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REPLIGEN CORP Form 8-K June 05, 2003

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) June 4, 2003

REPLIGEN CORPORATION

(Exact name of registration as specified in charter)

Delaware 0-14656 04-2729386 (State or other (Commission (IRS Employer jurisdiction of incorporation) Identification No.)

41 Seyon Street, Bldg. 1, Suite 100, Waltham, MA 02453 (Address of principal executive offices) (Zip Code)

Registrant's telephone number, including area code (781) 250-0111

(Former name or former address, if changed since last report.)

Item 5. Other Events and Required FD Disclosure.

In November 2000 and December 2000, Repligen entered into two License Agreements (the "UCSD License Agreements") with the University of California, San Diego ("UCSD") relating to certain patent applications pertaining to the use of uridine and uridine derivatives for the treatment of mitochondrial disease and purine autism. On June 21, 2001, Pro-Neuron, Inc. filed a complaint (the "Pro-Neuron Complaint") against the Regents of the University of California (the "Regents") and Repligen in the Superior Court of California, County of San Diego seeking to void the UCSD License Agreement relating to treatment of mitochondrial disease entered into between Repligen and the UCSD. Pro-Neuron subsequently made claims to the UCSD License Agreement related to purine autism, and claims for misappropriation of trade secrets.

On June 4, 2003 Repligen, the Regents and Pro-Neuron entered into a binding term sheet for settlement (the "Settlement") under which the Pro-Neuron complaint will be dismissed upon execution of definitive agreements between the parties. Under the terms of the Settlement Repligen will receive \$750,000. Repligen and the Regents agreed to restructure the UCSD Licensing Agreements to exclude the field of acylated pyrimidines, including triacetyluridine ("TAU"). Repligen will discontinue its clinical trial of TAU in mitochondrial disease and will continue its clinical trials of TAU in bipolar disorder/major depression and purine autism for up to two years. Repligen will assign to Pro-Neuron any inventions from these trials, for which it has rights, involving the use of acylated pyrimidines but will retain the rights to any inventions for all other chemical entities. Repligen may still direct future clinical trials and product development efforts to prodrugs or derivatives of uridine which are not acylated

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pyrimidines.

Item 7. Financial Statements, Pro Forma Financial Information and Exhibits.

(c) Exhibits.

None.

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.

REPLIGEN CORPORATION

Dated: June 5, 2003 By: /s/ Walter C. Herlihy

Walter C. Herlihy

Chief Executive Officer and President