EONIKO KENTPO A三IO＾OTHटH THE ПOIOTHTA乏 \＆TEXNONOTIA乏 ITHN YГEIA A．E．

NATIONAL EVALUATION CENTER
OF QUALITY \＆TECHNOLOGY
IN HEALTH S．A

## ПІІТОПОІНТIKO EK／EC CERTIFICATE $\triangle I A \Sigma Ф A \wedge I \Sigma H$ ПOIOTHTA乏 ПAPAГתГH乏I PRODUCTION QUALITY ASSURANCE








We hereby certify that the under mentioned manufacturer has established and maintains a quality assurance system according to the requirements of Directive 93／42／EEC，Annex V and its transposition in Greek legislation，
for the manufacture and final inspection of the products mentioned in this certificate．
The certificate is subject to terms and conditions overleaf．
Any significant changes in design or manufacture may render this certificate invalid．
Apı日нós Пıбтотоıптıкои́／Certificate Number：301011012CA

The present is issued to replace certificate Nr 301011012.
Kataøкєuaఠtís：MONADA BIOMHXANIK $\Omega$ N IATPIK $\Omega$ N AEPI $\Omega$ N KPHTH乏 －MOBIAK A．E．，＂MOBIAK A．E．＂．

Manufacturer：MOBIAK S．A．
EүKatáotaøף：KAOIANA AKP
Facility：KATHIANA AKROTIRIOU，CHANIA CRETE GREECE．
Проїо́vта：ОПЛ乏 ANAФEPONTAI $\Sigma T O ~ П А Р А Р Т Н M A . ~$
Products：AS LISTED IN ANNEX．
Kainүорıттоі́ŋбף Проїövтшv：
Devices Classification：

First issue date：
1： $\mathrm{Il}, 2$ ： $\mathrm{Il} a, 3: \mathrm{Ila}, 4: \mathrm{Ilb}, 5: \mathrm{Ib}$

03／11／2020

Current issue date：
22／09／2021

Valid until：
24／05／2024

Audit report：
200091012

 PIKROU－MORAIIAKI ELEF THERIA，President \＆Managing Director

EONIKO KENTPO A三IO＾OTH THE ПOIOTHTA乏 \＆TEXNO＾OГIA乏 ITHN YГEIA A．E．
NATIONAL EVALUATION CENTER OF QUALITY \＆TECHNOLOGY IN HEALTH S．A．

ПАРАРТНМА ТОҮ ҮП．АРІӨМ．301011012СА ПI乏ТОПОІНТІКОY． ANNEX No．301011012CA CERTIFICATE．

| ПPOİONTA | TYПОІ／EMПOPIKE乏 ONOMA乏IE乏 |
| :---: | :---: |
| 1．โYMחYKN （OXYGEN CONCENTRATORS） | FORCE，THORAX 5，IRENE． |
| 2．NEФEへOПOIHTE乏 （NEBULIZERS） | GEM |
|  （CONTINUOUS AND AUTOMATIC POSITIVE AIRWAY PRESSURE UNITS） | MORFEUS，MORFEUS SOFT，MORFEUS AUTO／ MORFEUS AUTO II． |
| 4．PYOMIइTE POH乏 IATPIK $\Omega$ DIA $\Omega \Omega N$ OEYFONOY （OXYGEN PRESSURE REGULATORS） | HERCULES，APHRODITE． |
| 5．ПААМIKO OЕYMETPO $\triangle A K T Y \wedge O Y ~$ （PULSE OXIMETER） | MY SPO2 |

## OPOI \＆ПPOŸПOOE



For class I sterile products，the certificate covers only the aspects of manufacture concerned with securing and maintaining sterile conditions．
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For Class I devices with a measuring function the certificate covers only the aspects of manufacture concerned with the conformity of the products with metrological requirements．


For class III products an additional Type Examination certificate is required according to the requirements of $93 / 42 / E E C$ ， Annex III．
 The certificate is valid only for the products and the facilities mentioned

 Periodical surveillance as referred in $93 / 42 / E E C$ will be held in order to verify that the manufacturer maintains and applies the quality system．

CE 0653 бта калитто́ $\mu \varepsilon v a$ тоїо́vта．
When meeting with the terms and conditions above，the manufaffurer may draw up an EC declaration of conformity and legally affix the CE 0653 mark．
 PIKROU－MORAITAKI ELEF THERIA，President \＆Managing Director

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    National Evaluation Center of Quality \＆Technology in Health S．A．（EKAPTY）is a Notified body according to Council Directive 93／42／EEC concerning medical devices，with identification number 0653.

