

REPUBLIC OF THE PHILIPPINES Department of Health

NATIONAL CENTER FOR MENTAL HEALTH





BIDS AND AWARDS COMMITTEE

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Section VI Schedule of Requirements

ITB No. 026-2022

The delivery schedule expressed as weeks/months stipulates hereafter a delivery date which is the date of delivery to the project site.

ITEM NO.	DESCRIPTION	QTY	TOTAL	Delivered, Weeks/Months
AUS22-01	Specifications:	6	646,800.00	60 to 90 calendar
AUS22-01	 Specifications: Power Supply Voltage: At least 127V – 230V Frequency: 50/60 HZ Power Consumption: At least 1400 – 1700W Electric shock protective type: Class one equipment applied parts Type B Mode of Operation: Continuous operation Protection against harmful water penetration or particulate materials: IPXO Sterilization pressure: At least 1.3 – 2.1 kg/cm2 Operation temperature: 122 C – 128 C (123 to 133 C) Sterilization Time: Can be selected at least 16 – 30 minutes Volume: At least 16 -23 liters Diameter of the chamber: At least 230 	6	646,800.00	60 to 90 calendar days
	 240 mm (diameter) 410 - 450 mm (dep.) 12. Water consumption per cycle 350 - 380 ml 			
	13. Net weight / Gross weight: 30kg/33kg 14. Fuse of Protection: F1 – 0.5a/f2 and F3 = 20a/f4 = 10a			
	Standard Requirements			
	A certification that the unit being offered must be brand new and not a			
	discontinued model.			

2. Current and valid Certificate Manufacturer's compliance with ISO and or CE certificate or its equivalent 3. Certification that the supplier has the capability for corrective and preventive maintenance of the unit. 4. Bidder's certificate that the parts shall be available at the authorized Philippine service center/s for a period of five (5) years after the warranty period. 5. Certification that the brand has been in the local market for at least five (5) years with at least 2 to 3 installations. 6. Certification that the supplier will be responsible for the notification, transportation to the site, delivery, installation and testing on the site (hospital/health facility) and expenses for such will be on the account of the supplier 7. Certification to provide users' Manual in English language: a. Operational Manual – 2 copies b. Service Manual – 2 copies 8. Certification that the supplier / bidder shall provide free installation of the equipment 9. Warranty Certificate: one (1) year for parts and two (2) years for service from the date of delivery, inspection and

CONFORME	D BY:
(Signature o	ver printed name)
DATE:	

acceptance



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ITEM NO.	DESCRIPTION	QTY	TOTAL	Delivered, Weeks/Months
UPU22-01	ULTRASOUND MACHINE WITH PORTABLE ULTRASOUND	1	7,700,00.00	60 to 90 calendar days
	System Standard Features			
	The equipment must be capable and with applications for Radiology and Obstetrics, Gynecology & Cardiology			
	The equipment must be console cart-type system			
	Power Supply			
	3. The equipment must have a: a. Voltage: 100 - 240 VAC at 50/60 Hz;			
	b. A power rating of 650 VA max – 450 VA min			
	4. The supplier must provide an online UPS 2KVA, AVR and isolation transformer for both equipment.			
	System Architecture Design			
	5. Height with Monitor at least 1600 mm			
	6. Width at least 500 mm			
	7. Depth at least 700 mm			
	8. Weight at least 50 kg (without peripherals) 9. Four (4) active transducer ports			
	10. Four (4) swivel and lockable caster wheels			
	11. Integrated image storage at least 500 GB 12. Integrated Digital B/W thermal printer			
	13. The system shall have at least 5 built-in Probe holders			
	14. At least two external USB port			
	15. The system shall have a built-in Gel Holder and warmer			
	16. User Interface Operator keyboard			
	Main Monitor			

- 17. At least Flat Panel Display, 21.5inch widescreen (LED/LCD) color with:
 - a. Brightness and contrast adjustment
 - b. Tilt/Rotate/Pan
 - c. Rotate, swivel and lift function
- 18. High resolution monitor display: with at least 1920 x 1080 pixels resolution

