

## Internal Audit Check list

## INTERNAL PROCESS VERIFICATION

Created:	17/May 1995	<b>Audit No 20</b> Sit With management and Complete this Form	
Revised:	06 August 2020		Page 1 of 14
Audit Date	9th Sept 20	Auditor <i>Helen Lamb</i> <i>Derek Lamb</i>	

Company / ISO Section	Criteria of ISO Section	Auditor Comments / Issues
VST Ltd ISO9001:201 5.5.1.1	<p><b>General</b></p> <p>Top management shall demonstrate leadership and commitment with respect to the quality management system by:</p> <ul style="list-style-type: none"> <li>a) taking accountability for the effectiveness of the quality management system;</li> <li>b) ensuring that the quality policy and quality objectives are established for the quality management system and are compatible with the context and strategic direction of the organization;</li> <li>c) ensuring the integration of the quality management system requirements into the organization's business processes;</li> <li>d) promoting the use of the process approach and risk-based thinking;</li> <li>e) ensuring that the resources needed for the quality management system are available;</li> <li>f) communicating the importance of effective quality management and of conforming to the quality management system requirements;</li> <li>g) ensuring that the quality management system achieves its intended results;</li> <li>h) engaging, directing and supporting persons to contribute to the effectiveness of the quality management system;</li> <li>i) promoting improvement;</li> <li>j) supporting other relevant management roles to demonstrate their leadership as it applies to their areas of responsibility.</li> </ul> <p>NOTE Reference to "business" in this International Standard can be interpreted broadly to mean those activities that are core to the purposes of the organization's existence, whether the organization is public, private, for profit or not for profit.</p>	<i>doc index</i> <i>intranet</i> <i>part map</i> <i>BS</i> <i>Required Reading</i> <i>Issues</i>
VST Ltd ISO9001:201 5.5.2.1	<p><b>Establishing the quality policy</b></p> <p>Top management shall establish, implement and maintain a quality policy that:</p> <ul style="list-style-type: none"> <li>a) is appropriate to the purpose and context of the organization and supports its strategic direction;</li> <li>b) provides a framework for setting quality objectives;</li> <li>c) includes a commitment to satisfy applicable requirements;</li> <li>d) includes a commitment to continual improvement of the quality management system.</li> </ul>	<i>intranet</i> <i>Doc index</i> <i>management plan</i>
VST Ltd ISO9001:201 5.6.2.2	<p>When planning how to achieve its quality objectives, the organization shall determine:</p> <ul style="list-style-type: none"> <li>a) what will be done;</li> </ul>	

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	b) what resources will be required; c) who will be responsible; d) when it will be completed; e) how the results will be evaluated.	Intrastarts
VST Ltd ISO9001:2015 7.5.1	<b>General</b> 7.5.1 General The organization's quality management system shall include: a) documented information required by this International Standard; b) documented information determined by the organization as being necessary for the effectiveness of the quality management system. NOTE The extent of documented information for a quality management system can differ from one organization to another due to: — the size of organization and its type of activities, processes, products and services; — the complexity of processes and their interactions; — the competence of persons.	Route map Doc index
Viamed Ltd ISO13485:2016 4.1.3	<b>Quality management system</b> 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 necessary to achieve planned results and maintain the effectiveness of these 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).	Intrastarts Doc index
Viamed Ltd ISO13485:2016 4.1.4	<b>Quality management system</b> For each quality management system process, the organization shall: The organization shall 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 c) controlled in accordance with the requirements of this International Standard and applicable regulatory requirements.	Route map Intrastarts
Viamed Ltd ISO13485:2016 4.2.1	<b>Documentation requirements</b> The quality management system documentation (see 4.2.4) shall include: a) documented statements of a quality policy and quality objectives;	

