Quality Management System Route Map to Documents and Procedures Viamed Ltd ISO13485:2016

Version Date: 02 Oct 2023

Listing of Current Sections Search for ** Double Asterix to see Updated Documents

Section	Documents related	Processes Direct Links
4 Quality ma	anagement syste	em
4.1	Top Level Document: QMS	
Quality management	Route Map Viamed Ltd	
system	ISO13485_2016	
	Revision Document	
	ID127784	
	**Date Revision 29 Aug	
	2023 Reviewed 29 Aug 2023	
	Top Level Document:	
	Viamed ISO 13485:2016	
	Scope Revision Document	
	ID70776	
	Date Revision 27 Sep 2021	
	Reviewed 13 Oct 2022	
	Top Level Document:	
	VM3COP02.01 Exclusions	
	to Viamed ISO13485:2016	
	boundaries of ISO	
	Revision Document	
	ID74571	
	Date Revision 10 Nov 2021	
	Reviewed 01 Aug 2023	
	Top Level Document:	
	VM3COP00.00 VOP00.00	
	Viamed Quality Statement	
	policy and objectives	
	Revision Document	
	ID22684	
	Date Revision 16 Oct 2017	
	Reviewed 24 Aug 2023	
	Top Level Document:	
	VM3COP02.02 Viamed	
	Company Responsibilitys	
	organisation chart	
	structure	
	Revision Document	
	ID27474	
	Date Revision 20 Sep 2018	
	Reviewed 08 Nov 2022	

BS5750 Viamed

Revision Document

ID21353

Date Revision 10 Aug 2017

Reviewed 10 Aug 2017

BS EN ISO 13485-2016

Revision Document

ID19400

Date Revision 27 Mar 2017

Reviewed 27 Mar 2017

Chart 40 Management

review plan Issues followup Revision Document

ID22458

Date Revision 05 Oct 2017

Reviewed 05 Oct 2017

Chart 42 Processes, Tasks and Audits Review

Revision Document

ID23559

Date Revision 28 Oct 2017

Reviewed 28 Oct 2017

Chart 43 Processes and

Intrastats

Revision Document

ID23561

Date Revision 28 Oct 2017

Reviewed 28 Oct 2017

Intrastats overview

Revision Document

ID23567

Date Revision 28 Oct 2017

Reviewed 28 Oct 2017

Issues Overview

Revision Document ID23112

Date Revision 22 Oct 2017

Reviewed 22 Oct 2017

Document Index Overview

Revision Document ID8047

Date Revision 17 Mar 2011

Reviewed 17 Mar 2011

VM3COP00.01 Company

objectives

Revision Document

ID22842

Date Revision 17 Oct 2017

Reviewed 17 Oct 2017

Need Risks and

Expectations of External

Parties Viamed

Revision Document

ID130432

**Date Revision 27 Sep

2023 Reviewed 27 Sep 2023

4.1.1

The organization shall document a quality

Top Level Document: VOP | Process: 7723 01 Documentation and Records, Control,

Audit 10b Process Verification Viamed 24 Aug 2016

management system and maintain its effectiveness in accordance with the requirements of this International Standard and applicable regulatory requirements.

The organization shall establish, implement and maintain any requirement, procedure, activity or arrangement required to be documented by this International Standard or applicable regulatory requirements.

applicable regulatory requirements.
The organization shall document the role(s) undertaken by the organization under the applicable regulatory requirements.
NOTE Roles undertaken by the organization can include manufacturer, authorized

representative, importer

Creation, Storage, Retrieval, Revision Contro and Online Records

Revision Document ID120321

Date Revision 01 Jun 2023 Reviewed 01 Jun 2023

Top Level Document: Viamed ISO 13485:2016 Scope

Revision Document ID70776

Date Revision 27 Sep 2021 Reviewed 13 Oct 2022

Audit 10 Documentation Control

Revision Document ID63807

Date Revision 30 Jun 2021 Reviewed 30 Jun 2021 Process: 41

Retrieval, Revision Control Responsibility Allocation : Documentation

Control 16 Feb 2016

Process: 9

Distribution Of Faxes 16 Feb 2016

Process: 10

Distribution Of Emails 16 Feb 2016

Process: 8025

Check We Do Not Require A EU European

Representatives 09 Mar 2023

4.1.2

processes.

or distributor.

The organization shall: a) determine the processes needed for the quality management system and the application of these processes throughout the organization taking into account the roles undertaken bv the organization; b) apply a risk based approach to the control of the appropriate processes needed for the quality management system; c) determine the sequence and interaction of these

Top Level Document: VM3COP02.02 Viamed

Company Responsibilitys organisation chart structure

Revision Document ID27474

Date Revision 20 Sep 2018 Reviewed 08 Nov 2022

Top Level Document: VOP 21 Risk, Risk Management and Risk Analysis

Revision Document ID75935

Date Revision 24 Nov 2021 Reviewed 24 Nov 2021

Explanation Employee Roles and Titles

Revision Document ID22144

Date Revision 20 Sep 2017 Reviewed 20 Sep 2017

Chart 00 System Model

Revision Document ID8674 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Chart 01 System and Documentation

Revision Document ID8675
Date Revision 12 Oct 2011

Process: 7743

Customer Complaints Paper File 26 Sep 2016

Process: 7723

Audit 10b Process Verification Viamed 24 Aug

2016

Process: 7725

Audit 12 CE Files Viamed 24 Aug 2016

Reviewed 12 Oct 2011

Chart 02 Resource

Management

Revision Document ID8676 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Chart 03 Customer

Requirements

Revision Document ID8677 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Chart 04 Design and Development

Revision Document ID8678 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Chart 05 Product

Realisation

Revision Document ID8679 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Chart 06 General Process Control

Revision Document ID8680 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Chart 07 Measurement and Analysis

Revision Document ID8681 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Chart 08 Correction and Prevention

Revision Document ID8682 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Chart 09 Management System

Revision Document ID8683 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Chart 10 Documentation

Revision Document ID8684 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Chart 11 Provision of

Resources

Revision Document ID8685 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Chart 12 Infrastructure and Environment

Revision Document ID8686 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Chart 13 Sales Orders

Revision Document ID8687 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Chart 15 Purchasing

Revision Document ID8688

Date Revision 12 Oct 2011

Reviewed 12 Oct 2011

Chart 16 Internal Audits

Revision Document ID8689

Date Revision 12 Oct 2011

Reviewed 12 Oct 2011

Chart 17 Design Repairs

Revision Document ID8690

Date Revision 12 Oct 2011

Reviewed 12 Oct 2011

Chart 18 Calibration

Revision Document ID8691

Date Revision 12 Oct 2011

Reviewed 12 Oct 2011

Chart 19 HSE Risk

Assesments

Revision Document ID8692

Date Revision 12 Oct 2011

Reviewed 12 Oct 2011

Chart 20 Production

Revision Document ID8693

Date Revision 12 Oct 2011

Reviewed 12 Oct 2011

Chart 21 Repairs

Revision Document ID8694

Date Revision 12 Oct 2011

Reviewed 12 Oct 2011

Chart 22 Stock Control

Revision Document ID8695

Date Revision 12 Oct 2011

Reviewed 12 Oct 2011

Chart 23 Picking and

Packing

Revision Document ID8696

Date Revision 12 Oct 2011

Reviewed 12 Oct 2011

Chart 24 Goods Inwards

Revision Document ID8697

Date Revision 12 Oct 2011

Reviewed 12 Oct 2011

Chart 25 Inspection and

Test

Revision Document ID8698

Date Revision 12 Oct 2011

Reviewed 12 Oct 2011

Chart 26 Data Analysis

Revision Document ID8699

Date Revision 12 Oct 2011

Reviewed 12 Oct 2011

Chart 27 Customer

Complaints Chart 27

Revision Document ID8700

Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Chart 28 Quarantine and

Hold

Revision Document ID8701 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Chart 29 Sales Acquisition Revision Document ID8702 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Chart 30 System Design Plan

Revision Document ID8703 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Chart 31 Chart Interfaces Revision Document ID8704 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Chart 32 Generic Sales Process

Revision Document ID8705 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Chart 33 Launch of a new product

Revision Document ID8706 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Chart 34 Process Teams Org Chart

Revision Document ID8707 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Audit 20 Process verification to Managment

Revision Document ID73324

Date Revision 26 Oct 2021 Reviewed 26 Oct 2021

4.1.3

For each quality management system process, the organization shall: a) determine criteria and methods needed to ensure that both the operation and control of these processes are effective; b) ensure the availability of resources and information necessary to support the operation and monitoring of these processes;

c) implement actions

results and maintain the

effectiveness of these

necessary to achieve planned

Top Level Document: VOP

13 Process Monitoring, System Reviews, Audits, **Management Reviews Analysis Data PMS Post**

Market

Revision Document ID75461

Date Revision 18 Nov 2021 Reviewed 18 Nov 2021

Explanation Employee **Roles and Titles**

Revision Document ID22144

Date Revision 20 Sep 2017 Reviewed 20 Sep 2017

VM3COP27.01 Searching Intrastats Issues

Revision Document ID6657

Process: 27

Management Reviews And Quality Audits 16 Feb 2016

Process: 7723

Audit 10b Process Verification Viamed 24 Aug 2016

Process: 7730

Audit 20 Process Verification To Managment Viamed 24 Aug 2016

Process: 5889

Responsibility Allocation: Audit And Task -Audit 24 Feb 2016

Process: 7714

Audit 01 Picking Packing Viamed 24 Aug

2016

Process: 7715

Audit 02 Contract Review Viamed 24 Aug

2016

Process: 7716

processes; d) monitor, measure as appropriate, and analyse these processes; e) establish and maintain records needed to demonstrate conformance to this International Standard and compliance with applicable regulatory

requirements (see 4.2.5).

Date Revision 02 Nov 2009 Reviewed 02 Nov 2009 VM3COP27.17 Complete

Auto calender Issues

Revision Document ID16995

Date Revision 26 May 2016 Reviewed 26 May 2016

Issues Overview

Revision Document ID23112 Process: 7720

Date Revision 22 Oct 2017 Reviewed 22 Oct 2017

Intrastats overview

Revision Document

ID23567

Date Revision 28 Oct 2017 Reviewed 28 Oct 2017

Employee Roles

Revision Document

ID20125

Date Revision 16 May 2017 Reviewed 16 May 2017

Employee roles Example

Process

Revision Document

ID20129

Date Revision 16 May 2017

Reviewed 16 May 2017

VM3COP27.02 Collecting **Emails and Distributing**

Revision Document

ID85362

Date Revision 22 Mar 2022

Reviewed 22 Mar 2022 Employee Roles Individual

Processes

Revision Document

ID20127

Date Revision 16 May 2017

Reviewed 16 May 2017

Audit 18 Management

Review

Revision Document

ID73320

Date Revision 26 Oct 2021 Reviewed 26 Oct 2021

Audit 20 Process

verification to Managment

Revision Document

ID73324

Date Revision 26 Oct 2021 Reviewed 26 Oct 2021

Audit 03 Design Control Viamed 24 Aug 2016

Process: 7717

Audit 05 Purchasing Suppliers Viamed 24 Aug

2016

Process: 7718

Audit 06 Calibration Viamed 24 Aug 2016

Process: 7719

Audit 07 Handling And Storage Viamed 24

Aug 2016

Audit 08 Training Viamed 24 Aug 2016

Process: 7721

Audit 09 Goods Inward And Product Identity

Viamed 24 Aug 2016

Process: 7722

Audit 10 Documentation Control Viamed 24

Aug 2016 Process: 7724

Audit 11 Repairs And Service Viamed 24 Aug

2016

Process: 7725

Audit 12 CE Files Viamed 24 Aug 2016

Process: 7726

Audit 14 Complaints And Corrective Actions

Viamed 24 Aug 2016

Process: 7727

Audit 15 Production Viamed 24 Aug 2016

Process: 7728

Audit 17 Internal Audits Viamed 24 Aug 2016

Process: 7729

Audit 19 Health And Saftey Viamed 24 Aug

2016

Process: 7731

Audit 21 Audit Of Audit Viamed 24 Aug 2016

Process: 7732

Audit 22 Post Market Survellance Viamed 24

Aug 2016

Process: 7733

Audit 23 Analysis Of Data Viamed 24 Aug

2016

Process: 26

Company Resources 16 Feb 2016

Process: 8025

Check We Do Not Require A EU European

Representatives 09 Mar 2023

Process: 8028

Viamed Shopify Sales Report Export 11 Apr

2023

4.1.4

For each quality

The organization shall

management system process, the organization shall:

Top Level Document: VOP Process: 7725 01 Documentation and Records, Control,

Creation, Storage, **Retrieval, Revision Control** Viamed 24 Aug 2016

Audit 12 CE Files Viamed 24 Aug 2016

Process: 7730

Audit 20 Process Verification To Managment

manage these quality
management system
processes in accordance with
the requirements of this
International Standard and
applicable regulatory
requirements. Changes to be
made to these processes shall
be:

- a) evaluated for their impact on the quality management system;
- b) evaluated for their impact on the medical devices produced under this quality management system
- with the requirements of this International Standard and applicable regulatory requirements.

c) controlled in accordance

and Online Records

Revision Document

Date Revision 01 Jun 2023 Reviewed 01 Jun 2023

Audit 20 Process

verification to Managment

Revision Document ID73324

Date Revision 26 Oct 2021 Reviewed 26 Oct 2021

Audit 18 Management Review

Revision Document ID73320

Date Revision 26 Oct 2021 Reviewed 26 Oct 2021

Issues Overview

Revision Document ID23112 Date Revision 22 Oct 2017 Reviewed 22 Oct 2017

Employee Roles

Revision Document ID20125

Date Revision 16 May 2017 Reviewed 16 May 2017

Employee roles Example Process

Revision Document

Revision Document ID20129

Date Revision 16 May 2017 Reviewed 16 May 2017

Employee Roles Individual

Processes

Revision Document ID20127

Date Revision 16 May 2017 Reviewed 16 May 2017

Explanation Employee

Roles and Titles

Revision Document

ID22144

Date Revision 20 Sep 2017

Reviewed 20 Sep 2017

Explanation Employee Roles Titles Responsibilitys

Processes and Repeating Tasks Monitoring

Revision Document

ID22287

Date Revision 27 Sep 2017 Reviewed 27 Sep 2017

Chart 43 Processes and

Intrastats

Revision Document

ID23561

Date Revision 28 Oct 2017 Reviewed 28 Oct 2017 Process: 7878

Review Possible Upcoming Regulation

Changes 22 Oct 2017

Process: 8025

Check We Do Not Require A EU European

Representatives 09 Mar 2023

Chart 42 Processes, Tasks and Audits Review

Revision Document ID23559

Date Revision 28 Oct 2017 Reviewed 28 Oct 2017

Chart 40 Management review plan Issues followup

Revision Document ID22458

Date Revision 05 Oct 2017 Reviewed 05 Oct 2017

VM3COP24.02 Document Change Performing a Risk Assessment

Revision Document ID75310

Date Revision 17 Nov 2021 Reviewed 17 Nov 2021

VM3COP24.01 Definitions of Risk

Revision Document ID75525

Date Revision 19 Nov 2021 Reviewed 19 Nov 2021

VM3COP24.00 Viamed **Overall Risk Analysis** Program Risk Register

Revision Document ID47771

Date Revision 12 Nov 2020 Reviewed 12 Nov 2020

4.1.5

For each quality management system process, the organization shall: When the organization chooses to outsource any process that affects product conformity to requirements, it shall monitor and ensure control over such processes. The organization shall retain responsibility of conformity to this International Standard and to customer and applicable regulatory requirements for outsourced processes. The controls shall be proportionate to the risk involved and the ability of the external party to meet the requirements in accordance with 7.4. The controls shall include

written quality agreements.

Top Level Document: VOP Process: 7717 05 Supplier Control,

Supplier Review, Purchase Orders, Supplier Returns and Rejection

Revision Document ID75847

Date Revision 23 Nov 2021 Reviewed 23 Nov 2021

Audit 05 Purchasing suppliers

Revision Document ID69314 Date Revision 09 Sep 2021

Reviewed 09 Sep 2021

Audit 05 Purchasing Suppliers Viamed 24 Aug 2016

Process: 7199

Non Conformities Review Viamed 09 Mar 2016

Process: 8025

Check We Do Not Require A EU European

Representatives 09 Mar 2023

4.1.6 For each quality management system process, the organization shall: The organization shall document procedures for the validation of the application of computer software used in the quality management system. Such software applications shall be validated prior to initial use and, as appropriate, after changes to such software or its application.

The specific approach and activities associated with software validation and revalidation shall be proportionate to the risk associated with the use of the Date Revision 16 May 2017 software.

Records of such activities shall be maintained (see 4.2.5).

Top Level Document: Audit 27 Software Validation

Revision Document ID113182

Date Revision 09 Mar 2023 Reviewed 09 Mar 2023

Top Level Document: VOP 27 Software Validation

Revision Document ID91486

Date Revision 10 Jun 2022 Reviewed 10 Jun 2022

Intrastats Amendment Log Revision Document ID20136

Date Revision 16 May 2017 Reviewed 16 May 2017

Validation of Intrastats Revision Document ID20140

Reviewed 16 May 2017

Process: 7850

Software Validation Scan Incorrect Product 01

Oct 2017 Process: 7851

Software Validation Scan Un-QA Product To

Order 01 Oct 2017 Process: 7852

Software Validation Expired Stock 01 Oct

2017

Process: 7853

Software Validation Non Sell Able Shelf 01

Oct 2017 Process: 7854

Software Validation In Production List 01 Oct

2017

Process: 7855

Software Validation - Production Lists 01 Oct

2017

Process: 7856

Software Validation Unchecked Orders 01 Oct

2017

Process: 7857

Software Validation Stock Tracking Check 01

Oct 2017

Process: 7858

Software Validation Attempt To QA Some

Stock 01 Oct 2017 Process: 7861

Software Validation Of Training Documents

Forced Reading 03 Oct 2017

Process: 7865

Software Validation Conflicting Audits 07 Oct

2017

Process: 7870

Software Validation Non Conformance Product Risk Feedback Loop 15 Oct 2017

4.2 Documentation requirements

Top Level Document: VOP 01 Documentation and Records, Control, Creation, Storage, Retrieval, Revision Control and Online Records

Revision Document ID120321

Date Revision 01 Jun 2023 Reviewed 01 Jun 2023

Audit 10 Documentation Control

Revision Document ID63807

Date Revision 30 Jun 2021 Reviewed 30 Jun 2021

Process: 23

Company Objectives 16 Feb 2016

Process: 22

Company Policys 16 Feb 2016

Process: 23

4.2.1

The quality management system documentation (see 4.2.4) shall include:

a) documented statements of

Top Level Document: VM3COP00.00 VOP00.00 Viamed Quality Statement policy and objectives Revision Document

a quality policy and quality objectives; b) a quality manual; c) documented procedures and records required by this International Standard; d) documents, including records, determined by the organization to be necessary to ensure the effective planning, operation. and control of its processes; e) other documentation

specified by applicable

regulatory requirements.

ID22684

Date Revision 16 Oct 2017 Reviewed 24 Aug 2023

Top Level Document: VM3COP00.00 VOP00.00 VST Quality Statement

policy and objectives Revision Document

ID22062

Date Revision 16 Sep 2017 Reviewed 24 Aug 2023

Top Level Document: VOP 01 Documentation and Records, Control, Creation, Storage,

Retrieval, Revision Control Process: 5877 and Online Records

Revision Document

ID120321

Date Revision 01 Jun 2023 Reviewed 01 Jun 2023

Explaination Quality Objectives

Revision Document

ID18483

Date Revision 18 Jan 2017 Reviewed 18 Jan 2017

Explanation Employee **Roles and Titles**

Revision Document ID22144

Date Revision 20 Sep 2017 Reviewed 20 Sep 2017

Audit 20 Process

verification to Managment

Revision Document

ID73324

Date Revision 26 Oct 2021 Reviewed 26 Oct 2021

Audit 10 Documentation Control

Revision Document

ID63807

Date Revision 30 Jun 2021 Reviewed 30 Jun 2021

VM3COP00.01 Company objectives

Revision Document

ID22842

Date Revision 17 Oct 2017 Reviewed 17 Oct 2017

Company Objectives 16 Feb 2016

Process: 7730

Audit 20 Process Verification To Managment

Viamed 24 Aug 2016

Process: 7723

Audit 10b Process Verification Viamed 24 Aug

2016

Process: 7834

Financial Review 20 Sep 2017

Process: 7862

Review The Audit Calender Screen 04 Oct

2017

Process: 27

Management Reviews And Quality Audits 16

Feb 2016

Review Company Data 17 Feb 2016

Process: 6861

Management Meeting Review Weekly Meeting

09 Mar 2016 Process: 7037

Responsibility Allocation: Responsibility, authority and communication 09 Mar 2016

Process: 7057

Responsibility Allocation: Complaints and

Vigilance Notifications 09 Mar 2016

Process: 7070

Management Review 09 Mar 2016

Process: 7713

Review Roles And Responsibilitys 17 Aug

2016

Process: 7830

Review Q.A. Failures Report 18 Sep 2017

Process: 7837

Review External Parties Influencing The QMS

VST / Viamed 23 Sep 2017

Process: 7838

Review VIAMED Feedback - Customer

Feedback Negative 23 Sep 2017

Process: 7839

Review VIAMED Feedback - Customer

Complaints 23 Sep 2017

Process: 7842

Review VIAMED Product Feedback Negative

23 Sep 2017

Process: 7845

7.1.4 Environment Of Operations 25 Sep 2017

Process: 7848

Review ISO Scopes 27 Sep 2017

Process: 7849

Review Product Failures New Codes 28 Sep

2017

Process: 7120

General Maintenance Requirements 09 Mar

2016

Process: 28

Supplier Review 16 Feb 2016

Process: 5887

Review ISO/EN Documents 24 Feb 2016

Process: 5889

Responsibility Allocation : Audit And Task -

Audit 24 Feb 2016 **Process: 6866**

Internal Process Verification Complete

Systems Review 09 Mar 2016

Process: 7199

Non Conformities Review Viamed 09 Mar

2016

Process: 7828

Review The Quality Policy Viamed 16 Sep

2017

Process: 6821

Responsibility Allocation: VIAMED

Management Meeting Supplier Review 09 Mar

2016

Process: 7697

Yearly Pricing Review 09 May 2016

Process: 57

Temporary Stock Notices 17 Feb 2016

Process: 8029

Send Intercompany Invoices To Jean 12 Apr

2023

4.2.2

The organization shall document a quality manual that includes:

- a) the scope of the quality management system, including details of and justification for any exclusion
- or non-application;
- b) the documented procedures for the quality management system, or reference to them;
- c) a description of the interaction between the processes of the quality management system. The quality manual shall outline the structure of the

documentation used in the quality management system.

Top Level Document:

VM3COP02.01 Exclusions to Viamed ISO13485:2016

boundaries of ISO

Revision Document ID74571

Date Revision 10 Nov 2021 Reviewed 01 Aug 2023

Top Level Document: VM3COP02.02 Viamed Company Responsibilitys

organisation chart structure

Revision Document

ID27474 Date Revision 20 Se

Date Revision 20 Sep 2018 Reviewed 08 Nov 2022

Top Level Document: Viamed ISO 13485:2016 Scope

Revision Document

ID70776

Date Revision 27 Sep 2021 Reviewed 13 Oct 2022

Structure of the documentation used in the quality management

system

Revision Document ID18487

Date Revision 18 Jan 2017 Reviewed 18 Jan 2017

Audit 20 Process

Process: 7723

Audit 10b Process Verification Viamed 24 Aug

2016

Process: 7730

Audit 20 Process Verification To Managment

Viamed 24 Aug 2016

verification to Managment **Revision Document** ID73324 Date Revision 26 Oct 2021 Reviewed 26 Oct 2021 **Audit 10 Documentation**

Control

Revision Document ID63807

Date Revision 30 Jun 2021 Reviewed 30 Jun 2021

4.2.3

For each medical device type **17 Design Research and** or medical device family, the organization shall establish and maintain one or more files either containing or referencing documents generated to demonstrate conformity with the requirement of this International Standard and compliance with applicable

- regulatory requirements. The content of the file(s) shall include, but is not limited to: a) general description of the
- medical device, intended use/purpose, and labelling, including any instructions for use;
- b) specifications for product;
- c) specifications or procedures for manufacturing, packaging, storage, handling and distribution;
- d) procedures for measuring and monitoring;
- e) as appropriate, requirements for installation; f) as appropriate, procedures

for servicing. Medical device file Documentation

requirements

4.2.4 Documents required by the quality management system shall be controlled. Records are a special type of document and shall be controlled according to the requirements given in 4.2.5. A documented procedure shall define the controls

Top Level Document: VOP | Process: 7716 Development

Revision Document ID25632 Date Revision 19 Mar 2018 Reviewed 19 Mar 2018

Route to Medical device files

Revision Document ID18495 Date Revision 18 Jan 2017 Reviewed 18 Jan 2017

Audit 03 Design Control Revision Document ID111315

Date Revision 17 Feb 2023 Reviewed 17 Feb 2023

Audit 03 Design Control Viamed 24 Aug 2016

Process: 7723

Audit 10b Process Verification Viamed 24 Aug

2016

Top Level Document: VOP 01 Documentation and

Records, Control, Creation, Storage, Retrieval, Revision Control and Online Records

Revision Document ID120321

Date Revision 01 Jun 2023 Reviewed 01 Jun 2023

Process: 7722

Audit 10 Documentation Control Viamed 24 Aug 2016

Process: 8032

Review Contact Documentation 22 Aug 2023

needed to:

- a) review and approve documents for adequacy prior to issue;
- b) review, update as necessary and re-approve documents:
- c) ensure that the current revision status of and changes to documents are identified;
- d) ensure that relevant versions of applicable documents are available at points of use;
- e) ensure that documents remain legible and readily identifiable;
- f) ensure that documents of external origin, determined by the organization to be necessary

for the planning and operation of the quality management system, are identified and their distribution controlled;

- g) prevent deterioration or loss of documents;
- h) prevent the unintended use of obsolete documents and apply suitable identification to them. The organization shall ensure that changes to documents are reviewed and approved either by the original approving function

or another designated function that has access to pertinent background information upon which to

base its decisions.
The organization shall define the period for which at least one copy of obsolete documents shall be retained. This period shall

ensure that documents to which medical devices have been manufactured and tested are available for at least the lifetime of the medical device as defined by the organization,

but not less than the retention period of any resulting record (see 4.2.5),

Explanation Control of documents

Revision Document ID21322

Date Revision 06 Aug 2017 Reviewed 06 Aug 2017

DO NOT USE VM3COP01

Document Updates / Amendment control

Revision Document ID22201

Date Revision 23 Sep 2017 Reviewed 23 Sep 2017

Audit 10 Documentation Control

Revision Document ID63807

Date Revision 30 Jun 2021 Reviewed 30 Jun 2021

DO NOT USE VM3COP14
Documentation

Revision Document ID9276 Date Revision 18 Oct 2011 Reviewed 18 Oct 2011

Audit 23 Analysis of Data Revision Document

ID67997

Date Revision 23 Aug 2021 Reviewed 23 Aug 2021

VM3COP10.02 Product Recall locate products out in the Field

Revision Document ID74788

Date Revision 12 Nov 2021 Reviewed 12 Nov 2021

or as specified by applicable
Control of documents
Documentation
requirements
4.2.5

Records shall be maintained

conformity to requirements

to provide evidence of

and of the effective operation of the quality

management system.

The organization shall

for the identification.

storage, security and

records.

time and disposition of

protecting confidential

contained in records in

accordance with the

applicable regulatory

readily identifiable and

record shall remain

retrievable. Changes to a

requirements.

identifiable.

health information

document procedures to

define the controls needed

integrity, retrieval, retention

The organization shall define and implement methods for

Records shall remain legible,

The organization shall retain

organization, or as specified

the records for at least the

lifetime of the medical

device as defined by the

by applicable regulatory

than two years from

requirements

requirements, but not less

the medical device release

by the organization. Control

of records Documentation

Top Level Document: VOP

01 Documentation and Records, Control, Creation, Storage,

and Online Records

Revision Document ID120321

Date Revision 01 Jun 2023 Reviewed 01 Jun 2023

DO NOT USE VM3COP01

Document Updates / Amendment control

Revision Document ID22201

Date Revision 23 Sep 2017 Reviewed 23 Sep 2017

VM3COP14.01 Disposition of Documents / Records.

Revision Document ID15464

Date Revision 14 Aug 2015 Reviewed 14 Aug 2015

Guide to Intrastats

Revision Document

ID24779

Date Revision 22 Dec 2017 Reviewed 22 Dec 2017

Intrastats overview

Revision Document

ID23567

Date Revision 28 Oct 2017 Reviewed 28 Oct 2017

DO NOT USE VM3COP14

Documentation

Revision Document ID9276 Date Revision 18 Oct 2011 Reviewed 18 Oct 2011

Audit 10 Documentation

Control

Revision Document ID63807

Date Revision 30 Jun 2021 Reviewed 30 Jun 2021

Process: 7722

Audit 10 Documentation Control Viamed 24

Aug 2016 Process: 7725

Retrieval, Revision Control Audit 12 CE Files Viamed 24 Aug 2016

Process: 8027

Update Pricing For Viamed Shopify 11 Apr

5 Management commitment

5.1

Top management shall provide evidence of its commitment to the development and

Top Level Document: VOP | Process: 7730 02 Personnel and

Responsibility , Staff and Staffing Issues, Training, **Roles and Tasks**

Audit 20 Process Verification To Managment

Viamed 24 Aug 2016

Process: 7715

Audit 02 Contract Review Viamed 24 Aug

implementation of the quality management system and maintenance of its effectiveness by:

a) communicating to the organization the importance of meeting customer as well as applicable

regulatory requirements;

- b) establishing the quality policy;
- c) ensuring that quality objectives are established;
- d) conducting management reviews;
- e) ensuring the availability of resources. **Management** commitment

Revision Document

ID93320

Date Revision 01 Jul 2022 Reviewed 01 Jul 2022

Top Level Document: VOP Process: 27 18 Maintenance Building, Fabric and Infrastructure

Revision Document ID119029

Date Revision 15 May 2023 Reviewed 15 May 2023

Top Level Document: VM3COP00.00 VOP00.00

Viamed Quality Statement policy and objectives

Revision Document ID22684

Date Revision 16 Oct 2017 Reviewed 24 Aug 2023

VM3COP02 Organisation **Responsibilities Viamed**

Revision Document ID17423

Date Revision 07 Sep 2016 Reviewed 07 Sep 2016

Chart 01 System and Documentation

Revision Document ID8675 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Chart 02 Resource Management

Revision Document ID8676 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

VM3COP19 Health and Safety

Revision Document ID21800

Date Revision 05 Sep 2017 Reviewed 05 Sep 2017

Audit 20 Process

verification to Managment

Revision Document

ID73324

Date Revision 26 Oct 2021 Reviewed 26 Oct 2021

Explaination Quality

Objectives

Revision Document

ID18483

Date Revision 18 Jan 2017

Reviewed 18 Jan 2017

Explanation Employee Roles and Titles

Revision Document

ID22144

Date Revision 20 Sep 2017

2016

Process: 7833

Importance Of Effective Quality Management 20 Sep 2017

Management Reviews And Quality Audits 16

Feb 2016 Process: 7070

Management Review 09 Mar 2016

Process: 7848

Review ISO Scopes 27 Sep 2017

Process: 23

Company Objectives 16 Feb 2016

Process: 7686

Thorough Checking Of Awaiting Action Tray -

Priority 8s 21 Apr 2016

Reviewed 20 Sep 2017 **Explanation Control of**

documents

Revision Document

ID21322

Date Revision 06 Aug 2017 Reviewed 06 Aug 2017

How to Hold Intrastat

Meetings

Revision Document ID8928 Date Revision 18 Oct 2011 Reviewed 18 Oct 2011

Chart 40 Management review plan Issues followup

Revision Document ID22458

Date Revision 05 Oct 2017 Reviewed 05 Oct 2017

Audit 18 Management

Review

Revision Document

ID73320

Date Revision 26 Oct 2021 Reviewed 26 Oct 2021

Viamed Top Level Quality

Objectives

Revision Document ID130426

**Date Revision 27 Sep 2023 Reviewed 27 Sep 2023

Top management shall ensure that customer requirements and applicable regulatory requirements are determined and met.

