

CARMAT announces its placement in receivership procedure

Resumption of CARMAT shares trading starting July 2, 2025, at stock market opening

Paris, July 1, 2025 – 4 pm CEST

CARMAT (FR0010907956, ALCAR), designer and developer of the world's most advanced total artificial heart, aiming to provide a therapeutic alternative for people suffering from advanced biventricular heart failure (the "**Company**" or "**CARMAT**"), today announces its placement in receivership procedure¹ as well as the resumption of CARMAT shares trading, starting July 2, 2025, at stock market opening.

Opening of receivership procedure

On June 30, 2025, CARMAT had announced in a press release filing for insolvency and requesting to be placed in receivership to the Versailles Economic Affairs Court (the "Court").

Following a hearing held on July 1, 2025, the Court has effectively decided to place CARMAT in receivership.

CARMAT's operations carry on during the observation period, in accordance with legal provisions. During this period, CARMAT will assess all options to ensure the continuation of its business activities, notably a disposal plan².

The Company also confirms that it will endeavor to provide continuous support to patients who currently benefit from its Aeson® artificial heart.

Press releases will be issued regularly as the Company's situation evolves and the proceedings progress.

Resumption of CARMAT shares trading (ISIN code: FR0010907956, Ticker: ALCAR)

CARMAT has asked Euronext to resume the trading of its shares starting July 2, 2025, at stock market opening.

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About CARMAT

CARMAT is a French MedTech that designs, manufactures and markets the Aeson[®] artificial heart. The Company's ambition is to make Aeson[®] the first alternative to a heart transplant, and thus provide a therapeutic solution to people suffering from end-stage biventricular heart failure, who are facing a well-known shortfall in available human grafts. The world's first physiological artificial heart that is highly hemocompatible, pulsatile and self-regulated, Aeson[®] could save, every year, the lives of thousands of patients waiting for a heart transplant. The device offers patients quality of life and mobility thanks to its ergonomic and portable external power supply system that is continuously connected to the implanted prosthesis. Aeson[®] is commercially available as a bridge to transplant in the European Union and other countries that recognize CE marking. Aeson[®] is also currently being assessed within the framework of an Early Feasibility Study (EFS) in the United States. Founded in 2008, CARMAT is based in the Paris region, with its head offices located in Vélizy-Villacoublay and its production site in Bois-d'Arcy. The Company can rely on the talent and

¹ « Redressement judiciaire »

² « Plan de cession »



expertise of a multidisciplinary team of circa 200 highly specialized people. CARMAT is listed on the Euronext Growth market in Paris (Ticker: ALCAR / ISIN code: FR0010907956).

For more information, please go to <u>www.carmatsa.com</u> and follow us on <u>LinkedIn</u>.

CARMAT Stéphane Piat Chief Executive Officer

Pascale d'Arbonneau Deputy Chief Executive Officer & Chief Financial Officer Tel.: +33 1 39 45 64 50 <u>contact@carmatsas.com</u> NewCap Press Relations

Nicolas Merigeau Arthur Rouillé Tel.: +33 1 44 71 94 98 carmat@newcap.eu



Name: CARMAT ISIN code: FR0010907956 Ticker: ALCAR

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NewCap Financial Communication & Investor Relations

Dusan Oresansky Jérémy Digel Tel.: +33 1 44 71 94 92 carmat@newcap.eu

Disclaimer

This press release and the information it contains do not constitute an offer to sell or subscribe, nor a solicitation of an offer to buy or subscribe, for CARMAT shares in any country.

This press release may contain forward-looking statements regarding the Company's objectives and outlook. These forward-looking statements are based on the current estimates and anticipations of the Company's management and are subject to risk factors and uncertainties, including those described in its Universal Registration Document filed with the French Financial Markets Authority (Autorité des marchés financiers) (the "AMF") under number D.25-0345 (the "**2024 Universal Registration Document**"), available free of charge on the websites of CARMAT (www.carmatsa.com/en/) and the AMF (www.amf-france.org).

Readers' attention is particularly drawn to the fact that the Company is currently placed in receivership (opened on July 1, 2025) and is facing a very high risk of default, including in the very short term. The Company is also exposed to other risks and uncertainties, such as its ability to implement its strategy, the pace of development of its production and sales, the progress and results of ongoing or planned clinical trials, technological developments, the competitive landscape, regulatory changes, industrial risks, and all risks related to the management of the Company's growth. Forward-looking statements mentioned in this press release may not be achieved due to these factors or other unknown risks and uncertainties, or risks that the Company does not currently consider to be material or specific.

Aeson® is an active implantable medical device commercially available in the European Union and other countries recognising the CE mark. The Aeson® total artificial heart is intended to replace the ventricles of the native heart and is indicated as a bridge to transplant in patients with end-stage biventricular heart failure (Intermacs classes 1-4) who cannot benefit from maximal medical therapy or a left ventricular assist device (LVAD) and who are likely to benefit from a heart transplant within 180 days of implantation. The decision to implant and the surgical procedure must be carried out by healthcare professionals trained by the manufacturer. The documentation (clinician's manual, patient's manual and alarm booklet) must be read carefully to learn about the characteristics of Aeson® and the information required for patient selection and proper use (contraindications, precautions, side effects) of Aeson®. In the United States, Aeson® is currently only available as part of a feasibility clinical trial approved by the Food & Drug Administration (FDA).