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General	<ul style="list-style-type: none"> <li>b) a quality manual;</li> <li>c) documented procedures and records required by this International Standard;</li> <li>d) documents, including records, determined by the organization to be necessary to ensure the effective planning, operation, and control of its processes;</li> <li>e) other documentation specified by applicable regulatory requirements.</li> </ul>	Doc index
Viamed Ltd ISO13485:20 16 4.2.2 Quality manual	<p><b>Documentation requirements</b></p> <p>The organization shall document a quality manual that includes:</p> <ul style="list-style-type: none"> <li>a) the scope of the quality management system, including details of and justification for any exclusion or non-application;</li> <li>b) the documented procedures for the quality management system, or reference to them;</li> <li>c) a description of the interaction between the processes of the quality management system.</li> </ul> <p>The quality manual shall outline the structure of the documentation used in the quality management system.</p>	Doc index VOP procedures
Viamed Ltd ISO13485:20 16 5.1	<p><b>Management commitment</b></p> <p>Top management shall provide evidence of its commitment to the development and implementation of the quality management system and maintenance of its effectiveness by:</p> <ul style="list-style-type: none"> <li>a) communicating to the organization the importance of meeting customer as well as applicable regulatory requirements;</li> <li>b) establishing the quality policy;</li> <li>c) ensuring that quality objectives are established;</li> <li>d) conducting management reviews;</li> <li>e) ensuring the availability of resources.</li> </ul>	Issues Intrastake Rante mgs Doc index
Viamed Ltd ISO13485:20 16 5.4.1	<p><b>Quality objectives</b></p> <p>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 quality objectives shall be measurable and consistent with the quality policy.</p>	Intrastake Roles & Responsibilities
Viamed Ltd ISO13485:20 16 5.4.2	<p><b>Quality management system planning</b></p> <p>Top management shall ensure that:</p> <ul style="list-style-type: none"> <li>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;</li> <li>b) the integrity of the quality management system is maintained when changes to the quality management system are planned and implemented.</li> </ul>	management meetings Rante map
Viamed Ltd	<b>Responsibility and authority</b>	

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ISO13485:20 16.5.5.1	Top management shall ensure that responsibilities and authorities are defined, documented and communicated within the organization. Top management shall document the interrelation of all personnel who manage, perform and verify work affecting quality and shall ensure the independence and authority necessary to perform these tasks.	<i>Required Reading Issues</i>
Viamed Ltd ISO13485:20 16.5.5.2	<b>Management representative</b> Top management shall appoint a member of management who, irrespective of other responsibilities, 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.	<i>MD</i>
Viamed Ltd ISO13485:20 16.5.6.3	<b>Review output</b> The output from management review shall be recorded (see 4.2.5) and include the input reviewed and 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.	<i>Handstabs Doc Index management area</i>
Viamed Ltd ISO13485:20 16.6.1	<b>Provision of resources</b> The organization shall determine and provide the resources needed to: a) implement the quality management system and to maintain its effectiveness; b) meet applicable regulatory and customer requirements.	<i>Handstabs</i>
Viamed Ltd ISO13485:20 16.8.2.4	<b>Internal audit</b> 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	<i>Audit Calendar Task + Audit System Doc Index</i>

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	<p>defined and recorded (see 4.2.5). The selection of auditors and conduct of audits shall ensure objectivity and impartiality of the audit 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 eliminate detected nonconformities and their causes. Follow-up activities shall include the verification of the actions taken and the reporting of verification results.</p> <p>NOTE Further information can be found in ISO 19011.</p>	ISSUES
Viamed Ltd ISO13485:20 16.8.3.4	<p><b>Rework</b></p> <p>The organization shall perform rework in accordance with documented procedures that takes into account the potential adverse effect of the rework on the product. These procedures shall undergo the same review and approval as the original procedure.</p> <p>After the completion of rework, product shall be verified to ensure that it meets applicable acceptance criteria and regulatory requirements.</p> <p>Records of rework shall be maintained (see 4.2.5).</p>	Procedures Intrastats Doc index
Viamed Ltd ISO13485:20 16.8.5.3	<p><b>Preventive action</b></p> <p>The organization shall determine action to eliminate the causes of potential nonconformities in order to prevent their occurrence. Preventive actions shall be proportionate to the effects of the potential problems.</p> <p>The organization shall document a procedure to describe requirements for:</p> <ul style="list-style-type: none"> <li>a) determining potential nonconformities and their causes;</li> <li>b) evaluating the need for action to prevent occurrence of nonconformities;</li> <li>c) planning and documenting action needed and implementing such action, including, as appropriate, updating documentation;</li> <li>d) verifying that the action does not adversely affect the ability to meet applicable regulatory requirements or the safety and performance of the medical device;</li> <li>e) reviewing the effectiveness of the preventive action taken, as appropriate.</li> </ul> <p>Records of the results of any investigations and of action taken shall be maintained (see 4.2.5).</p>	Intrastats Issues Doc index DRC

