

AN AGREEMENT is made the day of 2008

BETWEEN Viamed Limited whose registered office is situate at
15 Station Road, Crosshills, Keighley, West Yorkshire, BD20 7DT. United Kingdom.

AND
Saadat

WHEREAS:

1. This Agreement ensures that development design manufacture testing packaging approval release and delivery are all in accordance with the quality assurance measures specified herein
2. This Agreement assumes the mutual Trust of each party hereto and neither party shall utilise information in any way which it receives by virtue of this Agreement unless specifically authorised to do so by the other party hereto This clause shall have effect both during the currency of this Agreement and after termination
3. This Agreement relates to all articles manufactured and supplied by. Saadat as specified in the order from Viamed Saadat shall be responsible to Viamed for all sub-contractors on the same terms as pertain to Saadat under this Agreement notwithstanding that Viamed may have approved a sub-contractor and may have demanded that Saadat make purchases from a specified sub-contractor

IT IS HEREBY AGREED AS FOLLOWS

1. QUALITY CONTROL SYSTEM

Saadat must have an introduced efficient quality control system which complies with ISO 9001/7.94 and EN 4600/10.93 or ISO9001:2000 and ISO 13485:2003 and which has been third party certified and approved by a certifying body notified in the EU or alternatively have already commenced a process to introduce ISO 9001/9002 and has planned to comply within 12 months from the date hereof
Compliance with quality system requirements must exist each time an article is manufactured tested packaged and released

2. DEVELOPMENT AND DESIGN

Saadat shall be responsible for and pursue development and design work so that only fault-free articles supplied according to written specifications and intended operating reliability must continue after delivery by Saadat

Design shall be verified against functional specifications and written documents must be prepared for this purpose and comprise both method and measurement results

Such documents are to be kept on file in accordance with paragraph 10 hereof

3. MANUFACTURING DOCUMENTATION

Saadat shall assess and prepare all documentation required for manufacture testing approval packaging and release Saadat shall keep all submitted documentation on file in accordance with paragraph 10 hereof

4. MANUFACTURING MATERIALS AND MANUFACTURING COMPONENTS

Viamed shall be responsible for the purchase and arrival tests of those materials and components supplied by Third parties to Viamed prior to Viamed supplying Saadat

5. MANUFACTURING AND PROCESS VALIDATION

All manufacturing processes are to be validated and the methods and results are to be documented which documents must be kept on file in accordance with paragraph 10 hereof

For the assembly of parts of articles. Saadat may only use those sub-contractors who have an introduced efficient quality control system which complies with the ISO 9001 or 9002 or have already commenced a process to introduce ISO 9001/2 and who have planned to comply with such stipulations within 12 months

from the date hereof. 'Assembly' shall include pre-delivery assembly by sub-contractors of multiple components and manufacturing according to instruction by drawings. Manufacturing of pre-production samples must be in accordance with paragraph 9 hereof. Series production of new articles including when revisions have been made must be started in accordance with paragraph 6 hereof and parts of article must be made by the same sub-contractors who manufactured the same parts of the pre-production series. Only the specifications as approved by Viamed shall be used in respect of materials, process, measurement equipment, tools or quality control as in accordance with paragraph 6 hereof.

6. NEW ARTICLES AND REVISIONS

Before the manufacture of new articles commences, pre-production samples will be manufactured and validated with processes in accordance with approved documents. The pre-production series must be checked and approved in writing by Viamed. Where Saadat intends to make alterations to the agreed specifications in any manner, Viamed must be informed in writing immediately. Such changes in written specification of any article must be approved in writing by Viamed and previously issued documents must be replaced before any revised production (altered as necessary) in validated processes. Pre-production samples must be checked and approved in writing by Viamed. Saadat shall perform control measurements of pre-production samples and supply measurement protocols to Viamed showing measurement results confirming that submitted pre-production samples are in compliance with written specifications. Viamed reserves the right to waive the right to view such measurement protocols but where such waiver is invoked, Saadat remains liable for the supply of articles which comply with written specifications. All aforementioned written information/documentation prepared by Saadat, including measurement protocols and written approvals by Viamed, shall be kept on file in accordance with paragraph 10 hereof.

7. TESTS

Saadat shall perform tests in accordance with paragraph 3 hereof and protocols shall be kept showing that processes and tests were performed by Saadat in accordance with the documentation referred to in paragraph 3 hereof. Saadat must also assess quality control to identify defects and all defects must be eradicated immediately. Here, defects are discovered in final inspection, defective parts of any batch must be separated from the remainder of the batch immediately.

