

Internal Audit Check list			
Audit of Audits			
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Audit Date	6-12-22	Auditor DLAMTB	

Company / ISO Section	Criteria of ISO Section	Auditor Comments
VST Ltd ISO9001:2015 5.3	Organizational roles, responsibilities and authorities Top management shall ensure that the responsibilities and authorities for relevant roles are assigned, communicated and understood within the organization. Top management shall assign the responsibility and authority for: a) ensuring that the quality management system conforms to the requirements of this International Standard; b) ensuring that the processes are delivering their intended outputs; c) reporting on the performance of the quality management system and on opportunities for improvement (see 10.1), in particular to top management; d) ensuring the promotion of customer focus throughout the organization; e) ensuring that the integrity of the quality management system is maintained when changes to the quality management system are planned and implemented.	
VST Ltd ISO9001:2015 9.2.1	The organization shall conduct internal audits at planned intervals to provide information on whether the quality management system: a) conforms to: 1) the organization's own requirements for its quality management system; 2) the requirements of this International Standard; b) is effectively implemented and maintained.	
VST Ltd ISO9001:2015 9.2.2	The organization shall: a) plan, establish, implement and maintain an audit programme(s) including the frequency, methods, responsibilities, planning requirements and reporting, which shall take into consideration the importance of the processes concerned, changes affecting the organization, and the results of previous audits; b) define the audit criteria and scope for each audit; c) select auditors and conduct audits to ensure objectivity and the impartiality of the audit process; d) ensure that the results of the audits are reported to relevant management; e) take appropriate correction and corrective actions without undue delay; f) retain documented information as evidence of the implementation of the audit programme and the audit results. NOTE See ISO 19011 for guidance.	
Viamed Ltd ISO13485:2016 5.6.2	General The input to management review shall include, but is not limited to, information arising from:	

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Review input	<ul style="list-style-type: none"> 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; j) changes that could affect the quality management system; k) recommendations for improvement; l) applicable new or revised regulatory requirements. 	Management Review Risk, files Route Map Audit calendar Doc index QA system QI 21 forms
Viamed Ltd ISO13485:2016 8.2.4	Internal audit The organization shall conduct internal audits at planned intervals to determine whether the quality management system: <ul style="list-style-type: none"> 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 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. NOTE Further information can be found in ISO 19011.	Doc index Audit calendar Route Map
Viamed Ltd ISO13485:2016 8.5.1	General The organization shall identify and implement any changes necessary to ensure and maintain the continued suitability, adequacy and effectiveness of the quality management system as well as medical device safety and performance through the use of the quality policy, quality objectives, audit	Management Review meetings

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results, post market surveillance, analysis of data, corrective actions, preventive actions and management review.

TO BE COMPLETED LAST IN THE ANNUAL AUDIT CALENDAR

	QUESTION:	RESPONSE:	Y/N
1	Are there any audits outstanding.	2021 Completed 2022 on Schedule	Y
2	Are there any related issues outstanding to the audits.	2021 Completed 2022 Completed	Y
3	Are there any corrective actions not signed off.	No	Y
4	Are there any follow up actions not completed.	No	N
5	Is each audit properly numbered and dated.		Y
6	Has each audit got the current years processes linked to it. Are audit processes updated annually.		Y
7	Is each audit correctly signed off.	#279819 further Action Required	N
8	Have results of audits been brought to the attention of the person responsible where appropriate.		Y
9	Is there evidence that the frequency of audits should be changed.		N

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List Processes Per Title

Share Holder

Process Scope	Roll Task	Roll Audit	Risk	Action	Notes / Issues
PROCESSID 7862 Review The Audit Calendar Screen		173 Managing Director	Freq 1 Risk 1 Overall 1	Audit 12M	

Managing Director

Process Scope	Roll Task	Roll Audit	Risk	Action	Notes / Issues
PROCESSID 38 Management oversight of Internal Tasks and Audits Issue(s). Review the responses to Tasks and Audits. ensure they are being fulfilled and completed.	730 Managing Director		Freq 1 Risk 1 Overall 1	Task 12M	

ISO Controller

Process Scope	Roll Task	Roll Audit	Risk	Action	Notes / Issues
PROCESSID 7093 Review of outstanding Audits	725 Managing Director		Freq 1 Risk 1 Overall 1	Task 12M	

Humanmed Controller

Process Scope	Roll Task	Roll Audit	Risk	Action	Notes / Issues
PROCESSID 7670 Review of Humanmed sales and orders and clear any duplicates or problems.	611 Office Processes		Freq 3 Risk 1 Overall 3	Task 1M	

