

Medical Leaders Join BioSig Clinical Advisory Board

New Physician Board to Drive Adoption of PURE EP System

Santa Monica, CA, Nov. 28, 2018 (GLOBE NEWSWIRE) -- BioSig Technologies, Inc. (NASDAQ: BSGM), a medical device company developing a proprietary biomedical signal processing platform designed to address an unmet technology need for the \$4.6 billion electrophysiology (EP) marketplace, today announced that the Company formed a Clinical Advisory Board to complement existing scientific relationships with the nationwide key opinion leaders built during the R&D phase of product development.

The goal of the newly formed Board is to advise the Company on a range of aspects related to commercialization of BioSig's novel signal acquisition and processing technology, PURE EP(tm) System. Physicians from 17 leading medical centers in the United States will be gathering for the Board's first meeting on November 30, 2018. The agenda will include focused consultations on the human clinical data collection, commercial strategy for PURE EP(tm) System and potential clinical benefits of new technologies in the R&D pipeline.

"Our technology has been developed together with some of the most respected key opinion leaders in the electrophysiology industry. As we progress towards full commercial launch of our PURE EP(tm) System, it is essential to engage a wider group of talented physicians who can strengthen our product and launch plan. We are pleased that so many leading electrophysiologists agreed to dedicate their valuable time and provide input to help us better address unmet clinical needs in this very complex area of arrhythmia treatments," commented Kenneth L. Londoner, Chairman and CEO of BioSig Technologies, Inc.

BioSig received FDA 510(k) clearance for its first product, PURE EP(tm) System on August 8, 2018. The Company began trading on the Nasdaq Capital Markets on September 21, 2018 under the ticker symbol BSGM.

About BioSig Technologies

BioSig Technologies is a medical technology company developing a proprietary biomedical signal processing platform designed to improve the electrophysiology (EP) marketplace (www.biosigtech.com). Led by a proven management team and a veteran, independent Board of Directors, Los Angeles-based BioSig Technologies is preparing to commercialize its PURE EP[™] System. The technology has been developed to address an unmet need in a large and growing market.

The Company's first product, PURE EP™ System, is a novel cardiac signal acquisition and

display system which is engineered to assist electrophysiologists in clinical decision- making during procedures to diagnose and treat patients with abnormal heart rates and rhythms. BioSig's main goal is to deliver technology to improve upon catheter ablation treatments for the prevalent and potentially deadly arrhythmias, Atrial Fibrillation and Ventricular Tachycardia. BioSig has partnered with Minnetronix on technology development and has received FDA 510(k) clearance for the PURE EP[™] System in August 2018.

Forward-looking Statements

This press release contains "forward-looking statements." Such statements may be preceded by the words "intends," "may," "will," "plans," "expects," "anticipates," "projects," "predicts," "estimates," "aims," "believes," "hopes," "potential" or similar words. Forwardlooking statements are not guarantees of future performance, are based on certain assumptions and are subject to various known and unknown risks and uncertainties, many of which are beyond the Company's control, and cannot be predicted or quantified and consequently, actual results may differ materially from those expressed or implied by such forward-looking statements. Such risks and uncertainties include, without limitation, risks and uncertainties associated with (i) our inability to manufacture our products and product candidates on a commercial scale on our own, or in collaboration with third parties; (ii) difficulties in obtaining financing on commercially reasonable terms; (iii) changes in the size and nature of our competition; (iv) loss of one or more key executives or scientists; and (v) difficulties in securing regulatory approval to market our products and product candidates. More detailed information about the Company and the risk factors that may affect the realization of forward-looking statements is set forth in the Company's filings with the Securities and Exchange Commission (SEC), including the Company's Annual Report on Form 10-K and its Quarterly Reports on Form 10-Q. Investors and security holders are urged to read these documents free of charge on the SEC's website at http://www.sec.gov. The Company assumes no obligation to publicly update or revise its forward-looking statements as a result of new information, future events or otherwise.

Contact: Natasha Drapeau BioSig Technologies, Inc. Executive Vice President 12600 Hill Country Blvd R-275 Austin, TX 78738 ndrapeau@biosigtech.com 512-329-2643



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