

Risk Analysis Report

Date 6 January 1998

Product Tom Thumb Infant Resuscitator

Product Type :

Tom Thumb Series all models

MDD Class II

Model Type TT4

Product Description

Hand held gas powered resuscitator unit

Manufacturer Address

Viamed,

15 Station Road,

Crosshills

Keighley,

West Yorkshire BD20 7DT

We hereby declare that the statements made herein are correct and valid

Company: Viamed Ltd. 15, Station Road, Crosshills, Keighley, West Yorkshire, BD20 7DT.

Signature



Name J.S.Lamb Director

Date 6th Jan 1998

Level of Risk Product:

CEMARK\TOMTHUMB\RARTT

6

January 1998 Insignificant 4: Tolerable 3 : Significant 2: Catastrophic 1

TT20

Ref	Hazard	Part of Equipment which pose risks	I R	Design solution Adopted	R R	Document referenced
C	ENERGY					
C2.1	Electricity		4		4	
C2.2	Heat		4		4	
C2.3	Mechanical Force		4		4	
C2.4	Iononizing Radiation		4		4	
C2.5	Non Iononizing Radiation		4		4	
C2.6	Electromagnetic Fields		4		4	
C2.7	Moving Parts				4	
C2.8	Suspended Masses		4		4	
C2.9	Patient Support Failure		4		4	
C2.10	Pressure (Vessel rupture)		4		4	
C2.11	Acoustic pressure		4		4	
C2.12	Vibration		4		4	
C2.13	Magnetic Fields (MRI)		4		4	
C3	BIOLOGICAL		4		4	
C3.1	Bio-Burden		4		4	
C3.2	Bio-Contamination		4		4	
C3.3	Bio-Incompatibility		4		4	
C3.4	Incorrect Output (Substance/energy)		4		4	
C3.5	Incorrect Formulation (Chemical Composition)		4		4	
C3.6	Toxicity		4		4	
C3.7	Cross Infection		4		4	
C3.8	Pyrogenicity		4		4	
C3.9	Inability to Maintain Hygienic Standards		4		4	
C3.10	Degradation		4		4	
C4.	ENVIRONMENTAL		4		4	
C4.1	Electromagnetic Interference		4		4	
C4.2	Inadequate supply of Power or Coolant		4		4	
C4.3	Likelihood of Operation outside Prescribed Environmental Conditions		4		4	

Level of Risk Product:

January 1998 Insignificant 4: Tolerable 3 : Significant 2: Catastrophic 1

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6

TT 21

Ref.	Hazard	Part of Equipment which pose risks	I R	Design solution Adopted	R R	Document referenced
C 1	Incompatibility with other Devices		4		4	
C4.5	Accidental Mechanical Damage		4		4	
C4.6	Contamination due to Waste Products and or Device Disposal		4		4	
C5	DEVICE USE				4	
C5.1	Inadequate Labelling	No instructions on equipment	3	Instruction Leaflet	4	
C5.2	Inadequate Operating Instructions				4	
C5.3	Inadequate Specification of Accessories		4		4	
C5.4	Inadequate Specification of Pre-Use Checks		4		4	
C5.5	Over-complicated Operating Instructions		4		4	
C5.6	Unavailable or Separated Operating Instructions		4	Training required by operator	4	
C5.7	Use by Unskilled /untrained Personnel		4	Training required by operator	4	
C5.8	Reasonable Foreseeable Abuse	Over pressure	2	Two Blow off valves	4	
C5.9	Insufficient Warning of Side Effects		4		4	
C5.10	Inadequate Warnings of Hazards Likely With Re-use of Single Use Devices		4		4	
C5.11	Incorrect Measurement and other Metrological Aspects		4		4	
C5.12	Incorrect Diagnosis		4		4	
C5.13	Erroneous Data Transfer		4		4	
C5.14	Misrepresentation of Results		4		4	
			4		4	

Ref.	Hazard	Part of Equipment which pose risks	4	Design solution Adopted	4	Document referenced
C5	Incompatibility with Consumables /accessories / other Devices		4		4	
C6	FUNCTIONAL FAILURE MAINTENANCE and AGEING		4		4	
C6.1	Inadequacy of Performance Characteristics for the Intended Use		4		4	
C6.2	Lack of ,or Inadequate Specification for Maintenance including Post Maintenance Functional Tests		4		4	
C6.3	Inadequate maintenance		4		4	
C6.4	Lack of Adequate Determination of End of Device Life		4		4	
C6.5	Loss of Mechanical Integrity	Overpressure on start up	4	Instruction manual	4	
C6.6	Inadequate Packaging (contaminationand/or Deterioration of the Device)		4		4	
C6.7	Improper Use		4		4	
	See separate sheet					