Customer focus

Top Level Document: VOP | Process: 7 03 Contract Review,

Enquires, Office Processes

Revision Document ID77875

Date Revision 15 Dec 2021 Reviewed 15 Dec 2021

Top Level Document: VOP 19 Feedback Customer Complaints Vigilance and Notifications Viamed Ltd

Revision Document

ID75475

Date Revision 18 Nov 2021 Reviewed 18 Nov 2021

Top Level Document: VOP 07 Stock Control, Handling, Control of Labelling, Storage,

Movement

Revision Document

ID88809

Date Revision 06 May 2022 Reviewed 06 May 2022

Audit 02 Contract Review and Sales Order Processing Fax Paper 16 Feb 2016 Revision Document

Responsibility Allocation : Checking Of Sales

Orders 16 Feb 2016

Process: 11

Distribution Of Post 16 Feb 2016

Process: 5882

Responsibility Allocation: Send Post To

Humanmed 24 Feb 2016

Process: 2

Answering Telephones 16 Feb 2016

Process: 7715

Audit 02 Contract Review Viamed 24 Aug

2016

Process: 7743

Customer Complaints Paper File 26 Sep 2016

Process: 7716

Audit 03 Design Control Viamed 24 Aug 2016

Process: 7696

Send VIAMED Delivery Notifications 28 Apr

2016

Process: 6898

GHX Web Pricing 09 Mar 2016

Process: 19

Maintaining Leaflet Stocks 16 Feb 2016

Process: 14

Process: 15

12/10/2023, 11:03	Омз коите мар	viamed Ltd iSO13485:2016
	ID69328 Date Revision 09 Sep 2021 Reviewed 09 Sep 2021 Audit 16 Sales and Marketing Revision Document ID69457 Date Revision 10 Sep 2021 Reviewed 10 Sep 2021	Filing and Archiving 16 Feb 2016 Process: 10 Distribution Of Emails 16 Feb 2016 Process: 9 Distribution Of Faxes 16 Feb 2016 Process: 7996 Verification Repairs Older Repairs 07 Feb 2022 Process: 7934 Test Website Questions 02 May 2019
Top management shall ensure that the quality policy: a) is applicable to the purpose of the organization; b) includes a commitment to comply with requirements and to maintain the effectiveness of the quality management system; c) provides a framework for establishing and reviewing quality objectives; d) is communicated and understood within the organization; e) is reviewed for continuing suitability. Quality policy	Top Level Document: VM3COP00.00 VOP00.00 Viamed Quality Statement policy and objectives Revision Document ID22684 Date Revision 16 Oct 2017 Reviewed 24 Aug 2023 Top Level Document: VM3COP00.00 VOP00.00 VST Quality Statement policy and objectives Revision Document ID22062 Date Revision 16 Sep 2017 Reviewed 24 Aug 2023 VM3COP00.01 Company objectives Revision Document ID22842 Date Revision 17 Oct 2017 Reviewed 17 Oct 2017 Audit 18 Management Review Revision Document ID73320 Date Revision 26 Oct 2021 Reviewed 26 Oct 2021 Audit 20 Process verification to Managment Revision Document ID73324 Date Revision 26 Oct 2021 Reviewed 26 Oct 2021 Reviewed 26 Oct 2021 Reviewed 26 Oct 2021	Process: 23 Company Objectives 16 Feb 2016 Process: 22 Company Policys 16 Feb 2016 Process: 23 Company Objectives 16 Feb 2016 Process: 7723 Audit 10b Process Verification Viamed 24 Aug 2016 Process: 7833 Importance Of Effective Quality Management 20 Sep 2017 Process: 7828 Review The Quality Policy Viamed 16 Sep 2017 Process: 7827 Review The Quality Policy VST 16 Sep 2017
Planning		
Top management shall ensure that quality objectives, including those needed to meet applicable regulatory requirements and requirements for product, are established at relevant functions and levels within the organization. The	Top Level Document: VOP 07 Stock Control, Handling, Control of Labelling, Storage, Movement Revision Document ID88809 Date Revision 06 May 2022 Reviewed 06 May 2022 Top Level Document: VOP	Process: 7730 Audit 20 Process Verification To Managment Viamed 24 Aug 2016 Process: 7830 Review Q.A. Failures Report 18 Sep 2017 Process: 26 Company Resources 16 Feb 2016 Process: 5877 Review Company Data 17 Feb 2016

quality objectives shall be measurable and consistent with the quality policy.

Quality objectives

20 Goods in Purchases, Returns, Repairs, Inspection / Rejection

Revision Document ID75943

Date Revision 24 Nov 2021 Reviewed 24 Nov 2021

VM3COP18 Post Market Surveilance

Revision Document ID75985

Date Revision 24 Nov 2021 Reviewed 24 Nov 2021

Explanation Employee Roles and Titles

Revision Document

ID22144

Date Revision 20 Sep 2017 Reviewed 20 Sep 2017

Explaination Quality

Objectives

Revision Document

ID18483

Date Revision 18 Jan 2017

Reviewed 18 Jan 2017

Audit 20 Process

verification to Managment

Revision Document

ID73324

Date Revision 26 Oct 2021

Reviewed 26 Oct 2021

Viamed Top Level Quality Objectives

Revision Document

ID130426

**Date Revision 27 Sep 2023 Reviewed 27 Sep 2023

5.4.2

Top management shall ensure that:

a) the planning of the quality management system is carried out in order to meet the requirements given in 4.1, as well as the quality objectives;

b) the integrity of the quality management system is maintained when changes to the quality management system are

management system are planned and implemented.

Quality management system planning

Top Level Document:

VM3COP02.02 Viamed Company Responsibilitys organisation chart

structure
Revision Doc

Revision Document ID27474

Date Revision 20 Sep 2018

Reviewed 08 Nov 2022

Top Level Document:

VM3COP00.00 VOP00.00

VST Quality Statement policy and objectives

Revision Document ID22062

Date Revision 16 Sep 2017 Reviewed 24 Aug 2023

Top Level Document: VM3COP00.00 VOP00.00 Viamed Quality Statement Process: 11

Distribution Of Post 16 Feb 2016

Process: 5882

Responsibility Allocation : Send Post To

Humanmed 24 Feb 2016

Process: 7723

Audit 10b Process Verification Viamed 24 Aug

2016

Process: 7730

Audit 20 Process Verification To Managment

Viamed 24 Aug 2016

policy and objectives

Revision Document

ID22684

Date Revision 16 Oct 2017

Reviewed 24 Aug 2023

Top Level Document: VOP 21 Risk, Risk Management and Risk Analysis

Revision Document

ID75935

Date Revision 24 Nov 2021

Reviewed 24 Nov 2021

Explanation Employee

Roles and Titles

Revision Document

ID22144

Date Revision 20 Sep 2017

Reviewed 20 Sep 2017

Explaination Quality

Objectives

Revision Document

ID18483

Date Revision 18 Jan 2017

Reviewed 18 Jan 2017

Explanation Control of

documents

Revision Document

ID21322

Date Revision 06 Aug 2017

Reviewed 06 Aug 2017

Route to Medical device files

Revision Document

ID18495

Date Revision 18 Jan 2017

Reviewed 18 Jan 2017

VM3COP20.01 Post In Distributing the Post

Revision Document

ID103501

Date Revision 14 Nov 2022

Reviewed 14 Nov 2022

Audit 20 Process

verification to Managment

Revision Document

ID73324

Date Revision 26 Oct 2021

Reviewed 26 Oct 2021

Viamed Top Level Quality Objectives

Revision Document

ID130426

**Date Revision 27 Sep

2023 Reviewed 27 Sep 2023

VM3COP00.01 Company objectives

Revision Document

ID22842

	Date Revision 17 Oct 2017	
	Reviewed 17 Oct 2017	
5.5		
Responsibility, authority	Top Level Document: VOP 02 Personnel and	
and communication	Responsibility, Staff and	
	Staffing Issues, Training,	
	Roles and Tasks	
	Revision Document	
	ID93320	
	Date Revision 01 Jul 2022	
	Reviewed 01 Jul 2022	
	Top Level Document: QC	
	44 MHRA / CMDCAS Risk	
	Assessment Initial	
	Assessment form Revision Document	
	ID75549	
	Date Revision 19 Nov 2021	
	Reviewed 19 Nov 2021	
	Top Level Document: VOP	
	19 Feedback Customer	
	Complaints Vigilance and	
	Notifications Viamed Ltd	
	Revision Document	
	ID75475	
	Date Revision 18 Nov 2021	
	Reviewed 18 Nov 2021	
5.5.1		Process: 7720
Top management shall	02 Personnel and	Audit 08 Training Viamed 24 Aug 2016
ensure that responsibilities	Responsibility , Staff and	Process: 7730
and authorities are defined,	Staffing Issues, Training,	Audit 20 Process Verification To Managment
documented and communicated within the	Roles and Tasks Revision Document	Viamed 24 Aug 2016 Process: 7713
organization.	ID93320	Review Roles And Responsibilitys 17 Aug
Top management shall	Date Revision 01 Jul 2022	2016
document the interrelation of		Process: 6837
all personnel who manage,	Top Level Document:	Personnel Requirements and Training 09 Mar
perform and verify work	VM3COP02.02 Viamed	2016
affecting quality and shall	Company Responsibilitys	
ensure the independence and	organisation chart	
authority necessary to	structure	
perform these tasks.	Revision Document	
Responsibility and authority	ID27474 Date Revision 20 Sep 2018	
authority	Reviewed 08 Nov 2022	
	Explanation Employee	
	Roles and Titles	
	Revision Document	
	ID22144	
	Date Revision 20 Sep 2017	
	Reviewed 20 Sep 2017	
	VM3COP02 Organisation	
	Responsibilities Viamed	
	Revision Document ID17423	
	Date Revision 07 Sep 2016	
	Reviewed 07 Sep 2016	
II		

Chart 01 System and Documentation

Revision Document ID8675 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Chart 02 Resource Management

Revision Document ID8676 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Viamed Company Format Company format 1

Revision Document ID9039 Date Revision 18 Oct 2011 Reviewed 18 Oct 2011

Viamed Company Format Company format 2

Revision Document ID9040 Date Revision 18 Oct 2011 Reviewed 18 Oct 2011

Viamed Company Format Company format 3

Revision Document ID9041 Date Revision 18 Oct 2011 Reviewed 18 Oct 2011

Viamed Company Format Company format 4

Revision Document ID9042 Date Revision 18 Oct 2011 Reviewed 18 Oct 2011

Audit 08 Training, **Competence and Human** Resources

ID70147 Date Revision 20 Sep 2021

Reviewed 20 Sep 2021

Audit 20 Process

Revision Document

verification to Managment

Revision Document ID73324

Date Revision 26 Oct 2021

Reviewed 26 Oct 2021

Audit 19 Health and Safety, Working **Conditions and Building Fabric Issues**

Revision Document ID68045

Date Revision 24 Aug 2021 Reviewed 24 Aug 2021

5.5.2

Top management shall appoint a member of management who, irrespective of other responsibilities,

Top Level Document: VOP Process: 7730 02 Personnel and Responsibility, Staff and Staffing Issues, Training, **Roles and Tasks** Revision Document

Audit 20 Process Verification To Managment Viamed 24 Aug 2016

Process: 7833

Importance Of Effective Quality Management 20 Sep 2017

has responsibility and authority that includes: a) ensuring that processes needed for the quality management system are documented; b) reporting to top management on the effectiveness of the quality management system and any need for improvement; c) ensuring the promotion of awareness of applicable regulatory requirements and quality management system requirements throughout the organization. **Management** representative

ID93320
Date Revision 01 Jul 2022
Reviewed 01 Jul 2022
Top Level Document:
VM3COP02.02 VST
Company Responsibilitys
organisation chart
structure
Revision Document
ID29373
Date Revision 23 Apr 2019
Reviewed 08 Nov 2022

Top Level Document:
VM3COP02.02 Viamed
Company Responsibilitys
organisation chart
structure
Revision Document

ID27474 Date Revision 20 Sep 2018 Reviewed 08 Nov 2022

Explanation Employee
Roles and Titles
Revision Document
ID22144

Date Revision 20 Sep 2017 Reviewed 20 Sep 2017 **Audit 20 Process**

verification to Managment Revision Document

ID73324 Date Revision 26 Oct 2021 Reviewed 26 Oct 2021

VM3COP02 Organisation Responsibilities Viamed Revision Document

ID17423

Date Revision 07 Sep 2016 Reviewed 07 Sep 2016

VM3COP02 Organisation VST

Revision Document ID13954

Date Revision 19 May 2014 Reviewed 19 May 2014

5.5.3

Top management shall ensure that appropriate communication processes are established within the organization and that communication takes place regarding the effectiveness of the quality management system.

Internal communication

VM3COP27.01 Searching Intrastats Issues Revision Document ID6657

Date Revision 02 Nov 2009 Reviewed 02 Nov 2009

Intrastats overview
Revision Document

Revision Document ID23567

Date Revision 28 Oct 2017 Reviewed 28 Oct 2017

Issues Overview

Revision Document ID23112

Date Revision 22 Oct 2017 Reviewed 22 Oct 2017 **Overview Issues Meeting Headers List** Revision Document ID22169 Date Revision 22 Sep 2017 Reviewed 22 Sep 2017 Chart 42 Processes, Tasks and Audits Review **Revision Document** ID23559 Date Revision 28 Oct 2017 Reviewed 28 Oct 2017 **Chart 43 Processes and** Intrastats Revision Document ID23561 Date Revision 28 Oct 2017 Reviewed 28 Oct 2017 Chart 37 New Processes Revision Document ID23563 Date Revision 28 Oct 2017 Reviewed 28 Oct 2017

5.6 Management review

5.6.1

The organization shall document procedures for management review. Top management shall review the organization �s quality management system at documented planned intervals to ensure its continuing suitability, adequacy, and effectiveness. The review shall include assessing opportunities for improvement and the need for changes to the quality management system, including the quality policy and quality objectives. Records from management reviews shall be maintained General

Top Level Document: VOP 13 Process Monitoring, System Reviews, Audits,

Management Reviews **Analysis Data PMS Post** Market

Revision Document ID75461

Date Revision 18 Nov 2021 Reviewed 18 Nov 2021

How to Hold Intrastat Meetings

Revision Document ID8928 Date Revision 18 Oct 2011 Reviewed 18 Oct 2011

Audit 18 Management Review

Revision Document ID73320

Date Revision 26 Oct 2021 Reviewed 26 Oct 2021

Audit 10 Documentation Control

Revision Document ID63807

Date Revision 30 Jun 2021 Reviewed 30 Jun 2021

Management Review **Revision Document** ID30851

Process: 7846

ISO System Management Review Viamed 26 Sep 2017

Process: 27

Management Reviews And Quality Audits 16 Feb 2016

Process: 7070

Management Review 09 Mar 2016

Date Revision 18 Sep 2019 Reviewed 18 Sep 2019 Management reviews Revision Document ID19801 Date Revision 05 May 2017 Reviewed 05 May 2017

5.6.2

The input to management review shall include, but is not limited to, information arising from:

- a) feedback;
- b) complaint handling;
- c) reporting to regulatory authorities;
- d) audits;
- e) monitoring and

measurement of processes:

- f) monitoring and measurement of product;
- g) corrective action;
- h) preventive action;
- i) follow-up actions from previous management reviews:
- i) changes that could affect the quality management system;
- k) recommendations for improvement;
- l) applicable new or revised regulatory requirements.

General Review input

Top Level Document: VOP Process: 7743 19 Feedback Customer Complaints Vigilance and **Notifications Viamed Ltd**

Revision Document

ID75475

Date Revision 18 Nov 2021

Reviewed 18 Nov 2021

Top Level Document: VM3COP02.02 Viamed

Company Responsibilitys organisation chart

structure

Revision Document ID27474

Date Revision 20 Sep 2018 Reviewed 08 Nov 2022

Top Level Document: VOP 13 Process Monitoring. System Reviews, Audits, **Management Reviews Analysis Data PMS Post**

Market

Revision Document

ID75461

Date Revision 18 Nov 2021

Reviewed 18 Nov 2021

Chart 27 Customer Complaints Chart 27

Revision Document ID8700 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

VM3COP18 Post Market Surveilance

Revision Document ID75985

Date Revision 24 Nov 2021 Reviewed 24 Nov 2021

How to Hold Intrastat Meetings

Revision Document ID8928 Date Revision 18 Oct 2011 Reviewed 18 Oct 2011

Audit 18 Management Review

Revision Document

ID73320

Date Revision 26 Oct 2021 Reviewed 26 Oct 2021

Audit 21 Audit of Audit

Customer Complaints Paper File 26 Sep 2016

Process: 7743

Customer Complaints Paper File 26 Sep 2016

Process: 7743

Customer Complaints Paper File 26 Sep 2016

Process: 7838

Review VIAMED Feedback - Customer

Feedback Negative 23 Sep 2017

Process: 7839

Review VIAMED Feedback - Customer

Complaints 23 Sep 2017

Process: 7842

Review VIAMED Product Feedback Negative

23 Sep 2017 Process: 7846

ISO System Management Review Viamed 26

Sep 2017 Process: 7848

Review ISO Scopes 27 Sep 2017

Process: 7849

Review Product Failures New Codes 28 Sep

2017

Process: 7871

Review Exclusion From Viamed 13485:2016

And VST 9001:2015 15 Oct 2017

Process: 7837

Review External Parties Influencing The QMS

VST / Viamed 23 Sep 2017

Process: 7830

Review Q.A. Failures Report 18 Sep 2017

Process: 7741

Review Ethical Policy 14 Sep 2016

Process: 7713

Review Roles And Responsibilitys 17 Aug

2016

Process: 7070

Management Review 09 Mar 2016

Process: 6931

Customer Complaints 09 Mar 2016

Process: 7091

Calibration Index 09 Mar 2016

Process: 8014

Review VIAMED Product Feedback Positive

25 Jul 2022 Process: 8016

Review VIAMED Customer Feedback

Positive 25 Jul 2022

Revision Document ID77289 Date Revision 09 Dec 2021 Reviewed 09 Dec 2021 Audit 22 Post Market Survellance Revision Document ID120397 Date Revision 02 Jun 2023 Reviewed 02 Jun 2023 Audit 23 Analysis of Data Revision Document ID67997 Date Revision 23 Aug 2021 Reviewed 23 Aug 2021 Viamed Management **Review Blank Minutes** 20xxRevision Document ID126137 Date Revision 04 Aug 2023 Reviewed 04 Aug 2023 QC 21 Non Conformance

5.6.3

The output from management review shall be recorded (see 4.2.5) and include the input reviewed land any decisions and actions related to:

- a) improvement needed to maintain the suitability, adequacy, and effectiveness of the quality management system and its processes;
- b) improvement of product related to customer requirements;
- c) changes needed to respond to applicable new or revised regulatory requirements;
- d) resource needs. **Review** output

Top Level Document: QC Assessment Initial

Date Revision 11 Nov 2021 Reviewed 25 Nov 2022

Assessment form

Form

ID74728

Revision Document ID75549

Revision Document

Date Revision 19 Nov 2021 Reviewed 19 Nov 2021

Issues Overview

Revision Document ID23112 Date Revision 22 Oct 2017 Reviewed 22 Oct 2017

VM3COP27.01 Searching Intrastats Issues

Revision Document ID6657 Date Revision 02 Nov 2009 Reviewed 02 Nov 2009

Management Review

Revision Document ID30851

Date Revision 18 Sep 2019 Reviewed 18 Sep 2019

Management reviews

Revision Document ID19801

Date Revision 05 May 2017 Reviewed 05 May 2017

Management reviews minutes

Process: 7730

44 MHRA / CMDCAS Risk Audit 20 Process Verification To Managment Viamed 24 Aug 2016

Revision Document ID19803 Date Revision 05 May 2017 Reviewed 05 May 2017 **Audit 20 Process** verification to Managment Revision Document ID73324 Date Revision 26 Oct 2021 Reviewed 26 Oct 2021 **Audit 18 Management** Review Revision Document ID73320 Date Revision 26 Oct 2021 Reviewed 26 Oct 2021

6 Resource management

Resource management 6.1 Top Level Document: VOP Process: 7723 Audit 10b Process Verification Viamed 24 Aug The organization shall 02 Personnel and determine and provide the Responsibility , Staff and 2016 Staffing Issues, Training, resources needed to: Process: 7730 **Roles and Tasks** a) implement the quality Audit 20 Process Verification To Managment management system and to Revision Document Viamed 24 Aug 2016 maintain its effectiveness; ID93320 b) meet applicable regulatory Date Revision 01 Jul 2022 and customer requirements. Reviewed 01 Jul 2022 **Audit 20 Process** Provision of resources verification to Managment **Revision Document** ID73324 Date Revision 26 Oct 2021 Reviewed 26 Oct 2021 6.2 Top Level Document: VOP | Process: 7720 Personnel performing work 02 Personnel and Audit 08 Training Viamed 24 Aug 2016 affecting product quality Responsibility, Staff and shall be competent on the Staffing Issues, Training, basis of appropriate **Roles and Tasks** education, training, skills Revision Document and experience. ID93320 The organization shall Date Revision 01 Jul 2022 document the process(es) for Reviewed 01 Jul 2022 Top Level Document: VOP establishing competence, providing needed 12 Training training, and ensuring Revision Document awareness of personnel. ID31024 The organization shall: Date Revision 30 Sep 2019 Reviewed 30 Sep 2019 a) determine the necessary competence for personnel **Explanation Employee Roles and Titles** performing work affecting product quality; Revision Document b) provide training or take ID22144 other actions to achieve or Date Revision 20 Sep 2017

maintain the necessary competence; c) evaluate the effectiveness

of the actions taken:

d) ensure that its personnel are aware of the relevance and importance of their activities and how they contribute to the achievement of the quality objectives;

e) maintain appropriate records of education, training, skills and experience (see 4.2.5). NOTE The methodology used to check effectiveness is proportionate to the risk associated with the work for which the training or other action is being provided. **Human resources**

Reviewed 20 Sep 2017 Audit 08 Training,

Competence and Human Resources

Revision Document ID70147

Date Revision 20 Sep 2021 Reviewed 20 Sep 2021

Audit 19 Health and Safety, Working **Conditions and Building** Fabric Issues

Revision Document ID68045

Date Revision 24 Aug 2021 Reviewed 24 Aug 2021

Audit 07 Handling And Storage Viamed 24

Aug 2016 Process: 7721

Audit 09 Goods Inward And Product Identity

Viamed 24 Aug 2016

Process: 6855

Risk Assessment HSE 09 Mar 2016

Process: 6856

Fire Alarms 09 Mar 2016

Process: 54

Responsibility Allocation : Gents Toilets 17

Feb 2016 Process: 5907

Hoover Warehouse 03 Mar 2016

Process: 5908

Sweep Warehouse 03 Mar 2016

Process: 5909

Empty Warehouse Bins 03 Mar 2016

Process: 5911

Clear Cardboard 03 Mar 2016

Process: 5856

Cleaning The Kitchen 17 Feb 2016

Process: 7802

Clean Kitchen Sides 22 May 2017

Dishwashing 22 May 2017

Process: 7804

Sweep Kitchen Floor 22 May 2017

Process: 7805

Empty Kitchen Bins 22 May 2017

Process: 7806

Watering Plants 22 May 2017

Process: 56

Warehouse Outside Heating Guard 17 Feb

6.3

The organization shall document the requirements for the infrastructure needed to achieve conformity to product requirements, prevent product mix-up and ensure orderly handling of product. Infrastructure includes, as appropriate:

a) buildings, workspace and associated utilities;

b) process equipment (both hardware and software);

c) supporting services (such as transport, communication, or information systems).

The organization shall document requirements for the maintenance activities, including the interval

of performing the maintenance activities, when

such maintenance activities, or lack thereof, can affect product quality. As

appropriate, the requirements shall apply to equipment used in production, the

control of the work environment and monitoring

and measurement. Records of such maintenance Revision Document ID8713

Top Level Document: VOP Process: 7719 16 Health and Safety, **Company Personnel**

Manual

Revision Document ID31032

Date Revision 30 Sep 2019 Reviewed 30 Sep 2019

Top Level Document: VOP 18 Maintenance Building, Fabric and Infrastructure

Revision Document

ID119029

Date Revision 15 May 2023 Reviewed 15 May 2023

Top Level Document: VOP 06 Measurement Control Viamed VST, Calibration,

OA Stock

Revision Document

ID53615

Date Revision 11 Feb 2021 Reviewed 11 Feb 2021

Top Level Document: VOP 11 Equipment Control, Office, Warehouse, Pcs and Process: 7803

Equipment

Revision Document ID31008

Date Revision 30 Sep 2019 Reviewed 30 Sep 2019

DO NOT USE VM3COP11 Calibration

Date Revision 12 Oct 2011

shall be maintained Infrastructure

Reviewed 12 Oct 2011

HSE Fire / Exit Escape route Ground Floor plans Revision Document

ID127734

Date Revision 25 Aug 2023 Reviewed 25 Aug 2023

HSE Fire Exit / Escape **Route Ground Floor plans** Document

Revision Document ID2558 Date Revision 01 Aug 2007 Reviewed 01 Aug 2007

HSE Fire Risk Assessment Revision Document

ID21790

Date Revision 04 Sep 2017 Reviewed 04 Sep 2017

HSE Fire Safety Risk Assessment

Revision Document ID892 Date Revision 25 Oct 2006 Reviewed 25 Oct 2006

HSE Fire / Exit Escape route Basement floor plans

Revision Document ID127738

Date Revision 25 Aug 2023 Reviewed 25 Aug 2023

HSE Fire / Exit Escape route Ghyll House floor plans

Revision Document

ID95898

Date Revision 04 Aug 2022 Reviewed 04 Aug 2022

Ghyll House Fire Certificate

Revision Document

ID12303

Date Revision 15 Mar 2013 Reviewed 15 Mar 2013

CPM 21 Fire Exit / Escape **Route Procedures**

Revision Document ID21892

Date Revision 07 Sep 2017 Reviewed 07 Sep 2017

FIRE Report Premisis Revision Document

ID82517

Date Revision 15 Feb 2022 Reviewed 15 Feb 2022

VM3COP20.35 Ups

Calculator

Revision Document

ID88671

Date Revision 05 May 2022

 $\|2016\|$

Process: 5919

Check Out Side Drain 05 Mar 2016

Process: 5921

Clearing Water Downstairs 05 Mar 2016

Process: 7120

General Maintenance Requirements 09 Mar

2016

Process: 7742

Boiler Check 26 Sep 2016

Process: 7756

Carbon Monoxide Alarm 05 Jan 2017

Process: 7820

North Yorkshire Council Waste Tranfer 15 Jun

2017

Process: 7821

Controlled Waste Description And Transfer 15

Jun 2017

Process: 7835

Electrics Need Checking 20 Sep 2017

Process: 7836

Central Heating For Winter 20 Sep 2017

Process: 7713

Review Roles And Responsibilitys 17 Aug

2016

Process: 7845

7.1.4 Environment Of Operations 25 Sep 2017

Process: 45

Responsibility Allocation: Main Server Status

16 Feb 2016 Process: 48

Responsibility Allocation: Internet 16 Feb

2016

Process: 52

Software Verification Clear Down Backup

Emails 16 Feb 2016

Process: 5903

Responsibility Allocation: Weather Station 02

Mar 2016 Process: 5939

Responsibility Allocation: Email ISP Routing

05 Mar 2016 Process: 7121

Responsibility Allocation : General Computer

Maintenance 09 Mar 2016

Process: 7129

Intrastats Cross Reference Database Tables

Updates 09 Mar 2016

Process: 7672

Off Site Backup 09 Mar 2016

Process: 7704

Responsibility Allocation: Computer Failure

Diagnostics 24 May 2016

Process: 7850

Software Validation Scan Incorrect Product 01

Oct 2017

Process: 7851

Software Validation Scan Un-QA Product To

Reviewed 05 May 2022 VM3COP20.07 UPS

Procedures

Revision Document ID8722

Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

VM3COP03.05 Procedures Oct 2017 for customer returning

goods on our UPS account number

Revision Document

ID17155

Date Revision 05 Jul 2016 Reviewed 05 Jul 2016

Explanation Employee **Roles and Titles**

Revision Document

ID22144

Date Revision 20 Sep 2017 Reviewed 20 Sep 2017

Audit 07 Handling and

Storage

Revision Document

ID120355

Date Revision 02 Jun 2023 Reviewed 02 Jun 2023

Audit 19 Health and

Safety, Working **Conditions and Building**

Fabric Issues

Revision Document

ID68045

Date Revision 24 Aug 2021 Reviewed 24 Aug 2021

Audit 15 Production

Revision Document

ID119452

Date Revision 19 May 2023 Reviewed 19 May 2023

Order 01 Oct 2017

Process: 7852

Software Validation Expired Stock 01 Oct

2017

Process: 7853

Software Validation Non Sell Able Shelf 01

Process: 7854

Software Validation In Production List 01 Oct

2017

Process: 7855

Software Validation - Production Lists 01 Oct

2017

Process: 7856

Software Validation Unchecked Orders 01 Oct

2017

Process: 7857

Software Validation Stock Tracking Check 01

Oct 2017

Process: 7858

Software Validation Attempt To QA Some

Stock 01 Oct 2017

Process: 7861

Software Validation Of Training Documents

Forced Reading 03 Oct 2017

Process: 7832

Cleardown Emailed Invoices 20 Sep 2017

Process: 7755

Fast Hosts Invoice 08 Dec 2016

Process: 7739

Intrastats Amendment Log 12 Sep 2016

Process: 5853

Vacuuming Of The Office, Hall And Meeting

Room 17 Feb 2016

Process: 5878

Empty Office Bins 18 Feb 2016

Process: 5906

Empty Paper Bins 03 Mar 2016

Process: 5910

Clean Duckets 03 Mar 2016

Process: 7961

R D Room - Tidy, Empty Bins, Remove Cups. Caution Around Oxygen Supply 05 Oct 2020

Process: 7896

Tree In Car Park 22 Dec 2017

Process: 7864

ESD Work Stations 07 Oct 2017

Process: 46

Responsibility Allocation : Backup Server

Status 16 Feb 2016

Process: 44

Secure Socket Level Certificate 16 Feb 2016

Process: 49

Responsibility Allocation: Wifi 16 Feb 2016

Process: 50

Responsibility Allocation: Guest Access Wifi

16 Feb 2016 Process: 51

		Responsibility Allocation : Printers 16 Feb
		2016
		Process: 53
		Emails 16 Feb 2016
6.4	Top Level Document:	
Work environment and	VM3COP27.51 Incoming /	
contamination control	Goods in Contamination	
Contamination Control	Control	
	Revision Document	
	ID74855	
	Date Revision 12 Nov 2021	
	Reviewed 12 Nov 2021	
6.4.1		D 7710
II .	Top Level Document: VOP 16 Health and Safety,	Process: 7719
The organization shall document the requirements	Company Personnel	Audit 07 Handling And Storage Viamed 24
for the work environment	Manual	Aug 2016 Process: 7720
needed to achieve	Revision Document	
conformity to product	ID31032	Audit 08 Training Viamed 24 Aug 2016 Process: 7729
requirements.	Date Revision 30 Sep 2019	Audit 19 Health And Saftey Viamed 24 Aug
If the conditions for the work	II ±	2016
environment can have an		Process: 56
adverse effect on product	18 Maintenance Building,	Warehouse Outside Heating Guard 17 Feb
quality, the	Fabric and Infrastructure	2016
organization shall document	Revision Document	Process: 5919
the requirements for the	ID119029	Check Out Side Drain 05 Mar 2016
work environment and the	Date Revision 15 May 2023	Process: 5921
procedures to monitor	Reviewed 15 May 2023	Clearing Water Downstairs 05 Mar 2016
and control the work	CPM 15 Disciplinary	Process: 7120
environment.	Procedures	General Maintenance Requirements 09 Mar
The organization shall:	Revision Document	2016
a) document requirements	ID25502	Process: 7742
for health, cleanliness and	Date Revision 05 Mar 2018	Boiler Check 26 Sep 2016
clothing of personnel if	Reviewed 05 Mar 2018	Process: 7756
contact between such	CPM 16 Dress Code	Carbon Monoxide Alarm 05 Jan 2017
personnel and the product or	Revision Document ID7055	Process: 7820
work environment could	Date Revision 26 Apr 2010	North Yorkshire Council Waste Tranfer 15 Jun
affect medical device safety	Reviewed 22 Jul 2014	2017
or performance;		Process: 7821
b) ensure that all personnel	Policy Viamed	Controlled Waste Description And Transfer 15
who are required to work	Revision Document	Jun 2017
temporarily under special	ID14332	Process: 7835
environmental	Date Revision 25 Sep 2014	Electrics Need Checking 20 Sep 2017
conditions within the work	Reviewed 04 Sep 2017	Process: 7836
environment are competent	CPM 39 Smoking Policy	Central Heating For Winter 20 Sep 2017
or supervised by a competent		Process: 7864
person.	Date Revision 15 Feb 2010	ESD Work Stations 07 Oct 2017
NOTE Further information	Reviewed 15 Feb 2010	Process: 7873
can be found in ISO 14644	Audit 07 Handling and	On Site Environment Review 18 Oct 2017
and ISO 14698 Work	Storage Revision Document	Process: 54
environment	Revision Document ID120355	Responsibility Allocation : Gents Toilets 17 Feb 2016
	Date Revision 02 Jun 2023	Process: 5906
	Reviewed 02 Jun 2023	Empty Paper Bins 03 Mar 2016
	Audit 08 Training,	Process: 5907
	Competence and Human	Hoover Warehouse 03 Mar 2016
	Resources	Process: 5908
	Revision Document	Sweep Warehouse 03 Mar 2016
II		15p

ID70147
Date Revision 20 Sep 2021
Reviewed 20 Sep 2021
Audit 19 Health and
Safety, Working
Conditions and Building
Fabric Issues
Revision Document
ID68045
Date Revision 24 Aug 2021

Process: 5909
Empty Warehouse Bins 03 Mar 2016

Process: 5910

Clean Duckets 03 Mar 2016

Process: 5911

Clear Cardboard 03 Mar 2016

Process: 7698

Clean Toilets 17 May 2016

6.4.2

As appropriate, the organization shall plan and document arrangements for the control of contaminated or potentially contaminated product in order to prevent contamination of the work environment. personnel, or product. For sterile medical devices, the organization shall document requirements for control of contamination with microorganisms or particulate matter and maintain the required cleanliness during assembly or packaging processes.

