Scenable Access

We, **Enable Access Ltd**., as manufacturer, declare under our sole responsibility that the products listed below meet the provisions of **Regulation (EU) 2017/745**. The listed products are **Class I medical devices** and conformity with the requirements of the Regulation have been assessed following the procedure outlined in **Article 52 (section 7)** of the Regulation. The products meet the General Safety & Performance Requirements listed in Annex I of the Regulation and, in addition, meet the requirements of the following standard:

•BS6109: Part 2:1989, Appendix A.3

Product / code:

Doorline - Bridge: DB4, DB6, DB8, DB10 Butterfly Ramp: DVBM, DVBL Multi: DM5, DM7 Neatedge: DN2, DN3, DN4, DN5, DN6, DN15-9, DN25-9, DN35-9, DN50-9, DN75-9, DN100-9, DN125-9 Neatslope: DNS6, DNS6A, DNS10, DNS10A, DNS16, DNS16A Variwedge: DVWS, DVWM, DVWL, DVWXL Threshold Wedge: DTW1.5, DTW2, DTW3, DTW4, DTW5, DTW6

Technical documentation for the above products is kept by the manufacturer and can be made available by the Authorised Representative in the EU: Enable Access Global Ltd., 27 Phibsboro Place, Dublin 7, D07 V20, Republic of Ireland.

Signed:

Rhys Hibbert – Compliance Officer

Enable Access Ltd. Leighton Buzzard 11th April 2025