



DECLARATION OF CONFORMITY

Name of Manufacturer Liko AB
Address of Manufacturer Nedre vägen 100, 975 92 Luleå, SWEDEN
Name of Device Liko
Reference Viking
Model XS, S, M, L, XL
Product Number 2040007, 2040006, 2040015, 2040014, 2040013,
2040033, 2040034, 2040035

I undersigned Daniel Ahlqvist, General Manager for Liko AB, declare hereby that the medical devices specified above are of class I and conform to the Essential Requirements listed in the annex I of the EC Directive 93/42/EEC.

This Declaration is Supported by:

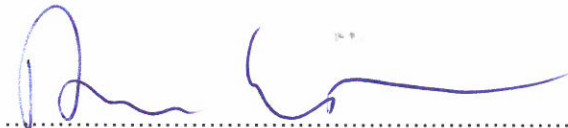
- *Quality Certificate Delivered by* INTERTEK SEMKO AB – Kista – SWEDEN
EN ISO 9001:2008
EN ISO 13485:2003
- *Certificates and Test Reports Delivered by* INTERTEK SEMKO AB – Kista – SWEDEN
EN ISO 10535:2006 (including EN 60601-1:2006 parts referred by EN ISO 10535:2006)
EN 60601-1-2:2007
- *Risk Analysis Carried out According to*
EN ISO 14971:2007

Technical file compiled by the manufacturer

In-house tests carried out by the manufacturer

Place Luleå, Sweden

Date 2013-02-26



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Daniel Ahlqvist