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<b>INTERNAL PROCESS VERIFICATION</b>			
A. Management System: B. Management Responsibility C. Resource Management D. Product Realisation E. Design & Development F. Product Provision G. Process Monitoring The following are questions that should be asked and answered either through Internal audits or at this meeting			
1	Review Last years Audit. Update processes if required. Are all follow on Issue resolved satisfactory.	#160654 <i>outstanding - now completed</i>	Y
<b><u>A – MANAGEMENT SYSTEM</u></b>			
2	Is the Quality Statement Policy and Objectives reviewed annually. ISO – Document Index Task ID (300). Search Issues and review.	#1961651	Y
3	Is the process manual up to date. ISO – Document Index Task ID (548). Search Issues and review.	#1961640	Y
4	Is documentation checked prior to formal approval and issue.		Y
5	Check that there is a system in operation for the request for amendments.	Issues	Y
6	Verify that amendments are updated electronically and old copies archived.		Y
7	Are sales orientated records filed and archived correctly in the ORD files, in the office and archiving.	<i>new system almost all now digital</i>	Y
8	Has organisation Chart changed. VM3COP02.02	<i>Chairman passed away new Chairman to be decided</i>	Y
9	Has personnel responsibility descriptions changed. Roles Titles Processes and Procedures ADMIN Over View for complete list	<i>Josh has left this week need to re assign jobs</i>	Y
10	Check that the CE files are maintained by sole responsibility.		Y
11	Check that the Notified body is informed of major changes to Documentation.		Y
12	Check that electronic documents are regularly backed up and secure off site. ISO – Document Index Task ID (452). Search Issues and review.	#196798	Y

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13	Is the management system applications a series of process controls and are they in place throughout the organisation. Are processes identified and are charts produced to this effect and are copies of these charts easily accessible for use by personnel.	Intrastats, Audit 10	Y
14	Check the documented system for its policies and objectives and its control of the above processes and procedures. Is the Process Manual up to date and does it indicates the company's objectives. Are procedures are in place Are they available to all personnel Are other company documents i.e. Technical Drawings, Standards; Operators Manuals etc. also available and controlled	Intrastats, Audit 10 Roles and Responsibilities.	Y
15	Are the latest revision of documents controlled by version and date status and are they easily accessible. Is the Managing Director or designate manager still giving final approval for document changes.	Intrastats, Audit 10	Y
16	Is the Managing Director or designate manager still giving final approval for document changes.	Yes for Controlled documents	Y 198273
17	Has the Business Continuity Plan has expired. ISO – Document Index Task 266. <i>still valid</i>	#165447	Y edit to add controlled
<b><u>B - MANAGEMENT RESPONSIBILITY</u></b>			
18	Is Top management showing full commitment to the overall system and are communication lines in place. Manage Review Task 290.	Intrastats, Director in control of QA system #197451	Y
19	Are all customer requirements defined and met.	Contract Review Audit 2	Y
20	Are all the processes and objectives, undertaken within the company, documented in Intrastats and have a procedure. Is it measurable. <i>114 as well</i> Check process for measurable. <i>ID114 Task 10 548</i> Documented In Staff – Audit of Roles, titles and procedures.		Y # new task 10 x1 198273
21	Does the person responsible for the management systems have the authority to implement actions and reports directly to top management with the need for these actions	Managing Director	Y
22	Are reviews of the management system undertaken regularly and the results and actions relayed throughout the organisation. Task 290 for weekly review #197451 Task 114 for bigger overview #195987 in terms Task 746 for total review # 161905	Issues, Message of Day, company meetings, management meetings, Management weekly reviews	Y

