

## SIGA Technologies to Host Business Update Call on March 2nd, 2023 Following Release of Fourth Quarter and Full Year 2022 Financial Results

February 23, 2023

NEW YORK, Feb. 23, 2023 (GLOBE NEWSWIRE) -- SIGA Technologies, Inc. (SIGA) (NASDAQ: SIGA), a commercial-stage pharmaceutical company, today announced that management will host a webcast and conference call to provide a business update at 4:30 P.M. ET on Thursday, March 2<sup>nd</sup>, 2023. Participating on the call will be Dr. Phil Gomez, Chief Executive Officer and Daniel Luckshire, Chief Financial Officer.

A live webcast of the call will also be available on the Company's website at <a href="www.siga.com">www.siga.com</a> under the 'Events & Presentations' tab in the Investor Relations section, or by <a href="clicking here">clicking here</a>. Please log in approximately 5-10 minutes prior to the scheduled start time.

Participants may access the call by dialing 1-877-425-9470 for domestic callers or 1-201-389-0878 for international callers.

A replay of the call will be available for two weeks by dialing 1-844-512-2921 for domestic callers or 1-412-317-6671 for international callers and using Conference ID: 13735849. The archived webcast will be available in the Events and Presentations section of the Company's website.

## ABOUT SIGA TECHNOLOGIES, INC. and TPOXX®

SIGA Technologies, Inc. is a commercial-stage pharmaceutical company focused on the health security market. Health security comprises countermeasures for biological, chemical, radiological and nuclear attacks (biodefense market), vaccines and therapies for emerging infectious diseases, and health preparedness. Our lead product is TPOXX®, also known as tecovirimat and ST-246®, an orally administered and IV formulation antiviral drug for the treatment of human smallpox disease caused by variola virus. TPOXX is a novel small-molecule drug and the US maintains a supply of TPOXX under Project BioShield. The oral formulation of TPOXX was approved by the FDA for the treatment of smallpox in 2018, and the IV formulation was approved for the same indication in 2022. The full label is available by clicking here. Oral tecovirimat received approval from the European Medicines Agency (EMA) and the Medicines and Healthcare Products Regulatory Agency (MHRA) in the United Kingdom in 2022. The EMA and UK approvals include labeling for oral tecovirimat indicating its use for the treatment of smallpox, monkeypox, cowpox, and vaccinia complications following vaccination against smallpox. The full label is available by clicking here. In September 2018, SIGA signed a contract with the Biomedical Advanced Research and Development Authority (BARDA), part of the office of the Assistant Secretary for Preparedness and Response within the U.S. Department of Health and Human Services, for additional procurement and development related to both oral and intravenous formulations of TPOXX. For more information about SIGA, please visit <a href="https://www.siga.com">www.siga.com</a>. The information contained on the websites referenced in this press release is not incorporated by reference herein.

## FORWARD-LOOKING STATEMENTS

This press release contains "forward-looking statements" within the meaning of the Private Securities Litigation Reform Act of 1995, as amended, including statements relating to the progress of SIGAs development programs and timelines for bringing products to market, as well as the impact of COVID-19 on SIGA's business. Forward-looking statements may be identified by words or phrases such as "believes," "estimates," "expects," "may," "will," "would," "can," "could," and similar words and phrases. Such forward-looking statements are based on current expectations and assumptions and subject to various known and unknown risks and uncertainties, and SIGA cautions you that any forward-looking information provided by or on behalf of SIGA is not a guarantee of future performance. SIGA's actual results could differ materially from those anticipated by such forward-looking statements due to a number of factors, some of which are beyond SIGAs control, including, but not limited to, (i) the risk that the U.S. Biomedical Advanced Research and Development Authority ("BARDA") elects, in its sole discretion as permitted under the BARDA Contracts (as defined below), not to exercise all, or any, of the remaining unexercised options under those contracts, (ii) the risk that SIGA may not complete performance under its contracts with BARDA (the "BARDA Contracts") on schedule or in accordance with contractual terms, (iii) the risk that the BARDA Contracts are modified or canceled at the request or requirement of the U.S. government, (iv) the risk that the nascent international biodefense market does not develop to a degree that allows SIGA to successfully market TPOXX internationally, (v) the risk that potential products, including potential alternative uses or formulations of TPOXX that appear promising to SIGA or its collaborators, cannot be shown to be efficacious or safe in subsequent pre-clinical or clinical trials, (vi) the risk that SIGA or its collaborators will not obtain appropriate or necessary governmental approvals to market these or other potential products or uses, (vii) the risk that SIGA may not be able to secure or enforce sufficient legal rights in its products, including intellectual property protection, (viii) the risk that any challenge to SIGAs patent and other property rights, if adversely determined, could affect SIGAs business and, even if determined favorably, could be costly, (ix) the risk that regulatory requirements applicable to SIGA's products may result in the need for further or additional testing or documentation that will delay or prevent SIGA from seeking or obtaining needed approvals to market these products, (x) the risk that the volatile and competitive nature of the biotechnology industry may hamper SIGAs efforts to develop or market its products, (xi) the risk that changes in domestic or foreign economic and market conditions may affect SIGAs ability to advance its research or may affect its products adversely, (xii) the effect of federal, state, and foreign regulation, including drug regulation and international trade regulation, on SIGAs businesses, (xiii) the risk that the COVID-19 pandemic could impact SIGAs operations by disrupting SIGAs supply chain for the manufacture of TPOXX, causing delays in SIGA's research and development activities, causing delays or the re-allocation of funding in connection with SIGA's government contracts,

or diverting the attention of government staff overseeing SIGA's government contracts, (xiv) the risk that the U.S. or foreign governments' responses (including inaction) to national or global economic conditions or infectious diseases such as COVID-19 are ineffective and may affect SIGA's business adversely, and (xv) other risk factors discussed in Item 1A. "Risk Factors" of SIGA's Annual Report on Form 10-K for the year ended December 31, 2021, and in SIGA's subsequent filings with the U.S. Securities and Exchange Commission. These documents are publicly available at the SEC's website at <a href="https://investor.siga.com">https://investor.siga.com</a>. Forward-looking statements are current only as of the date on which such statements were made, and except as may be otherwise required by law, we undertake no obligation to update publicly any forward-looking statements whether as a result of new information, future events, or otherwise.

The information contained in this press release does not necessarily reflect the position or the policy of the U.S. government and no official endorsement should be inferred. The information contained on the websites referenced herein is not incorporated by reference into this press release.

## **Investor Contacts:**

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