Contamination control

Top Level Document: VM3COP02.01 Exclusions to Viamed ISO13485:2016 boundaries of ISO

Reviewed 24 Aug 2021

Revision Document ID74571

Date Revision 10 Nov 2021 Reviewed 01 Aug 2023

Top Level Document: VOP 20 Goods in Purchases, Returns, Repairs, Inspection / Rejection

Revision Document ID75943

Date Revision 24 Nov 2021 Reviewed 24 Nov 2021

Top Level Document: VOP 09 Repairs and Servicing

Revision Document ID75927

Date Revision 24 Nov 2021 Reviewed 24 Nov 2021

Top Level Document: VOP 05 Supplier Control, Supplier Review, Purchase Orders, Supplier Returns and Rejection

Revision Document ID75847

Date Revision 23 Nov 2021 Reviewed 23 Nov 2021

Top Level Document: VM3COP27.51 Incoming / Goods in Contamination Control

Revision Document ID74855

Date Revision 12 Nov 2021 Reviewed 12 Nov 2021

Audit 09 Goods Inward and Product Identity

Revision Document ID55437

Date Revision 12 Mar 2021 Reviewed 09 Aug 2023

Process: 39

Environmental Policy Document Review 16 Feb

2016

Process: 7719

Audit 07 Handling And Storage Viamed 24 Aug 2016

Process: 7714

Audit 01 Picking Packing Viamed 24 Aug

2016

Process: 7721

Audit 09 Goods Inward And Product Identity

Viamed 24 Aug 2016

7 Product realization

Product realization 7.1 Top Level Document: VOP | Process: 7732 Audit 22 Post Market Survellance Viamed 24 08 Production, Reworks, The organization shall plan and develop the processes New Production Aug 2016 needed for product Revision Document Process: 7716 Audit 03 Design Control Viamed 24 Aug 2016 realization. Planning of ID31072 product realization shall be Date Revision 30 Sep 2019 Reviewed 30 Sep 2019 consistent with the requirements of the other Top Level Document: processes of the quality VM3COP27.11 Performing management system. a Technical File PMS and The organization shall risk assessment document one or more Revision Document processes for risk ID75465 management in product Date Revision 18 Nov 2021 realization. Reviewed 18 Nov 2021 Records of risk management VM3COP24.00 Viamed activities shall be maintained **Overall Risk Analysis** (see 4.2.5). Program Risk Register Revision Document In planning product ID47771 realization, the organization Date Revision 12 Nov 2020 shall determine the Reviewed 12 Nov 2020 following, as appropriate: VM3COP27.12 Clinical a) quality objectives and **Evaluation Risk** requirements for the product; assessment Technical Files b) the need to establish Revision Document processes and documents ID15453 (see 4.2.4) and to provide Date Revision 11 Aug 2015 resources specific to the Reviewed 11 Aug 2015 product, including Audit 22 Post Market infrastructure and work Survellance environment; Revision Document c) required verification, ID120397 validation, monitoring, Date Revision 02 Jun 2023 measurement, inspection and Reviewed 02 Jun 2023 test, handling, Audit 03 Design Control storage, distribution and Revision Document traceability activities specific ID111315 to the product together with Date Revision 17 Feb 2023 the criteria Reviewed 17 Feb 2023 for product acceptance; Audit 07 Handling and d) records needed to provide Storage **Revision Document** evidence that the realization processes and resulting ID120355 product meet Date Revision 02 Jun 2023 requirements (see 4.2.5). Reviewed 02 Jun 2023 The output of this planning Audit 23 Analysis of Data shall be documented in a Revision Document form suitable for the ID67997 organization s method of Date Revision 23 Aug 2021 Reviewed 23 Aug 2021 operations.

Audit 09 Goods Inward

2/10/2023, 11:03	QM3 Route Map	Viameu Ltu 15015465:2016
NOTE Further information	and Product Identity	
can be found in ISO 14971.	Revision Document	
Planning of product	ID55437	
realization	Date Revision 12 Mar 2021	
	Reviewed 09 Aug 2023	
	Audit 10 Documentation	
	Control	
	II I	
	Revision Document ID63807	
	II I	
	Date Revision 30 Jun 2021	
	Reviewed 30 Jun 2021	
7.2		
Customer-related processes		
	T II D	D 7722
7.2.1	Top Level Document:	Process: 7732
The organization shall	VM3COP03.07	Audit 22 Post Market Survellance Viamed 24
determine:	Humanmed Order	Aug 2016
a) requirements specified by	Checking	Process: 7715
the customer, including the	Revision Document	Audit 02 Contract Review Viamed 24 Aug
requirements for delivery	ID22266	2016
and postdelivery activities;	Date Revision 27 Sep 2017	Process: 7825
b) requirements not stated by	Reviewed 27 Sep 2017	Responsibility Allocation : Order Picking 06
the customer but necessary	Top Level Document:	Sep 2017
for specified or intended use,	VM3COP03.08	Process: 5
as known;	Humanmed Order	Responsibility Allocation: Processing Of Sales
c) applicable regulatory	Processing	Orders 16 Feb 2016
requirements related to the	Revision Document	Process: 7825
product;	ID24775	Responsibility Allocation : Order Picking 06
d) any user training needed	Date Revision 22 Dec 2017	Sep 2017
to ensure specified	Reviewed 22 Dec 2017	Process: 7825
performance and safe use of	Top Level Document:	Responsibility Allocation : Order Picking 06
the medical device;	VM3COP12.01 Viamed	Sep 2017
e) any additional	Policy on End User	Process: 7
requirements determined by	Training UK	Responsibility Allocation : Checking Of Sales
the organization	Revision Document	Orders 16 Feb 2016
Determination of	ID85827	Process: 7734
II.	Date Revision 29 Mar 2022	
requirements related to		Responsibility Allocation : Humanmed Order
product	Reviewed 29 Mar 2022	Processing 25 Aug 2016
	Top Level Document: VOP	
	03 Contract Review,	Responsibility Allocation : Processing Of Sales
	Enquires, Office Processes	Orders 16 Feb 2016
	Revision Document	Process: 7734
	ID77875	Responsibility Allocation : Humanmed Order
	Date Revision 15 Dec 2021	Processing 25 Aug 2016
	Reviewed 15 Dec 2021	Process: 7825
	Audit 22 Post Market	Responsibility Allocation : Order Picking 06
	Survellance	Sep 2017
	Revision Document	
	ID120397	
	Date Revision 02 Jun 2023	
	Reviewed 02 Jun 2023	
	Audit 02 Contract Review	
	and Sales Order Processing	
	Revision Document	
	ID69328	
	Date Revision 09 Sep 2021	
	Reviewed 09 Sep 2021	
	VM3COP20.31 Export	
		1

Order Processing

Revision Document

ID114755

Date Revision 28 Mar 2023

Reviewed 28 Mar 2023

VM3COP03.01 Order

Processing Priorities

Revision Document

ID20049

Date Revision 15 May 2017

Reviewed 15 May 2017

VM3COP20.30 UK Order

Processing

Revision Document

ID117589

Date Revision 28 Apr 2023

Reviewed 28 Apr 2023

VM3COP20.32 Order

Checking

Revision Document

ID34889

Date Revision 01 Apr 2020

Reviewed 01 Apr 2020

Infant Resuscitation

Cabinet - Training

Assessment Form

Revision Document

ID14334

Date Revision 25 Sep 2014

Reviewed 25 Sep 2014

Oxygen Sensor Training Powerpoint

Revision Document

ID15736

Date Revision 24 Sep 2015

Reviewed 25 Oct 2016

Oxygen Sensor Training Video

Revision Document

ID15737

Date Revision 24 Sep 2015

Reviewed 24 Sep 2015

Resuscitation Unit and

TC400 Training

Information Resuscitation

Cabinet Training

Revision Document ID4111

Date Revision 09 Jul 2008

Reviewed 09 Jul 2008

Resuscitation Unit

Maintenance Therapy

Equipment Suction

Controller Unit and TC400

Training Information

Therapy Workshop Inst.

Revision Document ID4122 Date Revision 09 Jul 2008

Reviewed 09 Jul 2008

Single Use Surgical Training Information certificates

Revision Document ID20220

Date Revision 19 May 2017 Reviewed 19 May 2017

SpO2 800 series Training Information

Revision Document

ID12687

Date Revision 02 Jul 2013 Reviewed 02 Jul 2013

TECcare Training Material

Revision Document ID11826

Date Revision 11 Jun 2012

Reviewed 11 Jun 2012

Temperature Probe

Training Material

Revision Document

ID18169

Date Revision 05 Dec 2016 Reviewed 05 Dec 2016

Tom Thumb Training

Information

Revision Document ID7880

Date Revision 07 Mar 2011

Reviewed 07 Mar 2011

Tom Thumb Training **Information 2009**

Revision Document

ID15644

Date Revision 16 Sep 2015

Reviewed 16 Sep 2015

Tom Thumb Training

Information Training

Manual Training

Information

Revision Document ID2973

Date Revision 31 Jan 2008

Reviewed 31 Jan 2008

Tom Thumb Training

Information Training V1.1

Revision Document

ID15641

Date Revision 16 Sep 2015

Reviewed 16 Sep 2015

Training information **Infant Resusitation Unit**

Revision Document ID8665

Date Revision 12 Oct 2011

Reviewed 12 Oct 2011

VM-2500 Product Training Materials - Frequently

Asked Questions

Revision Document ID6967

Date Revision 17 Mar 2010

Reviewed 17 Mar 2010

VM-2500 Product Training Materials Capnography **Product Application Notes**

Revision Document ID6749 Date Revision 08 Feb 2010 Reviewed 08 Feb 2010

VM-2500 Product Training Materials Capnography **Product Presentation** MASTER

Revision Document ID6750 Date Revision 08 Feb 2010 Reviewed 08 Feb 2010

VM-2500 Product Training Materials Mainstream or Sidestream Capnography

Revision Document ID6753 Date Revision 08 Feb 2010 Reviewed 08 Feb 2010

Audit 01 Picking packing

Revision Document

ID122441

Date Revision 26 Jun 2023 Reviewed 26 Jun 2023

Audit 16 Sales and Marketing

Revision Document ID69457

Date Revision 10 Sep 2021 Reviewed 10 Sep 2021

7.2.2

The organization shall review the requirements related to product. This review shall be conducted prior to the organization �s commitment to supply product to the customer (e.g. submission of tenders, acceptance of contracts or orders, acceptance of changes to contracts or orders) and shall ensure that: a) product requirements are defined and documented; b) contract or order requirements differing from those previously expressed

are resolved; c) applicable regulatory requirements are met; d) any user training identified in accordance with 7.2.1 is available or planned to be available:

e) the organization has the

Top Level Document: VOP Process: 7715 03 Contract Review, **Enquires, Office Processes**

Revision Document

ID77875

Date Revision 15 Dec 2021 Reviewed 15 Dec 2021

Audit 02 Contract Review and Sales Order Processing 2016

Revision Document ID69328

Date Revision 09 Sep 2021 Reviewed 09 Sep 2021

Audit 11 Repairs, Servicing and Returns

Revision Document ID124549

Date Revision 19 Jul 2023 Reviewed 19 Jul 2023

Audit 20 Process verification to Managment

Revision Document ID73324

Date Revision 26 Oct 2021 Reviewed 26 Oct 2021

Audit 10 Documentation

Audit 02 Contract Review Viamed 24 Aug 2016

Process: 7724

Audit 11 Repairs And Service Viamed 24 Aug

Process: 7723

Audit 10b Process Verification Viamed 24 Aug

Process: 7722

Audit 10 Documentation Control Viamed 24

Aug 2016 Process: 5871

Check Sale Or Returns 17 Feb 2016

Process: 5872

Check Sale Or Returns Export 17 Feb 2016

ability to meet the defined requirements. Records of the results of the review and actions arising from the review shall be maintained (see 4.2.5). When the customer provides no documented statement of requirement, the customer requirements shall be confirmed by the organization before acceptance. When product requirements

are changed, the organization shall ensure that relevant

amended and that relevant personnel are made aware of the changed requirements. **Review of requirements**

documents are

related to product

Control Revision Document ID63807 Date Revision 30 Jun 2021 Reviewed 30 Jun 2021

7.2.3

The organization shall plan and document arrangements for communicating with customers in relation to:

- a) product information;
- b) enquiries, contracts or order handling, including amendments;
- c) customer feedback, including complaints; d) advisory notices. The organization shall communicate with regulatory authorities in accordance with applicable regulatory requirements.

Communication

Top Level Document: VOP | Process: 2 03 Contract Review, **Enquires, Office Processes**

Revision Document ID77875

Date Revision 15 Dec 2021 Reviewed 15 Dec 2021

Top Level Document: VOP 19 Feedback Customer Complaints Vigilance and **Notifications Viamed Ltd**

Revision Document ID75475

Date Revision 18 Nov 2021 Reviewed 18 Nov 2021

VM3COP27.31 Processing Proforma Invoices and **Quotations**

Revision Document ID69812

Date Revision 15 Sep 2021 Reviewed 15 Sep 2021

VM3COP20.05 New Orders - How to enter into Opera Viamed

Revision Document ID13695

Date Revision 12 May 2014 Reviewed 12 May 2014

VM3COP20.32 Order

Checking

Revision Document ID34889

Date Revision 01 Apr 2020 Reviewed 01 Apr 2020

Answering Telephones 16 Feb 2016

Process: 7710

Responsibility Allocation: Proforma And

Quote Processing 29 Jun 2016

Process: 7825

Responsibility Allocation : Order Picking 06

Sep 2017 Process: 7743

Customer Complaints Paper File 26 Sep 2016

Process: 7743

Customer Complaints Paper File 26 Sep 2016

Process: 7726

Audit 14 Complaints And Corrective Actions

Viamed 24 Aug 2016

Process: 7715

Audit 02 Contract Review Viamed 24 Aug

2016

Process: 5943

Check Cardea And Multiquote 08 Mar 2016

Process: 7678

Check Catalog 360 Circle For Quotes And

Orders 08 Apr 2016

Process: 7758

Check For GHX Orders 17 Jan 2017

Process: 7760

Send Service Offers 31 Jan 2017

Process: 7670

Humanmed general Issues 09 Mar 2016

Process: 7782

Remove Started But Not Used Order Numbers

08 Feb 2017 Process: 7797

Check Order Are Being Picked In Priority

Order 10 May 2017

VM3COP20.49 Informing | Process: 7798 **Customers of Price** Amends

Revision Document

ID18357

Date Revision 05 Jan 2017 Reviewed 05 Jan 2017

VM3COP20.031 Viamed Repair Procedures Invoicing / customer

paperwork

Revision Document

ID24753

Date Revision 21 Dec 2017 Reviewed 21 Dec 2017 VM3COP20.22 Quoting

Customer Special prices.

Revision Document

ID15613

Date Revision 09 Sep 2015 Reviewed 09 Sep 2015

VM3COP10.02 Product Recall locate products out in the Field

Revision Document

ID74788

Date Revision 12 Nov 2021 Reviewed 12 Nov 2021

Audit 14 Complaints and **Corrective Actions** Revision Document

ID76091

Date Revision 25 Nov 2021 Reviewed 25 Nov 2021

Audit 02 Contract Review

Revision Document

ID69328

Date Revision 09 Sep 2021 Reviewed 09 Sep 2021

Audit 16 Sales and Marketing

Revision Document

ID69457

Date Revision 10 Sep 2021 Reviewed 10 Sep 2021

Audit 22 Post Market

Survellance

Revision Document

ID120397

Date Revision 02 Jun 2023 Reviewed 02 Jun 2023

Audit 01 Picking packing Revision Document

ID122441

Date Revision 26 Jun 2023 Reviewed 26 Jun 2023 Audit 04 Accounts and

Orders And Items Shipped Per Month 10 May

2017

Process: 7957

Warehouse Requests 29 May 2020

Process: 6959

Responsibility Allocation: Sales Forward

Orders Review 09 Mar 2016

Process: 6921

Responsibility Allocation: Customer pricing

agreements 09 Mar 2016

Process: 5876

E.Commerce Cardea And Multiquote 17 Feb

2016

Process: 7748

Check Repair Orders 10 Oct 2016

Process: 7860

Goods Out Picking 03 Oct 2017

Process: 5

Responsibility Allocation: Processing Of Sales

Orders 16 Feb 2016

Process: 6

Responsibility Allocation : Updating Contact

Management System 16 Feb 2016

Process: 7

Responsibility Allocation : Checking Of Sales

Orders 16 Feb 2016

Process: 8

Responsibility Allocation : Order And Status

Liaison With Customers 16 Feb 2016

Process: 9

Distribution Of Faxes 16 Feb 2016

Process: 10

Distribution Of Emails 16 Feb 2016

Process: 11

and Sales Order Processing Distribution Of Post 16 Feb 2016

Process: 12

Responsibility Allocation: Sales And

Technical Information Processing 16 Feb 2016

Process: 36

Emailing Of Invoices 16 Feb 2016

Process: 5850

Purchase Order Log 17 Feb 2016

Process: 5875

Check Paypal For Orders 17 Feb 2016

Process: 5857

Customer Service Logs 17 Feb 2016

Process: 5891

Processing Of Repair Quotes And Orders 25

Feb 2016

Process: 5892

Checking EBay And Amazon For Orders And

Messages 25 Feb 2016

Process: 5893

Answering Website Questions 25 Feb 2016

Process: 5899

Proforma And Quote Chasing 25 Feb 2016

Process: 5901

	Finance Revision Document ID63821 Date Revision 30 Jun 2021 Reviewed 30 Jun 2021	Link Call Log Contacts To The CRM 02 Mar 2016 Process: 5913 Check For Humanmed Orders In Logistics Mailbox 03 Mar 2016 Process: 6958 Responsibility Allocation: Shipped Order Queries 09 Mar 2016 Process: 7686 Thorough Checking Of Awaiting Action Tray-Priority 8s 21 Apr 2016 Process: 7734 Responsibility Allocation: Humanmed Order Processing 25 Aug 2016 Process: 7735 Ensure SOR's Are Followed Up 01 Sep 2016 Process: 7792 Shipped Order Success Report 13 Mar 2017
7.3 Design and development		
7.3.1	Top Level Document: VOP	Process: 7716
The organization shall document procedures for design and development General	17 Design Research and Development Revision Document ID25632 Date Revision 19 Mar 2018 Reviewed 19 Mar 2018 Revision Document ID111315 Date Revision 17 Feb 2023 Reviewed 17 Feb 2023 Reviewed 17 Feb 2023 Reviewed 17 Feb 2023 Reviewed 17 Feb 2023 Revision Document ID73324 Date Revision 26 Oct 2021 Revision Document ID73324 Date Revision 26 Oct 2021 Reviewed 26 Oct 2021 Reviewed 26 Oct 2021 Reviewed 27 Pec 2008 Revision Document ID4959 Date Revision 29 Dec 2008 Reviewed 29 Dec 2008 CE & Design files reorganisation Revision Document ID9085 Date Revision 18 Oct 2011 Reviewed 18 Oct 2011 Chart 04 Design and Development Revision Document ID8678 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011 Chart 17 Design Repairs Revision Document ID8690 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011	Audit 03 Design Control Viamed 24 Aug 2016 Process: 7723 Audit 10b Process Verification Viamed 24 Aug 2016

Chart 30 System Design Plan

Revision Document ID8703 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

New Project Design File Content

Revision Document ID9093 Date Revision 18 Oct 2011 Reviewed 18 Oct 2011

VM3COP16 Design and Design Changes Design requirements

Revision Document ID7396 Date Revision 10 Jan 2011 Reviewed 10 Jan 2011

Audit 12 CE Files

Revision Document ID63815

Date Revision 30 Jun 2021 Reviewed 30 Jun 2021

7.3.2

The organization shall plan and control the design and development of product. As appropriate,

design and development planning documents shall be maintained and updated as the design and development progresses.

During design and development planning, the

organization shall document:

- a) the design and development stages;
- b) the review(s) needed at each design and development stage;
- c) the verification,
 validation, and design
 transfer activities that are
 appropriate at each design
 and
- development stage;
- d) the responsibilities and authorities for design and development;
- e) the methods to ensure traceability of design and development outputs to design and development inputs; f) the resources needed
- including necessary competence of personnel

Top Level Document:

VM3COP27.11 Performing a Technical File PMS and risk assessment

Revision Document ID75465

Date Revision 18 Nov 2021 Reviewed 18 Nov 2021

Top Level Document: VOP 17 Design Research and Development

Revision Document ID25632

Date Revision 19 Mar 2018 Reviewed 19 Mar 2018

Top Level Document: VOP 02 Personnel and Responsibility, Staff and Staffing Issues, Training, Roles and Tasks

Revision Document ID93320

Date Revision 01 Jul 2022 Reviewed 01 Jul 2022

VM3COP16 Design and Design Changes Design requirements

Revision Document ID7396 Date Revision 10 Jan 2011 Reviewed 10 Jan 2011

VM3COP27.07 Project Manager

Revision Document

ID12734 Date Revision 11 Jul 2013

Reviewed 11 Jul 2013

Process: 7716

VM3COP27.11 Performing Audit 03 Design Control Viamed 24 Aug 2016

Process: 7723

Audit 10b Process Verification Viamed 24 Aug

2016

Process: 7720

Audit 08 Training Viamed 24 Aug 2016

Design and development planning

VM3COP27.12 Clinical **Evaluation Risk** assessment Technical Files

Revision Document ID15453

Date Revision 11 Aug 2015 Reviewed 11 Aug 2015

Audit 03 Design Control

Revision Document ID111315

Date Revision 17 Feb 2023 Reviewed 17 Feb 2023

Audit 20 Process

verification to Managment

Revision Document ID73324

Date Revision 26 Oct 2021 Reviewed 26 Oct 2021

Audit 08 Training,

Competence and Human

Resources

Revision Document

ID70147

Date Revision 20 Sep 2021

Reviewed 20 Sep 2021

Audit 12 CE Files

Revision Document

ID63815

Date Revision 30 Jun 2021

Reviewed 30 Jun 2021

QC 28B Design Changes

Revision Document

ID25508

Date Revision 05 Mar 2018 Reviewed 05 Mar 2018

Generic CE File Attached to All Assignment of responsibility Risk Management

Revision Document ID7742 Date Revision 02 Mar 2011 Reviewed 02 Mar 2011

7.3.3

Inputs relating to product requirements shall be determined and records maintained (see 4.2.5). These ID25632 inputs shall include:

- a) functional, performance, usability and safety requirements, according to the intended use;
- b) applicable regulatory requirements and standards;
- risk management;

c) applicable output(s) of d) as appropriate,

Top Level Document: VOP Process: 7716 17 Design Research and

Development

Revision Document

Date Revision 19 Mar 2018 Reviewed 19 Mar 2018

Audit 03 Design Control

Revision Document ID111315

Date Revision 17 Feb 2023 Reviewed 17 Feb 2023

Audit 20 Process verification to Managment Revision Document

Audit 03 Design Control Viamed 24 Aug 2016

Process: 7722

Audit 10 Documentation Control Viamed 24

Aug 2016

Process: 7723

Audit 10b Process Verification Viamed 24 Aug

2016

information derived from previous similar designs; e) other requirements essential for design and development of the product and processes. These inputs shall be reviewed for adequacy and approved. Requirements shall be complete, unambiguous, able to be verified or validated, and not in conflict with each other. NOTE Further information can be found in IEC 62366 1.	ID73324 Date Revision 26 Oct 2021 Reviewed 26 Oct 2021 Audit 12 CE Files Revision Document ID63815 Date Revision 30 Jun 2021 Reviewed 30 Jun 2021 Audit 23 Analysis of Data Revision Document ID67997 Date Revision 23 Aug 2021 Reviewed 23 Aug 2021	
Design and development inputs		
7.3.4 Design and development outputs shall: a) meet the input requirements for design and development; b) provide appropriate information for purchasing, production and service provision; c) contain or reference product acceptance criteria; d) specify the characteristics of the product that are essential for its safe and proper use. The outputs of design and development shall be in a form suitable for verification against the design and development inputs and shall be approved prior to release. Records of the design and development outputs shall be maintained (see 4.2.5). Design and development outputs	Top Level Document: VOP 17 Design Research and Development Revision Document ID25632 Date Revision 19 Mar 2018 Reviewed 19 Mar 2018 Audit 03 Design Control Revision Document ID111315 Date Revision 17 Feb 2023 Reviewed 17 Feb 2023 Audit 23 Analysis of Data Revision Document ID67997 Date Revision 23 Aug 2021 Reviewed 23 Aug 2021 Reviewed 23 Aug 2021 Audit 12 CE Files Revision Document ID63815 Date Revision 30 Jun 2021 Reviewed 30 Jun 2021	Process: 7716 Audit 03 Design Control Viamed 24 Aug 2016
7.3.5 Design and development review	Audit 12 CE Files Revision Document ID63815 Date Revision 30 Jun 2021	
	Reviewed 30 Jun 2021	
7.3.5 At suitable stages, systematic reviews of design	Top Level Document: VOP 17 Design Research and Development	Process: 7716 Audit 03 Design Control Viamed 24 Aug 2016

and development shall be performed in accordance with planned and documented arrangements

a) evaluate the ability of the results of design and development to meet

requirements; b) identify and propose necessary actions. Participants in such reviews shall include representatives of functions concerned with the design and development stage being reviewed, as well as other specialist personnel. Records of the results of the reviews and any necessary actions shall be maintained and include the identification of the design under review, the participants involved and the date of the review (see 4.2.5).

Revision Document ID25632

Date Revision 19 Mar 2018 Reviewed 19 Mar 2018

Audit 03 Design Control

Revision Document ID111315

Date Revision 17 Feb 2023 Reviewed 17 Feb 2023

Audit 12 CE Files

Revision Document

ID63815

Date Revision 30 Jun 2021 Reviewed 30 Jun 2021

7.3.6 Design and development verification shall be performed in accordance with planned and documented arrangements to ensure that the design and development outputs have met the design and development input requirements. The organization shall document verification plans that include methods, acceptance criteria and, as appropriate, statistical techniques with rationale for sample size. If the intended use requires that the medical device be connected to, or have an interface with, other medical device(s), verification shall include confirmation that the design outputs meet design inputs when so connected or interfaced. Records of the results and

conclusions of the

Top Level Document: VOP 17 Design Research and Development

Revision Document ID25632

Date Revision 19 Mar 2018 Reviewed 19 Mar 2018

Top Level Document: VOP 15 Data and Information Analysis

Revision Document ID98547

Date Revision 07 Sep 2022 Reviewed 07 Sep 2022

Audit 03 Design Control Revision Document

ID111315

Date Revision 17 Feb 2023 Reviewed 17 Feb 2023

Audit 12 CE Files

Revision Document ID63815

Date Revision 30 Jun 2021 Reviewed 30 Jun 2021

., 10, 1013, 11.03	Q. 15 Noute Hap	
verification and necessary		
actions shall be maintained		
(see 4.2.4 and 4.2.5). Design		
and development		
verification		
7.3.7	Audit 12 CE Files	
Design and development	Revision Document	
validation	ID63815	
	Date Revision 30 Jun 2021	
	Reviewed 30 Jun 2021	
	QC 30b Project	
	Verification & Validation	
	Summary Master	
	Revision Document	
	ID25482	
	Date Revision 01 Mar 2018	
	Reviewed 01 Mar 2018	
7.3.7	Top Level Document: VOP	Process: 7716
Design and development	17 Design Research and	Audit 03 Design Control Viamed 24 Aug 2016
validation shall be performed	11	Process: 7723
in accordance with planned	Revision Document	Audit 10b Process Verification Viamed 24 Aug
and documented	ID25632	2016
arrangements to ensure that	Date Revision 19 Mar 2018	
the resulting product is	Reviewed 19 Mar 2018	
capable of meeting the	Top Level Document: VOP	
requirements for the	15 Data and Information	
specified application or	Analysis	
intended use.	Revision Document	
The organization shall	ID98547	
document validation plans	Date Revision 07 Sep 2022	
that include methods,	Reviewed 07 Sep 2022	
acceptance criteria, and, as	Audit 03 Design Control	
appropriate, statistical techniques with rationale for	Revision Document	
sample size.	Date Revision 17 Feb 2023	
Design validation shall be	Reviewed 17 Feb 2023	
conducted on representative	Audit 12 CE Files	
product. Representative	Revision Document	
product includes	ID63815	
initial production units,	Date Revision 30 Jun 2021	
batches or their equivalents.	Reviewed 30 Jun 2021	
The rationale for the choice		
of product used for		
validation shall be recorded		
(see 4.2.5).		
As part of design and		
development validation, the		
organization shall perform		
clinical evaluations or		
performance evaluations of		
the medical device in		
accordance with applicable		
regulatory requirements. A medical device used for		
clinical evaluation or		
performance evaluation is		
not considered to be released		
line considered to be released	II	II

for use to the customer. If the intended use requires that the medical device be connected to, or have an interface with, other medical device(s), validation shall include confirmation that the requirements for the specified application or intended use have been met when so connected or interfaced. Validation shall be completed prior to release for use of the product to the customer. Records of the results and conclusion of validation and necessary actions shall be maintained (see 4.2.4 and 4.2.5).