Control Panel

- 19. at least 10 inch touch screen panel
- 20. at least with a 1024 x 800 resolution (Control Panel)

Scanning Parameters

- 21. Imaging frequency range from 2 to 16 MHz or wider range (probe dependent)
- 22. Gray Scale at least 256 shades
- 23. Dynamic Range at least 256 dB
- 24. Maximum Frame Rate of at least 1750 fps (Hz)
- 25. Displayed imaging shall have a depth range of at 2 33 cm or a wider range.
- 26. Transmission Focus selection: 1 8 focal points

System Standard Features

- 27. Will continuously optimize the brightness, contrast and uniformity of images when scanning different tissues.
- 28. Image and Cineloops storage
- 29. Cine playback function
- 30. The system shall provide post-processing functions in:
 - a. B-mode
 - b. Color Doppler Mode
 - c. Pulse wave Doppler
 - d. M-mode
 - e. Continuous Wave Doppler
 - f. Power Doppler
 - g. Anatomical M-Mode
 - h. Elastography
 - i. 4D
- 31. Digital Beamforming
- 32. A speckle reduction filter or application must be available
- 33. Image raw data analysis
- 34. Post-image optimization
- 35. Real-time automatic Doppler Calculations
- 36. Cardiac Measurements
- 37. Post measurement must be available
- 38. Auto IMT

- 39. Tissue Harmonic Imaging
- 40. Comprehensive measurements, body markers, calculations and application specific reports including Vascular, Cardiac, OB/GYN and General Imaging.
- 41. On screen annotation
- 42. Digital Calipers/Measurement
- 43. Panoramic imaging or extended view
- 44. Dual Image Cine Display
- 45. Quad Image Cine Display

Transducers/probes:

- 46. Convex array probe with Frequency range: 2 5 MHz or wider range
- 47. Linear array probe; Frequency at least: 4.5 12 MHz
- 48. Endocavity array probe; Frequency at least: 4 9 MHz
- 49. Phase array; Frequency at least: 2 4 MHz

Portable Ultrasound:

- 50. Physical Specification
 - a. Console: Laptop Style with Cart
 - b. At least 1 active probe port and at least 2 extension ports
 - c. At least 4 probe holders
 - d. Integrated external UPS
 - e. Built-in printer
 - f. Gel warmer
- 51. Electrical Specification
 - a. Input: 100-240 VAC, 50/60Hz
 - b. Output: not more than 200VA

Console

- 52. Storage: 256 GB
- 53. Windows 10 (64 bit)
- 54. At least 4 USB ports
- 55. 1 video output HDMI port
- 56. Lithium Battery at least 60 mins scanning time

Trolley Cart

- 57. Height at least 950 mm, Width at least 350 mm, Depth at least 300 mm
- 58. 4 Swivel wheel cart based type
- 59. Operation Mode
 - a. 2D Mode
 - b. Color Doppler Mode
 - c. Power Doppler
 - d. M-mode
 - e. PW Doppler
 - f. CW Doppler

- g. TDI
- h. Anatomical M Mode
- 60. Transducers
 - a. Convex Probe Frequency range of 2 5
 MHz or higher range
 - b. Endocavity Probe Frequency range of 49 MHz or higher range
 - Linear Probe Frequency range of 4.2 –
 13 MHz or higher range

Monitor

- 61. Display: at least 15 inch High Resolution LCD Monitor
- 62. Display Size: 1924 x 768
- 63. Brightness/contrast/color temperature adjustment

Data transfer

- 64. DICOM 3.0 compliant
- 65. Support digital data transfer via a DICOM network for both printing & storage.
- 66. Shall be capable of sending DICOM images to our PACS/RIS server
- 67. The system shall have a Open Modality Worklist Function to be connected to our existing PAC/RIS.

