Arbutus Announces New Data on AB-729 and AB-836 Programs with Presentation of Five Abstracts at the EASL International Liver Congress™ 2021 - All Selected for Best of ILC™

June 26, 2021

AB-729, dosed at 60 mg every 8 weeks, achieved a mean HBsAg decline of -1.87 log10 IU/mL at week 44, comparable to 60 mg dosed every 4 weeks.

**AB-729 resulted in HBsAg declines below 100 IU/ml in 75% of treated subjects, and increased HBV-specific immune responses in 3/5 evaluable subjects.**

**AB-729 resulted in decreases in HBV RNA and all HBsAg isoforms demonstrating broad target engagement.**

**AB-729 dosed for 48 weeks continued to demonstrate a favorable safety and tolerability profile.**

**AB-836 pre-clinical data suggest the potential for increased efficacy and an enhanced resistance profile relative to previous generation capsid inhibitors.**

Conference Call and Webcast Scheduled for 8:00 AM ET, Monday, June 28th, 2021

WARMINSTER, Pa., June 26, 2021 (GLOBE NEWSWIRE) -- Arbutus Biopharma Corporation (Nasdaq: ABUS), a clinical-stage biopharmaceutical company primarily focused on developing a cure for people with chronic hepatitis B virus (HBV) infection, as well as therapies to treat coronaviruses (including COVID-19), today announced the presentation of five abstracts at the European Association for the Study of the Liver (EASL) International Liver Congress™ (ILC). The abstracts included an oral late-breaker presentation (Presentation LBO-2764) by Professor Man-Fung Yuen, D.Sc., M.D., Ph.D., Deputy Head of Department, Chief of Division of Gastroenterology and Hepatology, Master of Lap Chee, University of Hong Kong, and lead investigator of AB-729’s Phase 1a/1b clinical trial, titled, “Repeat dosing of the GalNAc-siRNA AB-729 in subjects with chronic hepatitis B results in robust and sustained HBsAg suppression.”

As presented by Professor Yuen, AB-729 continues to demonstrate robust mean HBsAg reduction across all doses and dosing intervals with a favorable safety and tolerability profile, followed by a sustained plateau phase.

**Mean (range) change in HBsAg with repeat dosing of AB-729:**

<table>
<thead>
<tr>
<th>Visit</th>
<th>Cohort E AB-729 60 mg Q4W</th>
<th>Cohort F AB-729 60 mg Q8W</th>
<th>Cohort I AB-729 90 mg Q8W</th>
<th>p value between Cohorts</th>
</tr>
</thead>
<tbody>
<tr>
<td>Week 16</td>
<td>-1.44 (-0.71 to -1.95)</td>
<td>-1.39 (-1.61 to -1.08)</td>
<td>-1.63 (-0.89 to -2.44)</td>
<td>p ≥ 0.4</td>
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<tr>
<td>Week 24</td>
<td>-1.84 (-0.99 to -2.31)</td>
<td>-1.57 (-1.24 to -2.01)</td>
<td>-1.79 (-1.22 to -2.46)</td>
<td>p ≥ 0.2</td>
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<tr>
<td>Week 32</td>
<td>-1.84 (-0.94 to -2.36)</td>
<td>-1.68 (-1.37 to -2.15)</td>
<td>---</td>
<td>p = 0.5</td>
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<tr>
<td>Week 40</td>
<td>-1.84 (-0.88 to -2.47)</td>
<td>-1.78* (-1.40 to -2.14)</td>
<td>---</td>
<td>p = 0.7</td>
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<tr>
<td>Week 44</td>
<td>-1.81* (-0.93 to -2.43)</td>
<td>-1.87* (-1.32 to -2.34) [N=6]</td>
<td>---</td>
<td>p = 0.8</td>
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<tr>
<td>Week 48</td>
<td>-1.89* (-0.91 to -2.44)</td>
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† subjects switched to AB-729 60 mg Q12W after Week 20 dose

* Data updated since EASL ILC™ presentation

Professor Yuen stated, “These data continue to demonstrate that AB-729 delivers sustained and comparable HBsAg reduction across all doses and dose intervals. Importantly, AB-729 was generally safe and well tolerated. I believe these results support continued evaluation of AB-729 as a significant advancement in the future treatment of chronic HBV.”

Professor Yuen’s presentation can be accessed through the Investors section of Arbutus’ website under Events & Presentations at www.arbutusbio.com.

“The presented efficacy and safety data for AB-729, derived from up to one year of dosing, support our view that 60 mg every 8 weeks is an appropriate dose to move forward in our upcoming Phase 2a clinical trials,” stated Gaston Picchio, PhD, Chief Development Officer at Arbutus.

