Madrigal Pharmaceuticals Exceeds Target Enrollment in Phase 3 MAESTRO NAFLD-1 Trial

-- Completion of enrollment in MAESTRO-NAFLD-1 will enable reporting of topline 52-week data by the end of next year as planned.

-- Madrigal intends to present selected data from ongoing open label arm by the end of 2020

-- Recent presentations at EASL Digital ILC of secondary analysis of Ph2 data demonstrated that reduction in MRI-PDFF can predict improvement in all components of histologic response in NASH, including steatosis, ballooning, inflammation and fibrosis.

CONSHOHOCKEN, Pa., September 3, 2020 -- Madrigal Pharmaceuticals, Inc. (NASDAQ:MDGL) announced today that it has already exceeded the originally targeted enrollment of 700 patients in its MAESTRO NAFLD-1 clinical trial of resmetirom in patients with NASH and fibrosis that is diagnosed using non-invasive assessments. Resmetirom is the first orally administered, small-molecule, liver-directed, truly β-selective thyroid hormone receptor (THR) agonist currently in Phase 3 development for the treatment of NASH patients with fibrosis stage 2-3 (ClinicalTrials.gov NCT03900429 and ClinicalTrials.gov/NCT04197479).

MAESTRO-NAFLD-1, was originally planned to enroll 700 patients with non-alcoholic fatty liver disease (NAFLD), presumed NASH, randomized 1:1:1 to receive resmetirom 80 mg once a day, 100 mg once a day, or placebo, and included an 100 mg resmetirom open label arm in up to 100 patients. MAESTRO-NAFLD-1 enrollment has already exceeded the enrollment targets in the three double-blinded arms and in the open label arm. Although new screening of patients for the double-blind arms has ended, eligible patients who have already screened for the study will continue to enroll over the next few weeks.

Dr. Stephen Harrison, M.D., Medical Director for Pinnacle Clinical Research, San Antonio, Texas, and Visiting Professor of Hepatology, Oxford University, and Principal Investigator of the MAESTRO studies commented, “MAESTRO-NAFLD-1 has exceeded expectations in terms of enrollment, even during the COVID pandemic, that attests to the high prevalence of NASH and the enthusiasm of patients and investigators to participate in a Phase 3 clinical trial in which the NASH diagnosis is made without a liver biopsy. The ultimate goals of the biomarker tests and liver imaging, which have expanded rapidly in the past few years, are to diagnose NASH with fibrosis non-invasively in order to identify patients with high risk fatty liver disease.”
“We are pleased to have achieved our target enrollment in the MAESTRO-NAFLD-1 trial,” stated Paul Friedman, M.D., Madrigal’s Chief Executive Officer. “We plan to complete enrollment in the double-blind arms of this study near the beginning of October to enable us to report topline 52-week data by the end of next year as planned. We intend to present data from the ongoing open label arm before the end of 2020.”

Becky Taub, M.D., Chief Medical Officer and President of Research & Development of Madrigal, stated, “We are encouraged by patient participation in our Phase 3 MAESTRO-NASH and MAESTRO-NAFLD-1 studies of resmetirom, a once daily oral medication. We recently began and will continue to enroll patients with compensated NASH cirrhosis into the open label arm of MAESTRO-NAFLD-1 to collect exploratory efficacy and safety data in this important population. We believe the data from additional enrolled NASH patients in the double-blind arms and patients with NASH cirrhosis will reinforce the safety and efficacy of resmetirom and provide an even more robust safety data base for our Phase 3 NASH program.”

Dr. Taub continued, “As we have recently reported, including in presentations by NASH experts over the past week at the Digital International Liver Congress™ 2020 (EASL), secondary analyses of data from our Phase 2 NASH study demonstrate that liver fat reduction at three months after starting treatment has clear predictive power for NASH resolution and fibrosis reduction on subsequent liver biopsy. Further, once daily oral 80 mg and 100 mg Phase 3 doses of resmetirom deliver at least 50% reduction in liver fat, and, based on secondary analyses of Phase 2 data, are associated with a statistically significant reduction in all components of NASH, including 64% NASH resolution (p<0.0001), of which >60% had fibrosis reduction. Finally, data from these analyses demonstrate that resmetirom robustly and statistically significantly (p<0.001) reduces markers of net collagen deposition in the liver, supporting the anti-fibrotic action of resmetirom. The related presentations by NASH experts at EASL are available here: EASL Presentations by NASH Experts_August 2020.”

About Resmetirom (MGL-3196)
Thyroid hormone, through activation of its β-receptor in hepatocytes, plays a central role in liver function impacting a range of health parameters from levels of serum cholesterol and triglycerides to the pathological buildup of fat in the liver. Thyroid hormone receptor (THR)-β action in the liver is key to proper function of the liver, including regulation of mitochondrial activity such as breakdown of liver fat and control of the level of normal, healthy mitochondria. Patients with NASH have reduced levels of thyroid hormone activity in the liver with resultant impaired hepatic function, in part due to the inflamed state of the liver that causes degradation of thyroid hormone.

