



## **EU DECLARATION OF CONFORMITY**

Document number: ALDATK329101220708

Following the provisions of the medical device's regulation 2017/745(EU MDR).

We:

### **Legal Manufacturer**

CV. Anggrek-Liar  
Jalan Jomblang Barat 666 RT 09 RW 03  
Kelurahan Candi, Kecamatan Candisari  
Semarang 50257  
Central Java  
REPUBLIC OF INDONESIA

### **EU Authorized Representative**

Kaio-Dia Europe Sasu  
12, rue Bertrand  
31480 Lagraulet  
Saint Nicolas, France

Single Registration Number (SRN)  
ID-MF-000037819

Single Registration Number (SRN)  
FR-AR-000037267

### **Manufacturing Sites**

Anggrek-Liar / Kaio-Care  
Jalan Tumpang IV No. 38  
Kelurahan Bendan Ngisor  
Kecamatan Gajahmungkur  
Semarang 50233  
Central Java  
REPUBLIC OF INDONESIA

MAF Holding BV  
Laan van Leeuwesteyn 62  
2271 HL Voorburg, Nederland

Single Registration Number (SRN)  
NL-AR-000044388

Declare under our sole responsibility for the universal device fixation support:

**Kaio-Dia® Dia-T Kids©**  
Non-Invasive Top

**Basic UDI-DI:**  
8994228UW-DTKDY

**Identification number:**  
See Appendix 1

**SIGNATURE:**

Date: 24 January 2025

**Date of Issue :**  
**Place of Issue :** Semarang, Central Java, Indonesia  
**Name :** Anik Susilawatiningsih  
**Function :** Regulatory Affairs Manager

This Declaration of Conformity relating to Technical Documentation DOC329DTK82325 supersedes the previous Declaration signed 01-December-2021.



### Intended Purpose:

Kaio-Dia® Dia-T Kids®, a top wear are independently developed as universal after market accessories used to provide additional fixation support for a wide variety of automated insulin and medication on body delivery systems.

(see attachment **Appendix 1 – List of Supported Product version #1**)

Dia-T Kids® are available in pediatric sizes are non-sterile and may be reused (intended for single patient use only).

The devices are not designed or sold for use other than indicated.

### EMDN code and description:

**Z12040180** - GENERAL MEDICINE INSTRUMENTS FOR DIAGNOSIS AND MONITORING - HARDWARE

**Class: I**

**Classification rule (Annex VIII): 1**

to which this declaration relates is in conformity with the requirements of the medical devices regulation 2017/745 (EU MDR).

This conformity is based on the following elements:

- Technical Documentation reference: DOC329DTK82325, of the product to which this declaration relates.

As a Class I medical device, this product is self-certified in accordance with Regulation (EU) 2017/745. The conformity assessment has been carried out under the sole responsibility of the manufacturer, without the involvement of a Notified Body.

The undersigned, acting on behalf of CV. Anggrek-Liar, assumes full responsibility for ensuring that the product complies with all applicable European laws and regulations.

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Semarang, Central Java, Indonesia

Anik Susilawatiningsih

Regulatory Affairs Manager

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## Appendix 1 – List of Supported Product

Date: 24-01-2025 Version number #1

| Producer  | Product Name       |  |
|-----------|--------------------|--|
| Medtronic | Minimed™ 630g      |  |
| Medtronic | Minimed™ 770g      |  |
| Medtronic | Minimed™ 780g      |  |
| Tandem    | T:slim X2™         |  |
| Ypsomed   | Mylife™ YpsoPump®  |  |
| SOOIL     | Diabecare DANA-i   |  |
| SOOIL     | Dana Diabecare RS  |  |
| SOOIL     | Dana Diabecare R   |  |
| SOOIL     | Dana Diabecare IIS |  |

*End of Document*

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