

pharmaxis	FM-002-11	Date Effective: 11-Aug-23	Page 1 of 10
SUPPLIER APPROVAL QUESTIONNAIRE			

The purpose of this questionnaire is to assist Pharmaxis in assessing suppliers for inclusion in our approved supplier list for provision of material, service and computerised system providers which are essential to the quality of our products.


All information will be treated in the strictest confidence.

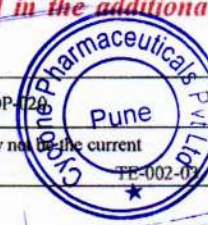
We would be grateful if you could take the time to complete this questionnaire, and return it as soon as possible to the Pharmaxis contact below.

SECTION A) TO BE COMPLETED BY THE PHARMAXIS REQUESTOR

Name of Pharmaxis Contact: Rachel Downing		Position: QA Officer	
Address	Pharmaxis Ltd. 20 Rodborough Road FRENCHS FOREST NSW 2086 AUSTRALIA		
Phone No.	02 9454 7214	Date	05Dec23
E-mail	rachel.d@amapharma.com		
Supplier Name	Cogniso Pty Ltd		
Supplier Address	Level 3, 476-478 George Street, Sydney NSW 2000		
Material / Service Required	Provision of IT services		
Specification	N/A		
Manufacturer/Principle Supplier Name and Address (if not the direct supplier)	N/A		

REQUESTOR NOTE. If the supplier is a consultant they must provide evidence to show they have adequate education, training, and experience, or any combination thereof, to advise on the subject for which they are engaged. A request for copies of these documents should be entered in the additional questions section of this form.

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pharmaxis	FM-002-11	Date Effective: 11-Aug-23	Page 2 of 10
SUPPLIER APPROVAL QUESTIONNAIRE			

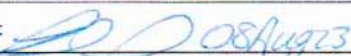
Prior to sending to the supplier ensure additional questions, if required, are entered in the allocated area at the end of this form.

Sections A and B is to accompany sections C or D as applicable.

Section C is to be completed by material and service providers.

Section D is to be completed computerised system providers.

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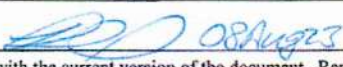


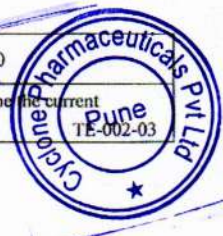
pharmaxis	FM-002-11	Date Effective: 11-Aug-23	Page 3 of 10
SUPPLIER APPROVAL QUESTIONNAIRE			

**SECTION B) TO BE COMPLETED BY THE MATERIAL/SERVICE
SUPPLIER/COMPUTERISED SYSTEM REPRESENTATIVE**

Supplier Name	Cogniso Pty Ltd		
Questionnaire Completed By	Name:	Ian McArthur	
	Position:	Managing Director	
	E-mail:	ian@cogniso.com.au	Phone No. 0439 452 981
	Date Completed:	02/01/2024	

LICENCING / CERTIFICATION
<p>What critical licencing or certification does your company require in order to supply the specified material, service or computerised system to Pharmaxis.</p> <p>Please provide the following details:</p> <p>Issuing Authority Name :</p> <p>Licence / Certificate No.:</p> <p>Scope of Licence / Certification :</p> <p>Please <u>attach</u> a copy of the certificate(s).</p> <p>If there is more than one critical licence /certification required please provide further details below:</p> <p>.....</p> <p>.....</p> <p>.....</p> <p>.....</p> <p>.....</p>
<p>What other licencing or certification has your company obtained.</p> <p>Please <u>attach</u> a copy of the certificate(s).</p>

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SUPPLIER APPROVAL QUESTIONNAIRE**SECTION C) TO BE COMPLETED BY THE MATERIAL/SERVICE SUPPLIER**

Complete each question by circling the applicable answer. If the question is not relevant to your business enter N/A and move on to the next question.

Please do not hesitate to contact the Pharmaxis Requestor via the contact details on page 1, should you have any queries regarding the questions.

Please attach scanned pdf copies of required documentation in your Email response.

