Internal Audit Check list					
Handling & Stock Control					
Created:	17/May 1995	Audit No 07			
Revised:	27 April 2022		Page 1 of 10		
Audit Date		Auditor			

Company / ISO Section	Criteria of ISO Section	Auditor Comments / Issues
VST Ltd ISO9001:201 5 7.1.4	Environment for the operation of processes The organization shall determine, provide and maintain the environment necessary for the operation of its processes and to achieve conformity of products and services. NOTE A suitable environment can be a combination of human and physical factors, such as: a) social (e.g. non-discriminatory, calm, non-confrontational); b) psychological (e.g. stress-reducing, burnout prevention, emotionally protective); c) physical (e.g. temperature, heat, humidity, light, airflow, hygiene, noise). These factors can differ substantially depending on the products and services provided.	
VST Ltd ISO9001:201 5 7.1.5.1	General 7.1.5.1 General The organization shall determine and provide the resources needed to ensure valid and reliable results when monitoring or measuring is used to verify the conformity of products and services to requirements. The organization shall ensure that the resources provided: a) are suitable for the specific type of monitoring and measurement activities being undertaken; b) are maintained to ensure their continuing fitness for their purpose. The organization shall retain appropriate documented information as evidence of fitness for purpose of the monitoring and measurement resources.	
VST Ltd ISO9001:201 5 8.1	Operational planning and control The organization shall plan, implement and control the processes (see 4.4) needed to meet the requirements for the provision of products and services, and to implement the actions determined in Clause 6, by: a) determining the requirements for the products and services; b) establishing criteria for: 1) the processes; 2) the acceptance of products and services; c) determining the resources needed to achieve conformity to the product and service requirements; d) implementing control of the processes in accordance with the criteria; e) determining, maintaining and retaining documented information to the extent necessary: 1) to have confidence that the processes have been carried out as planned; 2) to demonstrate the conformity of products and services to their requirements.	

The output of this planning shall be suitable for the organizations operations. The organization shall control planned changes and review the consequences of unintended changes, taking action to mitigate any adverse effects, as necessary. The organization shall ensure that outsourced processes are controlled (see 8.4). VST Ltd General ISO9001:201 The organization shall ensure that externally provided processes, products 5 8.4.1 and services conform to requirements. The organization shall determine the controls to be applied to externally provided processes, products and services when: a) products and services from external providers are intended for incorporation into the organization's own products and services; b) products and services are provided directly to the customer(s) by external providers on behalf of the organization; c) a process, or part of a process, is provided by an external provider as a result of a decision by the organization. The organization shall determine and apply criteria for the evaluation, selection, monitoring of performance, and re-evaluation of external providers, based on their ability to provide processes or products and services in accordance with requirements. The organization shall retain documented information of these activities and any necessary actions arising from the evaluations. VST Ltd Type and extent of control ISO9001:201 The organization shall ensure that externally provided processes, products and services do not adversely affect the organization's ability to 5 8.4.2 consistently deliver conforming products and services to its customers. The organization shall: a) ensure that externally provided processes remain within the control of its quality management system; b) define both the controls that it intends to apply to an external provider and those it intends to apply to the resulting output; c) take into consideration: 1) the potential impact of the externally provided processes, products and services on the organization's ability to consistently meet customer and applicable statutory and regulatory requirements; 2) the effectiveness of the controls applied by the external provider; d) determine the verification, or other activities, necessary to ensure that the externally provided processes, products and services meet requirements. VST Ltd Control of production and service provision ISO9001:201 The organization shall implement production and service provision under 5 8.5.1 controlled conditions. Controlled conditions shall include, as applicable: a) the availability of documented information that defines: 1) the characteristics of the products to be produced, the services to be provided, or the activities to be performed; 2) the results to be achieved; b) the availability and use of suitable monitoring and measuring resources;