Top Level Document: VOP 17 Design Research and

Development

Revision Document ID25632

Date Revision 19 Mar 2018 Reviewed 19 Mar 2018

Audit 03 Design Control **Revision Document**

as suitable for manufacturing ID111315

Date Revision 17 Feb 2023 Reviewed 17 Feb 2023

Audit 12 CE Files Revision Document

ID63815

Date Revision 30 Jun 2021 Reviewed 30 Jun 2021

Process: 7716

Audit 03 Design Control Viamed 24 Aug 2016

Process: 7722

Audit 10 Documentation Control Viamed 24

Aug 2016

7.3.9

7.3.8

The organization shall

transfer of design and

manufacturing. These

outputs are verified

and that production

requirements.

before becoming final production specifications

development outputs to

design and development

procedures shall ensure that

capability can meet product

Results and conclusions of

(see 4.2.5). **Design and** development transfer

the transfer shall be recorded

document procedures for

The organization shall document procedures to control design and development changes. The organization shall determine the significance of the change to function, performance, usability, safety and applicable regulatory requirements for the medical device and its intended use. Design and development changes shall be identified. Before implementation, the changes shall be:

Top Level Document: VOP 17 Design Research and Development

Revision Document ID25632

Date Revision 19 Mar 2018 Reviewed 19 Mar 2018

Audit 03 Design Control

Revision Document ID111315

Date Revision 17 Feb 2023 Reviewed 17 Feb 2023

Audit 12 CE Files Revision Document

ID63815 Date Revision 30 Jun 2021 Reviewed 30 Jun 2021

Process: 7716

Audit 03 Design Control Viamed 24 Aug 2016

Process: 7726

Audit 14 Complaints And Corrective Actions

Viamed 24 Aug 2016

a) reviewed;

b) verified;

c) validated, as appropriate;

d) approved.

The review of design and development changes shall include evaluation of the effect of the changes on constituent parts and product in process or already delivered, inputs or outputs of risk management and product realization processes. Records of changes, their

review and any necessary actions shall be maintained (see 4.2.5). **Control of** design and development

QC 28B Design Changes Revision Document

ID25508

Date Revision 05 Mar 2018 Reviewed 05 Mar 2018

7.3.10

changes

The organization shall maintain a design and development file for each medical device type or medical device family. This file shall include or reference records generated to demonstrate conformity to the requirements for design and development and records for design and development changes. **Design and** development files

Audit 03 Design Control

Revision Document ID111315

Date Revision 17 Feb 2023 Reviewed 17 Feb 2023

Audit 12 CE Files

Revision Document ID63815

Date Revision 30 Jun 2021 Reviewed 30 Jun 2021

Process: 7722

Audit 10 Documentation Control Viamed 24 Aug 2016

Process: 7716

Audit 03 Design Control Viamed 24 Aug 2016

7.4

Purchasing

DO NOT USE VM3COP04 Process: 5850

Purchasing / suppliers

Revision Document ID15473

Date Revision 14 Aug 2015 Reviewed 14 Aug 2015

VM3COP20.29 Checking the Purchase Order Log

Revision Document

ID73132 Date Revision 25 Oct 2021

Reviewed 25 Oct 2021

VM3COP27.34 Sending **Purchase Orders to**

Revision Document ID17070

Suppliers

Date Revision 22 Jun 2016 Reviewed 22 Jun 2016

VM3COP04.01 QC06 Supplier Questionnaire ISO Questionnaire Viamed

Purchase Order Log 17 Feb 2016

Process: 7707

Send Purchase Orders To Suppliers 13 Jun

2016

BlankRevision Document
ID21304
Date Revision 06 Aug 2017
Reviewed 06 Aug 2017

7.4.1

lbe:

The organization shall document procedures (see 4.2.4) to ensure that purchased product conforms to specified purchasing information.

The organization shall establish criteria for the evaluation and selection of suppliers. The criteria shall

- a) based on the supplier sability to provide product that meets the organizations requirements;
- b) based on the performance of the supplier;
- c) based on the effect of the purchased product on the quality of the medical device;
- d) proportionate to the risk associated with the medical device.

The organization shall plan the monitoring and re-evaluation of suppliers. Supplier performance in meeting requirements for the purchased product shall be monitored. The results of the monitoring shall provide an input into

shall provide an input into the supplier re-evaluation process.

Non-fulfilment of purchasing requirements shall be addressed with the supplier proportionate to the risk associated with the purchased product and compliance with applicable regulatory requirements. Records of the results of evaluation, selection, monitoring and re-evaluation of supplier capability or performance and any necessary actions arising

Top Level Document: VOP 05 Supplier Control, Supplier Review, Purchase

Supplier Review, Purchase Orders, Supplier Returns and Rejection

Revision Document ID75847

Date Revision 23 Nov 2021 Reviewed 23 Nov 2021

Top Level Document: VOP 20 Goods in Purchases, Returns, Repairs, Inspection / Rejection

Revision Document ID75943

Date Revision 24 Nov 2021 Reviewed 24 Nov 2021

Top Level Document: VOP 21 Risk, Risk Management and Risk Analysis

Revision Document ID75935

Date Revision 24 Nov 2021 Reviewed 24 Nov 2021

Audit 05 Purchasing suppliers

Revision Document ID69314

Date Revision 09 Sep 2021 Reviewed 09 Sep 2021

Audit 09 Goods Inward and Product Identity

Revision Document ID55437

Date Revision 12 Mar 2021 Reviewed 09 Aug 2023 Process: 7717

Audit 05 Purchasing Suppliers Viamed 24 Aug

2016

Process: 7725

Audit 12 CE Files Viamed 24 Aug 2016

Process: 5855

Purchase Order Requirements Teledyne 17 Feb

2016

Process: 8030

Purchase Order Invoice Review 23 Jun 2023

02/10/2023. 11:03 QMS Route Map Viamed Ltd ISO13485:2016 from these activities shall be maintained (see 4.2.5). Purchasing process 7.4.2 Top Level Document: VOP Process: 7717 Purchasing information shall 20 Goods in Purchases. Audit 05 Purchasing Suppliers Viamed 24 Aug describe or reference the Returns, Repairs, 2016 Inspection / Rejection Process: 6821 product to be purchased, including as appropriate: Revision Document Responsibility Allocation: VIAMED a) product specifications; ID75943 Management Meeting Supplier Review 09 Mar b) requirements for product Date Revision 24 Nov 2021 2016 Process: 6831 acceptance, procedures, Reviewed 24 Nov 2021 processes and equipment; Top Level Document: VOP Responsibility Allocation: VIAMED c) requirements for 05 Supplier Control, Management Meeting Supplier Review - Min / qualification of supplier Supplier Review, Purchase Max - Re-Orders 09 Mar 2016 personnel: Orders, Supplier Returns Process: 28 d) quality management and Rejection Supplier Review 16 Feb 2016 system requirements. Revision Document Process: 5868 The organization shall ID75847 Return Goods To Suppliers 17 Feb 2016 ensure the adequacy of Date Revision 23 Nov 2021 Process: 6829 specified purchasing Reviewed 23 Nov 2021 Supplier Review - Outstanding orders 09 Mar requirements prior to their Audit 05 Purchasing 2016 communication to the Process: 6832 suppliers supplier. Revision Document Supplier Review Future orders 09 Mar 2016 Purchasing information shall ID69314 Process: 7679 Date Revision 09 Sep 2021 include, as applicable, a Check Stock Requirements Supplier Teledyne written agreement that the Reviewed 09 Sep 2021 18 Apr 2016 supplier notify the Audit 09 Goods Inward Process: 7680 Check Stock Requirements Supplier Envited organization of changes in and Product Identity the purchased product prior Revision Document 18 Apr 2016 to implementation of any ID55437 Process: 7681 changes that affect Check Stock Requirements Supplier Posey 18 Date Revision 12 Mar 2021 the ability of the purchased Reviewed 09 Aug 2023 Apr 2016 product to meet specified Audit 23 Analysis of Data Process: 7682 purchase requirements. Revision Document Check Stock Requirements Supplier Bluepoint To the extent required for ID67997 18 Apr 2016 traceability given in 7.5.9, Date Revision 23 Aug 2021 Process: 7683 the organization shall Check Stock For Proforma 18 Apr 2016 Reviewed 23 Aug 2021 maintain relevant purchasing Process: 7784 information in the form of Check Returns Supplier Envitec 15 Feb 2017 documents (see 4.2.4) and Process: 7785 records (see 4.2.5). Check Returns Supplier Teledyne 15 Feb 2017 Purchasing information Process: 7786 Check Returns Supplier Maxtec 15 Feb 2017 Process: 7787 Check Returns All Supplier 15 Feb 2017 Process: 7826 Goods In Processes 06 Sep 2017 Process: 7923 Review Of Credits Received From Suppliers 08 Jan 2019 Process: 6819 Supplier Payments and Invoice processing 09 Mar 2016

Process: 7882

Process: 7933

Purchase Payments 23 Oct 2017

Purchasing Invoice Processing 22 Mar 2019

Process: 8030 Purchase Order Invoice Review 23 Jun 2023 Top Level Document: VOP || Process: 7717 7.4.3 The organization shall 07 Stock Control, Audit 05 Purchasing Suppliers Viamed 24 Aug establish and implement the Handling, Control of 2016 inspection or other activities Labelling, Storage, Process: 7721 Audit 09 Goods Inward And Product Identity necessary for ensuring Movement that purchased product meets Revision Document Viamed 24 Aug 2016 specified purchasing Process: 8030 ID88809 requirements. The extent of Purchase Order Invoice Review 23 Jun 2023 Date Revision 06 May 2022 verification activities Reviewed 06 May 2022 Top Level Document: VOP shall be based on the supplier evaluation results 06 Measurement Control and proportionate to the risks Viamed VST, Calibration. associated with the QA Stock Revision Document purchased product. When the organization ID53615 becomes aware of any Date Revision 11 Feb 2021 changes to the purchased Reviewed 11 Feb 2021 product, the organization Top Level Document: VOP shall 20 Goods in Purchases, determine whether these Returns, Repairs, changes affect the product Inspection / Rejection Revision Document realization process or the medical device. ID75943 When the organization or its Date Revision 24 Nov 2021 customer intends to perform Reviewed 24 Nov 2021 verification at the Audit 09 Goods Inward supplier �s premises, and Product Identity Revision Document the organization shall state ID55437 the intended verification Date Revision 12 Mar 2021 activities and method of Reviewed 09 Aug 2023 product release in the purchasing information. Records of the verification shall be maintained (see 4.2.5). Verification of purchased product 7.5 Production and service provision 7.5.1 Top Level Document: VOP Process: 7714 Production and service 22 Picking and Packing Audit 01 Picking Packing Viamed 24 Aug provision shall be planned, Dispatch and Goods Out 2016 carried out, monitored and Revision Document Process: 7719 controlled to ensure that ID31048 Audit 07 Handling And Storage Viamed 24 product conforms to Date Revision 30 Sep 2019 Aug 2016 Process: 7725 specification. As appropriate, Reviewed 30 Sep 2019 production controls shall Top Level Document: VOP Audit 12 CE Files Viamed 24 Aug 2016 include but are not limited 07 Stock Control, Process: 7727 Handling, Control of Audit 15 Production Viamed 24 Aug 2016 Labelling, Storage, a) documentation of Process: 7673 procedures and methods for Movement Check Expiry Dated Stock 09 Mar 2016 the control of production Revision Document Process: 6850 Current Stock Levels 09 Mar 2016 (see 4.2.4); ID88809 b) qualification of Process: 6838 Date Revision 06 May 2022

infrastructure;

c) implementation of monitoring and measurement of process parameters and product characteristics; d) availability and use of

monitoring and measuring equipment;

e) implementation of defined operations for labelling and packaging;

f) implementation of product release, delivery and postdelivery activities.

The organization shall establish and maintain a record (see 4.2.5) for each medical device or batch of medical devices that provides traceability to the extent specified in 7.5.9 and identifies the amount manufactured and amount lapproved for distribution. The record shall be verified and approved. **Control of** production and service

provision

Reviewed 06 May 2022

Top Level Document: VOP | Process: 5858 06 Measurement Control Viamed VST, Calibration. QA Stock

Revision Document ID53615

Date Revision 11 Feb 2021 Reviewed 11 Feb 2021

Top Level Document: VOP 08 Production, Reworks, New Production

Revision Document ID31072

Date Revision 30 Sep 2019 Reviewed 30 Sep 2019

Top Level Document: VOP 20 Goods in Purchases, Returns, Repairs, Inspection / Rejection Revision Document

Date Revision 24 Nov 2021 Reviewed 24 Nov 2021

VM3COP20.37 Generating a New Service Visit

Revision Document ID17116 Date Revision 28 Jun 2016 Reviewed 28 Jun 2016

Audit 06 Calibration

Revision Document

ID63048

ID75943

Date Revision 22 Jun 2021 Reviewed 22 Jun 2021

Audit 01 Picking packing

Revision Document ID122441

Date Revision 26 Jun 2023

Reviewed 26 Jun 2023

Audit 07 Handling and Storage

Revision Document

ID120355

Date Revision 02 Jun 2023 Reviewed 02 Jun 2023

Audit 15 Production

Revision Document

ID119452

Date Revision 19 May 2023 Reviewed 19 May 2023

Audit 24 Service Logs

Revision Document

ID68263

Date Revision 26 Aug 2021

Reviewed 26 Aug 2021

Audit 09 Goods Inward and Product Identity Revision Document

Opera Negative Stock 09 Mar 2016

Opera Stock Adjustments 17 Feb 2016

Process: 5935

Stock Allocations 05 Mar 2016

Process: 6945

Missing Stock or Adjustments 09 Mar 2016

Process: 6955

Production Requirements 09 Mar 2016

Process: 7689

Move Stock From QA Shelf To Stock Shelf

Monday 21 Apr 2016

Process: 7694

Move Stock From QA Shelf To Stock Shelf

Tuesday 28 Apr 2016

Process: 7695

Top Up Quick Shipping Shelves 28 Apr 2016

ID55437 Date Revision 12 Mar 2021 Reviewed 09 Aug 2023

7.5.2

The organization shall document requirements for cleanliness of product or contamination control of product if:

- a) product is cleaned by the organization prior to sterilization or its use;
- b) product is supplied nonsterile and is to be subjected to a cleaning process prior to sterilization or its use;
- c) product cannot be cleaned prior to sterilization or its use, and its cleanliness is of significance in use; d) product is supplied to be used non-sterile, and its cleanliness is of significance in use:
- e) process agents are to be removed from product during manufacture. If product is cleaned in accordance with a) or b) above, the requirements contained in 6.4.1 do not apply

prior to the cleaning process. Cleanliness of product

Top Level Document: VM3COP02.01 Exclusions to Viamed ISO13485:2016 boundaries of ISO

Revision Document ID74571

Date Revision 10 Nov 2021 Reviewed 01 Aug 2023

Audit 07 Handling and Storage

Revision Document

ID120355 Date Revision 02 Jun 2023 Reviewed 02 Jun 2023

Process: 7717

Audit 05 Purchasing Suppliers Viamed 24 Aug

Process: 7719

Audit 07 Handling And Storage Viamed 24

Aug 2016

7.5.3

The organization shall document requirements for medical device installation and acceptance criteria for verification of installation, as appropriate. If the agreed customer requirements allow installation of the medical device to be performed by an external party other than the organization or its supplier, the organization shall provide documented requirements for medical device installation and verification of installation. Records of medical device installation and verification of installation performed by

the organization or

Resuscitation Unit and TC400 Maintenance TC400 Installation Instructions

Revision Document ID8155 Date Revision 24 Mar 2011 Reviewed 24 Mar 2011

Resuscitation Unit Instructions for Use / Installation Ceratherm v3.01 Resuscitation Unit and TC400 Maintenance

Revision Document ID8178 Date Revision 24 Mar 2011 Reviewed 24 Mar 2011

Resuscitation Unit Instructions for Use / User Manual Nufer Wall Mount Installation

Revision Document ID1312 Date Revision 19 Mar 2007 Reviewed 19 Mar 2007

Process: 7717

Audit 05 Purchasing Suppliers Viamed 24 Aug 2016

its supplier shall be maintained (see 4.2.5). **Installation activities**

VM3COP51.20 Resuscitation Cabinet Installation Instructions

Revision Document ID18221

Date Revision 12 Dec 2016 Reviewed 12 Dec 2016

Audit 24 Service Logs

Revision Document ID68263

Date Revision 26 Aug 2021

Reviewed 26 Aug 2021

7.5.4

met.

If servicing of the medical device is a specified requirement, the organization shall document servicing procedures, reference materials, and reference measurements, as necessary, for performing servicing activities and verifying that product requirements are

The organization shall analyse records of servicing activities carried out by the organization or its supplier:

- a) to determine if the information is to be handled as a complaint;
- b) as appropriate, for input to the improvement process. Records of servicing activities carried out by the organization or its supplier shall be maintained (see 4.2.5). **Servicing activities**

Top Level Document: VM3COP50.13 Quality Control Tom Thumb

Revision Document ID3115 Date Revision 30 Sep 2019 Reviewed 30 Sep 2019

Top Level Document: VOP 09 Repairs and Servicing

Revision Document ID75927

Date Revision 24 Nov 2021 Reviewed 24 Nov 2021

VM3COP20.27 Annual Services for Resuscitation Cabinets

Revision Document ID24509

Date Revision 06 Dec 2017 Reviewed 06 Dec 2017

VM3COP20.37 Generating a New Service Visit

Revision Document ID17116 Date Revision 28 Jun 2016 Reviewed 28 Jun 2016

VM3COP50.12 Quality Control / Service Checks Tom Thumb

Revision Document

ID15367

Date Revision 05 Aug 2015 Reviewed 05 Aug 2015

Audit 24 Service Logs

Revision Document

ID68263

Date Revision 26 Aug 2021 Reviewed 26 Aug 2021

Audit 11 Repairs, Servicing and Returns

Revision Document ID124549

Date Revision 19 Jul 2023 Reviewed 19 Jul 2023

Audit 23 Analysis of Data

Revision Document ID67997

Process: 5857

Customer Service Logs 17 Feb 2016

Process: 7722

Revision Document ID31154 Audit 10 Documentation Control Viamed 24

Aug 2016

Date Revision 23 Aug 2021 Reviewed 23 Aug 2021 **Audit 14 Complaints and Corrective Actions** Revision Document ID76091 Date Revision 25 Nov 2021 Reviewed 25 Nov 2021

7.5.5

The organization shall maintain records of the sterilization process parameters used for each sterilization batch (see 4.2.5). Sterilization records shall be traceable to each production batch of medical devices. Particular requirements for sterile medical devices

Top Level Document: VM3COP02.01 Exclusions to Viamed ISO13485:2016 **boundaries of ISO**

Revision Document ID74571

Date Revision 10 Nov 2021 Reviewed 01 Aug 2023

Process: 7722

Audit 10 Documentation Control Viamed 24

Aug 2016 Process: 7717

Audit 05 Purchasing Suppliers Viamed 24 Aug

2016

7.5.6

The organization shall validate any processes for production and service provision where the resulting output cannot be or is not verified by subsequent monitoring or measurement and, as a consequence, deficiencies become apparent only after the product is in use or the service has been delivered. Validation shall demonstrate the ability of these processes to achieve planned results consistently.

The organization shall document procedures for validation of processes including:

- a) defined criteria for review and approval of the processes;
- b) equipment qualification and qualification of personnel;
- c) use of specific methods, procedures and acceptance criteria:
- d) as appropriate, statistical techniques with rationale for sample sizes
- e) requirements for records (see 4.2.5);
- f) revalidation, including criteria for revalidation;

Top Level Document: VOP 27 Software Validation

Revision Document ID91486

Date Revision 10 Jun 2022 Reviewed 10 Jun 2022

Top Level Document: VOP 15 Data and Information Analysis

Revision Document ID98547

Date Revision 07 Sep 2022 Reviewed 07 Sep 2022

VM3COP18 Post Market Surveilance

Revision Document ID75985

Date Revision 24 Nov 2021 Reviewed 24 Nov 2021

Audit 03 Design Control Revision Document

Date Revision 17 Feb 2023 Reviewed 17 Feb 2023

ID111315

Audit 24 Service Logs

Revision Document ID68263

Date Revision 26 Aug 2021 Reviewed 26 Aug 2021

Audit 11 Repairs, Servicing |2017|and Returns

Revision Document ID124549

Date Revision 19 Jul 2023 Reviewed 19 Jul 2023

Audit 10 Documentation Control

Process: 7849

Review Product Failures New Codes 28 Sep

2017

Process: 7870

Software Validation Non Conformance Product Risk Feedback Loop 15 Oct 2017

Process: 7879

Software Validation Scheduled Tasks And

Audits 22 Oct 2017 Process: 7850

Software Validation Scan Incorrect Product 01

Oct 2017 Process: 7851

Software Validation Scan Un-QA Product To

Order 01 Oct 2017 Process: 7852

Software Validation Expired Stock 01 Oct

2017

Process: 7853

Software Validation Non Sell Able Shelf 01

Oct 2017 Process: 7854

Software Validation In Production List 01 Oct

2017

Process: 7855

Software Validation - Production Lists 01 Oct 2017

Process: 7856

Software Validation Unchecked Orders 01 Oct

Process: 7857

Software Validation Stock Tracking Check 01 Oct 2017

Process: 7858

Software Validation Attempt To QA Some

Stock 01 Oct 2017 Process: 7861

g) approval of changes to the Revision Document processes.

The organization shall document procedures for the validation of the application of computer software used in production and service provision. Such software applications shall be validated prior to initial use and, as appropriate, after changes to such software or its application. The specific approach and activities associated with software validation and revalidation shall be proportionate to the risk associated with the use of the software including the effect on the ability of the product to conform to specifications. Records of the results and conclusion of validation and necessary actions from the validation shall be maintained (see 4.2.4 and 4.2.5). Validation of

processes for production and service provision

ID63807

Date Revision 30 Jun 2021 Reviewed 30 Jun 2021

Software Validation Of Training Documents Forced Reading 03 Oct 2017

Process: 7865

Software Validation Conflicting Audits 07 Oct 2017

Process: 7875

Software Validation Document Control 20 Oct 2017

Process: 7880

Software Validation Out Of Date Documents

22 Oct 2017 Process: 7881

Software Validation - Live Orders 22 Oct 2017

7.5.7

The organization shall document procedures (see 4.2.4) for the validation of processes for sterilization and sterile barrier systems. Processes for sterilization and sterile barrier systems shall be validated prior to implementation and following product or process changes, as appropriate. Records of the results and, conclusion of validation and necessary actions from the validation shall be maintained (see 4.2.4 and 4.2.5). NOTE Further information can be found in ISO 11607-1 and ISO 11607-2. Particular requirements for validation of processes for sterilization and sterile barrier systems

Top Level Document: VM3COP02.01 Exclusions to Viamed ISO13485:2016 boundaries of ISO

Revision Document ID74571 Date Revision 10 Nov 2021

Reviewed 01 Aug 2023

7.5.8 Top Level Document: VOP Process: 8024 The organization shall 07 Stock Control. Discontinue/Supersede Stock 01 Mar 2023 document procedures for Handling, Control of product identification and Labelling, Storage, identify product by suitable Movement means throughout product Revision Document realization. ID88809 The organization shall Date Revision 06 May 2022 identify product status with Reviewed 06 May 2022 respect to monitoring and **Top Level Document: VOP** 20 Goods in Purchases. measurement requirements throughout Returns, Repairs, product realization. **Inspection / Rejection** Identification of product **Revision Document** status shall be maintained ID75943 throughout production, Date Revision 24 Nov 2021 storage, installation and Reviewed 24 Nov 2021 servicing of product to Audit 07 Handling and ensure that only product that Storage has passed the required Revision Document inspections and tests or ID120355 released under an authorized Date Revision 02 Jun 2023 concession is dispatched, Reviewed 02 Jun 2023 used or installed. Audit 09 Goods Inward and Product Identity If required by applicable regulatory requirements, the Revision Document organization shall document ID55437 a system to assign Date Revision 12 Mar 2021 unique device identification Reviewed 09 Aug 2023 Audit 11 Repairs, Servicing to the medical device. The organization shall and Returns document procedures to Revision Document ensure that medical devices ID124549 returned to the Date Revision 19 Jul 2023 organization are identified Reviewed 19 Jul 2023 and distinguished from conforming product. Identification 7.5.9 VM3COP14.01 Disposition of Documents / Records. Traceability Revision Document ID15464 Date Revision 14 Aug 2015 Reviewed 14 Aug 2015 VM3COP14.01 Disposition 7.5.9.1 The organization shall of Documents / Records. document procedures for Revision Document traceability. These ID15464 procedures shall define the Date Revision 14 Aug 2015 Reviewed 14 Aug 2015 extent of traceability in accordance with applicable VM3COP23.00 EAN13 regulatory requirements and Barcodes to Stock and the the records to be **Online Databases** maintained (see 4.2.5). Revision Document General ID75624 Date Revision 22 Nov 2021 Reviewed 22 Nov 2021

Audit 07 Handling and Storage

Revision Document ID120355

Date Revision 02 Jun 2023 Reviewed 02 Jun 2023

Audit 10 Documentation Control

Revision Document ID63807

Date Revision 30 Jun 2021 Reviewed 30 Jun 2021

7.5.9.2

The records required for traceability shall include records of components, materials, and conditions for the work environment used. if these could cause the medical device not to satisfy its specified safety and performance requirements. The organization shall require that suppliers of distribution services or distributors maintain records

Top Level Document: VM3COP02.01 Exclusions to Viamed ISO13485:2016 boundaries of ISO

Revision Document ID74571

Date Revision 10 Nov 2021 Reviewed 01 Aug 2023

Audit 09 Goods Inward and Product Identity

Revision Document ID55437

Date Revision 12 Mar 2021 Reviewed 09 Aug 2023

Particular requirements for implantable medical devices

the distribution of medical devices to allow traceability and that these records are available for inspection. Records of the name and address of the shipping package consignee shall be maintained (see 4.2.5).

7.5.10

of

The organization shall identify, verify, protect, and safeguard customer property provided for use or incorporation into the product while it is under the organization s control or being used by the organization. If any customer property is lost, damaged or otherwise found to be unsuitable for use, the organization shall report this to the customer and maintain records (see 4.2.5).

Customer property

Top Level Document: VOP Process: 7684 09 Repairs and Servicing

Revision Document ID75927

Date Revision 24 Nov 2021 Reviewed 24 Nov 2021

DO NOT USE VM3COP09 Feb 2016 Repairs

Revision Document ID8712 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

VM3COP20.03 Repair Procedures Goods in

Revision Document ID13703

Date Revision 13 May 2014 Reviewed 13 May 2014 VM3COP20.031 Viamed

Repairs Ready For Quote 18 Apr 2016

Process: 7685

Repairs Ready For Invoice 18 Apr 2016

Process: 5891

Processing Of Repair Quotes And Orders 25

Process: 7693

Collect Repair Filing From Warehouse 22 Apr 2016

Process: 7863

Maintain Repair Codes List 05 Oct 2017

Process: 6847

Responsibility Allocation: Quarantine Repairs

09 Mar 2016 Process: 6862

Current Repairs 09 Mar 2016

Process: 7674

Repair Procedures Invoicing / customer paperwork

Revision Document ID24753

Date Revision 21 Dec 2017 Reviewed 21 Dec 2017

VM3COP20.47 Collecting Repair Paperwork

Revision Document ID17485

Date Revision 15 Sep 2016 Reviewed 15 Sep 2016

Audit 07 Handling and Storage

Revision Document ID120355

Date Revision 02 Jun 2023 Reviewed 02 Jun 2023

Audit 09 Goods Inward and Product Identity

Revision Document ID55437

Date Revision 12 Mar 2021 Reviewed 09 Aug 2023

Audit 11 Repairs, Servicing and Returns

Revision Document ID124549

Date Revision 19 Jul 2023 Reviewed 19 Jul 2023 Check Repairs Ready For Invoice List 10 Mar

2016

Process: 7897

Daily O2 Sensors Returns 04 Jan 2018

Process: 7944

Sealant, Glues, Greases, Sprays, Gases And Tapes You Use In Production, Service And Repairs For Viamed And VST 09 Oct 2019

Process: 7690

Ship Repairs 21 Apr 2016

Process: 7748

Check Repair Orders 10 Oct 2016

Process: 7749

Check Repair Quotes 10 Oct 2016

Process: 7752

SRS Folder 22 Nov 2016

7.5.11

The organization shall document procedures for preserving the conformity of product to requirements during processing, storage, handling, and distribution. Preservation shall apply to the constituent parts of a medical device. The organization shall protect product from alteration, contamination or damage when exposed to expected conditions and hazards during processing, storage, handling, and distribution by: a) designing and constructing suitable packaging and shipping containers: b) documenting requirements for special

conditions needed if

packaging alone cannot

Top Level Document: VOP 09 Repairs and Servicing

Revision Document ID75927

Date Revision 24 Nov 2021 Reviewed 24 Nov 2021

Top Level Document: VOP 07 Stock Control, Handling, Control of Labelling, Storage,

Movement

Revision Document ID88809

Date Revision 06 May 2022 Reviewed 06 May 2022

Top Level Document: VOP 20 Goods in Purchases, Returns, Repairs, Inspection / Rejection Revision Document

ID75943

Date Revision 24 Nov 2021 Reviewed 24 Nov 2021

VM3COP20.03 Repair Procedures Goods in Revision Document Process: 7684

Repairs Ready For Quote 18 Apr 2016

Process: 7685

Repairs Ready For Invoice 18 Apr 2016

Process: 5891

Processing Of Repair Quotes And Orders 25

Feb 2016 **Process: 7673**

Check Expiry Dated Stock 09 Mar 2016

provide preservation. If special conditions are required, they shall be controlled and recorded (see 4.2.5). Preservation of product

ID13703 Date Revision 13 May 2014 Reviewed 13 May 2014

VM3COP20.031 Viamed **Repair Procedures Invoicing / customer** paperwork

Revision Document ID24753

Date Revision 21 Dec 2017 Reviewed 21 Dec 2017

Audit 01 Picking packing

Revision Document ID122441

Date Revision 26 Jun 2023 Reviewed 26 Jun 2023

Audit 07 Handling and Storage

Revision Document ID120355 Date Revision 02 Jun 2023

Reviewed 02 Jun 2023

7.6

The organization shall determine the monitoring and measurement to be lundertaken and the monitoring and measuring equipment needed to provide evidence of conformity of broduct to determined requirements. The organization shall document procedures to ensure that monitoring and measurement can be carried out and are carried out in a manner that is consistent with the

monitoring and measurement requirements. As necessary to ensure valid results, measuring equipment Revision Document shall:

a) be calibrated or verified, or both, at specified intervals, or prior to use, against measurement standards traceable to international or national measurement standards: when no such standards exist, the basis used for calibration or verification shall be recorded (see 4.2.5); b) be adjusted or re-adjusted

as necessary: such

Top Level Document: VOP Process: 7048 06 Measurement Control Viamed VST, Calibration, OA Stock

Revision Document ID53615

Date Revision 11 Feb 2021 Reviewed 11 Feb 2021

DO NOT USE VM3COP11 Calibration

Revision Document ID8713 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Explanation Control of documents

Revision Document ID21322

Date Revision 06 Aug 2017 Reviewed 06 Aug 2017

Audit 06 Calibration

ID63048

Date Revision 22 Jun 2021 Reviewed 22 Jun 2021

Audit 23 Analysis of Data Revision Document

ID67997 Date Revision 23 Aug 2021 Reviewed 23 Aug 2021

Control of monitoring and measuring devices 09 Mar 2016

adjustments or readjustments shall be recorded (see 4.2.5); c) have identification in order to determine its calibration status; d) be safeguarded from adjustments that would invalidate the measurement result; e) be protected from damage and deterioration during handling, maintenance and storage. The organization shall perform calibration or verification in accordance with documented procedures. In addition, the organization shall assess and record the validity of the previous measuring results when the equipment is found not to conform to requirements. The organization shall take appropriate action in regard to the equipment and any product affected. Records of the results of calibration and verification shall be maintained (see 4.2.5). The organization shall document procedures for the validation of the application of computer software used for the monitoring and measurement of requirements. Such software applications shall be validated prior to initial use and, as appropriate, after changes to such software or its application. The specific approach and activities associated with software validation and revalidation shall be proportionate to the risk associated with the use of the software including the effect on the ability of the product to conform to specifications.

