

March 5, 2020



CNS Pharmaceuticals to Report Fourth Quarter 2019 Financial Results on March 12, 2020

Company to Host Business Update Conference Call at 4:30 pm ET

HOUSTON, March 5, 2020 /PRNewswire/ --**CNS Pharmaceuticals, Inc.** (NASDAQ: CNSP) ("CNS" or the "Company"), a biopharmaceutical company specializing in the development of novel treatments for primary and metastatic cancers of the brain and central nervous system, plans to report financial results for the three and 12 months ended December 31, 2019, after the close of the U.S. financial markets on Thursday, March 12, 2020.



CNS' senior management will provide a business update in a conference call and live audio webcast at 4:30 p.m. Eastern time on Thursday, March 12, 2020. The conference call dial-in and webcast information is as follows:


DOMESTIC DIAL-IN:	(844) 535-4071
INTERNATIONAL DIAL-IN:	(706) 679-2458
PASSCODE:	1254059
WEBCAST:	CNS Business Update Conference Call

For those unable to participate in the live conference call or webcast, a replay will be available beginning approximately two hours after the close of the conference call. To access the replay, dial 855-859-2056 or 404-537-3406. The replay passcode is 1254059. The replay can be accessed for a period of time on CNS website at [CNS Business Update Conference Call](#).

About CNS Pharmaceuticals, Inc.

CNS Pharmaceuticals is developing novel treatments for primary and metastatic cancers of the brain and central nervous system. Its lead drug candidate, Berubicin, is proposed for the

treatment of glioblastoma multiforme (GBM), an aggressive and incurable form of brain cancer. CNS holds a worldwide exclusive license to the Berubicin chemical compound and has acquired all data and know-how from Reata Pharmaceuticals, Inc. related to a completed Phase 1 trial with Berubicin in GBM which Reata conducted in 2006. In this trial, 44% of patients experienced a statistically significant improvement in progression-free survival. This 44% disease control rate was based on 11 patients (out of 25 evaluable patients) with stable disease, plus responders. One patient experienced a durable complete response and remains cancer-free as of February 20, 2020. In the second half of 2020, CNS expects to commence a Phase 2 clinical trial of Berubicin for the treatment of GBM in the U.S., while a sub-licensee partner undertakes a Phase 2 trial in adults and a first-ever Phase 1 trial in pediatric GBM patients in Poland. Its second drug candidate, WP1244, is a novel DNA binding agent that has shown in preclinical studies that it is 500-times more potent than the chemotherapeutic agent daunorubicin in inhibiting tumor cell proliferation.

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