Dr. Picchio added, “In addition, while only based on 3/5 evaluable subjects, long-term dosing with AB-729 demonstrated increased HBV-specific immune responses, providing support for combination therapy including immunomodulatory agents.”
In addition, Arbutus announced the following three posters and oral presentation were made during the EASL International Liver Congress™.

**AB-729 ILC™ 2021 Poster Presentations:**

**Poster 2823**
- Inhibition of hepatitis B surface antigen in chronic hepatitis B subjects by RNA interference therapeutic AB-729 is accompanied by upregulation of HBV-specific T cell activation markers; presenting author, Dr. Emily Thi, Ph.D., Director, Immunobiology and Biomarkers Research, Arbutus Biopharma Corp.
  - Key Findings -- AB-729 induced reductions in HBsAg are associated with increased HBV-specific immune responses in 3/5 evaluable subjects. These increases in HBV-specific immune responses were accompanied by mild to moderate ALT elevations. We believe these to be the first results in a setting of small interfering RNA (siRNA) therapy which strengthen the hypothesis that long-term HBV antigen suppression can promote immune reawakening in HBV subjects.

**Poster 2822**
- Inhibition of hepatitis B surface antigen by RNA interference therapeutic AB-729 in chronic hepatitis B subjects correlates with suppression of all HBsAg isoforms and HBV RNA; presenting author, Dr. Emily Thi, Ph.D., Director, Immunobiology and Biomarkers Research, Arbutus Biopharma Corp.
  - Key Findings -- AB-729 mediated total HBsAg decline correlated with decreases in circulating HBV RNA species and all HBsAg isoforms. Early reduction in HBV RNA was observed in both “slow” and “fast” responders, demonstrating broad target engagement by AB-729 in all HBV subjects. A sub-set of subjects receiving AB-729 experienced a consistent decline of HBsAg immune complex levels.

**Poster 2829**
- A single dose of the GalNAc-siRNA, AB-729, results in prolonged reductions in HBsAg, HBcrAg, HBV DNA and HBV RNA in the absence of nucleos(t)ide analogue (NA) therapy in HBeAg negative subjects with chronic hepatitis B infection; presenting author, Dr. Edward Gane, M.D., Professor of Medicine at the University of Auckland, New Zealand and Chief Hepatologist, Transplant Physician and Deputy Director of the New Zealand Liver Transplant Unit at Auckland City Hospital.
  - Key Findings -- In HBV DNA+ subjects, a single dose of AB-729 90 mg in the absence of NA therapy achieved potent and durable anti-viral responses demonstrating broad target engagement.

**AB-836 ILC™ 2021 Oral Presentation:**

**Presentation OS-595**
- Preclinical antiviral profile of AB-836, a potent highly selective hepatitis B virus capsid inhibitor, presented by Dr. Nagraj Mani, Ph.D., Research Fellow, Arbutus Biopharma Corp.
  - Key Findings -- AB-836’s pre-clinical data suggest the potential for increased efficacy and an enhanced resistance profile relative to previous generation capsid inhibitors. AB-836 is currently in a Phase 1a/1b clinical trial.

These posters and presentations can be accessed through the Investors section under Events & Presentations of Arbutus’ website at [www.arbutusbio.com](http://www.arbutusbio.com).

William Collier, President and Chief Executive Officer of Arbutus, stated, “Given our focus and commitment to the development of a cure for people living with HBV, we are gratified that all of our abstracts were selected for presentation at the Best of ILC™. In the second half of 2021, we look forward to the initiation of two Phase 2a proof of concept AB-729 clinical trials and to reporting initial data from both healthy volunteers and HBV subjects in our AB-836 Phase 1a/1b clinical trial. With the extensive amount of AB-729 data presented by Arbutus to date, we are confident that it has the potential to be a cornerstone drug in future HBV combination regimens.”

**Conference Call and Webcast:**
Arbutus will hold a conference call and webcast on Monday, June 28, 2021, at 8:00 AM Eastern Time to summarize the data presented at EASL. You can access a live webcast of the call, which will include presentation slides, through the Investors section of Arbutus’ website at [www.arbutusbio.com](http://www.arbutusbio.com) or directly at [Live Webcast](http://www.arbutusbio.com). Alternatively, you can dial (866) 393-1607 or (914) 495-8556 and reference conference ID 6857498.

An archived webcast will be available on the Arbutus website after the event. Alternatively, you may access a replay of the conference call by calling (855) 859-2056 or (404) 537-3406, and reference conference ID 6857498.

