DURECT CORP Form 8-K July 12, 2011

## 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

July 12, 2011

(July 11, 2011)

**Date of Report** 

(Date of earliest event reported)

## **DURECT CORPORATION**

(Exact name of Registrant as specified in its charter)

# Edgar Filing: DURECT CORP - Form 8-K

Delaware (State or other jurisdiction of	000-31615 (Commission	94-3297098 (I.R.S. Employer
incorporation or organization)	File Number) 2 Results Way	Identification No.)
Cupertino, CA 95014		
(Address of principal executive offices) (Zip code)		
(408) 777-1417		
(Registrant s telephone number, including area code)		
(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	2 under the Exchange Act (17 CFR 240.14a-12)	
" Pre-commencement communications pursu	uant to Rule 14d-2(b) under the Exchange Act (17	CFR 240.14d-2(b))
" Pre-commencement communications pursu	uant to Rule 13e-4(c) under the Exchange Act (17	CFR 240.13e-4(c))

#### Item 1.01 Entry into a Material Definitive Agreement

On July 11, 2011, Durect Corporation ( Durect ) and Zogenix, Inc., a Delaware corporation ( Zogenix ), entered into a Development and License Agreement (the License Agreement ). Under the License Agreement, Zogenix will be responsible for the clinical development and commercialization of a proprietary, long-acting injectable formulation of risperidone using Durect s SABER controlled-release formulation technology in combination with Zogenix s DosePro® needle-free, subcutaneous drug delivery system. Durect will be responsible for non-clinical, formulation and CMC development responsibilities. Durect will be reimbursed by Zogenix for its research and development efforts on the product.

Zogenix will pay a non-refundable upfront fee to Durect of \$2.25 million. Zogenix is obligated to pay Durect up to \$103 million in total future milestone payments with respect to the product subject to and upon the achievement of various development, regulatory and sales milestones. Zogenix is also required to pay a mid single-digit to low double-digit percentage patent royalty on annual net sales of the product determined on a jurisdiction-by-jurisdiction basis. The patent royalty term is equal to the later of the expiration of all Durect technology patents or joint patent rights in a particular jurisdiction, the expiration of marketing exclusivity rights in such jurisdiction, or 15 years from first commercial sale in such jurisdiction. After the patent royalty term, Zogenix will continue to pay royalties on annual net sales of the product at a reduced rate for so long as Zogenix continues to sell the product in the jurisdiction. Zogenix is also required to pay to Durect a tiered percentage of fees received in connection with any sublicense of the licensed rights.

Durect granted to Zogenix an exclusive worldwide license, with sub-license rights, to Durect intellectual property rights related to Durect s proprietary polymeric and non-polymeric controlled-release formulation technology to make and have made, use, offer for sale, sell and import risperidone products, where risperidone is the sole active agent, for administration by injection in the treatment of schizophrenia, bipolar disorder or other psychiatric related disorders in humans. Durect retains the right to supply Zogenix s Phase 3 clinical trial and commercial product requirements on the terms set forth in the License Agreement.

Durect retains the right to terminate the License Agreement with respect to specific countries if Zogenix fails to advance the development of the product in such country, either directly or through a sublicensee. In addition, either party may terminate the License Agreement upon insolvency or bankruptcy of the other party, upon written notice of a material uncured breach or if the other party takes any act impairing such other party s relevant intellectual property rights. Zogenix may terminate the License Agreement upon written notice if during the development or commercialization of the product, the product becomes subject to one or more serious adverse drug experiences or if either party receives notice from a regulatory authority, independent review committee, data safety monitory board or other similar body alleging significant concern regarding a patient safety issue. Zogenix may also terminate the License Agreement with or without cause, at any time upon prior written notice.

#### **Item 8.01 Other Events**

Durect issued a press release announcing this event on July 12, 2011, a copy of which is attached hereto as Exhibit 99.1.

#### Item 9.01. Financial Statements and Exhibits

(d) Exhibits

99.1 Press Release of DURECT Corporation dated July 12, 2011

## Edgar Filing: DURECT CORP - Form 8-K

#### **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.

### **DURECT Corporation**

By: /s/ Matt Hogan

Date: July 12, 2011

Matt Hogan

Chief Financial Officer