

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): March 21, 2018

**FENNEC PHARMACEUTICALS INC.**

(Exact name of registrant as specified in its charter)

001-32295

(Commission File Number)

British Columbia, Canada

(State or other jurisdiction of  
incorporation)

20-0442384

(I.R.S. Employer Identification No.)

PO Box 13628, 68 TW Alexander Drive,

Research Triangle Park, NC

(Address of principal executive offices)

27709

(Zip Code)

Registrant's telephone number, including area code: (919) 636-4530

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 (§230.405 of this chapter) or Rule 12b-2 of the Securities Exchange Act of 1934 (§240.12b-2 of this chapter).

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 8.01 Other Events.**

On March 21, 2018, Fennec Pharmaceuticals Inc. (“Fennec”) announced the U.S. Food and Drug Administration (FDA) has granted PEDMARK™ (a unique formulation of sodium thiosulfate) Fast Track designation for the prevention of cisplatin-related ototoxicity in pediatric patients with standard risk hepatoblastoma (SR-HB).

On March 21, 2018, Fennec issued a press release announcing this news as described above. A copy of the press release is filed as Exhibit 99.1 to this Current Report on Form 8-K.

**Item 9.01 Financial Statements and Exhibits.**

(d) Exhibits.

<u>Exhibit No.</u>	<u>Description</u>
<a href="#">Exhibit 99.1</a>	<a href="#">Press Release dated March 21, 2018</a>

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**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.

FENNEC PHARMACEUTICALS INC.

Date March 23, 2018

By: /s/ Rostislav Raykov  
Rostislav Raykov  
Chief Executive Officer

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## FENNEC PHARMACEUTICALS RECEIVES FAST TRACK DESIGNATION BY FDA FOR PEDMARK

**Research Triangle Park, NC, March 21, 2018** – Fennec Pharmaceuticals Inc. (NASDAQ:FENC; TSX: FRX), a specialty pharmaceutical company, today announced that the U.S. Food and Drug Administration (FDA) has granted PEDMARK (a unique formulation of sodium thiosulfate) Fast Track designation for prevention of cisplatin-related ototoxicity in pediatric patients with standard risk hepatoblastoma (SR-HB). There are currently no drugs approved in the US for this condition. Fast Track designation is intended to facilitate development and expedite review of drugs to treat serious and life-threatening conditions so that an approved product can reach the market expeditiously.

"We believe the receipt of Fast Track designation from the FDA highlights the serious nature of hearing loss that patients have following cisplatin chemotherapy and the current lack of safe and effective treatments," said Rosty Raykov, President and Chief Executive Officer of Fennec. "We look forward to the more frequent interactions with the Agency that the Fast Track designation provides, as we prepare for the NDA filing."

Fast Track designation allows for closer collaboration with the review team within the Oncology Division at FDA, with the goal of getting important new drugs to patients more rapidly. Through the Fast Track program, a product may be eligible for priority review at the time of a New Drug Application (NDA) filing and may also be eligible to submit completed sections of the NDA on a rolling basis, before the complete application is submitted.

### About PEDMARK™ (Sodium Thiosulfate (STS))

Cisplatin and other platinum compounds are essential chemotherapeutic components for many pediatric malignancies. Unfortunately, platinum-based therapies cause ototoxicity in many patients, and are particularly harmful to the survivors of pediatric cancer.

In the U.S. and Europe there is estimated that over 10,000 children may receive platinum based chemotherapy. The incidence of hearing loss in these children depends upon the dose and duration of chemotherapy, and many of these children require lifelong hearing aids. There is currently no established preventive agent for this hearing loss and only expensive, technically difficult and sub-optimal cochlear (inner ear) implants have been shown to provide some benefit. Infants and young children at critical stages of development lack speech language development and literacy, and older children and adolescents lack social-emotional development and educational achievement.

STS has been studied by cooperative groups in two Phase 3 clinical studies of survival and reduction of ototoxicity, The Clinical Oncology Group Protocol ACCL0431 and SIOPEL 6. Both studies are completed. The COG ACCL0431 protocol enrolled one of five childhood cancers typically treated with intensive cisplatin therapy for localized and disseminated disease, including newly diagnosed hepatoblastoma, germ cell tumor, osteosarcoma, neuroblastoma, and medulloblastoma. SIOPEL 6 enrolled only hepatoblastoma patients with localized tumors.

### About Fennec Pharmaceuticals

Fennec Pharmaceuticals, Inc., is a specialty pharmaceutical company focused on the development of Sodium Thiosulfate (STS) for the prevention of platinum-induced ototoxicity in pediatric patients. STS has received Orphan Drug Designation in the US in this setting. For more information, please visit [www.fennecpharma.com](http://www.fennecpharma.com).

### For further information, please contact:

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