

Galmed Announces Grant of New Patent for the Combination of Aramchol with Resmetirom (MGL-3196, REZDIFFRA) for the Treatment of NASH and Liver Fibrosis

TEL AVIV, Israel, March 15, 2024 /PRNewswire/ -- Galmed Pharmaceuticals Ltd. (Nasdaq: GLMD) ("Galmed" or the "Company"), a clinical-stage biopharmaceutical company for liver, metabolic and fibrotic diseases announced today the grant of a European patent related to the use of a combination of Aramchol and Resmetirom (MGL-3196, REZDIFFRA) for the treatment of NASH/MASH and liver fibrosis. The patent was granted in France, Germany, Italy, the Netherlands and the United Kingdom and the approval of the patent in the United States and other countries is pending. With this latest patent, Galmed is strengthening and extending the patent protection of its lead compound, Aramchol, until September 2039.

Previously, Galmed reported results from the Open-Label part of its Phase 3 NASH study, which demonstrated that treatment with Aramchol 300mg BID resulted in a high rate of subjects with fibrosis improvement. <https://galmedpharma.investorroom.com/2022-04-28-Galmed-reports-interim-results-from-the-Open-Label-part-of-the-ARMOR-study-with-Aramchol-showing-robust-fibrosis-improvement-across-multimodality-histological-assessment>

Galmed has long believed that the optimum treatment for NASH/MASH will be combination therapy. Aramchol has been shown to down-regulate the expression and activity of stearoyl-CoA desaturase-1 (SCD1) in hepatic stellate cells, resulting in a direct effect on fibrogenesis. Aramchol's unique mechanism of action differs from others in its competitive landscape, positioning it to work as a potent anti-fibrotic compound alongside effective treatments in both approved and pre-approval stages.

Allen Baharaff, CEO and President of Galmed Pharmaceuticals mentioned that "this new patent for the combination of Aramchol and Resmetirom reflects the same spirit of earlier patents which were granted to Galmed by the United States Patent and Trademark

Office (USPTO) for the treatment for hepatic fibrosis." Mr. Baharaff continued "It is clear today that NASH is a chronic condition with multiple liver pathologies. Among all pathologies,

excessive liver fat, high glycemic index and fibrosis are major treatment challenges. By combining Aramchol and Resmetirom, two distinct and selective compounds with complementary mechanisms, we believe this will provide a perfect treatment for NASH."

About Galmed Pharmaceuticals Ltd.

We are a clinical stage biopharmaceutical company focused on the development of Aramchol for liver and fibro-inflammatory diseases. We have focused almost exclusively on developing Aramchol for the treatment of NASH and are currently developing Aramchol for PSC and exploring the feasibility of developing Aramchol for other fibro-inflammatory indications outside of liver disease. We are also collaborating with the Hebrew University in the development of Amilo-5MER, a 5 amino acid synthetic peptide.

Forward-Looking Statements:

Forward-looking statements relate to anticipated or expected events, activities, trends or results as of the date they are made. Because forward-looking statements relate to matters that have not yet occurred, these statements are inherently subject to risks and uncertainties that could cause our actual results to differ materially from any future results expressed or implied by the forward-looking statements.

Forward-looking statements may include, but are not limited to, statements relating to our objectives plans and strategies, as well as statements, other than historical facts, that address activities, events or developments that we intend, expect, project, believe or anticipate will or may occur in the future. Many factors could cause our actual activities or results to differ materially from the activities and results anticipated in forward-looking statements, including, but not limited to, the timing and cost of any pre-clinical or clinical trial, for our product candidates; completion and receiving favorable results of any pre-clinical or clinical trial; regulatory action with respect to Aramchol or any other product candidate by the U.S. Food and Drug Administration, or the FDA, or the European Medicines Authority, or EMA, including but not limited to acceptance of an application for marketing authorization, review and approval of such application, and, if approved, the scope of the approved indication and labeling; the commercial launch and future sales of Aramchol and any future product candidates; our ability to comply with all applicable post-market regulatory requirements for Aramchol or any other product candidate in the countries in which we seek to market the product; our ability to achieve favorable pricing for Aramchol or any other product candidate; our expectations regarding the commercial market for non-alcoholic steato-hepatitis, or NASH, in patients or any other targeted indication; third-party payor reimbursement for Aramchol or any other product candidate; our estimates regarding anticipated capital requirements and our needs for additional financing; market adoption of Aramchol or any other product candidate by physicians and patients; the timing, cost or other aspects of the commercial launch of Aramchol or any other product candidate; our ability to obtain and maintain adequate protection of our intellectual property; the possibility that we may face third-party claims of intellectual property infringement; our ability to manufacture our product candidates in commercial quantities, at an adequate quality or at an acceptable cost; our ability to establish adequate sales, marketing and distribution channels; intense competition in our industry, with competitors having substantially greater financial, technological, research and development, regulatory and clinical, manufacturing, marketing and sales, distribution and personnel resources than we do; the development and approval of the use of Aramchol or any other product candidate for additional indications or in combination therapy; our ability to maintain the listing of our ordinary share on The Nasdaq Capital Market; and our expectations regarding licensing, acquisitions and

strategic operations. We believe these forward-looking statements are reasonable; however, these statements are only current predictions and are subject to known and unknown risks, uncertainties and other factors that may cause our or our industry's actual results, levels of activity, performance or achievements to be materially different from those anticipated by the forward-looking statements. We discuss many of these risks in our Annual Report on Form 20-F for the year ended December 31, 2022 filed with the SEC on March 29, 2023 in greater detail under the heading "Risk Factors." Given these uncertainties, you should not rely upon forward-looking statements as predictions of future events. All forward-looking statements attributable to us or persons acting on our behalf speak only as of the date hereof and are expressly qualified in their entirety by the cautionary statements included in this report. We undertake no obligations to update or revise forward-looking statements to reflect events or circumstances that arise after the date made or to reflect the occurrence of unanticipated events. In evaluating forward-looking statements, you should consider these risks and uncertainties.

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For further information: For further information: For further information: Guy Nehemya, Chief Operating Officer, Galmed Pharmaceuticals Ltd., investor.relations@galmedpharma.com, +972-3-693-8448

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