## Relmada Therapeutics Files Amended Complaint Against Laidlaw and Its Principals, Matthew Eitner and James Ahern

NEW YORK, Sept. 14, 2016 /PRNewswire/ -- Relmada Therapeutics, Inc. (OTCQB: RLMD), a clinical-stage company developing novel therapies for the treatment of central nervous system (CNS) diseases, today announced that it has filed a motion to amend its amended complaint against Laidlaw & Company (UK) Ltd. ("Laidlaw") in the U.S. District Court for the District of Nevada. The motion to file a second amended complaint is part of the lawsuit that the Company had previously filed in the Nevada District Court. The proposed second amended complaint, dated September 6, 2016 (the "Proposed Second Amended Complaint") alleges additional claims against the Defendants, including defamation/business disparagement, defamation per se, tortious interference with prospective economic advantage, violations of sections 1962(c) and 1962(d) of the Racketeer Influenced and Corrupt Organizations Act, and violation of Section 10(b) of the Securities Exchange Act of 1934 and Rule 10b-5 thereunder. Relmada believes that it has incurred substantial damages from Defendants' actions, which are addressed in the Proposed Second Amended Complaint.

Relmada noted that the Nevada court previously issued a temporary restraining order and associated injunction against Laidlaw and its principals, Matthew Eitner and James Ahern, as a result of their dissemination of false and misleading proxy materials. Relmada's Board believes Laidlaw must compensate the Company for the damages it has suffered as a result of Laidlaw's actions, and that Laidlaw must be prevented from harming the Company in the future.

A copy of the Proposed Second AmendedComplaints is attached as Exhibit 99.1 to the Company's recent Form 8-K.

## About Relmada Therapeutics, Inc.

Relmada Therapeutics is a clinical-stage, publicly traded biotechnology company developing novel versions of proven drug products together with new chemical entities that potentially address areas of high unmet medical need in the treatment of central nervous system (CNS) diseases. The Company has a diversified portfolio of four products at various stages of development, including d-Methadone (dextromethadone, REL-1017), an N-methyl-D-aspartate (NMDA) receptor antagonist for depression and neuropathic pain; LevoCap ER (REL-1015), an abuse resistant, sustained release dosage form of the opioid analgesic levorphanol; oral buprenorphine (BuTab, REL-1028), an oral dosage form of the opioid analgesic buprenorphine; and topical mepivacaine (MepiGel, REL-1021), an orphan drug designated topical formulation of the local anesthetic mepivacaine. The Company's product development efforts are guided by the internationally recognized scientific expertise of its

research team. The Company's approach is expected to reduce clinical development risks and costs while potentially delivering valuable products to address areas of high unmet medical needs. For more information, please visit Relmada's website at: <a href="https://www.relmada.com">www.relmada.com</a>.

## **Forward-Looking Statements**

The Private Securities Litigation Reform Act of 1995 provides a safe harbor for forwardlooking statements made by us or on our behalf. We may from time to time make written or oral statements in this letter, the proxy statements filed with the SEC communications to stockholders and press releases which constitute "forward-looking statements" within the meaning of Section 27A of the Securities Act of 1933 and Section 21E of the Securities Exchange Act of 1934. These forward-looking statements are based upon management's current expectations, estimates, assumptions and beliefs concerning future events and conditions and may discuss, among other things, anticipated future performance, expected product development, product potential, future business plans and costs. Any statement that is not historical in nature is a forward-looking statement and may be identified by the use of words and phrases such as "expects," "anticipates," "believes," "will," "will likely result," "will continue," "plans to" and similar expressions. No forward-looking statement can be guaranteed and actual results may differ materially from those projected. Relmada undertakes no obligation to publicly update any forward-looking statement, whether as a result of new information, future events, or otherwise. Readers are cautioned that it is not possible to predict or identify all of the risks, uncertainties and other factors that may affect future results and that the risks described herein should not be considered to be a complete list.

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SOURCE Relmada Therapeutics, Inc.