**About AB-729**
AB-729 is an RNA interference (RNAi) therapeutic targeted to hepatocytes using Arbutus’ novel covalently conjugated N-acetylgalactosamine...
(GalNAC) delivery technology that enables subcutaneous delivery. AB-729 inhibits viral replication and reduces all HBV antigens, including hepatitis B surface antigen in preclinical models. Reducing hepatitis B surface antigen is thought to be a key prerequisite to enable reawakening of a patient’s immune system to respond to the virus. Based upon clinical data generated thus far in an ongoing single- and multi-dose Phase 1a/1b clinical trial, AB-729 has demonstrated positive safety and tolerability data and meaningful reductions in hepatitis B surface antigen.

**About AB-836**

AB-836 is an oral HBV capsid inhibitor. HBV core protein assembles into a capsid structure, which is required for viral replication. The current standard-of-care therapy for HBV, primarily nucleos(t)ide analogues that work by inhibiting the viral polymerase, significantly reduce virus replication, but not completely. Capsid inhibitors inhibit replication by preventing the assembly of functional viral capsids. They also have been shown to inhibit the uncoating step of the viral life cycle thus reducing the formation of new covalently closed circular DNA (cccDNA), the genetic reservoir which the virus uses to replicate itself.

**About HBV**

Hepatitis B is a potentially life-threatening liver infection caused by HBV. HBV can cause chronic infection which leads to a higher risk of death from cirrhosis and liver cancer. Chronic HBV infection represents a significant unmet medical need. The World Health Organization estimates that over 250 million people worldwide suffer from chronic HBV infection, while other estimates indicate that approximately 2 million people in the United States suffer from chronic HBV infection. Approximately 900,000 people die every year from complications related to chronic HBV infection despite the availability of effective vaccines and current treatment options.

**About Arbutus**

Arbutus Biopharma Corporation is a publicly traded (Nasdaq: ABUS) biopharmaceutical company primarily focused on discovering, developing and commercializing a cure for people with chronic hepatitis B virus (HBV) infection. The Company is advancing multiple product candidates with distinct mechanisms of action that it believes have the potential to provide a new curative regimen for chronic HBV infection. Arbutus has also initiated a drug discovery and development effort for treating coronaviruses (including COVID-19). For more information, visit [www.arbutusbio.com](http://www.arbutusbio.com).

**Forward-Looking Statements and Information**

This press release contains forward-looking statements within the meaning of the Section 27A of the Securities Act of 1933 and Section 21E of the Securities Exchange Act of 1934, and forward-looking information within the meaning of Canadian securities laws (collectively, “forward-looking statements”). Forward-looking statements in this press release include, but may not be limited to, statements about Arbutus’ future development plans for AB-729 and AB-836, including the initiation of two Phase 2a proof of concept AB-729 clinical trials and the reporting of initial data from both healthy volunteers and HBV subjects in the AB-836 Phase 1a/1b clinical trial in the second half of this year; and Arbutus’ expectations regarding the potential for its product candidates to provide a curative regimen for chronic HBV infection.

With respect to the forward-looking statements contained in this press release, Arbutus has made numerous assumptions regarding, among other things: the effectiveness and timeliness of preclinical studies and clinical trials, and the usefulness of the data; the timeliness of regulatory approvals; the continued demand for Arbutus’ assets; and the stability of economic and market conditions. While Arbutus considers these assumptions to be reasonable, these assumptions are inherently subject to significant business, economic, competitive, market and social uncertainties and contingencies, including uncertainties and contingencies related to the ongoing COVID-19 pandemic.

Additionally, there are known and unknown risk factors which could cause Arbutus’ actual results, performance or achievements to be materially different from any future results, performance or achievements expressed or implied by the forward-looking statements contained herein. Known risk factors include, among others: anticipated pre-clinical studies and clinical trials may be more costly or take longer to complete than anticipated, and may never be initiated or completed, or may not generate results that warrant future development of the tested drug candidate; Arbutus may elect to change its strategy regarding its product candidates and clinical development activities; Arbutus may not receive the necessary regulatory approvals for the clinical development of Arbutus’ products; economic and market conditions may worsen; market shifts may require a change in strategic focus; and the ongoing COVID-19 pandemic could significantly disrupt Arbutus’ clinical development programs.

A more complete discussion of the risks and uncertainties facing Arbutus appears in Arbutus’ Annual Report on Form 10-K, Arbutus’ Quarterly Reports on Form 10-Q and Arbutus’ continuous and periodic disclosure filings, which are available at www.sedar.com and at sec.gov. All forward-looking statements herein are qualified in their entirety by this cautionary statement, and Arbutus disclaims any obligation to revise or update any such forward-looking statements or to publicly announce the result of any revisions to any of the forward-looking statements contained herein to reflect future results, events or developments, except as required by law.

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