

CERTIFICATE OF REGISTRATION

Fiscal Year 2022 This is to certify that KENMAK HASTANE MALZEMELERI VE ELEKTROSTATIK BOYA SAN. TIC. A.S. SARNIC YOLU UZERI NO:23 GAZIEMIR IZMIR, TR 35410

Has completed the annual registration with the U.S. Food and Drug Administration as required by Title 21, chapter I, subchapter H, part 807 of the United States Code of Federal Regulation.

Establishment Registration : 3016672059

Product Code : FNL

Regulation Number : 880.5100

Device Classification : 2

Device : AC-POWERED ADJUSTABLE HOSPITAL BED

U.S. Agent : HEALTHCARE INTERNATIONAL PARTNERS, LLC

2450 SW 145 AVENUE, SUITE 101

MIRAMAR, FL 33027

This certificate does not make representation or warranties to any person or entity other than the named certificate holder; it is issued for record-keeping purposes only. This certificate does not denote endorsement or approval of the certificate holder's facility or product by the U.S. Food and Drug Administration. Healthcare International Partners, LLC. Assumes no liability to any person or entity in connection with foregoing.

According to 21 CFR 807. 39 the assignment of a registration number or registration of a device facility does not in any way denote approval of the establishment or its products. Any representation that creates an impression of official approval because of registration or possession of a registration number is misleading and constitutes misbranding.

The U.S. Food and Drug Administration does not issue a certificate of registration, nor does the U.S. Food and Drug Administration recognize a certificate of registration. Healthcare International Partners, LLC. is not affiliated with the U.S. Food and Drug Administration.

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Dated: 01/07/2021 regulatory@hip-llc.com

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