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SCOLR Pharma, Inc. Form 8-K May 05, 2008

UNIT	TED STATES SECURITIES AT Washington,	ND EXCHANGE COMMISSION DC 20549
	Form	8-K
PURSUANT TO	CURRENT SECTION 13 OR 15(d) OF TH	REPORT IE SECURITIES EXCHANGE ACT OF 1934
	Date of report (date of ea May 5,	<u>*</u>
	SCOLR Pha (Exact name of registrant a	•
Delaware (State or other jurisdiction of incorporation)	001-31982 (Commission File No.)	91-1689591 (I.R.S. Employer Identification No.)
	3625 132nd Avenu Bellevue, W (Address of principal	/A 98006
	(425) 373 (Registrant's telephone num	
	low if the Form 8-K filing is in e following provisions (see Gen	tended to simultaneously satisfy the filing obligation of the struction A.2. below):
[] Written communications	pursuant to Rule 425 under the	Securities Act (17 CFR 230.425)
[] Soliciting material pursua	ant to Rule 14a-12 under the Ex	change Act (17 CFR 240.14a-12)
[] Pre-commencement com	munications pursuant to Rule 1-	4d-2(b) under the Exchange Act (17 CFR 240.14d-2(b))
Pre-commencement com	munications pursuant to Rule 1	3e-4(c) under the Exchange Act (17 CFR 240.13e-4(c))

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Item 7.01 Regulation FD Disclosure.

Representatives of SCOLR Pharma, Inc. will use the materials attached hereto as Exhibit 99.1 in investor presentations from time to time. We have also posted the presentation materials on our company website at www.scolr.com.

Please refer to page 2 of Exhibit 99.1 for a discussion of certain forward-looking statements included in the presentation materials. These forward-looking statements involve risks and uncertainties, including activities, events or developments that we expect, believe or anticipate will or may occur in the future. A number of factors could cause actual results to differ from those indicated in the forward-looking statements, including our ability to successfully develop new formulations and complete research and development, including pre-clinical and clinical studies, our ability to raise additional funds, the continuation of arrangements with our product development partners and customers, competition, government regulation and approvals, and general economic conditions. In order to obtain regulatory approvals for the commercial sale of potential products utilizing our CDT platform, we may be required to complete clinical trials in humans to demonstrate the safety and efficacy, or in certain cases, the bioequivalence, of the products. However, we may not be able to commence or complete these clinical trials in any specified time period, or at all, either because the appropriate regulatory agency objects or for other reasons. We plan to continue the costly process of simultaneously conducting clinical trials and preclinical research for multiple product candidates. Additional assumptions, risks and uncertainties are described in detail in our registration statements, reports and other filings with the Securities and Exchange Commission. Such filings are available on our website or at www.sec.gov.

Item 9.01 Financial Statements and Exhibits.

(d) Exhibits.

Exhibit No. Description

99.1 Presentation materials for the investor presentation by SCOLR Pharma, Inc.

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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 5, 2008

SCOLR PHARMA, INC.

By: /s/ Alan M. Mitchel

Alan M. Mitchel

Senior Vice President Business and Legal

Affairs

and Chief Legal Officer