

Sequana Medical announces results of Annual and Extraordinary General Meetings of Shareholders

Ghent, BELGIUM – 27 May 2021 – Sequana Medical NV (Euronext Brussels: SEQUA) (the "Company" or "Sequana Medical"), an innovator in the treatment of diuretic-resistant fluid overload in liver disease, malignant ascites and heart failure, today announces that all proposed resolutions submitted to the Annual and Extraordinary General Meeting of Shareholders were approved at the meetings held today at 09:00 am CEST.

The items on the agendas of the shareholders' meetings included the approval of a number of resolutions relating to the financial year ended on 31 December 2020, as well as the re-appointment of directors and the statutory auditor, the approval of a number of change of control clauses, the issuance of a new share option plan, and the renewal of the authorization to the Board of Directors to increase the share capital within the framework of the authorized capital.

The minutes of the shareholders' meetings, as well as the revised versions of the Company's articles of association, can be accessed on the [Company's website](#).

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About Sequana Medical

Sequana Medical is a commercial stage medical device company utilizing its proprietary **alfapump**[®] and **DSR**[®] (Direct Sodium Removal) technologies to develop innovative treatments for fluid overload in liver disease, malignant ascites and heart failure where diuretics are no longer effective. Fluid overload is a frequent complication of many large diseases including advanced liver disease driven by NASH (non-alcoholic steatohepatitis)-related cirrhosis and heart failure, with diuretic resistance being widespread. The U.S. market for the **alfapump** resulting from NASH-related cirrhosis is forecast to exceed €3 billion annually within the next 10-20 years. The heart failure market for **DSR** and the **alfapump DSR**[®] is estimated to be over €5 billion

annually in the U.S. and EU5 by 2026.

The **alfapump** is a unique, fully implanted wireless device that automatically pumps fluid from the abdominal cavity into the bladder, where it is naturally eliminated through urination. DSR is Sequana Medical's proprietary approach to managing sodium and fluid overload through use of a sodium-free infusate administered into the abdominal cavity.

In the U.S., the Company's key growth market, the **alfapump** has been granted breakthrough device designation by the FDA for recurrent or refractory ascites due to liver cirrhosis. Interim data from the ongoing North American pivotal study (POSEIDON) showed positive outcomes against all primary endpoints of the study. This study is intended to support a future marketing application of the **alfapump** in the U.S. and Canada. In Europe, the **alfapump** is CE-marked for the management of refractory ascites due to liver cirrhosis and malignant ascites and is included in key clinical practice guidelines. Over 850 **alfapump** systems have been implanted to date.

Sequana Medical has combined its proven **alfapump** and proprietary DSR therapy, and is developing the **alfapump** DSR, a breakthrough approach to fluid overload due to heart failure. RED DESERT, the repeated dose **alfapump** DSR study in diuretic-resistant heart failure patients has demonstrated that repeated DSR therapy is able to both manage the fluid and sodium balance of these patients as well as restore their diuretic response and improve their cardio-renal status.

Sequana Medical is headquartered in Ghent, Belgium. For further information, please visit www.sequanamedical.com.

Important Regulatory Disclaimers

*The **alfapump**[®] system is not currently approved in the United States or Canada. In the United States and Canada, the **alfapump**[®] system is currently under clinical investigation (POSEIDON Study) and is being studied in adult patients with refractory or recurrent ascites due to cirrhosis. For more information regarding the POSEIDON clinical study see www.poseidonstudy.com. The DSR[®] therapy is still in development and it should be noted that any statements regarding safety and efficacy arise from ongoing pre-clinical and clinical investigations which have yet to be completed. The DSR[®] therapy is not currently approved for clinical research in the United States or Canada. There is no link between the DSR[®] therapy and ongoing investigations with the **alfapump**[®] system in Europe, the United States or Canada.*

Note: **alfapump**[®] is a registered trademark. DSR[®] and **alfapump** DSR[®] are registered trademarks in the Benelux.