8. DELIVERIES, RELEASE AND CERTIFICATES

Prior to delivery, Saadat must approve and release articles such release to be based upon the results of documented approved tests of materials and components and on tests performed during the manufacturing process up to final inspection and including pre-packaging with markings and contents and according to ISO 9001 paragraph 4.10.4, 'Final Inspection and Testing'. Saadat shall be responsible for ensuring that supplied articles comply with any written specifications and also that certificates accompanying deliveries are in accordance with paragraph 10 hereof. Certificate contents are specified in Appendix A1 hereto.

9. TRACEABILITY

Saadat shall ensure traceability between received materials and components and Quality documents which lie on file in respect of articles delivered to Viamed.

10. FILING QUALITY CONTROL DOCUMENTS

Saadat shall file all documents as detailed in this Agreement under paragraphs 3, 5, 6, 7, 8 and 12 and additionally in accordance with ISO 9001 paragraph 4.10.5 'Inspection and Test Records' and paragraph 4.16 'Control of Quality Records'.

Should Viamed request any final test document which is held on file, copies of such documentation must be supplied to Viamed within 7 days from the date of request. All documents must be kept on file for 5 years after issue.

11. AUDITS

Viamed hereby reserves the right to visit Saadat together with any sub-contractors which have not been third party certified and approved by a certifying body notified within the EU.

Such visits are to enable Viamed to carry out a quality audit and check processes and documents which are on file. The right of access shall apply where Viamed is the subject of inspection by a public agency where such agency demands to inspect Saadat or a sub-contractor of Saadat.

The right of access hereby reserved shall at all times be preceded by written notice of an intended visit such notice to be 7 working days minimum

12. QUALITY INFORMATION

there must be a continuous exchange of quality information between Saadat . and Viamed such information being the basis for the introduction of requisite improvements to ensure that safety risks operational and production disruptions never occur Saadat must immediately advise Viamed in writing in respect of any technical problems and suspected defects in articles which. Saadat . has already delivered to Viamed Saadat must in such cases ensure that sub-contractors provide supporting information Viamed must immediately advise Saadat in writing of any discovery of defect Defective articles in respect of which Viamed submits claims or returns will be processed as complaints and both Viamed and Saadat shall process complaints in accordance with ISO9001 paragraph 8.5.2

Saadat must always and forthwith advise Viamed in writing of:

- i. the cause of any defect
- ii. the steps taken to prevent repetition of any defect in the future
- iii. an analysis of the risk of patient injury and functional disruption where appropriate
- iv. any other information requested by Viamed in any complaint matter
- v. Both parties hereto agree to inform the other in respect of anything which may impact upon quality and quality control including major changes in staff and organisation Where quality information is addressed to anyone other than the respective Contact Person (as detailed in paragraph 13 below) a copy of such information must always be sent in addition to the Contact Person

13. VALIDITY AND CONTACT PERSONS

This Agreement shall remain in force until terminated by either party hereto serving upon the other written notice of not less than 6 months

The parties hereto have nominated the following as Contact Persons who shall address technical matters or other matters arising out of this Agreement in the first instance

For Viamed: John S Lamb Tel: +44 1535 634542
Fax: +44 1535 635582

For...Saadat . :

SIGNED ON BEHALF OF VIAMED LIMITED:

Name :

Position :

SIGNED ON BEHALF OF.....

Name :

Position :

Appendix A1

Certificate contents

Deliveries must be accompanied by a certificate specifying that the supplied articles comply with all the written specifications.

Certificates shall be based on documented, approved results from:

The checks the supplier deems necessary to ensure that all delivered articles, including their marking and packaging, comply with specifications, e.g. on arrival inspections of materials and components, inspections during and after the manufacturing process and inspections of marking and packaging.

Certificates must contain the following information, where applicable:

The supplier's company name

Text certifying that all delivered articles comply with all the written specifications of the article.

Viamed's order number.

Viamed's article designation and article number.

The number of approved units

The date the certificate was issued.

A signature, name, tel. no., fax no., and department of the person responsible for ensuring that certificate information is correct.

A signature, name, tel. no., fax no-, and department of the person responsible for releasing the articles.

Deliveries without an accompanying certificate will be deemed defective as will deliveries containing defective parts which were approved by any unauthorized employee of Viamed

Any "approval" of deliveries by Viamed must be in writing, and a copy of same must in such instances accompany the certificate with the defect specified.

The policy of Viamed is to return deliveries deemed to be defective.