*Job done issue incomplete  
meeting very recently done*  
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23	Are all required actions are undertaken in a timely manner and closed where appropriate.	Intrastats Issues	Y
24	Are all output requirements in such a format that verification against inputs, is applicable and appropriate. Is fitness for Purpose validated and is it measurable.  Staff – Audit of Roles, titles and procedures – click into details - review Scope and Risks. To check relevance.  Staff – Audit of Roles, titles and procedures check down the page for gaps in the IP 1-6 (end tick boxes)		Y
25	Are actions recorded against verifications completed in a timely and responsible manner.	Intrastats Issues	Y
26	Are design changes recorded and all the relevant information filed in the appropriate places.	Design control Audit 3 Intrastats No Design	NA
<b>C - RESOURCE MANAGEMENT</b>			
27	Has top management established a mechanism for identifying and providing required resources, training etc.	Training Audit 8	Y
28	Does this include existing and new personnel.	Training Audit 8	Y
29	Has top management identified the competency levels and attributes required for existing and new personnel.	Training Audit 8	Y
30	Is the competency of personnel monitored, verified and the appropriate records maintained	Training Audit 8	Y
31	Are personnel responsibilities defined.	Roles and Responsibilities	Y
32	Do individuals know their responsibilities, reporting and communicating lines. Each employee has 'My Roles' Link Task 314	Intrastats communication #198273	Y
33	Verify that all procedures, detail who is responsible for it.	processes A	Y
34	Check that these responsibilities also cover personnel Health & Safety functions – Health and Safety Controller.		Y
35	Is the need for equipment, plant, services etc. identified and acted upon where necessary.	Production meetings, management meetings Health and Safety Questionnaire.	Y
36	Has the basic working infrastructure been planned with conformity to requirements in mind.	Health & safety Audit 19	Y
37	Check validations of unknown process control criteria. Are there any unknown process.		Y

# 198273

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38	Are there adequate mechanisms in place for the identification, handling etc. of product through all stages.	COP/07 procedures + Bar Codes	Y
39	Are the controls in place, to safeguard customer property, adequate for full protection against loss damage etc.	COP/09	Y
40	Is the process for monitoring and measurement of product in place at all stages throughout the production process.	Production COPs	Y
41	Is the process for control of measuring equipment adequate for the monitoring of product verifications.	Calibration Audit 06	Y
42	Are validity processes <del>are</del> in place to safeguard product integrity.	Bar coding traceability	Y
<b>D - PRODUCT REALISATION</b>		No New Products	NA
43	Is the planning process for the realisation of product undertaken at the relevant stages.	X <sup>2</sup>	NA
44	Does planning identify documentation, testing and other such activities as required and that all appropriate records are maintained.	X <sup>2</sup>	NA
45	Are all customer requirements being addressed, including statutory and regulatory and that the capabilities are identified to meet those requirements.	Contract Review Audit 02	NA
46	Establish that mechanisms are in place to review all customer requirements prior to any commitments by the organisation.	Contract Review Audit 02	NA
47	Check that there are adequate arrangements for customer communications and feedback.	Contract Review Audit 02	NA
48	Is collation and analysis of all relevant data determined and effective. Is corrective actions identified.	X <sup>2</sup>	NA
49	Are these actions completed in a timely and adequate manner and are these actions part of continual improvements.	X <sup>2</sup>	NA
50	Does the organisation have preventive measures in place to control potential non-conformities.	X <sup>2</sup>	NA
51	Are all the above actions are reviewed adequately.		NA
<b>E - DESIGN &amp; DEVELOPMENT</b>			
52	Are procedures in place to ensure adequate planning of product design and that all relevant stages are identified.	Design control Audit 3	NA
53	Are the interfaces and assignments of responsibilities identified.	Design control Audit 3	NA
54	Are all input requirements determined. Is the documentation identified.	Design control Audit 3	NA
55	Are all output requirements in such a format, that verification against inputs is applicable and appropriate. Is Fitness for Purpose validated.	Design control Audit 3	NA
56	Are actions recorded against verifications completed in a timely and responsible manner.	Design control Audit 3	NA
57	Are validation processes in place and are they determined in accordance with the relevant requirements.	Design control Audit 3	NA