Records of the results and
conclusion of validation and
necessary actions from the
validation shall be
maintained (see 4.2.4 and
4.2.5).
NOTE Further information
can be found in ISO 10012.
Control of monitoring and

measuring equipment

8 Measurement, analysis and improvement

0		
8		
Measurement, analysis and		
improvement		
8.1	Top Level Document:	Process: 7714
The organization shall plan	VM3COP27.11 Performing	Audit 01 Picking Packing Viamed 24 Aug
and implement the	a Technical File PMS and	2016
monitoring, measurement,	risk assessment	Process: 7715
analysis and improvement	Revision Document	Audit 02 Contract Review Viamed 24 Aug
processes needed to:	ID75465	2016
a) demonstrate conformity of	Date Revision 18 Nov 2021	Process: 7716
product;	Reviewed 18 Nov 2021	Audit 03 Design Control Viamed 24 Aug 2016
b) ensure conformity of the	Top Level Document: VOP	Process: 7717
quality management system;	13 Process Monitoring,	Audit 05 Purchasing Suppliers Viamed 24 Aug
c) maintain the effectiveness	System Reviews, Audits,	2016
of the quality management	Management Reviews	Process: 7718
system.	Analysis Data PMS Post	Audit 06 Calibration Viamed 24 Aug 2016
This shall include	Market	Process: 7720
determination of appropriate	Revision Document	Audit 08 Training Viamed 24 Aug 2016
methods, including statistical	ID75461	Process: 7719
techniques, and the	Date Revision 18 Nov 2021	Audit 07 Handling And Storage Viamed 24
extent of their use. General	Reviewed 18 Nov 2021	Aug 2016
	Top Level Document: VOP	Process: 7721
	15 Data and Information	Audit 09 Goods Inward And Product Identity
	Analysis	Viamed 24 Aug 2016
	Revision Document	Process: 7722
	ID98547	Audit 10 Documentation Control Viamed 24
	Date Revision 07 Sep 2022	Aug 2016
	Reviewed 07 Sep 2022	Process: 7724
	Explanation Employee	Audit 11 Repairs And Service Viamed 24 Aug
	Roles and Titles	2016
	Revision Document	Process: 7723
	ID22144	Audit 10b Process Verification Viamed 24 Aug
	Date Revision 20 Sep 2017	2016
	Reviewed 20 Sep 2017	Process: 7725
	Audit 22 Post Market	Audit 12 CE Files Viamed 24 Aug 2016
	Survellance	Process: 7726
	Revision Document	Audit 14 Complaints And Corrective Actions
	ID120397	Viamed 24 Aug 2016
	Date Revision 02 Jun 2023	Process: 7727
	Reviewed 02 Jun 2023	Audit 15 Production Viamed 24 Aug 2016
	Audit 23 Analysis of Data	Process: 7728
	Revision Document	Audit 17 Internal Audits Viamed 24 Aug 2016
	ID67997	Process: 7729
ıl —		

Reviewed 23 Aug 2021

DO NOT USE VM3COP13 Process: 7730

Audits

Revision Document ID8715 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Date Revision 23 Aug 2021 | Audit 19 Health And Saftey Viamed 24 Aug 2016

Audit 20 Process Verification To Managment

Viamed 24 Aug 2016

Process: 7731

Audit 21 Audit Of Audit Viamed 24 Aug 2016

Process: 7732

Audit 22 Post Market Survellance Viamed 24

Aug 2016 Process: 7733

Audit 23 Analysis Of Data Viamed 24 Aug

2016

Process: 7834

Financial Review 20 Sep 2017

Process: 7862

Review The Audit Calender Screen 04 Oct

2017

Process: 27

Management Reviews And Quality Audits 16

Feb 2016

Process: 5877

Review Company Data 17 Feb 2016

Process: 7070

Management Review 09 Mar 2016

Process: 7830

Review Q.A. Failures Report 18 Sep 2017

Process: 7837

Review External Parties Influencing The QMS

VST / Viamed 23 Sep 2017

Process: 7838

Review VIAMED Feedback - Customer

Feedback Negative 23 Sep 2017

Process: 7839

Review VIAMED Feedback - Customer

Complaints 23 Sep 2017

Process: 7840

Review VST Feedback - Customer Feedback

Negative 23 Sep 2017

Process: 7841

Review VST Feedback - Customer Complaints

23 Sep 2017

Process: 7842

Review VIAMED Product Feedback Negative

23 Sep 2017

Process: 7843

Review VST Product Feedback Negative 23 Sep 2017

Process: 7848

Review ISO Scopes 27 Sep 2017

Process: 7849

Review Product Failures New Codes 28 Sep

2017

Process: 7871

Review Exclusion From Viamed 13485:2016

And VST 9001:2015 15 Oct 2017

Process: 7874

Review For Latest Version Med Dev 2.12. 18

10/2023, 11:03	QM3 Route Map	Viamed Ltd ISO13485:2016
		Oct 2017
		Process: 7876
		Maintain Update Of ISO Route Maps 21 Oct
		2017
		Process: 7878
		Review Possible Upcoming Regulation
		Changes 22 Oct 2017
8.2		
Monitoring and		
measurement		
	m I In	D 7077
8.2.1	Top Level Document:	Process: 7877
As one of the measurements	VM3COP27.11 Performing	Disaster Planning 21 Oct 2017
of the effectiveness of the	a Technical File PMS and	Process: 5877
quality management system,	risk assessment	Review Company Data 17 Feb 2016
the organization	Revision Document	
shall gather and monitor	ID75465	
information relating to	Date Revision 18 Nov 2021	
whether the organization has	Reviewed 18 Nov 2021	
met customer	Top Level Document: VOP	
requirements. The methods	13 Process Monitoring,	
for obtaining and using this	System Reviews, Audits,	
information shall be	Management Reviews	
documented.	Analysis Data PMS Post	
The organization shall	Market	
document procedures for the	II I	
feedback process. This	ID75461	
	Date Revision 18 Nov 2021	
feedback process shall	II I	
include provisions to gather	Reviewed 18 Nov 2021	
data from production as well		
as post-production activities.		
The information gathered in	ID30851	
the feedback process shall	Date Revision 18 Sep 2019	
serve as potential input into	Reviewed 18 Sep 2019	
risk management	Management reviews	
for monitoring and	Revision Document	
maintaining the product	ID19801	
requirements as well as the	Date Revision 05 May 2017	
product realization or	Reviewed 05 May 2017	
improvement processes.	Audit 23 Analysis of Data	
If applicable regulatory	Revision Document	
requirements require the	ID67997	
organization to gain specific	Date Revision 23 Aug 2021	
experience from	Reviewed 23 Aug 2021	
postproduction activities, the	II - I	
review of this experience	Survellance	
shall form part of the	Revision Document	
feedback process. Feedback	II I	
recuback process. Feeuback	Date Revision 02 Jun 2023	
	II I	
	Reviewed 02 Jun 2023	
	Audit 14 Complaints and	
	Corrective Actions	
	Revision Document	
	ID76091	
	Date Revision 25 Nov 2021	
	Reviewed 25 Nov 2021	I .

8.2.2

The organization shall document procedures for timely complaint handling in accordance with applicable regulatory requirements. These procedures shall include at a minimum requirements and responsibilities for:

- a) receiving and recording information;
- b) evaluating information to determine if the feedback constitutes a complaint;
- c) investigating complaints;
- d) determining the need to report the information to the appropriate regulatory authorities;
- e) handling of complaintrelated product;
- f) determining the need to initiate corrections or corrective actions. If any complaint is not investigated, justification shall be documented. Any correction or corrective

action resulting from the complaint handling process shall be documented. If an investigation determines activities outside the organization contributed to the complaint, relevant information shall be exchanged between the organization and the external party involved.

Complaint handling records shall be maintained (see

4.2.5). Complaint handling

8.2.3

If applicable regulatory requirements require notification of complaints that meet specified reporting criteria of adverse events or issuance of advisory notices, the organization shall document procedures for providing notification to the appropriate regulatory authorities. Records of reporting to

Top Level Document: VOP Process: 7743 19 Feedback Customer **Complaints Vigilance and Notifications Viamed Ltd** Revision Document ID75475

Date Revision 18 Nov 2021 Reviewed 18 Nov 2021

Audit 14 Complaints and Corrective Actions

Revision Document ID76091

Date Revision 25 Nov 2021 Reviewed 25 Nov 2021

Customer Complaints Paper File 26 Sep 2016

Process: 7743

Customer Complaints Paper File 26 Sep 2016

Top Level Document: VOP

19 Feedback Customer Complaints Vigilance and Notifications Viamed Ltd

Revision Document ID75475

Date Revision 18 Nov 2021 Reviewed 18 Nov 2021

Audit 14 Complaints and Corrective Actions

Revision Document ID76091

Date Revision 25 Nov 2021

Process: 7743

Customer Complaints Paper File 26 Sep 2016

Process: 7743

Customer Complaints Paper File 26 Sep 2016

regulatory authorities shall be maintained (see 4.2.5). Reporting to regulatory authorities

Reviewed 25 Nov 2021 MHRA Correspondence / **RG2 Devices list**

Revision Document ID14763

Date Revision 12 Feb 2015 Reviewed 12 Feb 2015

MHRA Appendix A / Appendix B Class 1 Device Codes

Revision Document ID4798 Date Revision 24 Oct 2008 Reviewed 24 Oct 2008

CE Guidance 19 Own **Brand MHRA position obl** Revision Document ID3656 Date Revision 29 Apr 2008 Reviewed 29 Apr 2008

8.2.4

The organization shall conduct internal audits at planned intervals to determine whether the

quality management system: a) conforms to planned and documented arrangements, requirements of this International Standard, quality management system requirements established by the organization, and applicable

regulatory requirements;

b) is effectively implemented and maintained. The organization shall document a procedure to describe the responsibilities and requirements for planning and conducting audits and recording and reporting audit results. An audit program shall be planned, taking into consideration the status and importance of the processes and area to be audited, as well as the results of previous audits. The audit criteria, scope, interval and methods shall be defined and recorded (see 4.2.5). The selection of auditors and conduct of audits shall ensure objectivity and impartiality of the audit

Top Level Document: VOP

13 Process Monitoring, System Reviews, Audits, **Management Reviews Analysis Data PMS Post**

Market

Revision Document

ID75461 Date Revision 18 Nov 2021 Reviewed 18 Nov 2021

Audit 01 Picking packing

Revision Document ID122441

Date Revision 26 Jun 2023 Reviewed 26 Jun 2023

Audit 02 Contract Review and Sales Order Processing Process: 7720

Revision Document ID69328

Date Revision 09 Sep 2021 Reviewed 09 Sep 2021

Audit 06 Calibration

Revision Document ID63048

Date Revision 22 Jun 2021 Reviewed 22 Jun 2021

Audit 08 Training, Competence and Human

Resources

Revision Document

ID70147 Date Revision 20 Sep 2021

Reviewed 20 Sep 2021

Audit 09 Goods Inward and Product Identity Revision Document

ID55437

Date Revision 12 Mar 2021 Reviewed 09 Aug 2023

Process: 7714

Audit 01 Picking Packing Viamed 24 Aug

2016

Process: 7715

Audit 02 Contract Review Viamed 24 Aug

2016

Process: 7716

Audit 03 Design Control Viamed 24 Aug 2016

Process: 7717

Audit 05 Purchasing Suppliers Viamed 24 Aug

2016

Process: 7718

Audit 06 Calibration Viamed 24 Aug 2016

Process: 7719

Audit 07 Handling And Storage Viamed 24

Aug 2016

Audit 08 Training Viamed 24 Aug 2016

Process: 7721

Audit 09 Goods Inward And Product Identity

Viamed 24 Aug 2016

Process: 7722

Audit 10 Documentation Control Viamed 24

Aug 2016

Process: 7723

Audit 10b Process Verification Viamed 24 Aug

2016

Process: 7725

Audit 12 CE Files Viamed 24 Aug 2016

Process: 7724

Audit 11 Repairs And Service Viamed 24 Aug

2016

Process: 7726

Audit 14 Complaints And Corrective Actions

Viamed 24 Aug 2016

Process: 7727

Audit 15 Production Viamed 24 Aug 2016

Process: 7728

Audit 17 Internal Audits Viamed 24 Aug 2016

process. Auditors shall not audit their own work. Records of the audits and their results, including identification of the processes and areas audited and the conclusions, shall be maintained (see 4.2.5). The management responsible for the area being audited shall ensure that any necessary corrections and corrective actions are taken without undue delay to leliminate detected nonconformities and their causes. Follow-up activities shall include the verification of the actions taken and the reporting of verification results. NOTE Further information can be found in ISO 19011. Internal audit

Audit 10 Documentation Control

Revision Document ID63807

Date Revision 30 Jun 2021 Reviewed 30 Jun 2021

Audit 20 Process

verification to Managment Revision Document

ID73324

Date Revision 26 Oct 2021 Reviewed 26 Oct 2021

and Returns

Revision Document ID124549

Date Revision 19 Jul 2023 Reviewed 19 Jul 2023

Audit 15 Production

Revision Document ID119452

Date Revision 19 May 2023 Reviewed 19 May 2023

Audit 17 Internal Audits

Revision Document

ID77209

Date Revision 08 Dec 2021 Reviewed 08 Dec 2021

Audit 18 Management

Review

Revision Document ID73320

Date Revision 26 Oct 2021

Reviewed 26 Oct 2021

Audit 19 Health and Safety, Working

Conditions and Building

Fabric Issues

Revision Document

ID68045

Date Revision 24 Aug 2021 Reviewed 24 Aug 2021

Audit 21 Audit of Audit

Revision Document

ID77289

Date Revision 09 Dec 2021 Reviewed 09 Dec 2021

Audit 22 Post Market

Survellance

Revision Document

ID120397

Date Revision 02 Jun 2023

Reviewed 02 Jun 2023

Audit 23 Analysis of Data

Revision Document

ID67997

Date Revision 23 Aug 2021 Reviewed 23 Aug 2021

Process: 7729

Audit 19 Health And Saftey Viamed 24 Aug

2016

Process: 7730

Audit 20 Process Verification To Managment

Viamed 24 Aug 2016

Process: 7731

Audit 21 Audit Of Audit Viamed 24 Aug 2016

Process: 7732

Audit 22 Post Market Survellance Viamed 24

Aug 2016

Process: 7733

Audit 11 Repairs, Servicing Audit 23 Analysis Of Data Viamed 24 Aug

Audit 24 Service Logs **Revision Document** ID68263 Date Revision 26 Aug 2021 Reviewed 26 Aug 2021 **Explanation Employee Roles and Titles** Revision Document ID22144 Date Revision 20 Sep 2017 Reviewed 20 Sep 2017 DO NOT USE VM3COP13 Audits Revision Document ID8715 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011 Audit Schedule Revision Document ID23221 Date Revision 24 Oct 2017

8.2.5

8.2.6

The organization shall apply suitable methods for monitoring and, as appropriate, measurement of the quality management system processes. These methods shall demonstrate the ability of the processes to achieve planned results. When planned results are not achieved, correction and corrective action shall be taken, as appropriate.

Monitoring and measurement of processes

Top Level Document: VOP 13 Process Monitoring, System Reviews, Audits, **Management Reviews** Analysis Data PMS Post Market Revision Document ID75461 Date Revision 18 Nov 2021 Reviewed 18 Nov 2021 **Audit 23 Analysis of Data** Revision Document ID67997 Date Revision 23 Aug 2021 Reviewed 23 Aug 2021 **Audit 10 Documentation**

Reviewed 24 Oct 2017

Control Revision Document ID63807 Date Revision 30 Jun 2021

Reviewed 30 Jun 2021 DO NOT USE VM3COP11

The organization shall Calibration monitor and measure the Revision Document ID8713 characteristics of the product Date Revision 12 Oct 2011 to verify that product Reviewed 12 Oct 2011 requirements have been met. OLD DO NOT USE This shall be carried out at VM3COP29 Production applicable stages of the

Revision Document ID8727 Date Revision 12 Oct 2011 Reviewed 12 Oct 2011

Audit 07 Handling and Storage Revision Document

Evidence of conformity with the acceptance criteria shall

process in accordance with

documented procedures.

the planned and documented

product realization

arrangements and

ID120355 Date Revision 02 Jun 2023

Process: 27

Management Reviews And Quality Audits 16 Feb 2016

be maintained. The identity of the person authorizing release of product shall be recorded (see 4.2.5). As appropriate, records shall identify the test equipment used to perform measurement activities. Product release and service delivery shall not proceed until the planned and documented arrangements have been satisfactorily completed. For implantable medical devices, the organization shall record the identity of personnel performing any inspection or testing. Monitoring and measurement of product

Reviewed 02 Jun 2023 **Audit 15 Production** Revision Document ID119452 Date Revision 19 May 2023

Reviewed 19 May 2023

Process: 8024

Discontinue/Supersede Stock 01 Mar 2023

8.3 **Control of nonconforming** product

8.3.1

The organization shall ensure that product which does not conform to product requirements is identified and controlled to prevent its unintended use or delivery. The organization shall document a procedure to define the controls and related responsibilities and authorities for the identification, documentation, segregation, evaluation, and disposition of nonconforming product. The evaluation of nonconformity shall include a determination of the need for an investigation and notification of any external party responsible for the nonconformity. Records of the nature of the nonconformities and any subsequent action taken, including the evaluation, any investigation and the rationale for decisions shall

be maintained (see 4.2.5)

General

Top Level Document: VOP 19 Feedback Customer Complaints Vigilance and Notifications Viamed Ltd Revision Document

ID75475

Date Revision 18 Nov 2021 Reviewed 18 Nov 2021

Top Level Document: VOP 10 Non Conformance, **Corrective and Preventive** Actions

Revision Document ID124938

Date Revision 24 Jul 2023 Reviewed 24 Jul 2023

VM3COP10.02 Product Recall locate products out in the Field

Revision Document ID74788

Date Revision 12 Nov 2021 Reviewed 12 Nov 2021

Audit 07 Handling and Storage Revision Document

ID120355 Date Revision 02 Jun 2023 Reviewed 02 Jun 2023

Audit 09 Goods Inward and Product Identity **Revision Document**

Process: 7743

Customer Complaints Paper File 26 Sep 2016

Process: 7743

Customer Complaints Paper File 26 Sep 2016

	•	
	ID55437	
	Date Revision 12 Mar 2021	
	Reviewed 09 Aug 2023	
	Audit 23 Analysis of Data	
	Revision Document	
	ID67997	
	Date Revision 23 Aug 2021	
	Reviewed 23 Aug 2021	
3.3.2	Audit 07 Handling and	
The organization shall deal	Storage	
O	_{II} 5	(1

8.

with nonconforming product ||Revision Document by one or more of the following ways: a) taking action to eliminate

- the detected nonconformity; b) taking action to preclude its original intended use or application;
- c) authorizing its use, release or acceptance under concession.

The organization shall ensure that nonconforming product is accepted by concession only if the justification is provided, approval is obtained, and applicable regulatory requirements are met. Records of the acceptance by concession and the identity of the person authorizing the concession shall be maintained (see 4.2.5). Actions in response to

ID120355

Date Revision 02 Jun 2023 Reviewed 02 Jun 2023

8.3.3

When nonconforming product is detected after delivery or use has started, the organization shall take action appropriate to the effects, or potential effects, of the nonconformity. Records of actions taken shall be maintained (see 4.2.5).

nonconforming product detected before delivery

The organization shall document procedures for issuing advisory notices in accordance with applicable regulatory requirements. These procedures shall be capable of being put into effect at any time. Records of actions relating to the

Top Level Document: VOP 19 Feedback Customer Complaints Vigilance and Notifications Viamed Ltd

Revision Document ID75475

Date Revision 18 Nov 2021 Reviewed 18 Nov 2021

Audit 14 Complaints and Corrective Actions

Revision Document ID76091

Date Revision 25 Nov 2021 Reviewed 25 Nov 2021

2/10/2023, 11:03	QMS Route Map	Viamed Ltd ISO13485:2016
lissuance of advisory notices		
shall be maintained (see		
4.2.5). Actions in response		
to nonconforming product		
detected after delivery		
8.3.4	Top Level Document: VOP	
The organization shall	08 Production, Reworks,	
perform rework in	New Production	
accordance with documented	Revision Document	
procedures that takes into	ID31072	
account the potential adverse	Date Revision 30 Sep 2019	
effect of the rework on the	Reviewed 30 Sep 2019	
product. These procedures	Top Level Document: VOP	
shall undergo the	09 Repairs and Servicing	
same review and approval as	Revision Document	
the original procedure.	ID75927	
After the completion of	Date Revision 24 Nov 2021	
rework, product shall be	Reviewed 24 Nov 2021	
verified to ensure that it	Audit 20 Process	
meets applicable acceptance	verification to Managment	
criteria and regulatory	Revision Document	
requirements.	ID73324	
Records of rework shall be	Date Revision 26 Oct 2021	
maintained (see 4.2.5).	Reviewed 26 Oct 2021	
Rework	Audit 11 Repairs, Servicing	
	and Returns	
	Revision Document ID124549	
	Date Revision 19 Jul 2023	
	Reviewed 19 Jul 2023	
8.4	Top Level Document: VOP	l I
The organization shall	13 Process Monitoring,	Automotive Competitor Price Review 10 Mar
document procedures to	System Reviews, Audits,	2023
determine, collect and analyse appropriate data	Management Reviews Analysis Data PMS Post	
to demonstrate the	Market	
suitability, adequacy and	Revision Document	
effectiveness of the quality	ID75461	
management system. The	Date Revision 18 Nov 2021	
procedures shall include	Reviewed 18 Nov 2021	
determination of appropriate		
methods, including statistical		
techniques and	Supplier Review, Purchase	
the extent of their use.	Orders, Supplier Returns	
The analysis of data shall	and Rejection	
include data generated as a	Revision Document	
result of monitoring and	ID75847	
measurement and from	Date Revision 23 Nov 2021	
other relevant sources and	Reviewed 23 Nov 2021	
include, at a minimum, input	Top Level Document: VOP	
from:	15 Data and Information	
a) feedback;	Analysis	
b) conformity to product	Revision Document	
requirements;	ID98547	
c) characteristics and trends	Date Revision 07 Sep 2022	
of processes and product	II * I	I II
11 *	Reviewed 07 Sep 2022	
including opportunities for	II * I	

2/10/2023, 11:03	QM3 Route Map	Viamed Ltd ISO13485:2016
improvement;	Survellance	
d) suppliers;	Revision Document	
e) audits;	ID120397 Date Revision 02 Jun 2023	
f) service reports, as appropriate.	Reviewed 02 Jun 2023	
If the analysis of data shows	Audit 23 Analysis of Data	
that the quality management	Revision Document	
system is not suitable,	ID67997	
adequate or effective,	Date Revision 23 Aug 2021	
the organization shall use	Reviewed 23 Aug 2021	
this analysis as input for		
improvement as required in		
8.5.		
Records of the results of		
analyses shall be maintained		
(see 4.2.5). Analysis of data		
8.5		
Improvement		
8.5.1	Top Level Document: VOP	
The organization shall	10 Non Conformance,	
identify and implement any	Corrective and Preventive	
changes necessary to ensure	Actions	
and maintain the	Revision Document	
continued suitability,	ID124938	
adequacy and effectiveness of the quality management	Date Revision 24 Jul 2023 Reviewed 24 Jul 2023	
system as well as medical	Audit 06 Calibration	
device safety and	Revision Document	
performance through the use	ID63048	
of the quality policy, quality	Date Revision 22 Jun 2021	
objectives, audit results,	Reviewed 22 Jun 2021	
postmarket surveillance,	Audit 18 Management	
analysis of data, corrective	Review	
actions, preventive actions	Revision Document	
and management review.	ID73320	
General	Date Revision 26 Oct 2021	
	Reviewed 26 Oct 2021	
	Audit 22 Post Market	
	Survellance Revision Document	
	ID120397	
	Date Revision 02 Jun 2023	
	Reviewed 02 Jun 2023	
	Audit 23 Analysis of Data	
	Revision Document	
	ID67997	
	Date Revision 23 Aug 2021	
	Reviewed 23 Aug 2021	
	Audit 21 Audit of Audit	
	Revision Document	
	ID77289	
	Date Revision 09 Dec 2021 Reviewed 09 Dec 2021	
8.5.2		
The organization shall take	Top Level Document: VOP 10 Non Conformance,	
	Corrective and Preventive	

of nonconformities in order to prevent recurrence. Any necessary corrective actions shall be taken without undue delay. Corrective actions shall be proportionate to the effects of the nonconformities encountered. The organization shall document a procedure to

define requirements for: a) reviewing

nonconformities (including complaints);

b) determining the causes of nonconformities;

c) evaluating the need for action to ensure that nonconformities do not recur;

d) planning and documenting Reviewed 25 Nov 2021 action needed and implementing such action, including, as appropriate, updating documentation; e) verifying that the corrective action does not

adversely affect the ability to meet applicable regulatory requirements or the safety and performance of the medical device; f) reviewing the effectiveness of corrective action taken Records of the results of any

investigation and action taken shall be maintained

(see 4.2.5). Corrective action

8.5.3

The organization shall determine action to eliminate the causes of potential nonconformities in order to prevent their occurrence. Preventive lactions shall be proportionate to the effects of the potential problems. The organization shall document a procedure to describe requirements for: a) determining potential

Actions

Revision Document ID124938

Date Revision 24 Jul 2023 Reviewed 24 Jul 2023

Audit 20 Process

verification to Managment

Revision Document ID73324

Date Revision 26 Oct 2021 Reviewed 26 Oct 2021

Audit 10 Documentation Control

Revision Document ID63807

Date Revision 30 Jun 2021 Reviewed 30 Jun 2021

Audit 14 Complaints and Corrective Actions

Revision Document ID76091

Date Revision 25 Nov 2021

Top Level Document: VOP Process: 7839 10 Non Conformance, **Corrective and Preventive**

Actions

Revision Document ID124938

Date Revision 24 Jul 2023 Reviewed 24 Jul 2023

Audit 20 Process verification to Managment

Revision Document ID73324

Date Revision 26 Oct 2021 Reviewed 26 Oct 2021

Audit 14 Complaints and

Review VIAMED Feedback - Customer Complaints 23 Sep 2017

Process: 7838

Review VIAMED Feedback - Customer Feedback Negative 23 Sep 2017

Process: 7842

Review VIAMED Product Feedback Negative 23 Sep 2017

Process: 7849

Review Product Failures New Codes 28 Sep 2017

Process: 6866

Internal Process Verification Complete Systems Review 09 Mar 2016

nonconformities and their	Corrective Actions	Process: 7743
causes;	Revision Document	Customer Complaints Paper File 26 Sep 2016
b) evaluating the need for	ID76091	Process: 7199
action to prevent occurrence	Date Revision 25 Nov 2021	Non Conformities Review Viamed 09 Mar
of nonconformities;	Reviewed 25 Nov 2021	2016
c) planning and documenting		Process: 7671
action needed and		Humanmed Non Conformances 09 Mar 2016
implementing such action,		Process: 7091
including, as appropriate,		Calibration Index 09 Mar 2016
updating documentation;		Process: 7138
d) verifying that the action		Non Conformance Issues Any New QC21
does not adversely affect the		Forms 09 Mar 2016
ability to meet applicable		
regulatory		
requirements or the safety		
and performance of the		
medical device;		
e) reviewing the		
effectiveness of the		
preventive action taken, as appropriate.		
Records of the results of any		
investigations and of action		
taken shall be maintained		
(see 4.2.5). Preventive		
action		

Document ID	Sub Processes
ID70776	Viamed ISO 13485:2016 Scope Process: 5887 Review ISO/EN Documents 24 Feb 2016 Process: 7848 Review ISO Scopes 27 Sep 2017
ID74571	VM3COP02.01 Exclusions to Viamed ISO13485:2016 boundaries of ISO Process: 5887 Review ISO/EN Documents 24 Feb 2016 Process: 7871 Review Exclusion From Viamed 13485:2016 And VST 9001:2015 15 Oct 2017
ID22684	VM3COP00.00 VOP00.00 Viamed Quality Statement policy and objectives Process: 23 Company Objectives 16 Feb 2016 Process: 22 Company Policys 16 Feb 2016 Process: 7828 Review The Quality Policy Viamed 16 Sep 2017 Process: 7833 Importance Of Effective Quality Management 20 Sep 2017
ID27474	VM3COP02.02 Viamed Company Responsibilitys organisation chart structure Process: 5877 Review Company Data 17 Feb 2016
ID63807	Audit 10 Documentation Control Process: 10 Distribution Of Emails 16 Feb 2016 Process: 5939 Responsibility Allocation: Email ISP Routing 05 Mar 2016 Process: 5940 Thumb Nail Processor 07 Mar 2016 Process: 11 Distribution Of Post 16 Feb 2016 Process: 6 Responsibility Allocation: Updating Contact Management System 16 Feb 2016 Process: 52 Software Verification Clear Down Backup Emails 16 Feb 2016 Process: 53 Emails 16 Feb 2016 Process: 7672 Off Site Backup 09 Mar 2016 Process: 7700 Domain Name Management 19 May 2016 Process: 9 Distribution Of Faxes 16 Feb 2016 Process: 15 Filing and Archiving 16 Feb 2016 Process: 7711 Import Bank CSV 01 Jul 2016

Process: 7722 Audit 10 Documentation Control Viamed 24 Aug 2016 **Process: 7693** Collect Repair Filing From Warehouse 22 Apr 2016

Process: 12 Responsibility Allocation : Sales And Technical Information Processing 16 Feb

2016

Process: 16 Responsibility Allocation : Photocopying 16 Feb 2016 **Process: 5901** Link Call Log Contacts To The CRM 02 Mar 2016 **Process: 7699** Shred Sensitive Paperwork In JL Office 19 May 2016

Process: 7705 Checking For Uploaded Files 08 Jun 2016

Process: 7754

Process: 7770 Audit 10 Documentation Control VST 08 Feb 2017

Process: 6938 Responsibility Allocation : Customer Database Updates 09 Mar 2016 **Process: 6940** Responsibility Allocation : Customer Ongoing task List 09 Mar 2016

Process: 7090 Responsibility Allocation : Office Procedures 09 Mar 2016

Process: 7032 Responsibility Allocation : Document Requirements 09 Mar 2016

Process: 41 Responsibility Allocation : Documentation Control 16 Feb 2016

Process: 59 Out Of Date Documents 17 Feb 2016 **Process: 5851** Duplicate Documents 17 Feb 2016

Process: 5852 Responsibility Allocation: Retention Of Records 17 Feb 2016

Process: 7124 Responsibility Allocation : Intrastats 09 Mar 2016

Process: 7125 Responsibility Allocation: Intrastats Urgent Problems 09 Mar 2016

Process: 7126 Intrastats Requested Page updates 09 Mar 2016

Process: 7127 Responsibility Allocation : Intrastats Unfinished in progress Processes 09 Mar 2016

Process: 7128 Responsibility Allocation: Intrastats Future Features needed 09 Mar 2016

Process: 7129 Intrastats Cross Reference Database Tables Updates 09 Mar 2016

Process: 7130 Intrastats Information for Intrastats and L Drive 09 Mar 2016

Process: 7131 Responsibility Allocation : Intrastats Opera 09 Mar 2016

Process: 7133 Responsibility Allocation: Intrastats Contact Manager 09 Mar 2016

Process: 7739 Intrastats Amendment Log 12 Sep 2016

Process: 5877 Review Company Data 17 Feb 2016

Process: 44 Secure Socket Level Certificate 16 Feb 2016

Process: 5890 Check Website ISO Documents 24 Feb 2016 **Process: 7863** Maintain Repair Codes List 05 Oct 2017

Process: 7922 Back Up Emily's Accounts Docs 04 Jan 2019

Process: 7987 Sync External Telephone Logs 07 Feb 2022

Process: 7992 COSHH Datasheet Reminders 07 Feb 2022

Process: 8001 Verification Stock Linked To Documents 08 Feb 2022

Process: 8029 Send Intercompany Invoices To Jean 12 Apr 2023

Process: 8032 Review Contact Documentation 22 Aug 2023

ID120321 VOP 01 Documentation and Records, Control, Creation, Storage, Retrieval, Revision Control and Online Records