c) the implementation of monitoring and measurement activities at

appropriate stages to verify that criteria for control of processes or outputs,

	and acceptance criteria for products and services, have been met; d) the use of suitable infrastructure and environment for the operation of	
	processes; e) the appointment of competent persons, including any required qualification;	
	f) the validation, and periodic revalidation, of the ability to achieve planned results of the processes for production and service provision, where the	
	resulting output cannot be verified by subsequent monitoring or measurement;	
	g) the implementation of actions to prevent human error;h) the implementation of release, delivery and post-delivery activities	
VST Ltd ISO9001:201	Identification and traceability The organization shall use suitable means to identify outputs when it is	
5 8.5.2	necessary to ensure the conformity of products and services. The organization shall identify the status of outputs with respect to monitoring and measurement requirements throughout production and service provision.	
	The organization shall control the unique identification of the outputs when traceability is a requirement, and shall retain the documented information necessary to enable traceability.	
VST Ltd	Property belonging to customers or external providers	
ISO9001:201 5 8.5.3	The organization shall exercise care with property belonging to customers or external providers while it is under the organization's control or being used by the organization.	
	The organization shall identify, verify, protect and safeguard customers' or external providers' property provided for use or incorporation into the products and services.	
	When the property of a customer or external provider is lost, damaged or otherwise found to be unsuitable for use, the organization shall report this to the customer or external provider and retain documented information on	
	what has occurred. NOTE A customer's or external provider's property can include materials,	
	components, tools and equipment, premises, intellectual property and personal data.	
VST Ltd	Preservation	
ISO9001:201 5 8.5.4	The organization shall preserve the outputs during production and service provision, to the extent necessary to ensure conformity to requirements. NOTE Preservation can include identification, handling, contamination control, packaging, storage, transmission or transportation, and protection.	
VST Ltd ISO9001:201 5 8.7.1	The organization shall ensure that outputs that do not conform to their requirements are identified and controlled to prevent their unintended use or delivery.	
	The organization shall take appropriate action based on the nature of the nonconformity and its effect on the conformity of products and services. This shall also apply to nonconforming products and services detected after	
	delivery of products, during or after the provision of services. The organization shall deal with nonconforming outputs in one or more of the following ways:	
	a) correction;b) segregation, containment, return or suspension of provision of products	

	and services;	
	c) informing the customer;	
	d) obtaining authorization for acceptance under concession.	
	Conformity to the requirements shall be verified when nonconforming	
	outputs are corrected.	
VST Ltd	General	
ISO9001:201	The organization shall determine:	
5 9.1.1	a) what needs to be monitored and measured;	
	b) the methods for monitoring, measurement, analysis and evaluation	
	needed to ensure valid results;	
	c) when the monitoring and measuring shall be performed;	
	d) when the results from monitoring and measurement shall be analysed	
	and evaluated.	
	The organization shall evaluate the performance and the effectiveness of	
	the quality management system.	
	The organization shall retain appropriate documented information as	
	evidence of the results.	
Viamed Ltd	Infrastructure	
16 6.3	needed to achieve conformity to product requirements, prevent product	
10 0.5	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 shall be maintained	
Viamed Ltd	Work environment	
ISO13485:20		
16 6.4.1	environment needed to achieve conformity to product requirements.	
10 0.1.1	If the conditions for the work environment can have an adverse effect on	
	product quality, the organization shall document the requirements for the	
	work environment and the procedures to monitor and control the work	
	environment.	
	The organization shall:	
	a) document requirements for health, cleanliness and clothing of personnel	
	if contact between such personnel and the product or work environment	
	could affect medical device safety or performance;	
	b) ensure that all personnel who are required to work temporarily under	
	special environmental conditions within the work environment are	
	competent or supervised by a competent person.	
	NOTE Further information can be found in ISO 14644 and ISO 14698	
Viamed Ltd	Contamination control	
ISO13485:20		
15015405.20	rs appropriate, the organization shall plan and document arrangements for	

