

EC Declaration of Conformity
EG Konformitäts-Erklärung
Déclaration CE de Conformité
Declaración de Conformidad tipo EC

We, Datrend Systems Inc, declare in exclusive responsibility that the product:
Wir, Datrend Systems Inc, erklären in alleiniger Verantwortung, daß das Produkt:
Nous, Datrend Systems Inc, déclarons sous notre seule responsabilité que le produit:
Datrend Systems Inc declaramos en la responsabilidad exclusiva que el producto:

Model / Modell / Modèle / Modelo: **vPad-A1 Modular Patient Simulator¹**

to which this declaration relates is in conformity with the following Directives:
auf die sich diese Erklärung bezieht, steht im Einklang mit den folgenden Richtlinien:
auquel se réfère cette déclaration est conforme aux Directives suivantes:
a cuál se relaciona este declaración es conforme a las siguientes Directivas:

2014/30/EU - Electromagnetic Compatibility (EMC)
2014/35/EU - Low Voltage Directive (LVD)
2011/65/EU - RoHS Directive

Harmonized standards used:
Verwendete harmonisierte Standards:
Normes harmonisées utilisées:
Normas armonizados utilizados:

EN 61326-1:2013 *Electrical equipment for measurement, control and laboratory use - EMC requirements*
EN 61010-1:2010 *Safety requirements for electrical equipment for measurement, control and laboratory use*
EN 63000:2018 *Technical documentation for assessment of electrical and electronic products with respect to restriction of hazardous substances*

Signed for, and on behalf of, Datrend Systems Inc.



Keith Ching
Director of Manufacturing Technology

EU Authorised Representative:
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Place of Issue: Richmond BC Canada

¹ Comprising in combination the A1 Base Module and any one or more of the following simulation modules: vPad-PS Patient Simulator; vPad-O2 Pulse Oximeter Tester; and/or vPad-NIBP Non-Invasive Blood Pressure Simulator