Process: 5940 Thumb Nail Processor 07 Mar 2016

Process: 7827 Review The Quality Policy VST 16 Sep 2017

Process: 7828 Review The Quality Policy Viamed 16 Sep 2017

Process: 5934 Responsibility Allocation : Staff Training 05 Mar 2016

Process: 7032 Responsibility Allocation : Document Requirements 09 Mar 2016

Process: 41 Responsibility Allocation: Documentation Control 16 Feb 2016

Process: 59 Out Of Date Documents 17 Feb 2016

Process: 5851 Duplicate Documents 17 Feb 2016

Process: 5852 Responsibility Allocation: Retention Of Records 17 Feb 2016

Process: 7130 Intrastats Information for Intrastats and L Drive 09 Mar 2016

Process: 5890 Check Website ISO Documents 24 Feb 2016

Process: 7200 Responsibility Allocation: ISO Issues 09 Mar 2016

Process: 7744 FDA Device Establishment Registration And Listing 28 Sep 2016

Process: 7941 Check Leaflets, Letterhead And Other Paperwork To See If The Correct BSI

Logo Is In Use. Remove All Old If Found. 23 Sep 2019

Process: 7987 Sync External Telephone Logs 07 Feb 2022

/10/2023, 11:0	QMS Route Map Viamed Ltd ISO13485:2016
	Process: 7992 COSHH Datasheet Reminders 07 Feb 2022
	Process: 8001 Verification Stock Linked To Documents 08 Feb 2022
	Process: 8029 Send Intercompany Invoices To Jean 12 Apr 2023
	Process: 8032 Review Contact Documentation 22 Aug 2023
ID8700	Chart 27 Customer Complaints Chart 27
100/00	Process: 7743 Customer Complaints Paper File 26 Sep 2016
ID73324	Audit 20 Process verification to Managment
	Process: 7701 AWS Amazon Web Services 23 May 2016
	Process: 7723 Audit 10b Process Verification Viamed 24 Aug 2016
	Process: 7730 Audit 20 Process Verification To Managment Viamed 24 Aug 2016
	Process: 7827 Review The Quality Policy VST 16 Sep 2017
	Process: 7828 Review The Quality Policy Viamed 16 Sep 2017
	Process: 7771 Audit 10b Process Verification VST 08 Feb 2017
	Process: 7778 Audit 20 Process Verification To Managment VST 08 Feb 2017
	Process: 6866 Internal Process Verification Complete Systems Review 09 Mar 2016
	Process: 7755 Fast Hosts Invoice 08 Dec 2016
	Process: 7845 7.1.4 Environment Of Operations 25 Sep 2017
	Process: 7846 ISO System Management Review Viamed 26 Sep 2017
	Process: 7837 Review External Parties Influencing The QMS VST / Viamed 23 Sep 2017
	Process: 7832 Cleardown Emailed Invoices 20 Sep 2017
	Process: 7848 Review ISO Scopes 27 Sep 2017
	Process: 7851 Software Validation Scan Un-QA Product To Order 01 Oct 2017
	Process: 7852 Software Validation Expired Stock 01 Oct 2017
	Process: 7853 Software Validation Non Sell Able Shelf 01 Oct 2017
	Process: 7854 Software Validation In Production List 01 Oct 2017
	Process: 7855 Software Validation - Production Lists 01 Oct 2017
	Process: 7856 Software Validation Unchecked Orders 01 Oct 2017
	Process: 7857 Software Validation Stock Tracking Check 01 Oct 2017
	Process: 7858 Software Validation Attempt To QA Some Stock 01 Oct 2017
	Process: 7861 Software Validation Of Training Documents Forced Reading 03 Oct 2017
	Process: 7850 Software Validation Scan Incorrect Product 01 Oct 2017
	Process: 7871 Review Exclusion From Viamed 13485:2016 And VST 9001:2015 15 Oct 2017
	Process: 7865 Software Validation Conflicting Audits 07 Oct 2017
	Process: 7870 Software Validation Non Conformance Product Risk Feedback Loop 15 Oct
	2017
	Process: 7879 Software Validation Scheduled Tasks And Audits 22 Oct 2017
	Process: 7875 Software Validation Document Control 20 Oct 2017
	Process: 7880 Software Validation Out Of Date Documents 22 Oct 2017
	Process: 7881 Software Validation - Live Orders 22 Oct 2017
ID4 6005	
ID16995	VM3COP27.17 Complete Auto_calender Issues
	Process: 27 Management Reviews And Quality Audits 16 Feb 2016
ID85362	VM3COP27.02 Collecting Emails and Distributing
	Process: 10 Distribution Of Emails 16 Feb 2016
ID75461	VOP 13 Process Monitoring, System Reviews, Audits, Management Reviews Analysis
10/5401	Data PMS Post Market
	Process: 55 Business Continuity Plan 17 Feb 2016
	Process: 23 Company Objectives 16 Feb 2016
	Process: 27 Management Reviews And Quality Audits 16 Feb 2016
	Process: 27 Management Reviews And Quanty Addits 16 Feb 2016 Process: 7714 Audit 01 Picking Packing Viamed 24 Aug 2016
	Process: 7715 Audit 02 Contract Review Viamed 24 Aug 2016
	Process: 7716 Audit 03 Design Control Viamed 24 Aug 2016
	Process: 7717 Audit 05 Purchasing Suppliers Viamed 24 Aug 2016
	Process: 7718 Audit 06 Calibration Viamed 24 Aug 2016
	Process: 7719 Audit 07 Handling And Storage Viamed 24 Aug 2016
	Process: 7720 Audit 08 Training Viamed 24 Aug 2016
	Process: 7721 Audit 09 Goods Inward And Product Identity Viamed 24 Aug 2016
t .	III

Process: 7722 Audit 10 Documentation Control Viamed 24 Aug 2016 **Process: 7723** Audit 10b Process Verification Viamed 24 Aug 2016 **Process: 7724** Audit 11 Repairs And Service Viamed 24 Aug 2016

Process: 7725 Audit 12 CE Files Viamed 24 Aug 2016

Process: 7726 Audit 14 Complaints And Corrective Actions Viamed 24 Aug 2016

Process: 7727 Audit 15 Production Viamed 24 Aug 2016Process: 7728 Audit 17 Internal Audits Viamed 24 Aug 2016Process: 7729 Audit 19 Health And Saftey Viamed 24 Aug 2016

Process: 7730 Audit 20 Process Verification To Managment Viamed 24 Aug 2016

Process: 7731 Audit 21 Audit Of Audit Viamed 24 Aug 2016

Process: 7732 Audit 22 Post Market Survellance Viamed 24 Aug 2016

Process: 7733 Audit 23 Analysis Of Data Viamed 24 Aug 2016

Process: 6828

Process: 22 Company Policys 16 Feb 2016

Process: 7754

Process: 7762 Audit 01 Picking Packing VST 08 Feb 2017
Process: 7763 Audit 02 Contract Review VST 08 Feb 2017
Process: 7764 Audit 03 Design Control VST 08 Feb 2017
Process: 7765 Audit 05 Pagebooing Suppliers VST 08 Feb 201

Process: 7765 Audit 05 Purchasing Suppliers VST 08 Feb 2017

Process: 7766 Audit 06 Calibration VST 08 Feb 2017

Process: 7767 Audit 07 Handling And Storage VST 08 Feb 2017

Process: 7768 Audit 08 Training VST 08 Feb 2017

Process: 7769 Audit 09 Goods Inward And Product Identity VST 08 Feb 2017

Process: 7770 Audit 10 Documentation Control VST 08 Feb 2017 **Process: 7771** Audit 10b Process Verification VST 08 Feb 2017 **Process: 7772** Audit 11 Repairs And Service VST 08 Feb 2017

Process: 7773 Audit 12 CE Files VST 08 Feb 2017

Process: 7774 Audit 14 Complaints And Corrective Actions VST 08 Feb 2017

Process: 7775 Audit 15 Production VST 08 Feb 2017Process: 7776 Audit 17 Internal Audits VST 08 Feb 2017Process: 7777 Audit 19 Health And Saftey VST 08 Feb 2017

Process: 7778 Audit 20 Process Verification To Managment VST 08 Feb 2017

Process: 7779 Audit 21 Audit Of Audit VST 08 Feb 2017

Process: 7780 Audit 22 Post Market Survellance VST 08 Feb 2017

Process: 7781 Audit 23 Analysis Of Data VST 08 Feb 2017 **Process: 7808** Ensure All Invoice Correctly Tagged 02 Jun 2017

Process: 6886 Responsibility Allocation: VIAMED Sales And Marketing Sales Viamed

Medical Export 09 Mar 2016

Process: 6887 Responsibility Allocation : VIAMED Sales And Marketing Sales Viamed Automotive Export 09 Mar 2016

Process: 7204 Responsibility Allocation: VIAMED Board Directors Meeting Distributor Issues 09 Mar 2016

Process: 24 Responsibility Allocation: Compliance ISO Standards 16 Feb 2016

Process: 28 Supplier Review 16 Feb 2016

Process: 6865 Responsibility Allocation : Non Conformance Effectiveness 09 Mar 2016 **Process: 6866** Internal Process Verification Complete Systems Review 09 Mar 2016

Process: 7172 Responsibility Allocation : CE Technical Files 09 Mar 2016

Process: 7782 Remove Started But Not Used Order Numbers 08 Feb 2017

Process: 7090 Responsibility Allocation : Office Procedures 09 Mar 2016

Process: 7138 Non Conformance Issues Any New QC21 Forms 09 Mar 2016

Process: 57 Temporary Stock Notices 17 Feb 2016 **Process: 5854** Stock FAQ Admin List 17 Feb 2016

Process: 7043 Responsibility Allocation: Planning of product realization 09 Mar 2016

Process: 7045 Responsibility Allocation : Design and Development 09 Mar 2016

Process: 38 Audits Up to Date and Confirm next years Audit schedule 16 Feb 2016

Process: 5877 Review Company Data 17 Feb 2016

Process: 6904 Responsibility Allocation : Sales And Marketing Internal sales 09 Mar 2016

Process: 6944 Responsibility Allocation : Stock Meeting 09 Mar 2016

Process: 7846 ISO System Management Review Viamed 26 Sep 2017

Process: 7834 Financial Review 20 Sep 2017 **Process: 26** Company Resources 16 Feb 2016

Process: 7070 Management Review 09 Mar 2016

Process: 7837 Review External Parties Influencing The QMS VST / Viamed 23 Sep 2017

Process: 5887 Review ISO/EN Documents 24 Feb 2016

Process: 5889 Responsibility Allocation : Audit And Task - Audit 24 Feb 2016

Process: 7071 Post Market Surveillance 09 Mar 2016

Process: 7093 BSI Audits Calander 09 Mar 2016

Process: 7829

Process: 7670 Humanmed general Issues 09 Mar 2016

Process: 6821 Responsibility Allocation : VIAMED Management Meeting Supplier Review 09 Mar 2016

Process: 6831 Responsibility Allocation : VIAMED Management Meeting Supplier Review - Min / Max - Re-Orders 09 Mar 2016

Process: 6833 Responsibility Allocation : VIAMED Management Meeting MDA Recalls 09 Mar 2016

Process: 6834 Responsibility Allocation : VIAMED Management Meeting Additional Purchase Orders 09 Mar 2016

Process: 6836 Responsibility Allocation : VIAMED Management Meeting Research and Development rnd 09 Mar 2016

Process: 6920 Responsibility Allocation : VIAMED Sales And Marketing Price Lists UK 09 Mar 2016

Process: 6924 Responsibility Allocation : VIAMED Sales And Marketing Price Lists Export 09 Mar 2016

Process: 6935 Responsibility Allocation : VIAMED Sales And Marketing Products to be Marketed 09 Mar 2016

Process: 6936 Responsibility Allocation : VIAMED Sales And Marketing NHS Supplies Future Technology 09 Mar 2016

Process: 6941 Responsibility Allocation : VIAMED Sales And Marketing New Potential Products 09 Mar 2016

Process: 7039 Responsibility Allocation: Provision of Resources 09 Mar 2016

Process: 7187 Responsibility Allocation : VIAMED Board Directors Meeting Profiability 09 Mar 2016

Process: 7196 Responsibility Allocation : VIAMED Board Directors Meeting Stock Levels 09 Mar 2016

Process: 6871 ISO14001 Environmental management systems 09 Mar 2016

Process: 7830 Review Q.A. Failures Report 18 Sep 2017

Process: 7848 Review ISO Scopes 27 Sep 2017

Process: 7849 Review Product Failures New Codes 28 Sep 2017 **Process: 7862** Review The Audit Calender Screen 04 Oct 2017

Process: 7877 Disaster Planning 21 Oct 2017

Process: 7879 Software Validation Scheduled Tasks And Audits 22 Oct 2017

Process: 7876 Maintain Update Of ISO Route Maps 21 Oct 2017

Process: 7878 Review Possible Upcoming Regulation Changes 22 Oct 2017

Process: 7885 Audit 04 Accounts and Finance Viamed 23 Oct 2017

Process: 7886 Audit 18 Management Review Viamed 24 Oct 2017

Process: 7887 Audit 18 Management Review VST 24 Oct 2017

Process: 7889 Audit 24 Servicing Viamed 24 Oct 2017

Process: 7888 Review Processes Linked To VOPs And Audits 24 Oct 2017

Process: 7965 VST Feedback 29 Oct 2020

Process: 7964 Check Roles And Tasks For Incomplete Data 29 Oct 2020

Process: 7980 Review Gov Website For Applicable Required Standards ISO9001 15 Nov

Process: 7972 ISO System Management Review Vst 26 Oct 2021

Process: 7973 VST Product Performance - Customers 27 Oct 2021

Process: 7974 VST Product Performance - Suppliers 27 Oct 2021

Process: 7977 Review The Agenda For The Management Review / Board Meeting Prior To

The Annual Meeting 11 Nov 2021

Process: 7978 Regulatory Requirements and Review of QC21 form template 11 Nov 2021

Process: 7981 Review Process Updates For Risk To Systems 18 Nov 2021

Process: 8012 VAT Return Viamed Properties 06 Apr 2022

Process: 8014 Review VIAMED Product Feedback Positive 25 Jul 2022

Process: 8015 Review VST Product Feedback Positive 25 Jul 2022

Process: 8016 Review VIAMED Customer Feedback Positive 25 Jul 2022

Process: 8017 Review VST Customer Feedback Positive 25 Jul 2022

Process: 8018 Wednesday Meeting 09 Aug 2022

Process: 8019 Audit 04 Accounts And Finance VST 14 Sep 2022

ID73320 Audit 18 Management Review

Process: 55 Business Continuity Plan 17 Feb 2016

Process: 23 Company Objectives 16 Feb 2016

Process: 6813 Management Meeting Turnover Report 09 Mar 2016

Process: 27 Management Reviews And Quality Audits 16 Feb 2016

Process: 22 Company Policys 16 Feb 2016

Process: 7750 Meeting With Management 14 Oct 2016

Process: 7793 Team Review Meeting 16 Mar 2017

Process: 7753 Management Meeting Warehouse 22 Nov 2016

Process: 6861 Management Meeting Review Weekly Meeting 09 Mar 2016

Process: 7833 Importance Of Effective Quality Management 20 Sep 2017

Process: 7834 Financial Review 20 Sep 2017

Process: 26 Company Resources 16 Feb 2016

Process: 30 Responsibility Allocation: MHRA Licences And Notifications 16 Feb 2016

Process: 31 Responsibility Allocation : Notified Body Notifications 16 Feb 2016

Process: 32 MDALL Listings 16 Feb 2016

Process: 7057 Responsibility Allocation : Complaints and Vigilance Notifications 09 Mar 2016

Process: 7070 Management Review 09 Mar 2016

Process: 29 Responsibility Allocation: CMDCAS Updates And Licences 16 Feb 2016

Process: 5889 Responsibility Allocation : Audit And Task - Audit 24 Feb 2016

Process: 7744 FDA Device Establishment Registration And Listing 28 Sep 2016

Process: 7829

Process: 6871 ISO14001 Environmental management systems 09 Mar 2016

Process: 7874 Review For Latest Version Med Dev 2.12. 18 Oct 2017

Process: 7877 Disaster Planning 21 Oct 2017

Process: 7876 Maintain Update Of ISO Route Maps 21 Oct 2017

Process: 7878 Review Possible Upcoming Regulation Changes 22 Oct 2017

Process: 7886 Audit 18 Management Review Viamed 24 Oct 2017

Process: 7887 Audit 18 Management Review VST 24 Oct 2017

Process: 7890 New UPS Rates Needs Checking 24 Oct 2017

Process: 7888 Review Processes Linked To VOPs And Audits 24 Oct 2017

Process: 7895 FDA Device Establishment Registration 29 Oct 2017

Process: 7912 Review The Personel Information We Collect Or Store 20 Sep 2018

Process: 7913 Review Personnel Files 20 Sep 2018

Process: 7918 Backup Jeans Local Folder 08 Nov 2018

Process: 7964 Check Roles And Tasks For Incomplete Data 29 Oct 2020

Process: 7980 Review Gov Website For Applicable Required Standards ISO9001 15 Nov

Process: 7972 ISO System Management Review Vst 26 Oct 2021

Process: 7977 Review The Agenda For The Management Review / Board Meeting Prior To

The Annual Meeting 11 Nov 2021

Process: 7978 Regulatory Requirements and Review of QC21 form template 11 Nov 2021 **Process: 7979** Review The Template Of The QC 21 Form To Ensure It Is Current And Valid

12 Nov 2021

Process: 7981 Review Process Updates For Risk To Systems 18 Nov 2021

Process: 8018 Wednesday Meeting 09 Aug 2022

Process: 8026 Automotive Competitor Price Review 10 Mar 2023
Process: 8025 Check We Do Not Require A EU European Representatives 09 Mar 2023
VOP 05 Supplier Control, Supplier Review, Purchase Orders, Supplier Returns and
Rejection
Process: 6972 UPS Shipping Fuel Surcharge 09 Mar 2016
Process: 28 Supplier Review 16 Feb 2016
Process: 6960
Process: 7784 Check Returns Supplier Envitec 15 Feb 2017
Process: 7785 Check Returns Supplier Teledyne 15 Feb 2017
Process: 7786 Check Returns Supplier Maxtec 15 Feb 2017
Process: 7787 Check Returns All Supplier 15 Feb 2017
Process: 7975 Arrange Teledyne Returns 03 Nov 2021
Process: 7984 Check For Viking Invoices 19 Jan 2022
Process: 8009 Verification Stock Items And Locations 21 Feb 2022
Process: 7991 Verification Purchasing Documentation 07 Feb 2022
Process: 8002 Verification Todays Goods In 17 Feb 2022
Process: 8003 Verification Supplier Delivery Notes 17 Feb 2022
Process: 8030 Purchase Order Invoice Review 23 Jun 2023
Audit 05 Purchasing suppliers
Process: 7707 Send Purchase Orders To Suppliers 13 Jun 2016
Process: 6972 UPS Shipping Fuel Surcharge 09 Mar 2016
Process: 7717 Audit 05 Purchasing Suppliers Viamed 24 Aug 2016
Process: 5850 Purchase Order Log 17 Feb 2016
Process: 7751 VST Purchase Order Log 02 Nov 2016
Process: 7765 Audit 05 Purchasing Suppliers VST 08 Feb 2017
Process: 7794 V1000 Commissions Review 30 Mar 2017
Process: 7745 UPS Invoices Viamed 06 Oct 2016
Process: 7746 UPS Invoices VST 06 Oct 2016
Process: 7747 UPS Invoices Vandagraph 06 Oct 2016
Process: 7790 Humanmed Invoice them For Previous Month 10 Mar 2017
Process: 28 Supplier Review 16 Feb 2016
Process: 6960
Process: 5855 Purchase Order Requirements Teledyne 17 Feb 2016
Process: 5866 UPS Shipping Fuel Surcharge 17 Feb 2016
Process: 5868 Return Goods To Suppliers 17 Feb 2016
Process: 6829 Supplier Review - Outstanding orders 09 Mar 2016
Process: 6832 Supplier Review Future orders 09 Mar 2016
Process: 6848
Process: 6952 Responsibility Allocation : Lost in Shipping Claims 09 Mar 2016
Process: 6971 Responsibility Allocation : Freight Courier Cost Request 09 Mar 2016
Process: 7679 Check Stock Requirements Supplier Teledyne 18 Apr 2016
Process: 7680 Check Stock Requirements Supplier Envited 18 Apr 2016
Process: 7681 Check Stock Requirements Supplier Posey 18 Apr 2016
Process: 7682 Check Stock Requirements Supplier Bluepoint 18 Apr 2016
Process: 7784 Check Returns Supplier Envitec 15 Feb 2017
Process: 7785 Check Returns Supplier Teledyne 15 Feb 2017
Process: 7786 Check Returns Supplier Maxtec 15 Feb 2017
Process: 7787 Check Returns All Supplier 15 Feb 2017
Process: 34 Responsibility Allocation : Insurance Is Upto Date 16 Feb 2016
Process: 7683 Check Stock For Proforma 18 Apr 2016
<u>-</u>
Process: 7882 Purchase Payments 23 Oct 2017
Process: 7956 Teledyne Stock For Vandagraph 27 May 2020
Process: 7975 Arrange Teledyne Returns 03 Nov 2021
Process: 7984 Check For Viking Invoices 19 Jan 2022
Trucess. 7504 Check For Viking Invoices 15 Jun 2022
Process: 7991 Verification Purchasing Documentation 07 Feb 2022 Process: 8003 Verification Supplier Delivery Notes 17 Feb 2022

TD440400	A 11.00 C C XX 11.1
ID113182	Audit 27 Software Validation
	Process: 52 Software Verification Clear Down Backup Emails 16 Feb 2016
	Process: 7668 Responsibility Allocation : Upgrading Intrastats ISO Quality system 09 Mar
	2016
	Process: 7132 Responsibility Allocation : Intrastats Goldmine 09 Mar 2016
	Process: 7851 Software Validation Scan Un-QA Product To Order 01 Oct 2017
	Process: 7852 Software Validation Expired Stock 01 Oct 2017
	Process: 7853 Software Validation Non Sell Able Shelf 01 Oct 2017
	Process: 7854 Software Validation In Production List 01 Oct 2017
	Process: 7855 Software Validation - Production Lists 01 Oct 2017
	Process: 7856 Software Validation Unchecked Orders 01 Oct 2017
	Process: 7857 Software Validation Stock Tracking Check 01 Oct 2017
	Process: 7858 Software Validation Attempt To QA Some Stock 01 Oct 2017
	Process: 7861 Software Validation Of Training Documents Forced Reading 03 Oct 2017
	Process: 7850 Software Validation Scan Incorrect Product 01 Oct 2017
	Process: 7865 Software Validation Conflicting Audits 07 Oct 2017
	Process: 7870 Software Validation Non Conformance Product Risk Feedback Loop 15 Oct
	2017
	Process: 7879 Software Validation Scheduled Tasks And Audits 22 Oct 2017
	Process: 7875 Software Validation Document Control 20 Oct 2017
	Process: 7880 Software Validation Out Of Date Documents 22 Oct 2017
	Process: 7881 Software Validation - Live Orders 22 Oct 2017
	Process: 7892 Audit 27 Software Validation 26 Oct 2017
	Process: 7951 Server Review 05 Mar 2020
	Process: 8013 Software Validation Test Email System 29 Apr 2022
ID01.40C	
ID91486	VOP 27 Software Validation
	Process: 46 Responsibility Allocation : Backup Server Status 16 Feb 2016
	Process: 52 Software Verification Clear Down Backup Emails 16 Feb 2016
	Process: 7851 Software Validation Scan Un-QA Product To Order 01 Oct 2017
	Process: 7852 Software Validation Expired Stock 01 Oct 2017
	Process: 7853 Software Validation Non Sell Able Shelf 01 Oct 2017
	Process: 7854 Software Validation In Production List 01 Oct 2017
	Process: 7855 Software Validation - Production Lists 01 Oct 2017
	Process: 7856 Software Validation Unchecked Orders 01 Oct 2017
	Process: 7857 Software Validation Stock Tracking Check 01 Oct 2017
	Process: 7858 Software Validation Attempt To QA Some Stock 01 Oct 2017
	Process: 7861 Software Validation Of Training Documents Forced Reading 03 Oct 2017
	Process: 7850 Software Validation Scan Incorrect Product 01 Oct 2017
	Process: 7865 Software Validation Conflicting Audits 07 Oct 2017
	Process: 7870 Software Validation Non Conformance Product Risk Feedback Loop 15 Oct
	2017
	Process: 7879 Software Validation Scheduled Tasks And Audits 22 Oct 2017
	Process: 7875 Software Validation Document Control 20 Oct 2017
	Process: 7880 Software Validation Out Of Date Documents 22 Oct 2017
	Process: 7881 Software Validation - Live Orders 22 Oct 2017
	Process: 7892 Audit 27 Software Validation 26 Oct 2017
	Process: 8013 Software Validation Test Email System 29 Apr 2022
ID22062	VM3COP00.00 VOP00.00 VST Quality Statement policy and objectives
	Process: 23 Company Objectives 16 Feb 2016
	Process: 7827 Review The Quality Policy VST 16 Sep 2017
	Process: 7833 Importance Of Effective Quality Management 20 Sep 2017
ID25632	VOP 17 Design Research and Development
	Process: 42 Responsibility Allocation: Design Documentation 16 Feb 2016
	Process: 43 Responsibility Allocation: Product Post Market Survelance 16 Feb 2016
	Process: 6975 Responsibility Allocation : Projects 09 Mar 2016
	Process: 7045 Responsibility Allocation: Design and Development 09 Mar 2016
<u> </u>	1 10ccss. 7040 Incorporation of Americanian Persign and Development 03 Mai 2010

ID111315	Audit 03 Design Control
	Process: 7716 Audit 03 Design Control Viamed 24 Aug 2016
	Process: 42 Responsibility Allocation : Design Documentation 16 Feb 2016
	Process: 7764 Audit 03 Design Control VST 08 Feb 2017
	Process: 7043 Responsibility Allocation : Planning of product realization 09 Mar 2016
	Process: 7045 Responsibility Allocation : Design and Development 09 Mar 2016
	Process: 7047 Responsibility Allocation : Production and service provision 09 Mar 2016
	Process: 6942 Responsibility Allocation : Co ordination of Implementation 09 Mar 2016
	Process: 7173 Responsibility Allocation : Material Generation 09 Mar 2016
	Process: 5887 Review ISO/EN Documents 24 Feb 2016
	Process: 7919 Send Debtors Overview To Derek 06 Dec 2018
ID67997	Audit 23 Analysis of Data
	Process: 27 Management Reviews And Quality Audits 16 Feb 2016
	Process: 7733 Audit 23 Analysis Of Data Viamed 24 Aug 2016
	Process: 7781 Audit 23 Analysis Of Data VST 08 Feb 2017
	Process: 5877 Review Company Data 17 Feb 2016
	Process: 6931 Customer Complaints 09 Mar 2016
	Process: 7839 Review VIAMED Feedback - Customer Complaints 23 Sep 2017
	Process: 7838 Review VIAMED Feedback - Customer Feedback Negative 23 Sep 2017
	Process: 26 Company Resources 16 Feb 2016
	Process: 7070 Management Review 09 Mar 2016
	Process: 7713 Review Roles And Responsibilitys 17 Aug 2016
	Process: 7837 Review External Parties Influencing The QMS VST / Viamed 23 Sep 2017
	Process: 7840 Review VST Feedback - Customer Feedback Negative 23 Sep 2017
	Process: 7841 Review VST Feedback - Customer Complaints 23 Sep 2017
	Process: 7842 Review VIAMED Product Feedback Negative 23 Sep 2017
	Process: 7843 Review VST Product Feedback Negative 23 Sep 2017
	Process: 7071 Post Market Surveillance 09 Mar 2016
	Process: 7830 Review Q.A. Failures Report 18 Sep 2017
	Process: 7849 Review Product Failures New Codes 28 Sep 2017
	Process: 7862 Review The Audit Calender Screen 04 Oct 2017
	Process: 7930 Review Flow Of Data 12 Mar 2019
	Process: 7969 Weee Waste Reporting 23 Aug 2021
ID93320	VOP 02 Personnel and Responsibility , Staff and Staffing Issues, Training, Roles and
	Tasks
	Process: 39 Environmental Policy Document Review 16 Feb 2016
	Process: 7741 Review Ethical Policy 14 Sep 2016
	Process: 6839 Responsibility Allocation: Personnel Holidays and Time Adjustments 09 Mar
	2016
	Process: 5881 Training Records Review 18 Feb 2016
	Process: 5904 Taking On New Staff 02 Mar 2016
	Process: 6837 Personnel Requirements and Training 09 Mar 2016
	Process: 6877 Responsibility Allocation : Alarm Key Holders 09 Mar 2016
	Process: 6906 Responsibility Allocation : Time Working Away 09 Mar 2016
	Process: 6928 Responsibility Allocation : Staff 09 Mar 2016
	Process: 7074
	Process: 7042 Responsibility Allocation : Work Environment 09 Mar 2016
	Process: 5934 Responsibility Allocation : Staff Training 05 Mar 2016
	Process: 5874 Childcare Vouchers Edenred 17 Feb 2016
	Process: 7753 Management Meeting Warehouse 22 Nov 2016
	Process: 34 Responsibility Allocation: Insurance Is Upto Date 16 Feb 2016
	1
	Process: 5869 Responsibility Allocation: Legal Company Car Registration 17 Feb 2016
	Process: 6841 Responsibility Allocation : Grants 09 Mar 2016
	Process: 6843
	Process: 6861 Management Meeting Review Weekly Meeting 09 Mar 2016
	Process: 30 Responsibility Allocation : MHRA Licences And Notifications 16 Feb 2016
	Process: 31 Responsibility Allocation : Notified Body Notifications 16 Feb 2016
	II

2/10/2023, 11:0	5 QMS Route Map Viamed Ltd 15015465:2016
	Process: 32 MDALL Listings 16 Feb 2016 Process: 7022 Responsibility Allocation & Management commitment to ISO 00 May 2016
	Process: 7033 Responsibility Allocation : Management commitment to ISO 09 Mar 2016 Process: 7037 Responsibility Allocation : Responsibility, authority and communication 09 Mar 2016
	Process: 7057 Responsibility Allocation : Complaints and Vigilance Notifications 09 Mar 2016
	Process: 7713 Review Roles And Responsibilitys 17 Aug 2016 Process: 7837 Review External Parties Influencing The QMS VST / Viamed 23 Sep 2017 Process: 29 Responsibility Allocation : CMDCAS Updates And Licences 16 Feb 2016 Process: 7848 Review ISO Scopes 27 Sep 2017
	Process: 7891 Fire Alarm Evacuation Drill 25 Oct 2017 Process: 7908 Private Information Data 27 Jul 2018
	Process: 7907 Annual Review Doc Management 27 Jul 2018 Process: 7937 Diversity Impact Assessment 27 Jun 2019
	Process: 7961 R D Room - Tidy, Empty Bins, Remove Cups. Caution Around Oxygen Supply 05 Oct 2020
	Process: 7982 Check There Are No Changes To Employment Law 21 Nov 2021 Process: 7983 To Check On Line And See If There Have Been Any Changes To Gdpr We Need To Be Aware Of. 21 Nov 2021
ID17423	VM3COP02 Organisation Responsibilities Viamed Process: 6967 Responsibility Allocation: VIAMED Stock Meeting Repairs Review - Pulse Oximetry Sensors 09 Mar 2016 Process: 7000 Revel Mail Responsibilities Viamed Process: 7000 Revel Mail Responsibilities Viamed
	Process: 7900 Royal Mail - Mail Retention Form 29 Mar 2018
ID119029	VOP 18 Maintenance Building, Fabric and Infrastructure Process: 5856 Cleaning The Kitchen 17 Feb 2016
	Process: 5853 Vacuuming Of The Office, Hall And Meeting Room 17 Feb 2016 Process: 5900 Cleaning Of Office Windows 25 Feb 2016
	Process: 5878 Empty Office Bins 18 Feb 2016
	Process: 5912 Responsibility Allocation : Main Recycle Bins 03 Mar 2016
	Process: 5906 Empty Paper Bins 03 Mar 2016
	Process: 7805 Empty Kitchen Bins 22 May 2017
	Process: 5909 Empty Warehouse Bins 03 Mar 2016
	Process: 7706 Update Virus Software And Scan For Viruses 10 Jun 2016
	Process: 7802 Clean Kitchen Sides 22 May 2017
	Process: 7803 Dishwashing 22 May 2017
	Process: 7804 Sweep Kitchen Floor 22 May 2017
	Process: 7806 Watering Plants 22 May 2017 Process: 7807
	Process: 54 Responsibility Allocation : Gents Toilets 17 Feb 2016
	Process: 5907 Hoover Warehouse 03 Mar 2016
	Process: 5908 Sweep Warehouse 03 Mar 2016
	Process: 5910 Clean Duckets 03 Mar 2016
	Process: 5911 Clear Cardboard 03 Mar 2016
	Process: 7698 Clean Toilets 17 May 2016
	Process: 7131 Responsibility Allocation: Intrastats Opera 09 Mar 2016
	Process: 7133 Responsibility Allocation : Intrastats Contact Manager 09 Mar 2016 Process: 7132 Responsibility Allocation : Intrastats Goldmine 09 Mar 2016
	Process: 7896 Tree In Car Park 22 Dec 2017
ID21000	
ID21800	VM3COP19 Health and Safety Process: 6855 Risk Assessment HSE 09 Mar 2016
ID130426	Viamed Top Level Quality Objectives Process: 23 Company Objectives 16 Feb 2016
ID77875	VOP 03 Contract Review, Enquires, Office Processes
	Process: 5 Responsibility Allocation : Processing Of Sales Orders 16 Feb 2016
	Process: 10 Distribution Of Emails 16 Feb 2016
	Process: 36 Emailing Of Invoices 16 Feb 2016
	· ·