16 6.4.2	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.	
Viamed Ltd ISO13485:20	Planning of product realization The organization shall plan and develop the processes needed for product	
16 7.1	realization. Planning of product realization shall be consistent with the requirements of the other processes of the quality management system.	
	The organization shall document one or more processes for risk management in product realization.	
	Records of risk management activities shall be maintained (see 4.2.5). In planning product realization, the organization shall determine the	
	following, as appropriate:	
	a) quality objectives and requirements for the product;b) the need to establish processes and documents (see 4.2.4) and to provide resources specific to the product, including infrastructure and work	
	environment; c) required verification, validation, monitoring, measurement, inspection and test, handling, storage, distribution and traceability activities specific to	
	the product together with the criteria for product acceptance;	
	d) records needed to provide evidence that the realization processes and resulting product meet requirements (see 4.2.5).	
	The output of this planning shall be documented in a form suitable for the	
	organization's method of operations. NOTE Further information can be found in ISO 14971.	
Viamed Ltd	Control of production and service provision Production and service provision shall be planned, carried out, monitored	
16 7.5.1	and controlled to ensure that product conforms to specification. As	
	appropriate, production controls shall include but are not limited to: a) documentation of procedures and methods for the control of production	
	(see 4.2.4); b) qualification of 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 post-delivery 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 approved for distribution. The record shall be verified and	
	approved.	
Viamed Ltd ISO13485:20	Customer property The organization shall identify, verify, protect, and safeguard customer	
16 7.5.10	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).	
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Viamed Ltd ISO13485:20 16 7.5.11	Preservation of product 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 provide preservation. If special conditions are required, they shall be controlled and recorded (see 4.2.5).	
Viamed Ltd	Cleanliness of product	
ISO13485:20 16 7.5.2	contamination control of product if: a) product is cleaned by the organization prior to sterilization or its use; b) product is supplied non-sterile 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.	
Viamed Ltd	Identification	
ISO13485:20 16 7.5.8	The organization shall document procedures for product identification and identify product by suitable means throughout product realization. The organization shall identify product status with respect to monitoring and measurement requirements throughout product realization. Identification of product status shall be maintained throughout production, storage, installation and servicing of product to ensure that only product that has passed the required inspections and tests or released under an authorized concession is dispatched, used or installed. If required by applicable regulatory requirements, the organization shall document a system to assign unique device identification to the medical device. The organization shall document procedures to ensure that medical devices returned to the organization are identified and distinguished from conforming product.	
Viamed Ltd ISO13485:20 16 7.5.9.1	General The organization shall document procedures for traceability. These procedures shall define the extent of traceability in accordance with applicable regulatory requirements and the records to be maintained (see 4.2.5).	
Viamed Ltd	Monitoring and measurement of product	
ISO13485:20	The organization shall monitor and measure the characteristics of the	

16 8.2.6	product to verify that product requirements have been met. This shall be carried out at applicable stages of the product realization process in accordance with the planned and documented arrangements and documented procedures. Evidence of conformity with the acceptance criteria shall 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.	
Viamed Ltd ISO13485:20 16 8.3.1	General 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)	
Viamed Ltd ISO13485:20 16 8.3.2	Actions in response to nonconforming product detected before delivery The organization shall deal with nonconforming product 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).	

	QUESTION:	RESPONSE	Y/ N
1	Check all issues from the previous audit are completed.		
2	Check that incoming products are stored correctly on receipt.		
3	Check that the in-house stores area is adequate, safe and accessible.		

4	Verify that products for repair are suitably boxed prior to movement. i.e. In ducket with correct paperwork including SRS number.	
5	Verify that stock items are suitable packed and labelled for entry into stock.	
6	Check that gloves and or hand sanitiser is available and used, where necessary, when returns are received.	
7	Check in Intrastats that COSHH data sheets are available for all products.	
8	Check that items in a stock locations are correct to Intrastats. Verify that the quantity of an item in stock is correct to that in Intrastats. Check that the packing and labelling of the finished product is appropriate and will preserve quality to the end user. Check 5 items. 1. 2. 3. 4. 5.	
9	Check that demonstration and exhibition stock is separate from other stock, and areas labelled correctly.	
10	Verify that product in the non-conforming area can only be removed by authorised personnel. Verify that transfer of non-conformance stock is done by use form QC19.	
11	Verify that special requirement areas are available should the product require it.	
12	Check that completed products are adequately stored. List those checked. 1. 2. 3. 4. 5.	
13	Verify that there are adequate storage areas in the workshop for a working stock of assembly	

	components.	
14	Check that product movement around the workshop is by ducket only.	
15	Are stores and storage areas secure and suitably identified with signs. List problem areas.	
16	Are uncontrolled material and parts identified as such, and in the correct area. Check that items in Quarantine have HOLD labels with an issue number, date and initials.	
17	Check unentered and pre QA items have labels and/or are in the correct area.	
18	Are all parts in the warehouse properly identified with Viamed Location Tracking barcodes. Identify unmarked items.	