To exploit the thyroid hormone receptor (THR)-β pathway for therapeutic purposes in cardiometabolic and liver diseases, it is important to avoid activity at the THR-α receptor, the predominant systemic receptor for thyroid hormone that is responsible for activity outside the liver including in heart and bone. The lack of selectivity of older thyromimetic compounds, chemically-related toxicities and undesirable distribution in the body led to safety concerns. Madrigal recognized that greater selectivity for thyroid hormone receptor (THR)-β and liver
targeting might overcome these challenges and deliver the full therapeutic potential of THR-β agonism. Resmetirom has been shown to be highly selective based on 1) THR-β receptor functional selectivity based on both in vitro and in vivo assays 2) specific uptake into the liver, its site of action, virtually avoiding any uptake into tissues outside the liver. In short and long term human and animal studies, resmetirom has been confirmed to be safe and devoid of activity at the THR-α receptor and without impact on bone or cardiac parameters. Resmetirom does not impact the thyroid axis hormones, including the central thyroid axis. Madrigal believes that resmetirom is the first orally administered, small-molecule, liver-directed, truly β-selective THR agonist.

**About the Phase 3 Registration Program for the Treatment of NASH (Non-alcoholic steatohepatitis)**

Analyses from the resmetirom Phase 2 NASH study demonstrate that the magnitude of liver fat reduction accurately predicts NASH resolution and liver fibrosis reduction and, specifically, that the resmetirom doses being used in Madrigal’s Phase 3 MAESTRO-NASH trial could achieve the level of fat reduction predictive of NASH resolution and fibrosis reduction [Madrigal COVID and ABSTRACT Press Release_20200414].

The Phase 3 MAESTRO-NASH trial is expected to enroll 900 patients with biopsy-proven NASH (fibrosis stage 2 or 3), randomized 1:1:1 to receive resmetirom 80 mg once a day, 100 mg once a day, or placebo. After 52 weeks of treatment a second biopsy is performed. The primary surrogate endpoint on biopsy will be NASH resolution, with at least a 2-point reduction in NAS (NASH Activity Score), and with no worsening of fibrosis. Two key secondary endpoints are liver fibrosis improvement of at least one stage, with no worsening of NASH, and lowering of LDL-cholesterol [ClinicalTrials.gov/NCT03900429].

A second 52-week Phase 3 multi-center, double-blind, randomized, placebo-controlled study of resmetirom, MAESTRO-NAFLD-1, was initiated in December 2019 in 700 patients with non-alcoholic fatty liver disease (NAFLD), presumed NASH, randomized 1:1:1 to receive resmetirom 80 mg once a day, 100 mg once a day, or placebo. MAESTRO-NAFLD-1 also includes a 100 mg resmetirom open label arm in up to 100 patients. Unlike MAESTRO-NASH, MAESTRO-NAFLD-1 is a non-biopsy study and represents a “real-life” NASH study. NASH or presumed NASH is documented using historical liver biopsy or non-invasive techniques including fibroscan and MRI-PDFF. Using non-invasive measures, MAESTRO-NAFLD-1 is designed to provide incremental safety information to support the NASH indication as well as provide additional data regarding clinically relevant key secondary efficacy endpoints to better characterize the potential clinical benefits of resmetirom on cardiovascular and liver related endpoints. These key secondary endpoints include LDL-cholesterol, apolipoprotein B and triglyceride (TG) lowering; reduction of liver fat as determined by magnetic resonance imaging, proton density fat fraction (MRI-PDFF); and reduction of PRO-C3, a NASH fibrosis biomarker. [ClinicalTrials.gov/NCT04197479]

Additional secondary and exploratory endpoints will be assessed including reduction in liver enzymes, fibroscan scores and other fibrosis and inflammatory biomarkers.

These and other data, including safety parameters, form the basis for potential subpart H
submission to FDA for accelerated approval for the treatment of NASH. The original 900 patients in the MAESTRO-NASH study will continue on therapy after the initial 52-week treatment period; up to another 1,100 patients are to be added using the same randomization plan and the study is expected to continue for up to 54 months to accrue and measure clinical events, most relevantly progression to cirrhosis.

**About Resmetirom’s Potential to Confer Cardiovascular Risk Reduction in NASH patients**
Additionally, resmetirom lowers multiple atherogenic lipids, including LDL cholesterol, apolipoprotein B, triglycerides, and lipoprotein (a), as demonstrated in Phase 2, a key differentiating factor compared with other NASH therapeutics. The magnitude of reduction of these lipids support a potential indication for treatment of hyperlipidemia in NASH patients and predicts a potential for benefit on cardiovascular (CV) events in NASH patients who die most frequently of CV, not liver disease.