Process: 5892 Checking EBay And Amazon For Orders And Messages 25 Feb 2016

Process: 5894 Checking Of Active List 25 Feb 2016

Process: 7 Responsibility Allocation: Checking Of Sales Orders 16 Feb 2016

Process: 5943 Check Cardea And Multiquote 08 Mar 2016

Process: 5891 Processing Of Repair Quotes And Orders 25 Feb 2016

Process: 11 Distribution Of Post 16 Feb 2016 **Process: 2** Answering Telephones 16 Feb 2016

Process: 37 West Yorkshire Ambulance Stock 16 Feb 2016 **Process: 5948** Adding New Accounts To Opera 08 Mar 2016

Process: 5949 Filling Credit Card Slips 08 Mar 2016

Process: 6 Responsibility Allocation: Updating Contact Management System 16 Feb 2016

Process: 5895 Responsibility Allocation : Completing Office Job List 25 Feb 2016

Process: 5875 Check Paypal For Orders 17 Feb 2016

Process: 5944 Responsibility Allocation : Chasing Lost Customers 08 Mar 2016

Process: 3 Responsibility Allocation : Meeting And Greeting Visitors To The Company 16 Feb 2016

Process: 4 Responsibility Allocation: Assisting With Refreshments For Visitors 16 Feb 2016

Process: 7676 PDFing Of Invoices Viamed 17 Mar 2016

Process: 9 Distribution Of Faxes 16 Feb 2016

Process: 7696 Send VIAMED Delivery Notifications 28 Apr 2016

Process: 5857 Customer Service Logs 17 Feb 2016

Process: 5893 Answering Website Questions 25 Feb 2016

Process: 7678 Check Catalog 360 Circle For Quotes And Orders 08 Apr 2016

Process: 15 Filing and Archiving 16 Feb 2016

Process: 5899 Proforma And Quote Chasing 25 Feb 2016

Process: 7710 Responsibility Allocation: Proforma And Quote Processing 29 Jun 2016

Process: 7707 Send Purchase Orders To Suppliers 13 Jun 2016

Process: 14 Fax Paper 16 Feb 2016

Process: 5882 Responsibility Allocation : Send Post To Humanmed 24 Feb 2016 **Process: 7734** Responsibility Allocation : Humanmed Order Processing 25 Aug 2016

Process: 5850 Purchase Order Log 17 Feb 2016

Process: 7693 Collect Repair Filing From Warehouse 22 Apr 2016

Process: 7677

Process: 21 Office Sales Projects 16 Feb 2016

Process: 8 Responsibility Allocation : Order And Status Liaison With Customers 16 Feb 2016 **Process: 12** Responsibility Allocation : Sales And Technical Information Processing 16 Feb

2016

Process: 16 Responsibility Allocation : Photocopying 16 Feb 2016

Process: 17

Process: 20 Processing Of Mail Shots 16 Feb 2016

Process: 5896 Responsibility Allocation: Ensuring ORD's Are Taken To Goods Out And

Invoices Are Retrieved 25 Feb 2016

Process: 5897 Responsibility Allocation: Franking Mail 25 Feb 2016

Process: 5901 Link Call Log Contacts To The CRM 02 Mar 2016

Process: 5913 Check For Humanmed Orders In Logistics Mailbox 03 Mar 2016

Process: 5947 Responsibility Allocation : Search For Distributors 08 Mar 2016 **Process: 6958** Responsibility Allocation : Shipped Order Queries 09 Mar 2016

Process: 7686 Thorough Checking Of Awaiting Action Tray - Priority 8s 21 Apr 2016

Process: 7699 Shred Sensitive Paperwork In JL Office 19 May 2016

Process: 7705 Checking For Uploaded Files 08 Jun 2016

Process: 7709 Delivered not Invoiced 28 Jun 2016

Process: 7712 Review Inward Payments 01 Jul 2016

Process: 7735 Ensure SOR's Are Followed Up 01 Sep 2016

Process: 7751 VST Purchase Order Log 02 Nov 2016

Process: 7758 Check For GHX Orders 17 Jan 2017

Process: 7760 Send Service Offers 31 Jan 2017

Process: 7761 Send VST Delivery Notifications 01 Feb 2017

Process: 7783 PDF VST Invoices And Purchase Orders 10 Feb 2017

Process: 7792 Shipped Order Success Report 13 Mar 2017 **Process: 7795** Answering UK Web Questions 27 Apr 2017

Process: 7822 Review Oxylink Stock 26 Jul 2017

Process: 5876 E.Commerce Cardea And Multiquote 17 Feb 2016

Process: 5873 Distributor Contract Reviews 17 Feb 2016

Process: 5885 Responsibility Allocation : Monthly Reports 24 Feb 2016

Process: 6938 Responsibility Allocation: Customer Database Updates 09 Mar 2016 **Process: 6940** Responsibility Allocation: Customer Ongoing task List 09 Mar 2016

Process: 6956 Responsibility Allocation : Sales Order Issues 09 Mar 2016

Process: 5866 UPS Shipping Fuel Surcharge 17 Feb 2016

Process: 6952 Responsibility Allocation: Lost in Shipping Claims 09 Mar 2016 **Process: 6971** Responsibility Allocation: Freight Courier Cost Request 09 Mar 2016

Process: 7692 Responsibility Allocation: Take Complete Repair Paperwork To Office 22 Apr

2016

Process: 7796 Review Franking Label Errors 08 May 2017

Process: 6916 Responsibility Allocation: Service exisiting 09 Mar 2016 **Process: 6917** Responsibility Allocation : Service extension 09 Mar 2016

Process: 7863 Maintain Repair Codes List 05 Oct 2017

Process: 7872 Embargo Countries NOT Allowed To Sell To 16 Oct 2017

Process: 7890 New UPS Rates Needs Checking 24 Oct 2017

Process: 7893 VST Price Lists 28 Oct 2017

Process: 7894 VST Customer Agreements 28 Oct 2017 **Process: 7901** UPS Exceptions Checkup 20 Apr 2018

Process: 7957 Warehouse Requests 29 May 2020

Process: 7959 Audit 16 Sales And Marketing Viamed 28 Sep 2020 **Process: 7970** Proforma And Quote Chasing Ryan 31 Aug 2021

Process: 7971 Proforma And Quote Chasing Steve Hardaker 31 Aug 2021

Process: 7988 Verification Contact Details Internal CRM 07 Feb 2022

Process: 7989 Verification Contact Details Accounts 07 Feb 2022

Process: 7990 Verification Invoice Details Accounts 07 Feb 2022

Process: 8020 Checking Proformas And Quotes Vandagraph To The Bank 05 Dec 2022 **Process: 8023** Vandagraph Check Shopify Order Delivery Notifications 17 Feb 2023

Process: 8026 Automotive Competitor Price Review 10 Mar 2023

ID69328 Audit 02 Contract Review and Sales Order Processing

Process: 5 Responsibility Allocation: Processing Of Sales Orders 16 Feb 2016

Process: 36 Emailing Of Invoices 16 Feb 2016

Process: 5892 Checking EBay And Amazon For Orders And Messages 25 Feb 2016

Process: 5894 Checking Of Active List 25 Feb 2016

Process: 7 Responsibility Allocation: Checking Of Sales Orders 16 Feb 2016

Process: 5943 Check Cardea And Multiquote 08 Mar 2016

Process: 5891 Processing Of Repair Quotes And Orders 25 Feb 2016

Process: 2 Answering Telephones 16 Feb 2016

Process: 37 West Yorkshire Ambulance Stock 16 Feb 2016

Process: 5945 Responsibility Allocation : Sending Samples 08 Mar 2016

Process: 5946 Responsibility Allocation : Sending Sale Or Returns 08 Mar 2016

Process: 5948 Adding New Accounts To Opera 08 Mar 2016

Process: 5949 Filling Credit Card Slips 08 Mar 2016

Process: 5895 Responsibility Allocation : Completing Office Job List 25 Feb 2016

Process: 5875 Check Paypal For Orders 17 Feb 2016

Process: 7675 Responsibility Allocation: Ordering Demo Stock For Humanmed Reps 11 Mar 2016

Process: 5944 Responsibility Allocation : Chasing Lost Customers 08 Mar 2016

Process: 3 Responsibility Allocation: Meeting And Greeting Visitors To The Company 16

Process: 4 Responsibility Allocation: Assisting With Refreshments For Visitors 16 Feb 2016

Process: 7676 PDFing Of Invoices Viamed 17 Mar 2016

Process: 7696 Send VIAMED Delivery Notifications 28 Apr 2016

Process: 5893 Answering Website Questions 25 Feb 2016

Process: 7678 Check Catalog 360 Circle For Quotes And Orders 08 Apr 2016

Process: 5899 Proforma And Quote Chasing 25 Feb 2016

Process: 7710 Responsibility Allocation: Proforma And Quote Processing 29 Jun 2016

Process: 14 Fax Paper 16 Feb 2016

Process: 5882 Responsibility Allocation : Send Post To Humanmed 24 Feb 2016

Process: 7715 Audit 02 Contract Review Viamed 24 Aug 2016

Process: 7734 Responsibility Allocation: Humanmed Order Processing 25 Aug 2016

Process: 7677

Process: 6954 Back Orders Review - By Customer 09 Mar 2016

Process: 8 Responsibility Allocation : Order And Status Liaison With Customers 16 Feb 2016

Process: 5896 Responsibility Allocation : Ensuring ORD`s Are Taken To Goods Out And

Invoices Are Retrieved 25 Feb 2016

Process: 5897 Responsibility Allocation: Franking Mail 25 Feb 2016

Process: 5913 Check For Humanmed Orders In Logistics Mailbox 03 Mar 2016

Process: 5947 Responsibility Allocation : Search For Distributors 08 Mar 2016

Process: 6958 Responsibility Allocation : Shipped Order Queries 09 Mar 2016

Process: 7686 Thorough Checking Of Awaiting Action Tray - Priority 8s 21 Apr 2016

Process: 7709 Delivered not Invoiced 28 Jun 2016

Process: 7712 Review Inward Payments 01 Jul 2016

Process: 7735 Ensure SOR`s Are Followed Up 01 Sep 2016

Process: 7758 Check For GHX Orders 17 Jan 2017

Process: 7761 Send VST Delivery Notifications 01 Feb 2017

Process: 7783 PDF VST Invoices And Purchase Orders 10 Feb 2017

Process: 7795 Answering UK Web Questions 27 Apr 2017

Process: 7822 Review Oxylink Stock 26 Jul 2017

Process: 7791 Price List Check 10 Mar 2017

Process: 7763 Audit 02 Contract Review VST 08 Feb 2017

Process: 7808 Ensure All Invoice Correctly Tagged 02 Jun 2017

Process: 5872 Check Sale Or Returns Export 17 Feb 2016

Process: 5871 Check Sale Or Returns 17 Feb 2016

Process: 5876 E.Commerce Cardea And Multiquote 17 Feb 2016

Process: 7782 Remove Started But Not Used Order Numbers 08 Feb 2017

Process: 6956 Responsibility Allocation : Sales Order Issues 09 Mar 2016

Process: 6921 Responsibility Allocation: Customer pricing agreements 09 Mar 2016

Process: 6922

Process: 6959 Responsibility Allocation: Sales Forward Orders Review 09 Mar 2016

Process: 7801 VST Price Review 17 May 2017

Process: 5905 Responsibility Allocation : Price Checking 02 Mar 2016

Process: 6950

Process: 7697 Yearly Pricing Review 09 May 2016

Process: 7670 Humanmed general Issues 09 Mar 2016

Process: 7872 Embargo Countries NOT Allowed To Sell To 16 Oct 2017

Process: 7893 VST Price Lists 28 Oct 2017

Process: 7894 VST Customer Agreements 28 Oct 2017

Process: 7936 B2B Router / Peppol Responsibilitys 19 Jun 2019

Process: 7941 Check Leaflets, Letterhead And Other Paperwork To See If The Correct BSI

Logo Is In Use. Remove All Old If Found. 23 Sep 2019

Process: 7953 Vandagraph Delivery Notifications 26 May 2020

Process: 7954 Vandagraph Email Of Invoices 26 May 2020

Process: 7955 Vandagraph Shipper SignOff Collection 26 May 2020

Process: 7970 Proforma And Quote Chasing Ryan 31 Aug 2021

Process: 7971 Proforma And Quote Chasing Steve Hardaker 31 Aug 2021

Process: 8005 Verification Of SRS Information added 17 Feb 2022

Process: 7988 Verification Contact Details Internal CRM 07 Feb 2022

Process: 7989 Verification Contact Details Accounts 07 Feb 2022

Process: 8020 Checking Proformas And Quotes Vandagraph To The Bank 05 Dec 2022

Process: 8023 Vandagraph Check Shopify Order Delivery Notifications 17 Feb 2023

10/2023, 11:	US QIMS Route Map Viamed Ltd 15013463:2016
	Process: 8027 Update Pricing For Viamed Shopify 11 Apr 2023
	Process: 8028 Viamed Shopify Sales Report Export 11 Apr 2023
ID75475	VOP 19 Feedback Customer Complaints Vigilance and Notifications Viamed Ltd
	Process: 7743 Customer Complaints Paper File 26 Sep 2016
	Process: 7671 Humanmed Non Conformances 09 Mar 2016
	Process: 6931 Customer Complaints 09 Mar 2016
	Process: 7839 Review VIAMED Feedback - Customer Complaints 23 Sep 2017
	Process: 7838 Review VIAMED Feedback - Customer Feedback Negative 23 Sep 2017
	Process: 7070 Management Review 09 Mar 2016
	Process: 7840 Review VST Feedback - Customer Feedback Negative 23 Sep 2017
	Process: 7841 Review VST Feedback - Customer Complaints 23 Sep 2017
	Process: 7842 Review VIAMED Product Feedback Negative 23 Sep 2017
	Process: 7843 Review VST Product Feedback Negative 23 Sep 2017
	Process: 7174
	Process: 7175
	Process: 7179
	Process: 7874 Review For Latest Version Med Dev 2.12. 18 Oct 2017
	Process: 7954 Vandagraph Email Of Invoices 26 May 2020
	Process: 7979 Review The Template Of The QC 21 Form To Ensure It Is Current And Valid
	12 Nov 2021
ID69457	Audit 16 Sales and Marketing
	Process: 21 Office Sales Projects 16 Feb 2016
	Process: 17
	Process: 40 Responsibility Allocation : Calender 16 Feb 2016
	Process: 5870 Book Arab Health 17 Feb 2016
	Process: 19 Maintaining Leaflet Stocks 16 Feb 2016
	Process: 20 Processing Of Mail Shots 16 Feb 2016
	Process: 5873 Distributor Contract Reviews 17 Feb 2016
	Process: 5885 Responsibility Allocation : Monthly Reports 24 Feb 2016
	Process: 5883 **Responsibility Allocation : Monthly Sales Report 11 Sep 2023
	Process: 6888 Viamed Automotive UK 09 Mar 2016
	Process: 6898 GHX Web Pricing 09 Mar 2016
	Process: 5884 Responsibility Allocation : Monthly Report 24 Feb 2016
	Process: 5886 Responsibility Allocation : Monthly Report 24 Feb 2016
	Process: 6891 Responsibility Allocation : Exhibitions Co-ordinator 09 Mar 2016
	Process: 7909 EAN GTIN Online Database 06 Aug 2018
	Process: 7920 Sales Warnings 20 Dec 2018
	Process: 7927 Contract Pricing Review 14 Feb 2019
	Process: 7926 Sales Forecasts Export 22 Jan 2019
	Process: 7921 VST Bags And Grey Sensor 03 Jan 2019
	Process: 7925 Providing Ebay Feedback 16 Jan 2019
	Process: 7916 Google Webmaster Tools 16 Oct 2018
	Process: 7931 Competitor Pricing 14 Mar 2019
	Process: 7949 **Sales Projects Send To Sales Team 05 Sep 2023 Process: 7047 9010004 J. L. C.C.P. Oxygon Sensor Ordons 04 Mar 2020
	Process: 7947 8010004 - JJ-CCR Oxygen Sensor Orders 04 Mar 2020
	Process: 7948 8010006 - REVo Oxygen Sensor Orders 04 Mar 2020
	Process: 7950 Envited Oxygen Sensor Parts Stock Check 05 Mar 2020 Process: 7050 Audit 16 Salas And Marketing Viernad 28 San 2020
	Process: 7959 Audit 16 Sales And Marketing Viamed 28 Sep 2020
	Process: 7960 Audit 16 Sales And Marketing VST 28 Sep 2020 Process: 9021 Tendors Povious 02 Aug 2022
	Process: 8031 Tenders Review 02 Aug 2023
ID88809	VOP 07 Stock Control, Handling, Control of Labelling, Storage, Movement
	Process: 6973 Responsibility Allocation : Stock Transfers. (QC19) 09 Mar 2016
	Process: 7675 Responsibility Allocation : Ordering Demo Stock For Humanmed Reps 11 Ma
	2016
	Process: 5872 Check Sale Or Returns Export 17 Feb 2016
	Process: 5871 Check Sale Or Returns 17 Feb 2016
	Process: 5855 Purchase Order Requirements Teledyne 17 Feb 2016

Process: 5858 Opera Stock Adjustments 17 Feb 2016 **Process: 5868** Return Goods To Suppliers 17 Feb 2016

Process: 5935 Stock Allocations 05 Mar 2016

Process: 6829 Supplier Review - Outstanding orders 09 Mar 2016

Process: 6832 Supplier Review Future orders 09 Mar 2016

Process: 6840 Process: 6848

Process: 6850 Current Stock Levels 09 Mar 2016

Process: 6945 Missing Stock or Adjustments 09 Mar 2016

Process: 6955 Production Requirements 09 Mar 2016

Process: 7046 Responsibility Allocation : Stock Purchasing 09 Mar 2016

Process: 7051 Responsibility Allocation: Control of nonconforming product 09 Mar 2016

Process: 7673 Check Expiry Dated Stock 09 Mar 2016

Process: 7679 Check Stock Requirements Supplier Teledyne 18 Apr 2016 **Process: 7680** Check Stock Requirements Supplier Envitec 18 Apr 2016 **Process: 7681** Check Stock Requirements Supplier Posey 18 Apr 2016 **Process: 7682** Check Stock Requirements Supplier Bluepoint 18 Apr 2016

Process: 7687 Vandagraph Duckets 21 Apr 2016

Process: 7688

Process: 7689 Move Stock From QA Shelf To Stock Shelf Monday 21 Apr 2016 **Process: 7694** Move Stock From QA Shelf To Stock Shelf Tuesday 28 Apr 2016

Process: 7695 Top Up Quick Shipping Shelves 28 Apr 2016

Process: 7708 Acorn 0014904 17 Jun 2016

Process: 7798 Orders And Items Shipped Per Month 10 May 2017

Process: 6961 Responsibility Allocation : VIAMED Stock Meeting Purchase Order

Requirements 09 Mar 2016

Process: 7683 Check Stock For Proforma 18 Apr 2016

Process: 6968 Responsibility Allocation : VIAMED Stock Meeting Repairs Review - General 09 Mar 2016

Process: 6949 Responsibility Allocation : VIAMED Stock Meeting QA Processing 09 Mar 2016

Process: 6948 Responsibility Allocation : VIAMED Stock Meeting Stock Processing 09 Mar 2016

Process: 6947 Responsibility Allocation : VIAMED Stock Meeting Stock Queries 09 Mar 2016

Process: 7830 Review Q.A. Failures Report 18 Sep 2017

Process: 7864 ESD Work Stations 07 Oct 2017

Process: 7873 On Site Environment Review 18 Oct 2017

Process: 7866 Oxygen Cylinder Check 13 Oct 2017

Process: 7897 Daily O2 Sensors Returns 04 Jan 2018

Process: 7909 EAN GTIN Online Database 06 Aug 2018

Process: 7943 Review Stocks Of 8000004 01 Oct 2019

Process: 7944 Sealant, Glues, Greases, Sprays, Gases And Tapes You Use In Production,

Service And Repairs For Viamed And VST 09 Oct 2019

Process: 7962 VST Supplier QA Results 28 Oct 2020

Process: 7967 VST Stock Count For End April 01 Jul 2021

Process: 7969 Weee Waste Reporting 23 Aug 2021

Process: 8006 Verification Warehouse Unidentified Stock 17 Feb 2022 **Process: 8008** Verification Warehouse Hand Sanitiser 21 Feb 2022

Process: 8009 Verification Stock Items And Locations 21 Feb 2022

Process: 8010 Verification Of Ebay Stock 21 Feb 2022

Process: 8011 Verification Of Demo Stock 21 Feb 2022

Process: 7996 Verification Repairs Older Repairs 07 Feb 2022

Process: 8002 Verification Todays Goods In 17 Feb 2022

Process: 8004 Verification Of Non Conforming Products 17 Feb 2022

Process: 8024 Discontinue/Supersede Stock 01 Mar 2023

ID75943	VOP 20 Goods in Purchases, Returns, Repairs, Inspection / Rejection
	Process: 5938 Responsibility Allocation : Receive Goods 05 Mar 2016
	Process: 5898 Processing Depleted Sensors 25 Feb 2016
	Process: 5879 Responsibility Allocation : Customer Returning Goods On Our UPS Account
	18 Feb 2016
	Process: 7826 Goods In Processes 06 Sep 2017
	Process: 7859 Check POR Files For Items Delivered But Not Removed From File 02 Oct
	2017
	Process: 7976 Decontamination Of Incoming Products And Repairs 08 Nov 2021
ID103501	VM3COP20.01 Post In Distributing the Post
1000001	Process: 11 Distribution Of Post 16 Feb 2016
	Process: 5882 Responsibility Allocation : Send Post To Humanmed 24 Feb 2016
ID701.47	
ID70147	Audit 08 Training, Competence and Human Resources
	Process: 7720 Audit 08 Training Viamed 24 Aug 2016
	Process: 6839 Responsibility Allocation: Personnel Holidays and Time Adjustments 09 Mar
	2016 Process F001 Training Decords Device v 10 Feb 2016
	Process: 5881 Training Records Review 18 Feb 2016
	Process: 5904 Taking On New Staff 02 Mar 2016
	Process: 5936 Wages Calculations 05 Mar 2016 Process: 6937 Personnal Paguiroments and Training 09 Mar 2016
	Process: 6837 Personnel Requirements and Training 09 Mar 2016
	Process: 6851 Review Accident Book 09 Mar 2016
	Process: 6877 Responsibility Allocation: Alarm Key Holders 09 Mar 2016
	Process: 6906 Responsibility Allocation: Time Working Away 09 Mar 2016
	Process: 6928 Responsibility Allocation : Staff 09 Mar 2016 Process: 7074
	Process: 7759 Health Declaration Sheet 23 Jan 2017
	Process: 7768 Audit 08 Training VST 08 Feb 2017
	Process: 5934 Responsibility Allocation : Staff Training 05 Mar 2016 Process: 38 Audits Up to Date and Confirm next years Audit schedule 16 Feb 2016
	Process: 6841 Responsibility Allocation : Grants 09 Mar 2016
	Process: 7070 Management Review 09 Mar 2016
	Process: 7713 Review Roles And Responsibilitys 17 Aug 2016
	Process: 7883 Appraisal 23 Oct 2017
	Process: 7884 Pay Review 23 Oct 2017
	Process: 7908 Private Information Data 27 Jul 2018
	Process: 7907 Annual Review Doc Management 27 Jul 2018
	Process: 7937 Aintual Review Doc Management 27 Jun 2019
	Process: 7951 Server Review 05 Mar 2020
	Process: 7982 Check There Are No Changes To Employment Law 21 Nov 2021
	Process: 7983 To Check On Line And See If There Have Been Any Changes To Gdpr We
	Need To Be Aware Of. 21 Nov 2021
IDC0045	
ID68045	Audit 19 Health and Safety, Working Conditions and Building Fabric Issues
	Process: 5941 Responsibility Allocation: Replace Main Server 07 Mar 2016
	Process: 45 Responsibility Allocation: Main Server Status 16 Feb 2016
	Process: 46 Responsibility Allocation: Backup Server Status 16 Feb 2016
	Process: 7704 Responsibility Allocation : Computer Failure Diagnostics 24 May 2016 Process: 5856 Cleaning The Kitchen 17 Feb 2016
	Process: 7729 Audit 19 Health And Saftey Viamed 24 Aug 2016
	Process: 5853 Vacuuming Of The Office, Hall And Meeting Room 17 Feb 2016 Process: 5900 Cleaning Of Office Windows 25 Feb 2016
	Process: 39 Environmental Policy Document Review 16 Feb 2016
	Process: 7741 Review Ethical Policy 14 Sep 2016
	Process: 7741 Review Editical Policy 14 Sep 2016 Process: 5878 Empty Office Bins 18 Feb 2016
	Process: 5076 Empty Office Bins 16 Feb 2016 Process: 5912 Responsibility Allocation : Main Recycle Bins 03 Mar 2016
	Process: 5912 Responsibility Anocation: Main Recycle Bins 03 Mar 2016 Process: 7821 Controlled Waste Description And Transfer 15 Jun 2017
	Process: 7821 Controlled Waste Description And Transfer 15 Jun 2017 Process: 7820 North Yorkshire Council Waste Transfer 15 Jun 2017
	Process: 5906 Empty Paper Bins 03 Mar 2016

/10/2023, 11:0	QMS Route Map Viamed Ltd ISO13485:2016
	Process: 7805 Empty Kitchen Bins 22 May 2017
	Process: 5909 Empty Warehouse Bins 03 Mar 2016
	Process: 7042 Responsibility Allocation : Work Environment 09 Mar 2016
	Process: 7706 Update Virus Software And Scan For Viruses 10 Jun 2016
	Process: 7802 Clean Kitchen Sides 22 May 2017
	Process: 7803 Dishwashing 22 May 2017
	Process: 7804 Sweep Kitchen Floor 22 May 2017
	Process: 7806 Watering Plants 22 May 2017
	Process: 7807
	Process: 7777 Audit 19 Health And Saftey VST 08 Feb 2017
	Process: 54 Responsibility Allocation : Gents Toilets 17 Feb 2016
	Process: 5907 Hoover Warehouse 03 Mar 2016
	Process: 5908 Sweep Warehouse 03 Mar 2016
	Process: 5910 Clean Duckets 03 Mar 2016
	Process: 5911 Clear Cardboard 03 Mar 2016
	Process: 7687 Vandagraph Duckets 21 Apr 2016
	Process: 7698 Clean Toilets 17 May 2016
	Process: 6849 First Aid 09 Mar 2016
	Process: 6855 Risk Assessment HSE 09 Mar 2016
	Process: 6856 Fire Alarms 09 Mar 2016
	Process: 7092
	Process: 56 Warehouse Outside Heating Guard 17 Feb 2016
	Process: 5919 Check Out Side Drain 05 Mar 2016
	Process: 5921 Clearing Water Downstairs 05 Mar 2016
	Process: 7120 General Maintenance Requirements 09 Mar 2016
	Process: 7742 Boiler Check 26 Sep 2016
	Process: 7756 Carbon Monoxide Alarm 05 Jan 2017
	Process: 48 Responsibility Allocation : Internet 16 Feb 2016
	Process: 49 Responsibility Allocation : Wifi 16 Feb 2016
	Process: 50 Responsibility Allocation : Guest Access Wifi 16 Feb 2016
	Process: 51 Responsibility Allocation : Printers 16 Feb 2016
	Process: 5903 Responsibility Allocation : Weather Station 02 Mar 2016
	Process: 7121 Responsibility Allocation : General Computer Maintenance 09 Mar 2016
	Process: 7178 Responsibility Allocation : Systems Innovation 09 Mar 2016
	Process: 6843
	Process: 7835 Electrics Need Checking 20 Sep 2017
	Process: 7836 Central Heating For Winter 20 Sep 2017
	Process: 7847 Health And Safety Review 26 Sep 2017
	Process: 7864 ESD Work Stations 07 Oct 2017
	Process: 7867 Bandsaw Checklist 13 Oct 2017
	Process: 7868 Pillar Drill Checklist 13 Oct 2017
	Process: 7869 Hand Drill Checklist 13 Oct 2017
	Process: 7891 Fire Alarm Evacuation Drill 25 Oct 2017
	Process: 7896 Tree In Car Park 22 Dec 2017
	Process: 7910 Review CCTV Warning Signs 20 Sep 2018
	Process: 7928 Fire Test Points Checking 21 Feb 2019
	Process: 7929 Emergency Lighting And Fire Extinguishers 21 Feb 2019
	Process: 7911 Review Security Of The Special Category Personal Data 20 Sep 2018
	Process: 7961 R D Room - Tidy, Empty Bins, Remove Cups. Caution Around Oxygen Supply
	05 Oct 2020
	Process: 7982 Check There Are No Changes To Employment Law 21 Nov 2021
	Process: 7999 Building Risk Assesments 08 Feb 2022
ひついつつつ	VM3COP02.02 VST Company Responsibilitys organisation chart structure
ID29373	Process: 5877 Review Company Data 17 Feb 2016
ID29373 ID77289	Process: 5877 Review Company Data 17 Feb 2016 Audit 21 Audit of Audit Process: 7731 Audit 21 Audit Of Audit Viamed 24 Aug 2016