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58	Are design changes recorded and all the relevant information filed in the appropriate places.	Design control Audit 3	NA
<b>F - PRODUCT PROVISION</b>			
59	Are supplier profiles adequate and appropriate for the organisation. Are they monitored, for their ability to provide the requirements, is this monitoring controlled.	Purchasing Controls (Supplier Performance) Audit 5	Y
60	Is all the required information necessary, forwarded to suppliers in the correct format. Is this authorised prior to order placement.	Purchasing Controls (Supplier Performance) Audit 5 <i>Supply Review</i>	Y
61	Are goods and services received correct to the requirements stipulated.	Goods Inward Audit 9	Y
62	Are the provisions available, suitable for control of production and service, including procedures and equipment etc.	Production Audit 15	Y
63	Are there adequate mechanisms in place for the identification, handling etc. of product through all stages.	Production Audit 15	Y
64	Are the controls in place to safeguard customer property adequate for full protection against loss damage etc.	Production Audit 15	Y
65	Is the process for monitoring and measurement of products in place at all stages throughout the production process.	Production Audit 15	Y
66	Is the process, for control of measuring equipment, adequate for the monitoring of product verifications.	Calibration Audit 6	Y
<b>G - PROCESS MONITORING</b>			
67	Are mechanisms in place to monitor all relevant processes, including customer satisfaction. Are these verified against known criteria. Check process ID 114 / 548	# 198273	Y
68	Are controls in place for non-conforming product and processes. Are adequate to prevent unintended uses.	Goods Inward Audit 9 <i>Barcode</i>	Y
69	Where non-conforming product / process have been detected is appropriate action taken.	Goods Inward Audit 9 <i>Supplier Returns</i>	Y
70	Is collation and analysis of all relevant data determined and effective Is corrective actions identified.	Intrastats	Y
71	Are these actions completed in a timely and adequate manner. Are these actions part of continual improvements.	Issues	Y
72	Are all the above actions are reviewed adequately. Check process ID 114 / 548	# 198273 Annually	Y
73	Are regular analyses undertaken to identify any outstanding requirements. 548	# 198273 Intrastats	Y
74	Are necessary changes implemented where and when required.	<i>See Any follow up</i>	Y
75	Is any outsourcing done.		NA
76	Check the documented system for its policies, objectives and its control of the above processes and procedures.	Intrastats <i>doc 22429</i>	Y

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Intrastats – document index – VM3COP00.00 / VM3COP00.01.  
Check documents for location of objectives and policies.

Y

77 Are records of inspections filed.

## Audits

v

## **Sub Processes Linked to Audit**

Review the below processes tasks and audits and ensure they are completed in a timely manner.

### List Processes Per Title

## Managing Director

Process Scope	Roll Task	Roll Audit	Risk	Action	Notes / Issues
PROCESSID 7837 To Review the External Parties Influencing The QMS VST / Viamed Checked the Scopes and Risks, Review the Underlining Processes and Tasks	743 Managing Director	784 Company Secretary	Freq 1 Risk 1 Overall 1	Task 12M Audit 12M	#155651 ✓ #157833
PROCESSID 7845 Determine, provide and maintain the environment necessary for the operation of its processes and to achieve conformity of products and services.	745 Managing Director		Freq 1 Risk 1 Overall 1	Task 12M	#155251
PROCESSID 7846 To Comply with Top Level Re-authorise the Current Audits for next 12 Months Cover the Agenda as Per VOP13	746 Managing Director		Freq 1 Risk 1 Overall 1	Task 12M	#161905 and onwards

## IT Controller

Process Scope	Roll Task	Roll Audit	Risk	Action	Notes / Issues
<b>PROCESSID 7701</b> Amazon Web Services, is an online service, which basically simply provides a Linux PC out on the Web. Viamed uses this, for Web development of Websites: It hosts a working backup of many websites. Viamed / vst / vandagraph etc..	511	#196016	Freq 3 Risk 1 Overall 3	Task 1M	
<b>PROCESSID 7755</b> To Send Invoice for online services to Helen	597	#196213	Freq 3 Risk 1	Task 1M	

