K filing is intended to simultaneously satisfy the filing obligation of the registrant under any of the following provisions (see General Instruction A.2. below):

[ ] Written communications pursuant to Rule 425 under the Securities Act (17 CFR 230.425)
[ ] Soliciting material pursuant to Rule 14a-12 under the Exchange Act (17 CFR 240.14a-12)
[ ] Pre-commencement communications pursuant to Rule 14d-2(b) under the Exchange Act (17 CFR 240.14d-2(b))
[] Pre-commencement communications pursuant to Rule 13e-4(c) under the Exchange Act (17 CFR 240.13e-4(c))

#### **Section 2 - Financial Information**

#### Item 2.02. Results of Operations and Financial Condition.

On May 6, 2013, NewLink Genetics Corporation, a Delaware corporation (the "Company"), issued a press release reporting financial results for the first quarter ended March 31, 2014.

The press release is attached hereto as Exhibit 99.1, which is furnished under Item 2.02 of this report and shall not be deemed to be "filed" for purposes of Section 18 of the Securities Exchange Act of 1934 (the "Exchange Act") or otherwise subject to the liabilities of that section, nor shall it be deemed incorporated by reference in any filing under the Securities Act of 1933 or the Exchange Act, regardless of any general incorporation language in such filing.

#### Section 9 - Financial Statements and Exhibits

#### Item 9.01. Financial Statements and Exhibits.

(d) Exhibits.

Exhibit Number	Description				
99.1	Press Release, dated May 6, 2014, entitled "NewLink Genetics Corporation Reports First Quarter 2014 Financial Results"				

#### **SIGNATURES**

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

Dated: May 6, 2014

#### **NewLink Genetics Corporation**

By: /s/ Gordon H. Link, Jr.
Gordon H. Link, Jr.
Its: Chief Financial Officer

#### INDEX TO EXHIBITS

Exhibit Number	Description				
	Press Release, dated May 6, 2014, entitled "NewLink Genetics Corporation Reports First Quarter 2014 Financial				
99.1	Results"				



Contact:
Gordon Link
Chief Financial Officer
515-598-2925
glink@linkp.com

FOR IMMEDIATE RELEASE

#### **NewLink Genetics Corporation Reports First Quarter 2014 Financial Results**

AMES, IA -- 05/06/14 -- NewLink Genetics Corporation (NASDAQ: NLNK), today reported consolidated financial results for the first quarter of 2014 and progress in its development programs.

"In 2013, we achieved a major milestone when we completed patient enrollment in our algenpantucel-L pivotal Phase 3 IMPRESS study," commented Dr. Charles Link, Chairman and Chief Executive Officer of NewLink. Following completion of the first interim analysis in March of this year, the NewLink Genetics' Independent Data Safety Monitoring Committee (DSMC) reviewed patient data and, as anticipated, recommended study continuation without modification. A second interim analysis is planned upon reaching 333 patient events and, if needed, a final analysis is planned at 444 patient events. "We continue to look forward to the second interim analysis of our IMPRESS study near the end of this year and, assuming positive data, we plan to file a BLA for algenpantucel-L in 2015," said Dr. Link.

During 2014, the Company plans to continue advancing product development efforts across both HyperAcute and IDO pathway inhibitor platforms. Currently NewLink has six HyperAcute vaccines in various stages of clinical development for multiple indications (pancreas, lung, melanoma, prostate, breast and renal). In 2014, the Company continued expanding the breadth and depth of its IDO pathway inhibitor program. This included additional clinical development for its lead product candidate, indoximod, and also initiation of patient enrollment in a first-in-human clinical study of NLG919, its second compound from this platform.

"During the first quarter of 2014 at AACR we presented promising pre-clinical data demonstrating the synergistic anti-tumor activity of our IDO pathway inhibitors indoximod and NLG919 in combination with other checkpoint inhibitors and cancer immunotherapies," commented Dr. Nicholas Vahanian, President and Chief Medical Officer of NewLink. "At the same meeting, we also presented pre-clinical data on a novel class of anti-cancer agents called TDO inhibitors, which are structurally and functionally related to IDO."

