UNITED STATES SECURITIES AND EXCHANGE COMMISSION

		Washington, D.C. 205	49	
		FORM 8-K		
	Dursuant to Section	Current Report n 13 or 15(d)of the Secur		ue Act of 1934
		rt (Date of earliest event report	_	
	TF	AVERE THERAPEUT	TICS, INC.	
	(Exa	et name of registrant as specific	ed in its charter)	
	Delaware (State or other jurisdiction of incorporation)	001-36257 (Commission File Numbe	er)	27-4842691 (I.R.S. Employer Identification No.)
	(Address	3611 Valley Centre Drive, S San Diego, CA 9213 of Principal Executive Offices,	0	de)
	(Regis	(888) 969-7879 trant's Telephone Number, inclu	uding Area Code)	
	(Former Na	Not Applicable ne or Former Address, if Chanç	ged Since Last R	eport)
	eck the appropriate box below if the Form 8-K filing owing provisions:	is intended to simultaneously s	satisfy the filing o	bligation of the registrant under any of the
	Written communications pursuant to Rule 425 und	er the Securities Act (17 CFR 23	30.425)	
	Soliciting material pursuant to Rule 14a-12 under	the Exchange Act (17 CFR 240.1	.4a-12)	
	Pre-commencement communications pursuant to	Rule 14d-2(b) under the Exchanç	ge Act (17 CFR 24	40.14d-2(b))
	Pre-commencement communications pursuant to	Rule 13e-4(c) under the Exchanç	ge Act (17 CFR 24	40.13e-4(c))
	cate by check mark whether the registrant is an er pter) or Rule 12b-2 of the Securities Exchange Ac			of the Securities Act of 1933 (§230.405 of th
Em	erging growth company \square			
new	n emerging growth company, indicate by check may or revised financial accounting standards provide	d pursuant to Section 13(a) of t		
Sec	eurities registered pursuant to Section 12(b) of the	Act:		
	Title of each class	Trading Symbol(s)	Name o	f each exchange on which registered

Title of each class	Trading Symbol(s)	Name of each exchange on which registered
Common Stock, par value \$0.0001 per share	TVTX	The Nasdaq Global Market

Item 8.01 Other Events.

On September 7, 2021, Travere Therapeutics, Inc. (the "Company") provided a regulatory update on the Company's development program for sparsentan in focal segmental glomerulosclerosis ("FSGS") following a Type A meeting with the U.S. Food and Drug Administration (FDA) during which the Company and the FDA reached alignment on a pathway for the Company to proceed with a submission for accelerated approval, pending additional eGFR data. The Company intends to provide the FDA with additional eGFR data from the ongoing DUPLEX Study in the first half of 2022, and if such data are supportive, to submit an application for accelerated approval in the U.S. in mid-2022. The DUPLEX Study is continuing as planned with no changes to the statistical analysis plan, and patients will proceed in a blinded manner to assess the treatment effect on eGFR slope over 108 weeks in the confirmatory endpoint analysis.

Forward-Looking Statements

This Current Report on Form 8-K contains "forward-looking statements" as that term is defined in the Private Securities Litigation Reform Act of 1995. Without limiting the foregoing, these statements are often identified by the words "may", "might", "believes", "thinks", "anticipates", "plans", "expects", "intends" or similar expressions. In addition, expressions of our strategies, intentions or plans are also forward-looking statements. Such forward-looking statements include, but are not limited to, references to the Company's expected timing and plan for submitting additional eGFR data from the DUPLEX Study to the FDA; the expectation of submitting an application for accelerated approval of sparsentan for FSGS in mid-2022, pending additional supportive eGFR data; and expectations regarding the future conduct of the ongoing DUPLEX study. Such forward-looking statements are based on current information available to the Company and involve inherent risks and uncertainties, including factors that could delay, divert or change any such forward-looking statements, and could cause actual outcomes and results to differ materially from current expectations. No forward-looking statement can be guaranteed. Among the factors that could cause actual results to differ materially from those indicated in the forward-looking statements are risks and uncertainties associated with the regulatory review and approval process, including the Subpart H accelerated approval pathway in the United States, and the risk that the additional eGFR data will not support an accelerated approval submission. There is no guarantee that the FDA will accept for filing a NDA for sparsentan for FSGS under the Subpart H approval pathway, that the FDA will grant accelerated approval of sparsentan for FSGS or that sparsentan will be approved at all. The Company faces the risk that the Phase 3 DUPLEX Study of sparsentan in FSGS will not demonstrate that sparsentan is safe or effective or serve as a basis for accelerated approval of sparsentan as planned and risk that ongoing clinical trials may not succeed or may be delayed for safety, regulatory or other reasons. In addition, such risks and uncertainties may include those described in the Company's annual, quarterly and current reports (i.e., Form 10-K, Form 10-Q and Form 8-K) as filed or furnished with the SEC, which are available at the Company's website (www.travere.com) under "Investors & Media". You are cautioned not to place undue reliance on any forward-looking statements as there are important factors that could cause actual results to differ materially from those in any forward-looking statements, many of which are beyond our control. Except to the extent required by law, the Company undertakes no obligation to publicly update any forward-looking statement.

Item 9.01 Financial Statements and Exhibits.

(d) Exhibits

104 Cover Page Interactive Data File (embedded within the Inline XBRL document).

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: September 7, 2021

TRAVERE THERAPEUTICS, INC.

By: /s/ Elizabeth E. Reed

Name: Elizabeth E. Reed

Title: Senior Vice President, General Counsel and Secretary