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		Overall 3		
PROCESSID 7832	Backup of all Sent Emails sent to External Address for Verification	731 #191334 Managing Director	Freq 4 Risk 1 Overall 4	Task 2W
PROCESSID 7850	Test the Goods out process disabling picking of items Goods Out not relating to an order	752 #195444 Managing Director	Freq 3 Risk 2 Overall 6	Task 3M Audit 12M
PROCESSID 7851	To test intrastats does not allow picking of unprocessed products to live customer orders	754 #186913 Goods Out	Freq 3 Risk 4 Overall 12	Task 6M Audit 12M
PROCESSID 7852	To attempt to Scan a product that has gone past its expire date.	756 #156616 Goods Out	Freq 3 Risk 2 Overall 6	Task 12M Audit 12M
PROCESSID 7853	Warehouse shelves can be tagged as sellable stock / unsellable stock. Either for quarantine purposes or holding items for other customer orders.	759 #195445 Goods Out	Freq 3 Risk 3 Overall 9	Task 12M Audit 12M
	Test that Order picking cannot pick unsellable stock locations to an Order	#159799		
PROCESSID 7854	Software Validation of the production lists.	761 #189070 Goods In	Freq 2 Risk 2 Overall 4	Task 3M Audit 6M
	By confirming no extra production jobs are stuck in the system, and all listed production jobs are found. the production tracking is validated	#175534		
PROCESSID 7855	Software Validation - Production Lists	761 #189070 Goods In	Freq 2 Risk 2 Overall 4	Task 3M Audit 6M
	Review the current active production lists in intrastats to the actual in progress production lists	#175534		
PROCESSID 7856	To check order picking cannot pick against an unchecked order	764 #161540 Office Processes	Freq 2 Risk 2 Overall 4	Task 12M Audit 12M
PROCESSID 7857	To confirm Software Validation Stock Tracking Check, is functioning as expected	763 #172867 Goods In	Freq 2 Risk 1 Overall 2	Task 6M
PROCESSID 7858	Test the QA System that Staff not trained for QA are unable to QA a Product.	766 #185657 Office Processes	Freq 3 Risk 3 Overall 9	Task 6M
PROCESSID 7861	Software Validating Of Training Documents via Forced Required Reading	768 #155911 Managing Director	Freq 1 Risk 2 Overall 2	Task 12M
PROCESSID 7865	Software Validation of the system: To check all process(s) tasks and audits are not clashed with the same person doing the Task as the	779 #156187 Managing Director	Freq 1 Risk 1 Overall 1	Task 12M Audit 12M
		#156187		

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#### PROCESSID 7870

Scope to check the automatic system of tagging product non conformance and other issues to the post market surveillance review report.

789

Managing

Director

# 156791

Freq 1

Risk 1

Overall 1

Task 12M

#### PROCESSID 7875

To test document control is working as intended.

802

Managing

Director

# 151063

803

Company

Secretary

# 157854

Freq 1

Risk 1

Overall 1

Task 12M

Audit

12M

#### PROCESSID 7879

To check the Scheduled Tasks and Audits is working as Intended.

808

Managing

Director

# 162935

809

Company

Secretary

# 15036

Freq 1

Risk 4

Overall 4

Task 12M

Audit

6M

#### PROCESSID 7880

To confirm the out of documents computer software functions as expected flagging out of date items on the list

808

Managing

Director

# 162935

809

Company

Secretary

# 15036

Freq 1

Risk 1

Overall 1

Task 12M

Audit

6M

#### PROCESSID 7881

To compare Opera Live Orders to Intrastats Back order Active List

810

Managing

Director

# 157161

Freq 1

Risk 3

Overall 3

Task 12M

#### ISO Controller

### Process Scope

#### PROCESSID 6866

Review the Internal Process and Verification's are suitable for the current standards

Roll Task

55

Managing

Director

# 155516

Roll Audit

/

Overall 1

Risk

Task 12M

Freq 1

Risk 1

Overall 1

#### PROCESSID 7827

To review the Quality policy and check it is still valid and upto date.

301

Managing

Director

# 194653

Freq 1

Risk 1

Overall 1

Task 12M

#### PROCESSID 7828

To review the Quality policy and check it is still valid and upto date.

723

Managing

Director

# 194706

Freq 1

Risk 1

Overall 1

Task 12M

### Audits

#### Process Scope

#### PROCESSID 7723

To carry out Audit 10b Process Verification Viamed

Roll Task

3

Managing

Director

Roll Audit

/

Overall 2

Risk

Task 12M

Freq 1

Risk 2

Overall 2

#### Now Defunct - See Audit 20

#### PROCESSID 7730

To carry out Audit 20 Process Verification To Management Viamed

172

Company

Secretary

# 157177

Freq 1

Risk 2

Overall 2

Audit

12M

#### PROCESSID 7771

To carry out Audit 10b Process Verification VST

177

Managing

Director

#

Freq 1

Risk 2

Overall 2

# Internal Audit Check list

## INTERNAL PROCESS VERIFICATION

Created:	17/May 1995	<b>Audit No 20</b> Sit With management and Complete this Form	
Revised:	06 August 2020		Page 14 of 14
Audit Date		Auditor	

Now Defunct - See Audit 20

PROCESSID 7778

To carry out Audit 20 Process Verification To Management VST

181	Freq 1	Audit
Company Secretary	Risk 2	12M
	Overall 2	

# 157782