

## **CENTOGENE Extends Partnership With Takeda**

April 14, 2021

CAMBRIDGE, Mass., ROSTOCK, Germany and BERLIN, April 14, 2021 (GLOBE NEWSWIRE) -- Centogene N.V. (Nasdaq: CNTG), a commercial-stage company focused on rare diseases that transforms real-world clinical and genetic data into actionable information for patients, physicians, and pharmaceutical companies, announced today that it has extended its partnership with Takeda Pharmaceutical Company Limited (TSE: 4502/NYSE: TAK) to diagnose patients with certain genetic disorders.

As part of the agreement, which has been extended until March 2022, CENTOGENE will continue providing access to genetic testing to patients around the world.

Andrin Oswald, M.D., CEO of CENTOGENE, said, "We are pleased to be extending our partnership with Takeda, which will allow us to continue diagnosing and connecting rare disease patients globally. There is still a lot of work to be done, but with each step, we are closer to bringing patients the life-saving medical solutions they need."

In January 2015, CENTOGENE originally entered into an agreement with Shire Pharmaceuticals, which is now a subsidiary of Takeda Pharmaceutical Company Limited, to provide diagnostic testing capability to enhance early diagnosis of patients suffering from genetic rare diseases.

## **About CENTOGENE**

CENTOGENE engages in diagnosis and research around rare diseases transforming real-world clinical and genetic data into actionable information for patients, physicians, and pharmaceutical companies. Our goal is to bring rationality to treatment decisions and to accelerate the development of new orphan drugs by using our extensive rare disease knowledge, including epidemiological and clinical data, as well as innovative biomarkers. CENTOGENE has developed a global proprietary rare disease platform based on our real-world data repository with over 3.6 billion weighted data points from approximately 595,000 patients representing over 120 different countries as of September 30, 2020.

The Company's platform includes epidemiologic, phenotypic, and genetic data that reflects a global population, and also a biobank of these patients' blood samples. CENTOGENE believes this represents the only platform that comprehensively analyzes multi-level data to improve the understanding of rare hereditary diseases, which can aid in the identification of patients and improve our pharmaceutical partners' ability to bring orphan drugs to the market. As of September 30, 2020, the Company collaborated with over 40 pharmaceutical partners covering over 45 different rare diseases.

## **Important Notice and Disclaimer**

This press release contains statements that constitute "forward-looking statements" as that term is defined in the United States Private Securities Litigation Reform Act of 1995, including statements that express the Company's opinions, expectations, beliefs, plans, objectives, assumptions or projections regarding future events or future results, in contrast with statements that reflect historical facts. Examples include discussion of our strategies, financing plans, growth opportunities, and market growth. In some cases, you can identify such forward-looking statements by terminology such as "anticipate," "intend," "believe," "estimate," "plan," "seek," "project" or "expect," "may," "will," "would," "could," or "should," the negative of these terms or similar expressions. Forward-looking statements are based on management's current beliefs and assumptions and on information currently available to the Company. However, these forward-looking statements are not a guarantee of our performance, and you should not place undue reliance on such statements. Forward-looking statements are subject to many risks, uncertainties, and other variable circumstances, such as negative worldwide economic conditions and ongoing instability and volatility in the worldwide financial markets, the effects of the COVID-19 pandemic on our business and results of operations, possible changes in current and proposed legislation, regulations and govern-mental policies, pressures from increasing competition and consolidation in our industry, the expense and uncertainty of regulatory approval, including from the U.S. Food and Drug Administration, our reliance on third parties and collaboration partners, including our ability to manage growth and enter into new client relationships, our dependency on the rare disease industry, our ability to manage international expansion, our reliance on key personnel, our reliance on intellectual property protection, fluctuations of our operating results due to the effect of exchange rates, or other factors. Such risks and uncertainties may cause the statements to be inaccurate and readers are cautioned not to place undue reliance on such statements. Many of these risks are outside of the Company's control and could cause its actual results to differ materially from those it thought would occur. The forward-looking statements included in this press release are made only as of the date hereof. The Company does not undertake, and specifically declines, any obligation to update any such statements or to publicly announce the results of any revisions to any such statements to reflect future events or developments, except as required by

For further information, please refer to the Risk Factors section in our Annual Report for the year ended December 31, 2019, on Form 20-F filed with the SEC on April 23, 2020, Form 6-K containing our financial results for the three months ended March 31, 2020, furnished to the SEC on June 15, 2020, Form 6-K containing our financial results for the three and nine months ended September 30, 2020, furnished to the SEC on December 16, 2020, and other current reports and documents furnished to or filed with the U.S. Securities and Exchange Commission (SEC). You may get these documents by visiting EDGAR on the SEC website at <a href="https://www.sec.gov">www.sec.gov</a>.

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