## UNITED STATES SECURITIES AND EXCHANGE COMMISSION Washington, D.C. 20549

#### Form 6-K

# REPORT OF FOREIGN PRIVATE ISSUER PURSUANT TO RULE 13a-16 OR 15d-16 UNDER THE SECURITIES EXCHANGE ACT OF 1934

For the month of April 2015 Commission File Number 0-30314

# PORTAGE BIOTECH INC.

(Translation of registrant's name into English)

**47 Avenue Rd., Suite 200, Toronto, Ontario, Canada M5R 2G3** (Address of principal executive office)

| Indicate by check mark whether the registrant files or will file annual reports under cover Form 20-F or Form 40-F.  Form 20-FX Form 40-F                                                                                                      |
|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Indicate by check mark if the registrant is submitting the Form 6-K in paper as permitted by Regulation S-T Rule 101(b)(1):                                                                                                                    |
| Indicate by check mark if the registrant is submitting the Form 6-K in paper as permitted by Regulation S-T Rule 101(b)(7):                                                                                                                    |
| Indicate by check mark whether the registrant by furnishing the information contained in this Form is also thereby furnishing the information to the Commission pursuant to Rule 12g3-2(b) under the Securities Exchange Act of 1934.  Yes NoX |
| If "Yes" is marked, indicate below the file number assigned to the registrant in connection with Rule 12g3-2(b): 82                                                                                                                            |
|                                                                                                                                                                                                                                                |

### **SIGNATURES**

Pursuant to the requirements of the Securities Exchange Act of 1934, the registrant has duly caused this report to be signed on its behalf by the undersigned, thereunto duly authorized.

Dated: April 1, 2015

PORTAGE BIOTECH INC.

By: /s/ Kam Shah Kam Shah Chief Financial Officer

#### PORTAGE'S BIOHAVEN ANNOUNCES AGREEMENT WITH

#### **INVENTIV HEALTH**

Biohaven and inVentiv Health Enter into Agreement for Operational Execution of Biohaven's Lead Compound Phase 1 Clinical Trial.

**Toronto, Ontario, April 1, 2015** – Portage Biotech Inc. ("Portage") **(OTC Market: PTGEF, Canadian Securities Exchange: PBT.U),** is pleased to announce that Biohaven Pharmaceutical Holding Company Limited (Biohaven), which Portage holds 54% equity, has entered into an agreement with inVentiv Health for the operational execution of Biohaven's first clinical trial with their lead compound (BHV-0223).

Biohaven is engaged in the identification and development of clinical stage neuroscience compounds targeting the glutamatergic system. Clinical drug supply manufacturing has begun for BHV-0223, Biohaven's exciting novel formulation lead compound. Biohaven plans to begin a Phase 1 pharmacokinetic and biomarker study by 3Q2015 to confirm optimized drug exposure levels of its novel formulation. in Ventiv Health will oversee the study execution of this clinical trial.

"inVentiv Health is a global provider of best-in-class clinical development and comprehensive commercialization services. Biohaven is excited to have inVentiv assisting with strategic preparation and clinical trial execution of our development program for BHV-0223," commented Kimberly Gentile, Biohaven's Vice President of Clinical Operations.

BHV-0223 is a glutamate modulating agent being developed using Section 505(b)(2) of FDA guidelines. Section 505(b)(2) permits approval of new drug applications based, in part, upon prior findings of safety and/or effectiveness from a previously approved drug product.

#### About Biohaven

Biohaven is a privately-held biopharmaceutical company engaged in the identification and development of clinical stage neuroscience compounds targeting the glutamatergic system. Biohaven founders were among the first researchers at Yale University to discover the therapeutic potential of the NMDA antagonist ketamine and other glutamate modulating agents in the treatment of neuropsychiatric disorders. Biohaven has a worldwide license from Yale University to use intellectual property relating to the use of certain glutamate modulating agents in the treatment of neuropsychiatric disorders. The company's first drug candidate, BHV-0223, is a reformulated glutamate modulating agent being developed for treatment-resistant mood and anxiety disorders.

#### About Portage:

Portage is engaged in researching and developing pharmaceutical and biotech products through to clinical "proof of concept" with an initial focus on unmet clinical needs. Following proof of concept, Portage will look to sell or license the products to large pharmaceutical companies for further development and commercialization.

Portage is seeking discovery and co-development partners in areas such as certain inherited diseases, inflammatory and autoimmune disease, cancer, infectious disease, neurology and psychiatry developing novel targeted therapies, and even older marketed products that have been found to have novel patentable characteristics that bring new value to patients.

Portage seeks to work with a wide range of partners, in all phases of development through in-licensing or other types of alliances. The collaboration may include direct funding or investing human capital from our extensive pool of talented scientists and physicians. Specifically, Portage will invest sweat equity as well as, or instead of, capital. This internal pool of drug developers, financiers, scientists and physicians will provide unique value-add for our partners including but not limited to mitigating risks, clinical trial design, regulatory expertise and maximizing the rewards.

Apart from Biohaven, Portage also has fully owned subsidiary, Portage Pharmaceuticals Limited (PPL). PPL has successfully validated a new proprietary cell permeable peptide platform technology derived from human genes. This proprietary platform technology has been shown to efficiently deliver an active pharmacological agent or cargo into a cell without disrupting the cell membrane. The platform has favourable pharmaceutical properties simplifying formulation development for systemic and locally administered conjugates which will allow more rapid development of drug products. PPL has converted its previously filed provisional patent application for this delivery system to an international patent application that includes a variety of structures utilizing cargos that address important areas of medical need.

For further information, contact Dr. Greg Bailey, the Chairman at <u>gb@portagebiotech.com</u> or Kam Shah, Chief Financial Officer, at <u>(416) 929-1806</u> or <u>ks@portagebiotech.com</u> or visit our website at <u>www.portagebiotech.com</u>.

#### **Forward-Looking Statements**

This news release includes forward-looking statements within the meaning of the U.S. federal and Canadian securities laws. Any such statements reflect Portage's current views and assumptions about future events and financial performance. Portage cannot assure that future events or performance will occur. Important risks and factors that could cause actual results or events to differ materially from those indicated in our forward-looking statements.

Portage assumes no obligation and expressly disclaims any duty to update the information in this News Release.