Audits

Process Scope	Roll Task	Roll Audit	Risk	Action	Notes / Issues
PROCESSID 7731 To carry out Audit 21 Audit Of + Audit Viamed	270304 Managing Director	173	Freq 1 Risk 2 Overall 2	Audit 12M	
PROCESSID 7779 To carry out Audit 21 Audit Of + Audit VST	279306 Managing Director	192	Freq 1 Risk 2 Overall 2	Audit 12M	

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Month	Description	TaskID	ProcessID	Date	Signed off	Issues outstanding
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Jan

Jan	Audit 03 Design Control Viamed	TaskID 22	ProcessID 7716			
Jan	Audit 01 Picking Packing Viamed	TaskID 24	ProcessID 7860			
Jan	Audit 03 Design Control VST	TaskID 193	ProcessID 7764			
Jan	Audit 01 Picking Packing VST	TaskID 194	ProcessID 7762			

Feb

Feb	Audit 10b Process Verification Viamed	TaskID 3	ProcessID 7723			
Feb	Audit 10b Process Verification VST	TaskID 177	ProcessID 7771			
Feb	Audit 27 Software Validation	TaskID 821	ProcessID 7892			

Mar

Mar	Audit 09 Goods Inward And Product Identity Viamed	TaskID 170	ProcessID 7721			
Mar	Audit 12 CE Files Viamed	TaskID 16	ProcessID 7725			
Mar	Audit 09 Goods Inward And Product Identity VST	TaskID 174	ProcessID 7769			
Mar	Audit 12 CE Files VST	TaskID 176	ProcessID 7773			

Apr

Apr	Audit 22 Post Market Surveillance Viamed	TaskID 14	ProcessID 7732			
Apr	Audit 07 Handling And Storage Viamed	TaskID 25	ProcessID 7719			

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Apr Audit 07 Handling And Storage TaskID ProcessID 7767
VST 178

Apr Audit 22 Post Market Surveillance TaskID ProcessID 7780
VST 180

May

May Audit 06 Calibration Viamed TaskID ProcessID 7718
20

May Audit 15 Production Viamed TaskID ProcessID 7727
28

May Audit 15 Production VST TaskID ProcessID 7775
175

May Audit 06 Calibration VST TaskID ProcessID 7766
182

Jun

Jun Audit 08 Training Viamed TaskID ProcessID 7720
10

Jun Audit 10 Documentation Control TaskID ProcessID 7722
Viamed 27

Jun Audit 10 Documentation Control TaskID ProcessID 7770
VST 183

Jun Audit 08 Training VST TaskID ProcessID 7768
184

Jul

Jul Audit 23 Analysis Of Data TaskID ProcessID 7733
Viamed 43

Jul Audit 11 Repairs And Service TaskID ProcessID 7724
Viamed 171

Jul Audit 11 Repairs And Service TaskID ProcessID 7772
VST 179

Jul Audit 23 Analysis Of Data VST TaskID ProcessID 7781
185

Aug

Aug Audit 19 Health And Saftey TaskID ProcessID 7729
Viamed 13

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Aug Audit 19 Health And Saftey VST TaskID 186 ProcessID 7777

Aug Audit 24 Due Servicing TaskID 288 ProcessID 7889

Sep

Sep Audit 02 Contract Review TaskID 36 ProcessID 7715
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Sep Audit 05 Purchasing Suppliers TaskID 37 ProcessID 7717
Viamed

Sep Audit 02 Contract Review VST TaskID 187 ProcessID 7763

Sep Audit 05 Purchasing Suppliers TaskID 190 ProcessID 7765
VST

Oct

Oct Audit 18 Management Review TaskID 21 ProcessID 7886
Viamed

Oct Audit 18 Management Review TaskID 188 ProcessID 7887
VST

Oct Audit 04 Accounts TaskID 817 ProcessID 7885

Nov

Nov Audit 14 Complaints And TaskID 30 ProcessID 7726
Corrective Actions Viamed

Nov Audit 20 Process Verification To TaskID 172 ProcessID 7730
Managment Viamed

Nov Audit 20 Process Verification To TaskID 181 ProcessID 7778
Managment VST

Nov Audit 14 Complaints And TaskID 189 ProcessID 7774
Corrective Actions VST

Dec

Dec Audit 17 Internal Audits Viamed TaskID 11 ProcessID 7728

Dec Audit 21 Audit Of Audit Viamed TaskID 173 ProcessID 7862

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Dec Audit 17 Internal Audits VST TaskID 191 ProcessID 7776

Dec Audit 21 Audit Of Audit VST TaskID 192 ProcessID 7779