/10/2023, 11:0	QMS Route Map Viamed Ltd ISO13485:2016
	Process: 38 Audits Up to Date and Confirm next years Audit schedule 16 Feb 2016
	Process: 7093 BSI Audits Calander 09 Mar 2016
	Process: 7670 Humanmed general Issues 09 Mar 2016
	Process: 7862 Review The Audit Calender Screen 04 Oct 2017
ID120397	Audit 22 Post Market Survellance
	Process: 7732 Audit 22 Post Market Survellance Viamed 24 Aug 2016
	Process: 43 Responsibility Allocation : Product Post Market Survelance 16 Feb 2016
	Process: 7780 Audit 22 Post Market Survellance VST 08 Feb 2017
	Process: 6889 Responsibility Allocation : Post Market Surveilance 09 Mar 2016
	Process: 7809 Pro-Active Marketing 06 Jun 2017
	Process: 7810 Research Activities 06 Jun 2017
	Process: 5863 Responsibility Allocation : Sales Meetings UK 17 Feb 2016
	Process: 5864 Responsibility Allocation : Sales Meeting EX 17 Feb 2016
	Process: 7973 VST Product Performance - Customers 27 Oct 2021
	Process: 7974 VST Product Performance - Suppliers 27 Oct 2021
	Process: 8014 Review VIAMED Product Feedback Positive 25 Jul 2022
	Process: 8015 Review VST Product Feedback Positive 25 Jul 2022
	Process: 8016 Review VIAMED Customer Feedback Positive 25 Jul 2022
	Process: 8017 Review VST Customer Feedback Positive 25 Jul 2022
D126137	Viamed Management Review Blank Minutes 20xx
.D120157	Process: 7846 ISO System Management Review Viamed 26 Sep 2017
D74720	
D74728	QC 21 Non Conformance Form
	Process: 7138 Non Conformance Issues Any New QC21 Forms 09 Mar 2016
	Process: 7979 Review The Template Of The QC 21 Form To Ensure It Is Current And Valid
	12 Nov 2021
D31024	VOP 12 Training
	Process: 7750 Meeting With Management 14 Oct 2016
	Process: 7793 Team Review Meeting 16 Mar 2017
	Process: 5934 Responsibility Allocation : Staff Training 05 Mar 2016
	Process: 7833 Importance Of Effective Quality Management 20 Sep 2017
	Process: 7845 7.1.4 Environment Of Operations 25 Sep 2017
	Process: 7883 Appraisal 23 Oct 2017
D14696	
.D14050	Process: 6972 UPS Shipping Fuel Surcharge 09 Mar 2016
D17155	
D17155	VM3COP03.05 Procedures for customer returning goods on our UPS account number
	Process: 5879 Responsibility Allocation: Customer Returning Goods On Our UPS Account
	18 Feb 2016
D31032	VOP 16 Health and Safety, Company Personnel Manual
	Process: 7821 Controlled Waste Description And Transfer 15 Jun 2017
	Process: 7820 North Yorkshire Council Waste Tranfer 15 Jun 2017
	Process: 6851 Review Accident Book 09 Mar 2016
	Process: 7759 Health Declaration Sheet 23 Jan 2017
	Process: 6849 First Aid 09 Mar 2016
	Process: 6855 Risk Assessment HSE 09 Mar 2016
	Process: 6856 Fire Alarms 09 Mar 2016
	Process: 7092
	Process: 56 Warehouse Outside Heating Guard 17 Feb 2016
	Process: 5919 Check Out Side Drain 05 Mar 2016
	Process: 5921 Clearing Water Downstairs 05 Mar 2016
	Process: 7120 General Maintenance Requirements 09 Mar 2016
	Process: 7742 Boiler Check 26 Sep 2016
	Process: 7756 Carbon Monoxide Alarm 05 Jan 2017
	Process: 7835 Electrics Need Checking 20 Sep 2017
	Process: 7836 Central Heating For Winter 20 Sep 2017
	Process: 7847 Health And Safety Review 26 Sep 2017
	Terrent 20 Sep 2017
	TI CONTRACTOR OF THE PROPERTY

Process: 7867 Bandsaw Checklist 13 Oct 2017 Process: 7868 Pillar Drill Checklist 13 Oct 2017 Process: 7869 Hand Drill Checklist 13 Oct 2017 Process: 7928 Fire Test Points Checking 21 Feb 2019 Process: 7999 Building Risk Assesments 08 Feb 2022 ID120355 Audit 07 Handling and Storage Process: 6973 Responsibility Allocation: Stock Transfers. (QC19) 09 Mar 201 Process: 7719 Audit 07 Handling And Storage Viamed 24 Aug 2016 Process: 7767 Audit 07 Handling And Storage VST 08 Feb 2017 Process: 5858 Opera Stock Adjustments 17 Feb 2016 Process: 5935 Stock Allocations 05 Mar 2016 Process: 6840 Process: 6850 Current Stock Levels 09 Mar 2016 Process: 6945 Missing Stock or Adjustments 09 Mar 2016 Process: 7046 Responsibility Allocation: Stock Purchasing 09 Mar 2016 Process: 7051 Responsibility Allocation: Control of nonconforming product 0 Process: 7673 Check Expiry Dated Stock 09 Mar 2016 Process: 7688 Process: 7689 Move Stock From QA Shelf To Stock Shelf Monday 21 Apr 201 Process: 7694 Move Stock From QA Shelf To Stock Shelf Tuesday 28 Apr 201 Process: 7695 Top Up Quick Shipping Shelves 28 Apr 2016 Process: 7873 On Site Environment Review 18 Oct 2017 Process: 7866 Oxygen Cylinder Check 13 Oct 2017 Process: 7903 Empty Warehouse Depleted Sensor Bin 17 Jul 2018 Process: 7904 Check Weeee Waste Pallet And Sensor Bin 17 Jul 2018 Process: 7902 Empty Depleted Sensor Bin From The Offic 17 Jul 2018	9 Mar 2016 16
Process: 7869 Hand Drill Checklist 13 Oct 2017 Process: 7928 Fire Test Points Checking 21 Feb 2019 Process: 7999 Building Risk Assesments 08 Feb 2022 ID120355 Audit 07 Handling and Storage Process: 6973 Responsibility Allocation: Stock Transfers. (QC19) 09 Mar 201 Process: 7719 Audit 07 Handling And Storage Viamed 24 Aug 2016 Process: 7767 Audit 07 Handling And Storage VST 08 Feb 2017 Process: 5858 Opera Stock Adjustments 17 Feb 2016 Process: 5935 Stock Allocations 05 Mar 2016 Process: 6840 Process: 6945 Missing Stock or Adjustments 09 Mar 2016 Process: 7046 Responsibility Allocation: Stock Purchasing 09 Mar 2016 Process: 7051 Responsibility Allocation: Control of nonconforming product 0 Process: 7673 Check Expiry Dated Stock 09 Mar 2016 Process: 7688 Process: 7689 Move Stock From QA Shelf To Stock Shelf Monday 21 Apr 201 Process: 7694 Move Stock From QA Shelf To Stock Shelf Tuesday 28 Apr 201 Process: 7695 Top Up Quick Shipping Shelves 28 Apr 2016 Process: 7873 On Site Environment Review 18 Oct 2017 Process: 7866 Oxygen Cylinder Check 13 Oct 2017 Process: 7903 Empty Warehouse Depleted Sensor Bin 17 Jul 2018 Process: 7904 Check Weeee Waste Pallet And Sensor Bin 17 Jul 2018	9 Mar 2016 16
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Service And Repairs For Viamed And VST 09 Oct 2019 Process: 8008 Verification Warehouse Hand Sanitiser 21 Feb 2022	
Process: 8002 Verification Todays Goods In 17 Feb 2022	
Process: 8004 Verification Of Non Conforming Products 17 Feb 2022 Process: 8024 Discontinue/Supersede Stock 01 Mar 2023	
ID53615 VOP 06 Measurement Control Viamed VST, Calibration, QA Stock	
Process: 7718 Audit 06 Calibration Viamed 24 Aug 2016	
Process: 7091 Calibration Index 09 Mar 2016	
Process: 7998 Verification Calibrated Equipment 08 Feb 2022	
ID119452 Audit 15 Production	
Process: 7727 Audit 15 Production Viamed 24 Aug 2016	
Process: 7736 Production Start Job List 03 Sep 2016	
Process: 7737 Production In Production List 03 Sep 2016	
Process: 7738 Production Statistics 03 Sep 2016	
Process: 7775 Audit 15 Production VST 08 Feb 2017	
Process: 6845 Responsibility Allocation : Quarantine Production 09 Mar 2016	
Process: 6955 Production Requirements 09 Mar 2016	
Process: 7169 Responsibility Allocation : Production 09 Mar 2016	
Process: 7170 Responsibility Allocation : Production Production Schedule 09 1	Mar 2016
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Process: 7072 Responsibility Allocation : Manufacturing Processes 09 Mar 20	
Process: 8000 Verification Production Paperwork 08 Feb 2022	
ID31008 VOP 11 Equipment Control, Office, Warehouse, Pcs and Equipment	
Process: 5939 Responsibility Allocation : Email ISP Routing 05 Mar 2016	
Process: 5959 Responsibility Allocation: Enfail 15P Routing 05 Mai 2016 Process: 5941 Responsibility Allocation: Replace Main Server 07 Mar 2016	
Process: 3941 Responsibility Allocation : Replace Main Server 07 Mai 2016 Process: 45 Responsibility Allocation : Main Server Status 16 Feb 2016	
Process: 45 Responsibility Allocation : Main Server Status 16 Feb 2016 Process: 46 Responsibility Allocation : Backup Server Status 16 Feb 2016	
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Process: 52 Software Verification Clear Down Backup Emails 16 Feb 2016

Process: 53 Emails 16 Feb 2016

Process: 7672 Off Site Backup 09 Mar 2016

Process: 6813 Management Meeting Turnover Report 09 Mar 2016

Process: 7700 Domain Name Management 19 May 2016 **Process: 7701** AWS Amazon Web Services 23 May 2016

Process: 7704 Responsibility Allocation: Computer Failure Diagnostics 24 May 2016

Process: 48 Responsibility Allocation : Internet 16 Feb 2016 **Process: 49** Responsibility Allocation : Wifi 16 Feb 2016

Process: 50 Responsibility Allocation: Guest Access Wifi 16 Feb 2016

Process: 51 Responsibility Allocation : Printers 16 Feb 2016

Process: 5903 Responsibility Allocation: Weather Station 02 Mar 2016

Process: 6838 Opera Negative Stock 09 Mar 2016

Process: 7121 Responsibility Allocation: General Computer Maintenance 09 Mar 2016

Process: 7124 Responsibility Allocation : Intrastats 09 Mar 2016

Process: 7125 Responsibility Allocation: Intrastats Urgent Problems 09 Mar 2016

Process: 7126 Intrastats Requested Page updates 09 Mar 2016

Process: 7127 Responsibility Allocation: Intrastats Unfinished in progress Processes 09 Mar

2016

Process: 7128 Responsibility Allocation: Intrastats Future Features needed 09 Mar 2016

Process: 7129 Intrastats Cross Reference Database Tables Updates 09 Mar 2016 **Process: 7178** Responsibility Allocation : Systems Innovation 09 Mar 2016

Process: 7739 Intrastats Amendment Log 12 Sep 2016

Process: 7755 Fast Hosts Invoice 08 Dec 2016

Process: 44 Secure Socket Level Certificate 16 Feb 2016

Process: 7668 Responsibility Allocation : Upgrading Intrastats ISO Quality system 09 Mar

2016

Process: 7832 Cleardown Emailed Invoices 20 Sep 2017

Process: 7823 Saftey Tester Data 02 Aug 2017

ID55437 Audit 09 Goods Inward and Product Identity

Process: 5938 Responsibility Allocation: Receive Goods 05 Mar 2016

Process: 7721 Audit 09 Goods Inward And Product Identity Viamed 24 Aug 2016

Process: 7826 Goods In Processes 06 Sep 2017

Process: 7792 Shipped Order Success Report 13 Mar 2017

Process: 7769 Audit 09 Goods Inward And Product Identity VST 08 Feb 2017

Process: 6969 Responsibility Allocation: VIAMED Stock Meeting `Goods In` Review 09

Mar 2016

Process: 57 Temporary Stock Notices 17 Feb 2016

Process: 5854 Stock FAQ Admin List 17 Feb 2016

Process: 7181 Responsibility Allocation: Product Catagories 09 Mar 2016

Process: 6894 Product Cross References 09 Mar 2016

Process: 6838 Opera Negative Stock 09 Mar 2016

Process: 7830 Review Q.A. Failures Report 18 Sep 2017

Process: 7859 Check POR Files For Items Delivered But Not Removed From File 02 Oct

2017

Process: 7897 Daily O2 Sensors Returns 04 Jan 2018

Process: 7898 Stamp Deliveries 30 Jan 2018

Process: 7903 Empty Warehouse Depleted Sensor Bin 17 Jul 2018

Process: 7914 Proofs of Delivery 02 Oct 2018

Process: 7915 Reserve Stock Review 02 Oct 2018

Process: 7917 Human Med Purchase Order 18 Oct 2018

Process: 7923 Review Of Credits Received From Suppliers 08 Jan 2019

Process: 7943 Review Stocks Of 8000004 01 Oct 2019

Process: 7957 Warehouse Requests 29 May 2020

Process: 7962 VST Supplier QA Results 28 Oct 2020

Process: 7967 VST Stock Count For End April 01 Jul 2021

Process: 7976 Decontamination Of Incoming Products And Repairs 08 Nov 2021

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	Process: 8009 Verification Stock Items And Locations 21 Feb 2022
	Process: 8010 Verification Of Ebay Stock 21 Feb 2022
	Process: 8011 Verification Of Demo Stock 21 Feb 2022
ID75007	
ID75927	VOP 09 Repairs and Servicing
	Process: 7684 Repairs Ready For Quote 18 Apr 2016
	Process: 7685 Repairs Ready For Invoice 18 Apr 2016
	Process: 7690 Ship Repairs 21 Apr 2016
	Process: 7752 SRS Folder 22 Nov 2016
	Process: 6847 Responsibility Allocation : Quarantine Repairs 09 Mar 2016
	Process: 6862 Current Repairs 09 Mar 2016
	Process: 7048 Control of monitoring and measuring devices 09 Mar 2016
	Process: 7674 Check Repairs Ready For Invoice List 10 Mar 2016
	Process: 7814 Responsibility Allocation : Viamed Repairs 06 Jun 2017
	Process: 7811 Responsibility Allocation : General Area 06 Jun 2017
	Process: 7812 Responsibility Allocation : Vandagraph Repairs 06 Jun 2017
	Process: 7813 Responsibility Allocation : VST Repairs 06 Jun 2017
	Process: 7815 Responsibility Allocation : Product Types To Relevant Person 06 Jun 2017
	Process: 7942 Do We Have Service Manual / QA For All Our Stock Coming In. 23 Sep 2019
	Process: 7940 Review The Tom Thumb Grease Date 18 Sep 2019
	Process: 7985 OverDue Servicing 03 Feb 2022
	Process: 7993 Verification Warranty Repairs Customer Approval 07 Feb 2022
	Process: 7994 Verification Repairs Paperwork Completed 07 Feb 2022
	Process: 7995 Verification Visual Check Repair Shelf 07 Feb 2022
	Process: 7996 Verification Repairs Older Repairs 07 Feb 2022
	Process: 7997 Verification Repair Qa Reports 07 Feb 2022
	Process: 8005 Verification Of SRS Information added 17 Feb 2022
	Process: 8022 Vandagraph Repair Review 06 Feb 2023
ID31072	VOP 08 Production, Reworks, New Production
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ID114755 ID20049 ID117589 ID22266 ID24775	Process: 7169 Responsibility Allocation: Production 09 Mar 2016 Process: 7170 Responsibility Allocation: Production Production Schedule 09 Mar 2016 Process: 7171 Responsibility Allocation: Production Production Problems 09 Mar 2016 Process: 7072 Responsibility Allocation: Manufacturing Processes 09 Mar 2016 Process: 6962 Responsibility Allocation: VIAMED Stock Meeting Returns Overview 09 Ma 2016 Process: 8000 Verification Production Paperwork 08 Feb 2022 VM3COP20.31 Export Order Processing Process: 7825 Responsibility Allocation: Order Picking 06 Sep 2017 VM3COP03.01 Order Processing Priorities Process: 5 Responsibility Allocation: Order Picking 06 Sep 2017 VM3COP20.30 UK Order Processing Process: 7825 Responsibility Allocation: Order Picking 06 Sep 2017 VM3COP03.07 Humanmed Order Checking Process: 7 Responsibility Allocation: Checking Of Sales Orders 16 Feb 2016 Process: 7734 Responsibility Allocation: Humanmed Order Processing 25 Aug 2016 VM3COP03.08 Humanmed Order Processing Process: 5 Responsibility Allocation: Processing Of Sales Orders 16 Feb 2016

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ID34889	VM3COP20.32 Order Checking
	Process: 7825 Responsibility Allocation : Order Picking 06 Sep 2017
ID122441	Audit 01 Picking packing
	Process: 7714 Audit 01 Picking Packing Viamed 24 Aug 2016
	Process: 7825 Responsibility Allocation : Order Picking 06 Sep 2017
	Process: 5859 Review Un-shipped Parcels 17 Feb 2016
	Process: 6970
	Process: 7691 Ship Sale Or Returns 21 Apr 2016
	Process: 7762 Audit 01 Picking Packing VST 08 Feb 2017
	Process: 7796 Review Franking Label Errors 08 May 2017
	Process: 7797 Check Order Are Being Picked In Priority Order 10 May 2017
	Process: 7798 Orders And Items Shipped Per Month 10 May 2017
	Process: 7860 Goods Out Picking 03 Oct 2017
	Process: 8027 Update Pricing For Viamed Shopify 11 Apr 2023
ID124549	Audit 11 Repairs, Servicing and Returns
	Process: 5898 Processing Depleted Sensors 25 Feb 2016
	Process: 5879 Responsibility Allocation : Customer Returning Goods On Our UPS Account
	18 Feb 2016
	Process: 5857 Customer Service Logs 17 Feb 2016
	Process: 7724 Audit 11 Repairs And Service Viamed 24 Aug 2016
	Process: 7684 Repairs Ready For Quote 18 Apr 2016
	Process: 7685 Repairs Ready For Invoice 18 Apr 2016
	Process: 7690 Ship Repairs 21 Apr 2016
	Process: 7748 Check Repair Orders 10 Oct 2016
	Process: 7749 Check Repair Quotes 10 Oct 2016
	Process: 7752 SRS Folder 22 Nov 2016
	Process: 7760 Send Service Offers 31 Jan 2017
	Process: 7772 Audit 11 Repairs And Service VST 08 Feb 2017
	Process: 6847 Responsibility Allocation : Quarantine Repairs 09 Mar 2016
	Process: 6862 Current Repairs 09 Mar 2016
	Process: 7138 Non Conformance Issues Any New QC21 Forms 09 Mar 2016
	Process: 7674 Check Repairs Ready For Invoice List 10 Mar 2016
	Process: 7692 Responsibility Allocation: Take Complete Repair Paperwork To Office 22 Apr
	2016
	Process: 6916 Responsibility Allocation : Service exisiting 09 Mar 2016
	Process: 6917 Responsibility Allocation : Service extension 09 Mar 2016
	Process: 7823 Saftey Tester Data 02 Aug 2017
	Process: 7905 Generate RMA Box, Link Items And Add Faults 17 Jul 2018
	Process: 7906 Request RMA Based On The RMA Boxes 17 Jul 2018
	Process: 7993 Verification Warranty Repairs Customer Approval 07 Feb 2022
	Process: 7994 Verification Repairs Paperwork Completed 07 Feb 2022
	Process: 7995 Verification Visual Check Repair Shelf 07 Feb 2022
	Process: 7996 Verification Repairs Older Repairs 07 Feb 2022
	Process: 7996 Verification Repairs Older Repairs 07 Feb 2022 Process: 7997 Verification Repair Qa Reports 07 Feb 2022
	Process: 7997 Verification Repair Qa Reports 07 Feb 2022 Process: 8022 Vandagraph Repair Review 06 Feb 2023
ID69812	VM3COP27.31 Processing Proforma Invoices and Quotations Process: 7710 Responsibility Allocation: Proforma And Quote Processing 29 Jun 2016
ID13695	VM3COP20.05 New Orders - How to enter into Opera Viamed Process: 7936 B2B Router / Peppol Responsibilitys 19 Jun 2019
ID21314	Process: 6828
ID76091	Audit 14 Complaints and Corrective Actions
דבטטיים:	Process: 7726 Audit 14 Complaints And Corrective Actions Viamed 24 Aug 2016
	Process: 7/26 Addit 14 Complaints And Corrective Actions viamed 24 Aug 2016 Process: 6828
	Process: 7743 Customer Complaints Paper File 26 Sep 2016
	Process: 7774 Audit 14 Complaints And Corrective Actions VST 08 Feb 2017

Process: 6865 Responsibility Allocation: Non Conformance Effectiveness 09 Mar 2016

Process: 7199 Non Conformities Review Viamed 09 Mar 2016

Process: 7671 Humanmed Non Conformances 09 Mar 2016

Process: 6931 Customer Complaints 09 Mar 2016

Process: 7839 Review VIAMED Feedback - Customer Complaints 23 Sep 2017

Process: 7838 Review VIAMED Feedback - Customer Feedback Negative 23 Sep 2017

Process: 7840 Review VST Feedback - Customer Feedback Negative 23 Sep 2017

Process: 7841 Review VST Feedback - Customer Complaints 23 Sep 2017

Process: 7842 Review VIAMED Product Feedback Negative 23 Sep 2017

Process: 7843 Review VST Product Feedback Negative 23 Sep 2017

Process: 7849 Review Product Failures New Codes 28 Sep 2017

Process: 7934 Test Website Questions 02 May 2019

Process: 7965 VST Feedback 29 Oct 2020

Process: 7264 Responsibility Allocation: VST Management Meeting Non Conformance

Issues 09 Mar 2016

ID63821 Audit 04 Accounts and Finance

Process: 7702 Responsibility Allocation: Vandagraph Pay Pay Issue Refund 23 May 2016

Process: 7703 Vandagraph Pay Pal Retrieve Funds 23 May 2016

Process: 5915 Opera Sales Ledger Close 05 Mar 2016

Process: 7740 Weights Per Region Needed To Submit EC Sales List 13 Sep 2016

Process: 5929 HMRC Intrastats Sales Data 05 Mar 2016

Process: 7799 Opera Purchase Ledger Close 11 May 2017

Process: 7800 Opera Nominal Ledger Close 11 May 2017

Process: 5937 Review the Delivered Not Invoiced Reports 05 Mar 2016

Process: 5865 Vandagraph Loan 17 Feb 2016

Process: 5867 Accounts On Stop 17 Feb 2016

Process: 5874 Childcare Vouchers Edenred 17 Feb 2016

Process: 5914 End Of Year Reports For Accountants 04 Mar 2016

Process: 5916 Bank Details Opera reports entered Intrastats 05 Mar 2016

Process: 5917 Fill in Cashbook / Bank Rec for previous Month 05 Mar 2016

Process: 5918 Journals for the End of Month accounts 05 Mar 2016

Process: 5920 Responsibility Allocation : Cheques To Bank - Fill in Paying in Book 05 Mar

2016

Process: 5922 Credit Cards Expenses Calculations 05 Mar 2016

Process: 5923 Credits Note Processing 05 Mar 2016

Process: 5924 Export Cheques sent by Currency Lodgement 05 Mar 2016

Process: 5925 Customs Clearance 05 Mar 2016

Process: 5926 Responsibility Allocation : Petty Cash Expenses receipts and cash 05 Mar 2016

Process: 5927 Responsibility Allocation : Accounts Filing 05 Mar 2016

Process: 5928 Responsibility Allocation: Filing Cabinets 05 Mar 2016

Process: 5930 VAT Return Viamed 05 Mar 2016

Process: 5931 Purchase Invoices in to Opera 05 Mar 2016

Process: 5932 Remit Processing and entry into Opera 05 Mar 2016

Process: 5933 Responsibility Allocation : Sales Accounts Reminders 05 Mar 2016

Process: 5942 Chase the Debtors viamed 08 Mar 2016

Process: 6819 Supplier Payments and Invoice processing 09 Mar 2016

Process: 6822

Process: 6876 Issues for Accountants - P11D Form re Benefits to Revenue and Customs 09

Mar 2016

Process: 6946 Accounts Debtors Review - Export 09 Mar 2016

Process: 6951 Accounts Debtors Review - UK 09 Mar 2016

Process: 7192

Process: 7084 Responsibility Allocation : Accounts Issues 09 Mar 2016

Process: 7195 Responsibility Allocation: Loans between companies 09 Mar 2016

Process: 7788 Petty Cash Reconciliation 02 Mar 2017

Process: 7789 Withdraw Funds From Paypal 02 Mar 2017

Process: 7817 Issues For Accountants - Check suggested invoice report in operas 13 Jun 2017

/10/2023, 11:0	US QMS Route Map Viamed Ltd 15015465:2016
	Process: 7818 Issues For Accountants - Check Purchasing Journals to see if VAT handled
	correctly Previous Month 13 Jun 2017
	Process: 7819 Issues For Accountant - Check Contra account 8000 and clear it 13 Jun 2017
	Process: 7824 Chase The Debtors VST 27 Aug 2017 Process: 7708 Acorn 0014904 17 Jun 2016
	Process: 5869 Responsibility Allocation: Legal Company Car Registration 17 Feb 2016
	Process: 7831 Intrastats Debtors And Creditor Figures 18 Sep 2017 Process: 7885 Audit 04 Accounts and Finance Viamed 23 Oct 2017
	Process: 7899 Region Checker 06 Feb 2018
	Process: 7900 Royal Mail - Mail Retention Form 29 Mar 2018
	Process: 7901 UPS Exceptions Checkup 20 Apr 2018
	Process: 7920 Sales Warnings 20 Dec 2018
	Process: 7927 Contract Pricing Review 14 Feb 2019
	Process: 7919 Send Debtors Overview To Derek 06 Dec 2018
	Process: 7924 PDFing Of Invoices Vandagraph 11 Jan 2019
	Process: 7932 Check Debtors Report 15 Mar 2019
	Process: 7933 Purchasing Invoice Processing 22 Mar 2019
	Process: 7935 PCI DSS Compliance 03 Jun 2019
	Process: 7938 VAT Return Vandagraph 22 Jul 2019
	Process: 7939 VAT Return VST 22 Jul 2019
	Process: 7945 Xero Review Sales Contacts 05 Feb 2020
	Process: 7946 Xero Merge Customers That Are Duplicates 05 Feb 2020
	Process: 7952 Check Xero To Barclays Bank Statements End On Month GBP, USD And Euro
	Viamed 06 Mar 2020
	Process: 7958 Exchange Rate In To Intrastats 03 Sep 2020
	Process: 7966 Xero Sync 10 Mar 2021
	Process: 7968 Shred CC Slips 06 Aug 2021
	Process: 7984 Check For Viking Invoices 19 Jan 2022
	Process: 8007 Verification Credit Notes 17 Feb 2022
	Process: 7986 Check Creditors 03 Feb 2022
	Process: 7990 Verification Invoice Details Accounts 07 Feb 2022
	Process: 8012 VAT Return Viamed Properties 06 Apr 2022
	Process: 8019 Audit 04 Accounts And Finance VST 14 Sep 2022
	Process: 8021 Check Xero Bank For The Year To The Barclays Bank Account 06 Jan 2023
ID63815	Audit 12 CE Files
	Process: 7725 Audit 12 CE Files Viamed 24 Aug 2016
	Process: 7773 Audit 12 CE Files VST 08 Feb 2017
	Process: 24 Responsibility Allocation : Compliance ISO Standards 16 Feb 2016
	Process: 7172 Responsibility Allocation : CE Technical Files 09 Mar 2016 Process: 7071 Post Market Surveillance 09 Mar 2016
ID73132	VM3COP20.29 Checking the Purchase Order Log Process: 5850 Purchase Order Log 17 Feb 2016
ID17070	
ID17070	VM3COP27.34 Sending Purchase Orders to Suppliers
	Process: 7707 Send Purchase Orders To Suppliers 13 Jun 2016
ID63048	Audit 06 Calibration
	Process: 7718 Audit 06 Calibration Viamed 24 Aug 2016
	Process: 7766 Audit 06 Calibration VST 08 Feb 2017
	Process: 7048 Control of monitoring and measuring devices 09 Mar 2016
	Process: 7091 Calibration Index 09 Mar 2016 Process: 7009 Varification Calibrated Equipment 08 Feb 2022
	Process: 7998 Verification Calibrated Equipment 08 Feb 2022
ID68263	Audit 24 Service Logs
	Process: 5857 Customer Service Logs 17 Feb 2016
	Process: 7760 Send Service Offers 31 Jan 2017
	Process: 7889 Audit 24 Servicing Viamed 24 Oct 2017
	Process: 7985 OverDue Servicing 03 Feb 2022

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ID31048	VOP 22 Picking and Packing Dispatch and Goods Out
	Process: 5945 Responsibility Allocation : Sending Samples 08 Mar 2016
	Process: 5946 Responsibility Allocation : Sending Sale Or Returns 08 Mar 2016
	Process: 7825 Responsibility Allocation : Order Picking 06 Sep 2017
	Process: 5859 Review Un-shipped Parcels 17 Feb 2016
	Process: 6954 Back Orders Review - By Customer 09 Mar 2016
	Process: 6970
	Process: 7691 Ship Sale Or Returns 21 Apr 2016
	Process: 7748 Check Repair Orders 10 Oct 2016
	Process: 7749 Check Repair Quotes 10 Oct 2016
	Process: 7797 Check Order Are Being Picked In Priority Order 10 May 2017
	Process: 6969 Responsibility Allocation : VIAMED Stock Meeting `Goods In` Review 09
	Mar 2016
	Process: 7860 Goods Out Picking 03 Oct 2017
ID24509	VM3COP20.27 Annual Services for Resuscitation Cabinets
	Process: 5857 Customer Service Logs 17 Feb 2016
ID75624	VM3COP23.00 EAN13 Barcodes to Stock and the Online Databases
	Process: 7909 EAN GTIN Online Database 06 Aug 2018
ID8712	DO NOT USE VM3COP09 Repairs
	Process: 7684 Repairs Ready For Quote 18 Apr 2016
	Process: 7685 Repairs Ready For Invoice 18 Apr 2016
	Process: 7814 Responsibility Allocation : Viamed Repairs 06 Jun 2017
ID13703	VM3COP20.03 Repair Procedures Goods in
	Process: 5891 Processing Of Repair Quotes And Orders 25 Feb 2016
ID17485	VM3COP20.47 Collecting Repair Paperwork
	Process: 7693 Collect Repair Filing From Warehouse 22 Apr 2016
ID77209	Audit 17 Internal Audits
1111//4UJ	Audit 1/ Internal Audits
10//200	
110//200	Process: 7728 Audit 17 Internal Audits Viamed 24 Aug 2016 Process: 7776 Audit 17 Internal Audits VST 08 Feb 2017
10//203	Process: 7728 Audit 17 Internal Audits Viamed 24 Aug 2016
ID124938	Process: 7728 Audit 17 Internal Audits Viamed 24 Aug 2016 Process: 7776 Audit 17 Internal Audits VST 08 Feb 2017 Process: 7972 ISO System Management Review Vst 26 Oct 2021
	Process: 7728 Audit 17 Internal Audits Viamed 24 Aug 2016 Process: 7776 Audit 17 Internal Audits VST 08 Feb 2017
	Process: 7728 Audit 17 Internal Audits Viamed 24 Aug 2016 Process: 7776 Audit 17 Internal Audits VST 08 Feb 2017 Process: 7972 ISO System Management Review Vst 26 Oct 2021 VOP 10 Non Conformance, Corrective and Preventive Actions
	Process: 7728 Audit 17 Internal Audits Viamed 24 Aug 2016 Process: 7776 Audit 17 Internal Audits VST 08 Feb 2017 Process: 7972 ISO System Management Review Vst 26 Oct 2021 VOP 10 Non Conformance, Corrective and Preventive Actions Process: 7199 Non Conformities Review Viamed 09 Mar 2016
	Process: 7728 Audit 17 Internal Audits Viamed 24 Aug 2016 Process: 7776 Audit 17 Internal Audits VST 08 Feb 2017 Process: 7972 ISO System Management Review Vst 26 Oct 2021 VOP 10 Non Conformance, Corrective and Preventive Actions Process: 7199 Non Conformities Review Viamed 09 Mar 2016 Process: 7069 Responsibility Allocation : Corrective Actions 09 Mar 2016
	Process: 7728 Audit 17 Internal Audits Viamed 24 Aug 2016 Process: 7776 Audit 17 Internal Audits VST 08 Feb 2017 Process: 7972 ISO System Management Review Vst 26 Oct 2021 VOP 10 Non Conformance, Corrective and Preventive Actions Process: 7199 Non Conformities Review Viamed 09 Mar 2016 Process: 7069 Responsibility Allocation : Corrective Actions 09 Mar 2016 Process: 7849 Review Product Failures New Codes 28 Sep 2017