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): December 15, 2023

ARCUTIS BIOTHERAPEUTICS, INC.

(Exact name of registrant as specified in its charter)

001-39186

(Commission

File Number)

Delaware (State or other jurisdiction

of incorporation)

81-2974255

(IRS Employer

Identification Number)

V	7 Townsgate Road, Suite 30 Westlake Village, CA 91361 rincipal executive offices, including	
Registrant's telephon	ne number, including area co	de: (805) 418-5006
theck the appropriate box below if the Form 8-K filing is intended bllowing provisions (see General Instructions A.2. below):	led to simultaneously satisfy t	he filing obligation of the registrant under any of the
Written communications pursuant to Rule 425 under the Se	curities Act (17 CFR 230.425	
Soliciting material pursuant to Rule 14a-12 under the Excha	ange 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))
ecurities registered pursuant to Section 12(b) of the Act:		
Title of each class	Trading Symbol(s)	Name of each exchange on which registered
Common Stock, par value \$0.0001 per share	ARQT	The Nasdaq Global Select Market
ndicate by check mark whether the registrant is an emerging gro- hapter) or Rule 12b-2 of the Securities Exchange Act of 1934 (see Fig. 2) an emerging growth company, indicate by check mark if the re- revised financial accounting standards provided pursuant to S	§ 240.12b-2 of this chapter). It egistrant has elected not to use	Emerging growth company \square the extended transition period for complying with any new

Item 8.01 Other Events.

FDA	Ap	prova	l
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On December 15, 2023, Arcutis Biotherapeutics, Inc. (the "Company") announced that the U.S. Food and Drug Administration (the "FDA") approved the new drug application for ZORYVE (roflumilast) topical foam, 0.3% for the treatment of seborrheic dermatitis in individuals 9 years of age and older. Another formulation of ZORYVE, roflumilast cream 0.3% is approved by the FDA for the topical treatment of plaque psoriasis, including intertriginous areas, in individuals 6 years of age and older.

SIGNATURES

Pursuant to the requirements of the Securities Exchange Act of 1934, as amended, the registrant has duly caused this report to be signed on its behalf by the undersigned hereunto duly authorized.

ARCUTIS BIOTHERAPEUTICS, INC.

By: /s/ Todd Franklin Watanabe

Date: December 18, 2023

Todd Franklin Watanabe

President and Chief Executive Officer