Because of their diabetes, dyslipidemia, hypertension, obesity in concert with an inflamed, fatty liver, NASH patients, particularly those with advanced fibrosis, are at a substantially increased CV risk compared to the general population. Resmetirom’s ability to decrease liver fat, which is an independent risk factor for CV events, and resmetirom’s effect to reduce atherogenic lipids are being further evaluated in several key secondary endpoints in both MAESTRO Phase 3 clinical studies.

**About Madrigal Pharmaceuticals**
Madrigal Pharmaceuticals, Inc. (Nasdaq: MDGL) is a clinical-stage biopharmaceutical company pursuing novel therapeutics that target a specific thyroid hormone receptor pathway in the liver, which is a key regulatory mechanism common to a spectrum of cardio-metabolic and fatty liver diseases with high unmet medical need. Madrigal’s lead candidate, resmetirom, is a first-in-class, orally administered, small-molecule, liver-directed, thyroid hormone receptor (THR)-β selective agonist that is currently in two Phase 3 clinical studies, MAESTRO-NASH and MAESTRO-NAFLD-1, designed to demonstrate multiple benefits across a broad spectrum of NASH (non-alcoholic steatohepatitis) and NAFLD (non-alcoholic fatty liver disease) patients. For more information, visit [www.madrigalpharma.com](http://www.madrigalpharma.com).

**Forward-Looking Statements**
This communication contains “forward-looking statements” made pursuant to the safe harbor provisions of the Private Securities Litigation Reform Act of 1995, that are based on our beliefs and assumptions and on information currently available to us, but are subject to factors beyond our control. Forward-looking statements include but are not limited to statements or references concerning: our clinical trials; research and development activities; the timing and results associated with the future development of our lead product candidate, MGL-3196 (resmetirom); our primary and secondary study endpoints for resmetirom and the potential for achieving such endpoints and projections; optimal dosing levels for resmetirom; projections regarding potential future NASH resolution, safety, fibrosis treatment, cardiovascular effects, lipid treatment or biomarker effects with resmetirom; the predictive power of liver fat reduction on NASH resolution with fibrosis reduction or improvement; the achievement of enrollment objectives
concerning patient number, safety database and/or timing for our studies; potential NASH or
NAFLD patient risk profile benefits with resmetirom; and our possible or assumed future results
of operations and expenses, business strategies and plans, capital needs and financing plans,
trends, market sizing, competitive position, industry environment and potential growth
opportunities, among other things. Forward-looking statements: reflect management’s current
knowledge, assumptions, judgment and expectations regarding future performance or events;
include all statements that are not historical facts; and can be identified by terms such as
“anticipates,” “be,” “believes,” “continue,” “could,” “demonstrates,” “design,” “estimates,”
“expects,” “forecasts,” “future,” “goal,” “hopeful,” “intends,” “may,” “might,” “plans,”
“potential,” “predicts,” “predictive,” “projects,” “seeks,” “should,” “will,” “would” or similar
expressions and the negatives of those terms. Although management presently believes that
the expectations reflected in such forward-looking statements are reasonable, it can give no
assurance that such expectations will prove to be correct and you should be aware that actual
results could differ materially from those contained in the forward-looking statements.

Forward-looking statements are subject to a number of risks and uncertainties including, but
not limited to: our clinical development of resmetirom; enrollment uncertainties, generally and
in relation to COVID-19 shelter-in-place and social distancing measures and individual
precautionary measures that may be implemented or continued for an uncertain period of time;
outcomes or trends from competitive studies; the risks of achieving potential benefits in studies
that includes substantially more patients than our prior studies; the timing and outcomes of
clinical studies of resmetirom; and the uncertainties inherent in clinical testing. Undue reliance
should not be placed on forward-looking statements, which speak only as of the date they are
made. Madrigal undertakes no obligation to update any forward-looking statements to reflect
new information, events or circumstances after the date they are made, or to reflect the
occurrence of unanticipated events. Please refer to Madrigal’s filings with the U.S. Securities and
Exchange Commission for more detailed information regarding these risks and uncertainties and
other factors that may cause actual results to differ materially from those expressed or implied.
We specifically discuss these risks and uncertainties in greater detail in the section entitled "Risk
Factors" in our Annual Report on Form 10-K for the year ended December 31, 2019 and our
Quarterly Report on Form 10-Q for the period ended June 30, 2020, as well as in our other filings
with the SEC.

**Investor Contact:**
Marc Schneebaum, Madrigal Pharmaceuticals, Inc. IR@madrigalpharma.com

**Media Contact:**
Mike Beyer, Sam Brown Inc. mikebeyer@sambrown.com 312 961 2502