# Outlook Therapeutics® to Present at the BTIG Virtual Biotechnology Conference 2023

August 1, 2023

#### Live webcast fireside chat on Monday, August 7th at 1:00 PM ET

ISELIN, N.J., Aug. 01, 2023 (GLOBE NEWSWIRE) -- <u>Outlook Therapeutics</u>, Inc. (Nasdaq: OTLK), a biopharmaceutical company working to achieve FDA approval for the first ophthalmic formulation of bevacizumab for the treatment of wet AMD, today announced that <u>Russell Trenary</u>. <u>President and Chief Executive Officer</u> of Outlook Therapeutics will participate in a fireside chat at the BTIG Virtual Biotechnology Conference 2023 on Monday, August 7, 2023 at 1:00 PM ET.

In addition to the fireside chat, management will be available to participate in virtual one-on-one meetings with qualified members of the investor community who are registered to attend the conference.

A <u>live video webcast</u> of the fireside chat will be accessible on the <u>Events</u> page in the <u>Investors</u> section of the Company's website (<u>outlooktherapeutics.com</u>). The webcast replay will be archived for 90 days following the event.

### About Outlook Therapeutics, Inc.

Outlook Therapeutics is a biopharmaceutical company working to achieve FDA approval for the launch of ONS-5010/ LYTENAVA TM (bevacizumab-vikg) as the first FDA-approved ophthalmic formulation of bevacizumab for use in retinal indications, including wet AMD, DME and BRVO. The FDA accepted Outlook Therapeutics' BLA submission for ONS-5010 to treat wet AMD with a PDUFA goal date of August 29, 2023. The submission is supported by Outlook Therapeutics' wet AMD clinical program, which consists of three clinical trials: NORSE ONE, NORSE TWO, and NORSE THREE. If ONS-5010 ophthalmic bevacizumab is approved, Outlook Therapeutics expects to commercialize it as the first and only FDA-approved ophthalmic formulation of bevacizumab for use in treating retinal diseases in the United States, United Kingdom, Europe, Japan, and other markets. As part of the Company's multi-year commercial planning process, and in anticipation of potential FDA approval in August 2023, Outlook Therapeutics and AmerisourceBergen have entered into a strategic commercialization agreement to expand the Company's reach for connecting to retina specialists and their patients. AmerisourceBergen will provide third-party logistics (3PL) services and distribution, as well as pharmacovigilance services and other services in the United States. For more information, please visit <a href="https://www.outlooktherapeutics.com">www.outlooktherapeutics.com</a>.

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Source: Outlook Therapeutics, Inc.