

August 22, 2017



Moleculin Announces Meeting with European Medicines Agency

HOUSTON, TX -- (Marketwired) -- 08/22/17 -- Moleculin Biotech, Inc. (NASDAQ: MBRX) ("Moleculin" or the "Company"), a preclinical pharmaceutical company focused on the development of anti-cancer drug candidates, some of which are based on license agreements with The University of Texas System on behalf of the M.D. Anderson Cancer Center, today announced it will meet with the European Medicines Agency (EMA) on Wednesday, August 30, 2017 to discuss Clinical Trial Authorization for the study of Annamycin for the treatment of acute myeloid leukemia.

The European Medicines Agency is the European equivalent to the US Food and Drug Administration (FDA) and oversees the approval of new drugs for the European Union.

"With our planned clinical trial for Annamycin in the US and Poland which may begin shortly, dependent upon the filing and allowance of an IND with the FDA," commented Walter Klemp, Chairman and CEO of Moleculin, "we want to be in a position to move quickly with Annamycin in the rest of Europe. This meeting will lay the groundwork for expanding Annamycin's market throughout the European Union."

About Moleculin Biotech, Inc.

Moleculin Biotech, Inc. is a preclinical stage pharmaceutical company focused on the development of anti-cancer drug candidates, some of which are based on discoveries made at M.D. Anderson Cancer Center. Our lead product candidate is Annamycin, an anthracycline being studied for the treatment of relapsed or refractory acute myeloid leukemia, more commonly referred to as AML. We also have two preclinical small molecule portfolios, one of which is focused on the modulation of hard-to-target tumor cell signaling mechanisms and the recruitment of the patient's own immune system. The other portfolio targets the metabolism of tumors.

For more information about the Company, please visit <http://www.moleculin.com>.

Forward-Looking Statements

Some of the statements in this release are forward-looking statements within the meaning of Section 27A of the Securities Act of 1933, Section 21E of the Securities Exchange Act of 1934 and the Private Securities Litigation Reform Act of 1995, which involve risks and uncertainties. Forward-looking statements in this press release include, without limitation, the ability of Annamycin to begin clinical trials. These statements relate to future events, future expectations, plans and prospects. Although Moleculin Biotech believes that the expectations reflected in such forward-looking statements are reasonable as of the date made, expectations may prove to have been materially different from the results expressed

or implied by such forward-looking statements. Moleculin Biotech has attempted to identify forward-looking statements by terminology including "believes," "estimates," "anticipates," "expects," "plans," "projects," "intends," "potential," "may," "could," "might," "will," "should," "approximately" or other words that convey uncertainty of future events or outcomes to identify these forward-looking statements. These statements are only predictions and involve known and unknown risks, uncertainties, and other factors, including those discussed under Item 1A. "Risk Factors" in our most recently filed Form 10-K filed with the Securities and Exchange Commission ("SEC") and updated from time to time in our Form 10-Q filings and in our other public filings with the SEC. Any forward-looking statements contained in this release speak only as of its date. We undertake no obligation to update any forward-looking statements contained in this release to reflect events or circumstances occurring after its date or to reflect the occurrence of unanticipated events.

Contacts

PCG Advisory Group

Investors:

Kirin M. Smith

Chief Operating Officer

D: 646.863.6519

E: ksmith@pcgadvisory.com

Source: Moleculin Biotech, Inc.