



NERVGEN PHARMA ENGAGES CRO FOR PHASE 1 CLINICAL TRIAL

Vancouver, Canada. December 15, 2020 – **NervGen Pharma Corp. (TSX-V: NGEN) (OTCQX: NGENF)** (“NervGen” or the “Company”), a biotech company dedicated to creating innovative solutions for the treatment of nerve damage and neurodegenerative diseases, today announced that it has engaged Novotech (Australia) Pty Limited (“Novotech”), a leading full-service contract research organization (“CRO”) in Asia-Pacific, for its upcoming Phase 1 clinical trial for NVG-291. NervGen currently plans to initiate its Phase 1 clinical trial in Australia starting in Q1 2021.

“We are very pleased to bring on one of Asia-Pacific’s foremost CROs to support our first-in-human clinical study,” stated Paul Brennan, NervGen’s President & CEO. “With Novotech’s experience in conducting early-stage clinical trials, we are optimally positioned to execute the Phase 1 program in Australia where we expect to have a reduced risk of impact from COVID-19, as well as the ability to conduct this phase of our development quickly and cost-effectively.”

About Novotech

Novotech is internationally recognized as the leading regional full-service contract research organization. Novotech has been instrumental in the success of over a thousand Phase I - IV clinical trials for biotechnology companies. Novotech was established in 1996, with offices in 11 locations across the region, and site partnerships with major health institutions.

Novotech provides clinical development services across all clinical trial phases and therapeutic areas including: feasibility assessments; ethics committee and regulatory submissions, data management, statistical analysis, medical monitoring, safety services, central lab services, report write-up to ICH requirements, project and vendor management.

Novotech is a subsidiary of Novotech Health Holdings, the leading Asian biotech CRO.

About NervGen

NervGen is restoring life’s potential by creating innovative solutions for the treatment of nerve damage and neurodegenerative diseases. The Company is developing drugs for the treatment of multiple sclerosis, spinal cord injury and Alzheimer’s disease. NervGen’s platform technology targets protein tyrosine phosphatase sigma (“PTP σ ”), a neural receptor that impedes nerve repair. Inhibition of the PTP σ receptor has been shown to promote regeneration and remyelination of damaged nerves, as well as improvement of nerve function in animal models for various medical conditions.

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Follow NervGen on Twitter (@NervgenC) and LinkedIn (NervGen Pharma Corp.) for the latest news on the Company.

Neither the TSX Venture Exchange nor its Regulation Services Provider (as that term is defined in the policies of the TSX Venture Exchange) accepts responsibility for the adequacy or accuracy of this release.

Cautionary Note Regarding Forward-Looking Statements

This news release may contain “forward-looking information” and “forward-looking statements” within the meaning of applicable Canadian and United States securities legislation. Such forward-looking statements and information herein include, but are not limited to, the Company’s current and future plans, expectations and intentions, results, levels of activity, performance, goals or achievements, or any other future events or developments constitute forward-looking statements, and the words “may”, “will”, “would”, “should”, “could”, “expect”, “plan”, “intend”, “trend”, “indication”, “anticipate”, “believe”, “estimate”, “predict”, “likely” or “potential”, or the negative or other variations of these words or other comparable words or phrases, are intended to identify forward-looking statements. Forward-looking statements include, without limitation, statements relating to: the clinical development of NVG-291 for multiple sclerosis and spinal cord injuries, both sub-acute and chronic; steps taken to minimize the impact of the COVID-19 pandemic on our operations; our belief that we can quickly and cost-effectively conduct our Phase 1 study in Australia; and the creation of innovative solutions for the treatment of nerve damage and neurodegenerative diseases.

Forward-looking statements are based on estimates and assumptions made by the Company in light of management’s experience and perception of historical trends, current conditions and expected future developments, as well as other factors that we believe are appropriate and reasonable in the circumstances. In making forward-looking statements, the Company has relied on various assumptions, including, but not limited to: the Company’s ability to manage the effects of the COVID-19 pandemic; the accuracy of the Company’s financial projections; the Company obtaining positive results in its clinical and other trials; the Company obtaining necessary regulatory approvals; and general business, market and economic conditions.

Many factors could cause our actual results, level of activity, performance or achievements or future events or developments to differ materially from those expressed or implied by the forward-looking statements, including without limitation, a lack of revenue, insufficient funding, the impact of the COVID-19 pandemic, reliance upon key personnel, the uncertainty of the clinical development process, competition, and other factors set forth in the “Risk Factors” section of the Company’s Annual Information Form, Amended and Restated Prospectus Supplement, financial statements and Management Discussion and Analysis which can be found on SEDAR.com. All clinical development plans are subject to additional funding.

Readers should not place undue reliance on forward-looking statements made in this news release. Furthermore, unless otherwise stated, the forward-looking statements contained in this news release are made as of the date of this news release, and we have no intention and undertake no obligation to update or revise any forward-looking statements, whether as a result of new information, future events or otherwise, except as required by applicable law. The forward-looking statements contained in this news release are expressly qualified by this cautionary statement.