UNITED STATES SECURITIES AND EXCHANGE COMMISSION

WASHINGTON, D.C. 20549

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 15, 2023

Apellis Pharmaceuticals, Inc.

(Exact Name of Registrant as Specified in its Charter)

Delaware (State or Other Jurisdiction of Incorporation)

001-38276 (Commission File Number)

27-1537290 (IRS Employer Identification No.)

100 Fifth Avenue Waltham, MA (Address of Principal Executive Offices)

02451 (Zip Code)

Registrant's telephone number, including area code: (617) 977-5700

Not applicable

	(Former Name o	or Former Address, if Changed Since Last I	Report)	
Charle the appropriate have below if t		anded to simultaneously satisfy the fi	ling obligation of the registrant under any of the	
following provisions (see General In:	O	ended to simultaneously satisfy the fi	ing obligation of the registrant under any of the	
☐ Written communication	Written communications pursuant to Rule 425 under the Securities Act (17 CFR 230.425)			
☐ Soliciting material purs	Soliciting material pursuant to Rule 14a-12 under the Exchange Act (17 CFR 240.14a-12)			
☐ Pre-commencement con	Pre-commencement communications pursuant to Rule 14d-2(b) under the Exchange Act (17 CFR 240.14d-2(b))			
☐ Pre-commencement con	Pre-commencement communications pursuant to Rule 13e-4(c) under the Exchange Act (17 CFR 240.13e-4(c))			
Securities registered pursuant to Sect	ion 12(b) of the Act:			
Title of each class		Trading Symbol(s)	Name of each exchange on which registered	
Common Stock		APLS	Nasdaq Global Select Market	
Indicate by check mark whether the r chapter) or Rule 12b-2 of the Securit			405 of the Securities Act of 1933 (§230.405 of this	
			Emerging growth company \Box	

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. \Box

Item 7.01 Regulation FD Disclosure.

On March 14, 2023, Apellis Pharmaceuticals, Inc. (the "Company") received notification from the U.S. Food and Drug Administration (the "FDA") that the FDA is extending the Prescription Drug User Fee Act ("PDUFA") goal date for the supplemental New Drug Application (the "sNDA") for the EMPAVELI Injector. The FDA stated in writing that "the labeling is still under review at this time and [the FDA] will miss the goal date of March 15, 2023." The timing of the new PDUFA goal date is still being determined.

The EMPAVELI Injector is an on-body device designed to enhance self-administration of EMPAVELI® (pegcetacoplan), which is approved for adults with paroxysmal nocturnal hemoglobinuria ("PNH").

The information in this Item 7.01 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.

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.

Date: March 15, 2023

Apellis Pharmaceuticals, Inc.

By: /s/ Timothy Sullivan

Timothy Sullivan Chief Financial Officer