Standard Accessories

- 68. Ultrasound bed
- 69. Bar stool with height adjustment and footstool
- 70. At least five rolls of Thermal Papers
- 71. 3 gallon ultrasonic gel
- 72. 6 bottles of probe cleaner
- 73. Multipurpose Ultrasound Phantom

Inclusions

- 74. Two desktop computers with at least;
 - a. Processor: Equivalent to 11th Generation Intel Core i7
 - b. Memory: 8 GB RAM
 - c. Hard Disk: 500 GB HDD
 - d. Operating System: Windows 10 OS and license Microsoft Office
 - e. Video cards compatible to the existing PACS/RIS.
- 75. Each computer will be supplied with keyboard, mouse, 24" LED HD monitors and 650 VA UPS
- 76. All-in-one printer with ink tank (Toner Type)
- 77. 3TR Floor Mounted Air conditioner

Warranty

- 78. With three (3) years warranty for parts and services with free semi-annual preventive maintenance after end user acceptance/authorized hospital personnel including all transducers. Warranty shall commence upon the acceptance of the end user. The winning bidder shall issue a certificate reflecting the duration of warranty period.
- 79. Bidder's certificate guaranteeing the availability on the supply of spare parts ten (10) years from end of production. A certification must be issued by the manufacturer for the bidder.
- 80. With quarterly preventive maintenance and calibration
- 81. Certification in case that the equipment will be for repair or maintenance within the warranty period, it should be done onsite. Otherwise, a service unit with equivalent or higher specification than the supplied machine shall be provided.

Standard Requirements

- 82. The supplier shall secure a certificate from the manufacturer stating that the equipment is brand new, unused and not a discontinued model or was listed in the market recall. The certificate reflect the brand, model, serial number of the machine, and the date and place it was manufactured.
- 83. The bidder must provide the current and valid certificate of Manufacturer's compliance with ISO 13485.
- 84. The supplier shall secure a certificate from the manufacturer stating that both equipment is at least of mid-range level. The certificate must be issued by the manufacturer for the bidder.
- 85. Delivery of equipment and all accessories within 90 calendar days upon receipt of Notice to Proceed (NTP).
- 86. The machine should have a US FDA Approval or CE Mark Approval. The bidder shall provide the manufaturer's approved US FDA Premarket Notification (PMN) or 510(k) certificate or CE Mark Approval certificate.
- 87. Certification that the supplier has the capability for corrective and preventive

- maintenance of the unit. The certificate must be issued by the manufacturer for the bidder.
- 88. Certification from the manufacturer authenticated by the Philippine consulate from the country of origin of the unit that the warranty should not be affected with a change of distributor.
- 89. Certification that the brand must be in the local market for a minimum of ten (10) years with at least 3 installations in both private and government facilities.
- 90. Certification that the bidder is existing in the local market for a minimum of ten (10) years.
- 91. Certification from the supplier will be responsible for the notification, transportation, delivery, installation, acceptance testing and commissioning and expenses for such will be on the account of the supplier.
- 92. Certification that the supplier/bidder shall provide applications training for users and maintenance personnel of the hospital.
- 93. On-house training with certificate for the End-User Familiarization of the operating procedures of the equipment
- 94. Certification to provide user's Manuals:
 - Two (2) sets of Service Manual in English
 Language upon delivery of the equipment
 - b. Two (2) sets of User Manual in English
 Language upon delivery of the equipment
- 95. Certification that the suppliers/bidder shall provide free installation of the equipment.
- 96. Current and Valid Certificate of Authorized or Exclusive Distributorship
- 97. The manufacturer must have a local office and service center in the Philippines for at least 10 years. Provide a certification from manufacturer.
- 98. 24/7 Technical Support.
- 99. Both equipment should be of the same brand.
- 100. A certification of good performance from at least three (3) tertiary hospital stating the name of facility, address, type of equipment and its model, and the date of delivery must be provided.

CONFORMED BY:
(Signature over printed name)
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