NewLink reported a net loss of \$9.2 million or (\$.33) per share for the first quarter of 2014 compared to a net loss of \$7.9 million or (\$.33) per share for the comparable period in 2013.

Research and development expense in the first quarter of 2014 was \$6.4 million compared to \$6.3 million during the comparable period in 2013. The increase was primarily due to an increase in personnel-related expenses, offset by a decrease in contract research, manufacturing and consulting fees.

General and administrative expense in the first quarter of 2014 was \$3.3 million compared to \$2.0 million during the comparable period in 2013. The increase was primarily due to an increase in share-based compensation expense.

NewLink ended the quarter on March 31, 2014 with cash, cash equivalents, and certificates of deposit totaling \$84.0 million and expects to end the year with approximately \$40 million in cash, cash equivalents and

marketable securities. NewLink received gross proceeds from sales under its ATM of approximately \$28.3 million in the first quarter of 2014. NewLink ended the first quarter of 2014 with 27,862,390 shares outstanding.

#### **Recent Accomplishments**

- <u>HyperAcute Platform.</u> Completed first interim analysis for Phase 3 IMPRESS clinical trial and DSMC recommended study continuation without modification. Continued advancing the platform across multiple indications including pancreas, lung, melanoma and renal cancer.
- <u>IDO Inhibitors.</u> Presented preclinical data at the American Association for Cancer Research (AACR) 2014 Annual Meeting demonstrating that combining multiple checkpoint inhibitors that target the IDO (indoleamine-(2,3)-dioxygenase) pathway is effective in reducing local tumor-mediated immunosuppression and providing potential for enhanced anti-tumor activity. These data demonstrated the synergistic effects of combining NLG919, indoximod and anti-PD-1/PD-L1/PD-L2 antibodies to block both the IDO and PD pathways resulting in enhanced anti-tumor effects compared to blocking each pathway independently. This synergy was demonstrated in the context of established tumors treated with otherwise ineffective chemo-immunotherapy regimens.
- <u>TDO Inhibitors.</u> A novel class of compounds that mediate TDO (*tryptophan-2,3-dioxygenase*) activity were also presented at the AACR meeting. These data showed novel compounds with potent and selective TDO inhibition as well as IDO-specific inhibition and dual inhibition of TDO and IDO.

#### **About NewLink Genetics Corporation**

NewLink is a biopharmaceutical company focused on discovering, developing and commercializing novel immunotherapeutic products to improve treatment options for patients with cancer. NewLink's portfolio includes biologic and small molecule immunotherapy product candidates intended to treat a wide range of oncology indications. NewLink's product candidates are designed to harness multiple components of the immune system to combat cancer without significant incremental toxicity, either as a monotherapy or in combination with other treatment regimens. For more information please visit <a href="http://www.linkp.com">http://www.linkp.com</a>.

By leveraging its dual cancer immunotherapy platforms, which are designed to harness multiple components of the immune system to combat cancer, NewLink is well positioned to establish a leadership position in immuno-oncology. NewLink's HyperAcute® immunotherapy platform uniquely stimulates the patient's immune system to recognize and attack cancer cells, while its IDO pathway inhibitor platform technology targets a key immune checkpoint and disrupts mechanisms by which tumors evade the patient's immune system. NewLink's broad product pipeline includes biologic and small molecule immunotherapy product candidates designed to treat a wide range of oncology indications either as monotherapy or in combination with other treatment regimens. NewLink's most advanced product candidates include algenpantucel-L and tergenpumatucel-L HyperAcute immunotherapies, currently in Phase 3 clinical development for pancreatic cancer and Phase 2b/3 for non-small cell lung cancer, respectively. The IDO pathway inhibitor platform has two drug candidates currently in development. The first, indoximod, is currently in Phase 2 development for a range of solid tumor cancers. NewLink's second IDO pathway inhibitor, NLG919, is currently in Phase 1 development for advanced solid tumors. By targeting multiple immune system deficits, NewLink's product pipeline offers a broad approach to immuno-oncology.

