

## EC-declaration of conformity

As manufacturer confirms

**Permobil AB**

**Box 120**

**861 23 Timrå, Sweden**

that below specified product is in conformity with the

**Directive 93/42/EEC**

**Medical Devices**

**implemented in Swedish law 1993:584**

<b>General description</b>	Electrically powered wheelchair ISO code 12 23 06 under ISO 9999 and its accessories. All the devices are classified as Class 1 devices. The trade name of the wheelchair is: <b>M400 Corpus HD</b> All models are manufactured by Permobil.
<b>Product class:</b>	Class I (according to rule 12)
<b>Conformity procedure used</b>	Annex VII
<b>Design drawings</b>	Required design drawings, circuit diagrams and other manufacture documentation is filed in the Master Device File for each device.
<b>Descriptions</b>	Descriptions, explanations and operation descriptions will be seen from the manufacture documents, where appropriate.
<b>Risk analysis</b>	Where appropriate, the result of the risk analysis is declared in the Device Master File. The risk analyses are, where applicable, in compliance with EN ISO 14971
<b>EMC</b>	All devices confirm to the requirements specified in EN 12184.
<b>Standards</b>	All devices confirm to the appropriate parts of EN 12184.
<b>User manual</b>	User manuals and labeling are defined in the Master Device File. User manuals are for some devices made as assembly instructions. Devices that can be used safely without user manual/assembly instruction is delivered without such instructions.
<b>Corrective action</b>	All reported malfunctions and incidents/accidents are filed and reviewed and where appropriate reported to the competent authority.
<b>Drawn up</b>	2013-04-12
<b>Updated</b>	2018-06-07



Rikard Rönn

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