



Tricida to Present at the Cowen and Company 39th Annual Health Care Conference

February 27, 2019

SOUTH SAN FRANCISCO, Calif.--(BUSINESS WIRE)--Feb. 27, 2019-- Tricida, Inc. (Nasdaq: TCDA) today announced that it will present at the Cowen and Company 39th Annual Health Care Conference on Tuesday, March 12, 2019 at 8:20 am PT / 11:20 am ET. Geoff Parker, Tricida's Chief Financial Officer, will provide a company overview, business update and progress on the company's key initiatives.

A live webcast of the presentation will be accessible on the Tricida website at IR.Tricida.com. An archive of the webcast will be available for 90 days following the presentation.

About Tricida

Tricida, Inc. is a pharmaceutical company focused on the development and commercialization of its drug candidate, TRC101, a non-absorbed, orally-administered polymer designed to treat metabolic acidosis in patients with chronic kidney disease (CKD). Metabolic acidosis is a condition commonly caused by CKD that is believed to accelerate the progression of kidney deterioration. Tricida has successfully completed a Phase 3, double-blind, placebo-controlled trial of TRC101 in patients with CKD and metabolic acidosis. The results of this Phase 3 trial, along with results from a successful double-blind, randomized, placebo-controlled Phase 1/2 trial and an ongoing safety extension trial, TRCA-301E, are intended to serve as the basis for the submission of an NDA for TRC101 under the FDA's Accelerated Approval Program.

For more information about Tricida, please visit www.Tricida.com.

Cautionary Note on Forward-Looking Statements

This press release includes forward-looking statements, including for example, statements about our ability to submit an NDA for TRC101 under the FDA's Accelerated Approval Program. Forward -looking statements involve known and unknown risks, uncertainties, assumptions and other factors that may cause our actual results, performance or achievements to be materially different from any future results, performance or achievements expressed or implied by the forward-looking statements. These risks and uncertainties include, among others, timing and results of clinical trials and other studies; that many drug candidates that have completed Phase 3 trials do not become approved drugs on a timely or cost effective basis or at all; there can be no assurance that the FDA would approve an NDA under the Accelerated Approval Program, or at all, and even if approval for a drug is obtained, there can be no assurance that it will be adopted in the market or accepted as a benefit to patients and healthcare providers; possible safety and efficacy concerns; and that we completely rely on third-party suppliers to manufacture our clinical drug supply. The forward-looking statements contained in this press release reflect Tricida's current views with respect to future events, and Tricida does not undertake and specifically disclaims any obligation to update any forward-looking statements.

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