#### Cautionary Note Regarding Forward-Looking Statements

This press release contains forward-looking statements of NewLink that involve substantial risks and uncertainties. All statements, other than statements of historical facts, contained in this press release are forward-looking statements, within the meaning of The Private Securities Litigation Reform Act of 1995. The words "anticipate," "believe," "estimate," "expect," "intend," "may," "plan," "target," "potential," "will," "could," "should," "seek," or the negative of these terms or other similar expressions are intended to identify forward-looking statements, although not all forward-looking statements contain these identifying words. These forward-looking statements include, among others, statements about: NewLink's financial guidance for 2014; enrollment in its clinical trials for product candidates based on NewLink's HyperAcute and IDO platform technologies; its timing of release of clinical data from ongoing clinical studies; its plans related to moving additional indications into clinical development; NewLink's future financial performance, results of operations, cash position and sufficiency of capital resources to fund its operating requirements; and any other statements other than statements of historical fact. Actual results or events could differ materially from the plans, intentions and expectations disclosed in the forward-looking

statements that NewLink makes due to a number of important factors, including those risks discussed in "Risk Factors" and elsewhere in NewLink's Annual Report on Form 10-K for the year ended December 31, 2013, Form S-3 Registration Statement filed December 28, 2012 and in its other filings with the Securities and Exchange Commission. The forward-looking statements in this press release represent NewLink's views as of the date of this press release. NewLink anticipates that subsequent events and developments will cause its views to change. However, while it may elect to update these forward-looking statements at some point in the future, it specifically disclaims any obligation to do so. You should, therefore, not rely on these forward-looking statements as representing NewLink's views as of any date subsequent to the date of this press release.

## NewLink Genetics Corporation Condensed Consolidated Statement of Operations (unaudited)

(In thousands, except share and per share amounts)

### Three Months Ended

	March 31,			
	 2014		2013	
Grant Revenue	\$ 334	\$	302	
Operating expenses:				
Research and development	6,387		6,343	
General and administrative	3,251		2,001	
Loss from operations	(9,304)		(8,042)	
Other (expense) income, net	68		108	
Net loss	\$ (9,236)	\$	(7,934)	
Net loss per common share, basic and diluted	\$ (0.33)	\$	(0.33)	
Weighted average common shares outstanding	 27,605,910		23,860,469	

# NewLink Genetics Corporation Condensed Consolidated Balance Sheets (unaudited)

(In thousands, except share and per share data)

	March 31, 2014		December 31, 2013	
Assets				
Current assets:				
Cash, cash equivalents and certificates of deposit	\$	83,962	\$	61,540
Prepaid expenses and other current assets		1,347		2,430
Total current assets		85,309		63,970
Property and equipment, net		6,434		6,587
Total assets	\$	91,743	\$	70,557
Liabilities and Equity				
Current liabilities:				
Accounts payable and accrued expenses	\$	3,358	\$	3,603
Deferred rent		84		84
Other current liabilities		190		189
Total current liabilities		3,632		3,876
Long-term liabilities:				
Royalty obligation payable		6,000		6,000
Notes payable and obligations under capital leases		986		1,033
Deferred rent		1,300		1,321
Total long-term liabilities		8,286		8,354
Total liabilities		11,918	'	12,230
Stockholder's equity:			,	
Preferred stock		_		_
Common stock		279		266
Additional paid-in capital, net		224,941		194,038
Treasury Stock, at cost		(182)		_
Deficit accumulated during the development stage		(145,213)		(135,977)
Total equity		79,825		58,327
Commitments		_		_
Total liabilities and equity	\$	91,743	\$	70,557