
**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): **February 11, 2019**

Progenics Pharmaceuticals, Inc.

(Exact Name of Registrant as Specified in Charter)

Delaware
(State or other jurisdiction
of incorporation)

000-23143
(Commission File Number)

13-3379479
(IRS Employer
Identification No.)

One World Trade Center, 47th Floor, New York, New York 10007
(Address of Principal Executive Offices) (Zip Code)

Registrant's telephone number, including area code: **(646) 975-2500**

Not Applicable
(Former name or former address, if changed since last report)

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:

- 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))

Indicate by check mark whether the registrant is an emerging growth company as defined in Rule 405 of the Securities Act of 1933 (17 CFR §230.405) or Rule 12b-2 of the Securities Exchange Act of 1934 (17 CFR §240.12b-2).

Emerging growth company

If an emerging growth company, indicate by check mark if the registrant has elected not to use the extended transition period for complying with any new or revised financial accounting standards provided pursuant to Section 13(a) of the Exchange Act.

Item 7.01. Regulation FD Disclosure.

On February 11, 2019, Progenics Pharmaceuticals, Inc. (“Progenics” or the “Company”) issued a press release announcing acquisition of the Somerset, NJ manufacturing facility for AZEDRA[®] (ioberguane I 131) for cash consideration of \$8.0 million. AZEDRA is the first and only FDA-approved radiopharmaceutical indicated for the treatment of pheochromocytoma and paraganglioma, ultra-rare cancers. A copy of the foregoing press release is furnished as Exhibit 99.1 to this Current Report on Form 8-K and is incorporated in this Item 7.01 by reference.

The information in this Item 7.01, including Exhibit 99.1 attached hereto, is being furnished and shall not be deemed “filed” for purposes of Section 18 of the Securities Exchange Act of 1934, as amended (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, as amended, or the Exchange Act, except as expressly set forth by specific reference in such a filing.

Item 9.01 Financial Statements and Exhibits

- (d) Exhibits.

Exhibit

No.	Description
99.1	<u>Press Release announcing acquisition of the Somerset, NJ manufacturing facility, dated February 11, 2019.</u>

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.

PROGENICS PHARMACEUTICALS, INC.

By: /s/ Patrick Fabbio
Patrick Fabbio
Senior Vice President and Chief Financial Officer
(Principal Financial and Accounting Officer)

Date: February 11, 2019



Progenics Pharmaceuticals, Inc.
One World Trade Center
47th Floor, Suite J
New York, New York 10007
(646) 975-2500
www.progenics.com

Contact: Melissa Downs
Investor Relations
(646) 975-2533
mdowns@progenics.com

Progenics Acquires AZEDRA[®] (iobenguane I 131) Radiopharmaceutical Manufacturing Facility

NEW YORK, NY, February 11, 2019 – Progenics Pharmaceuticals, Inc. (NASDAQ:PGNX), an oncology company developing innovative medicines and imaging analysis technology for targeting and treating cancer, today announced that it has acquired the Somerset, NJ manufacturing facility for AZEDRA[®] (iobenguane I 131) for cash consideration of \$8.0 million. AZEDRA is the first and only FDA-approved radiopharmaceutical indicated for the treatment of pheochromocytoma and paraganglioma, ultra-rare cancers.

This Somerset site serves as the launch facility for AZEDRA and will also provide manufacturing support for the Company's development stage radiopharmaceuticals, including 1095. The production of AZEDRA uses a proprietary Ultratrace[®] process which concentrates the MIBG targeted radiolytic activity by eliminating non-therapeutic "cold" MIBG molecules, giving AZEDRA a uniquely high specific activity.

Progenics has also secured the long-term supply of iodine necessary for the production of both AZEDRA and 1095.

"This strategic transaction extends our leadership position in radiopharmaceuticals, establishing the infrastructure and manufacturing capabilities to label multiple types of isotopes, including iodine-131," stated Mark Baker, CEO of Progenics. "With this transaction, we are building the capabilities to ensure the supply of AZEDRA."

About Progenics

Progenics develops innovative medicines and other technologies to target and treat cancer, including: therapeutic agents designed to treat cancer (AZEDRA[®], 1095, and PSMA TTC); prostate-specific membrane antigen ("PSMA") targeted imaging agents for prostate cancer (PyLTM); and imaging analysis technology (aBSI and PSMA AI). Progenics has two commercial products, AZEDRA, for the treatment of patients with unresectable, locally advanced or metastatic pheochromocytoma or paraganglioma (rare neuroendocrine tumors of neural crest origin) who require systemic anticancer therapy; and RELISTOR[®] (methylnaltrexone bromide) for the treatment of opioid-induced constipation, which is partnered with Bausch Health Companies Inc.

This press release contains "forward-looking statements" regarding future events. Statements contained in this communication that refer to Progenics' estimated or anticipated future results or other non-historical facts are forward-looking statements that reflect Progenics' current expectations regarding the AZEDRA manufacturing site and Progenics' current perspective of existing trends and information as of the date of this communication. Forward looking statements are generally accompanied by words such as "anticipate," "believe," "plan," "could," "should," "estimate," "expect," "forecast," "outlook," "guidance," "intend," "may," "might," "will," "possible," "potential," "predict," "project," or other similar words, phrases or expressions. Such statements are predictions only and are subject to risks and uncertainties that could cause actual events or results to differ materially. These risks and uncertainties include, among others, market acceptance for approved products; the cost, timing and unpredictability of results of clinical trials and other development activities and collaborations; our ability to successfully develop and commercialize products, such as 1095; our ability to operate the AZEDRA manufacturing site in a manner consistent with our current intentions; the effectiveness of the efforts of our partners to market and sell products on which we collaborate and the royalty revenue generated thereby; generic and other competition; the possible impairment of, inability to obtain and costs of obtaining intellectual property rights; possible product safety or efficacy concerns, general business, financial, regulatory and accounting matters, litigation and other risks. More information concerning Progenics and such risks and uncertainties is available on its website, and in its press releases and reports it files with the U.S. Securities and Exchange Commission, including those risk factors included in its Annual Report on Form 10-K for the fiscal year ended December 31, 2017, as updated in its subsequent Quarterly Reports on Form 10-Q. Progenics is providing the information in this press release as of its date and, except as expressly required by law, Progenics disclaims any intent or obligation to update or revise any forward-looking statements, whether as a result of new information, future events or circumstances or otherwise.

Additional information concerning Progenics and its business may be available in press releases or other public announcements and public filings made after this release. For more information, please visit www.progenics.com. Information on or accessed through our website or social media sites is not included in the company's SEC filings.

(PGNX-F)