QUESTIONNAIRE	Circle Applicable Answer
GENERAL	
Can you manufacture / supply the material or service required as defined in Section A) meeting all specified requirements?	<input checked="" type="radio"/> Yes / No
Would you have to subcontract any or all of the material manufacture /service being supplied to Pharmaxis. If yes Please provide details:.....	Yes / <input checked="" type="radio"/> No
Do you have a procedure for the selection of subcontractors	Yes / No / <input checked="" type="radio"/> N/A
Do you agree to obtain written consent from Pharmaxis via the Pharmaxis contact before subcontracting any work relating to this material/service?	Yes / No / <input checked="" type="radio"/> N/A
Does your company have a Quality department. If yes, how many Quality staff: Ref. No. TR/001	<input checked="" type="radio"/> Yes / No / N/A
Does your company have training records for Quality department staff.	<input checked="" type="radio"/> Yes / No / N/A
CALIBRATION PROVIDERS	
If you provide calibration services, do you have applicable certification to perform these services? (Please attach certification)	Yes / <input checked="" type="radio"/> No / N/A
If you are not certified to perform calibration, how are the reference standards maintained used for calibration of clients equipment.	
EVIDENCE OF QUALITY SYSTEM	
Do you have current, documented work instructions for each process associated with the material / service required?	<input checked="" type="radio"/> Yes / No / N/A
Do you have a documented system for control, calibration and maintenance of measuring and test equipment?	<input checked="" type="radio"/> Yes / No / N/A

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08 Aug 23

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
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SUPPLIER APPROVAL QUESTIONNAIRE

QUESTIONNAIRE	Circle Applicable Answer
Do you operate documented procedures for investigating non-conformities (including customer complaints) and taking corrective action?	<input checked="" type="radio"/> Yes / No / N/A
Do you have documented procedures for handling deviations	<input checked="" type="radio"/> Yes / No / N/A
Do you have documented procedures for change management	<input checked="" type="radio"/> Yes / No / N/A
Do you have documented procedures for storage and maintenance of records?	<input checked="" type="radio"/> Yes / No / N/A
Do you retain records for a minimum retention period of 5 years.	<input checked="" type="radio"/> Yes / No / N/A
Is there a procedure for creation, review, approval and maintaining procedures.	<input checked="" type="radio"/> Yes / No / N/A
Do you have an internal audit program	Yes / No / N/A
Contact name and position for Pharmaxis to contact when requesting for current documentation (e.g. licences, certifications, Agreements)	
Name:	Position:
SUPPLIER SPECIFIC QUESTIONS	
	Yes / No / N/A
	Yes / No / N/A
	Yes / No / N/A
	Yes / No / N/A

End of Section C.

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
pharmaxis	FM-002-11	Date Effective: 11-Aug-23	Page 6 of 10
SUPPLIER APPROVAL QUESTIONNAIRE			

We appreciate you taking the time to complete this questionnaire. Your responses assist us in upholding the quality of our products and the safety of our patients.

Supplier please ensure you attach required documentation before returning back to the Pharmaxis requestor.

Questions have been appropriately completed by the supplier.

Sign and Date Pharmaxis Requestor:.....

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SUPPLIER APPROVAL QUESTIONNAIRE

SECTION D) TO BE COMPLETED BY THE COMPUTERISED SYSTEM SUPPLIER REPRESENTATIVE

This section of the questionnaire is only to be sent to computerised system providers.

QUESTIONNAIRE – COMPUTERISED SYSTEMS	Circle Applicable Answer
Software Development	
Does your company have a written policy/procedure for the Software Development Life Cycle (SDLC) methodology (i.e. Project Initiation → Project Planning → Requirements Gathering → System Design → System Development → System Testing → System Release) followed within your company?	<input checked="" type="radio"/> Yes / No / N/A
If yes, provide details on governing Procedure/ Document: Ref. SOP No - SOP/IT/019/00 Ref. SOP No. - SOP/IT/020/00 Ref. SOP No - SOP/IT/022/00	
Software Quality Policies, Standards, and Procedures	
Is the development & testing of systems performed & documented at multiple layers (e.g. module/ code review/integration /IQ/OQ).	<input checked="" type="radio"/> Yes / No / N/A
Is there a source code listing?	<input checked="" type="radio"/> Yes / No / N/A
If yes, provide details Ref. SOP No. - SOP/IT/011/00 Ref. SOP No. - SOP/IT/006/00	
Does your company have written procedures for the system testing strategy at each level of development (such as unit/module testing (UT), system testing (ST), integration testing (IT), stress/ performance testing and user acceptance testing (UAT)?	<input checked="" type="radio"/> Yes / No / N/A
Related document/ procedure:	
Does your company have a written procedure for executing tests (such as defining acceptance criteria/expected results, retaining raw data, recording test results, handling and resolving test failures (variances), and reviewing test results)?	<input checked="" type="radio"/> Yes / No / N/A
Related document/ procedure:	
Does your company have a written procedure for verifying the traceability between requirements/design specification and executed tests?	Yes / No / N/A
Ref. SOP No. - SOP/IT/018/00	
If yes, please attach documented evidence that traceability verification was performed for the system/ product, under consideration.	