Sub Processes Linked to Audit 07

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

List Processes Per Title

Product Controller					
Process Scope	Roll Task	Roll Audit	Risk	Action	Notes / Issues
PROCESSID 7873	800		Freq 2	Task	
Review the Highs and Lows in	Director 3		Risk 2	1	
Temperature of stored stock and	(Steve)		Overal		
products.			1 4		
Warehouse Team Leader					
Process Scope	Roll Task	Roll Audit	Risk	Action	Notes / Issues
PROCESSID 5858	110	261	Freq 4		
Opera Counts bulk stock in and issues			Risk 1		
stock out against orders.			Overal		
Multiple processes cause stock to be			1 4		
used internally,					
Opera requires a weekly update to					
bring the stock count into line with					
whats been used outside the invoicing					

systems					
NO LONGER REQUIRED, New system live counts these now					
PROCESSID 5935 To allocate stock that has not automatically be linked to a repair or invoice.	447 Company Secretary		Freq 4 Risk 1 Overal 1 4	I I	
PROCESSID 6850 Review current stock levels	615 Goods In	778 Managing Director	Freq 4 Risk 1 Overal 1 4	2W	
PROCESSID 6945 To synchronise Opera stock Count against Intrastats internal movement of stock, E.G. Items that wont uniquely appear on an opera order - such as production parts.	110	783 Managing Director	Freq 4 Risk 2 Overal 18		
PROCESSID 7673 To check that all the stock on the selves are within their use by dates.	294 Goods In	477 Managing Director	Freq 3 Risk 2 Overal 16		
PROCESSID 7689 Move Stock From QA Shelf To Stock Shelf	545 Goods In		Freq 4 Risk 1 Overal 1 4	Task 1W	
PROCESSID 7694 Move Stock From QA Shelf To Stock Shelf	544 Goods In	782	Freq 4 Risk 1 Overal 14	1W	
PROCESSID 7695 Move Stock From QA Shelf To Quick Shipping Shelves	495 Goods In		Freq 4 Risk 1 Overal 14	Task 1W	
PROCESSID 7866 Ensure we do not run out of oxygen	785 Production Processes		Freq 2 Risk 1 Overal 12	I I	
PROCESSID 7902 Empty depleted sensor bin from the office	876	877	Freq 1 Risk 1 Overal 1 1		
PROCESSID 7903 Empty Warehouse depleted sensor bin into Bin in cage and record weights in document in T Drive	878 Goods In	879	Freq 1 Risk 1 Overal 1 1	Task 1M Task 3W Audit	

				6M	
PROCESSID 7904	880	881	Freq 1	Task	
Check Weeee waste pallet and sensor	Goods In	001	Risk 1		
bin,			Overal		
arrange collection if FULL			1 1	12M	
PROCESSID 7942	1036	1037	Freq 1	Task	
To make sure we have a QA procedure	Company	Managing	Risk 3		
or service manual in place for all our	Secretary	Director	Overal		
stock coming through Viamed and			13	12M	
VST.					
Some may just say check packaging					
and barcode and other may need to go					
further in depth. With testing					
procedures.					
Those who do not require testing					
should state this in the procedure.					
Audits					
Process Scope	Roll Task	Roll Audit	Risk	Action	Notes / Issues
PROCESSID 7719		25	Freq 1	Audit	
To carry out Audit Audit 07 Handling		Company	Risk 2		
And Storage Viamed		Secretary	Overal		
		1.0	12		
PROCESSID 7767		178	Freq 1		
To carry out Audit 07 Handling And Storage VST		Company Secretary	Risk 2 Overal	12M	
Storage VS1		Secretary	12		
Production Processes					
Process Scope	Roll Task	Roll Audit	Risk	Action	Notes / Issues
PROCESSID 7940	1003	1004	Freq 1	Task	1100057 155005
To check the date of the grease used in		Managing	Risk 1	1	
the production and servicing of the	Secretary	Director	Overal	Audit	
Tom Thumb. To see if it needs to be			1 1	12M	
removed. Look at date purchased then					
add 4 years to the date.					
Dispose of this when it goes beyond					
this date.					
PROCESSID 7944	1011	1012		Task	
To check the use by date or	Managing	Company	Risk 1		
_ ·	Director	Secretary	1		
			1 1	1 2 MI	
of.					
Dispose of this when it goes beyond this date. PROCESSID 7944 To check the use by date or manufacturers life span, of any Sealant, Glues, Greases, Sprays, Gases And Tapes You Use In Production, Service And Repairs For Viamed And VST. To see if it needs to be disposed				6M	

Dispose of and where needed re order			
new, when it goes beyond this date.			