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SUPPLIER APPROVAL QUESTIONNAIRE

QUESTIONNAIRE – COMPUTERISED SYSTEMS	Circle Applicable Answer
Personnel/ Supplier / Subcontractor Qualifications and Performance	
Is your company's staff qualified & trained appropriately to develop/configure/customize the application software/computer system? Is the staff trained on regulatory requirement such as 21 CFR Part 11/Annexure 11 Ref. Document Name - Key Technical Staff (TS/001) Provide details as an attachment.	<input checked="" type="radio"/> Yes / No / N/A
Has the application been tested for compliance to regulatory requirements such as 21 CFR Part 11 / Annexure 11 ?	<input checked="" type="radio"/> Yes / No / N/A
Is any of the software used to provide the service to Pharmaxis purchased from a third party. If yes, has it been subjected to regulatory compliance checks such as 21 CFR Part 11?	Yes / No / N/A lan
Software Code Control / Security & Maintenance Practices	
Is access to your company's system development environment or Data/Infrastructure center (in case of cloud based systems) restricted? Related procedure/ document: Ref. SOP. NO. - SOP/IT/007/00 Ref. SOP NO. - SOP/IT/016/00 Ref. SOP. NO. - SOP/IT/016/00	<input checked="" type="radio"/> Yes / No / N/A N/A
Does your company have a written procedure for backup and recovery (including secure storage and onsite and offsite handling of media) of software code/ system data (in case of cloud based systems)? Related procedure/ document: Ref. SOP. NO - SOP/IT/010/00 Venkat Ref. SOP NO. - SOP/IT/007/00 Venkat Ref. SOP. NO - SOP/IT/008/00	<input checked="" type="radio"/> Yes / No / N/A Yes
Has your company performed disaster recovery testing of software code or system data/ network infrastructure (in case of cloud based systems)? SOP/IT/018/00 Yes Mamatha/ Venkat	<input checked="" type="radio"/> Yes / No / N/A Yes Mamatha/ Venkat
Does the system provide for archival and retrieval of records? Related procedure/ document: Yes Mamatha/ Venkat	<input checked="" type="radio"/> Yes / No / N/A
In case of cloud products/services, has the aspects such as network infrastructure physical/ logical security framework and data security measures have been tested and documented?	<input checked="" type="radio"/> Yes / No / N/A N/A

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pharmaxis	FM-002-11	Date Effective: 11-Aug-23	Page 9 of 10
SUPPLIER APPROVAL QUESTIONNAIRE			

SECTION D) CONTINUED

QUESTIONNAIRE- COMPUTERISED SYSTEMS	Circle Applicable Answer
Software Release / Customer Support Practices	
Does your company have a reporting system for communicating to customer's upgrades/patches and/or new releases to the system? Related procedure/ document: <i>Ref. SOP/IT/017/00</i> <i>Ref SOP/IT/016/00</i>	<input checked="" type="radio"/> Yes / No / N/A
Does your company have written procedure for bug fixes and patch management? Can you describe how customers are alerted: <i>Ref. SOP/IT/027/00</i>	<input checked="" type="radio"/> Yes / No / N/A
Documentation Support	
Does your company have documented evidence that the system has been tested for the base or out-of-box functionality? If yes, please attach a Test Summary Report/ Qualification Certificate for the system. <i>(It is part of validation)</i>	<input checked="" type="radio"/> Yes / No / N/A
Supplier Specific Questions	
	Yes / No / N/A <i>Yes Mamatha/ Venkat</i>
	Yes / No / N/A
	Yes / No / N/A

End of Section D.

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pharmaxis	FM-002-11	Date Effective: 11-Aug-23	Page 10 of 10
SUPPLIER APPROVAL QUESTIONNAIRE			

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
Supplier please ensure you attach required documentation before returning back to the Pharmaxis requestor.

Questions have been appropriately completed by the supplier.

Sign and Date Pharmaxis Requestor:.....

Sign and Date Pharmaxis Computer Systems Engineer:.....

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