

Republic of the Philippines Department of Health METRO MANILA CENTER FOR HEALTH DEVELOPMENT



SUPPLEMENTAL/ BID BULLETIN NO. 1

IB No. 2025-010 Procurement of Glucometer Strips (EARLY PROCUREMENT ACTIVITY)

This Supplemental/Bid Bulletin No. 1 is being issued to revise provisions/specifications in the Bidding Documents for a forecited project:

Query during Pre-bidding Conference:								
Technical Specification	Query	Response of the End User Unit						
9,130 boxes	228,250 strips with one (1) blood lancet per strip, automatic lancing device	228,250 strips with one (1) blood lancet per strip, automatic lancing device						
Battery: Lithium coin cell battery	Battery: Lithium coin cell battery	Battery: Lithium coin cell battery GRANTED						

Furthermore, this is to inform bidders that the bidding will be moved on the **29th of November**, **2024** instead of November 27, 2024 due to lack of quorum

Bidders are advised to use the following attached forms and submit them together with all required documents for the submission of bids on the 29th day of November 2024, 9:00 AM:

This Supplemental/Bid Bulletin No. 1 shall be integral to the Bidding Documents. All other provisions indicated in the bidding documents not affected by this Supplemental/Bid Bulletin No. 1 shall remain in effect.

For guidance and information of all concerned.

Issued this 20th day of November 2024 in MMCHD

Approved by: JEREMIAS FRANCIS Y. CHAN, MD Licensing Officer V / BAC Chairperson

	Republic of the Philippi	nes						
	Department of Healt							
	Metro Manila Center for Health Development TECHNICAL SPECIFICATIONS							
ltem	Glucometer Strips	Qty./Unit	228,250 strips					
No. 1								
Name of	Manufacturer:	Country of Origin (if applicable)						
Brand:		Model: (if applicable)						
ABC: P 1,	369,500.00							
PURCHAS	SER'S SPECIFICATION	STATEMENT OF COMPLIANCE						
SPECIFIC	ATIONS:							
	strips with one (1) blood lancet per strip, ic lancing device							
Test Rang	ge: 20-600mg/dl (1.1-33.3mml/L							
Test Bloo	d Volume: 0.5-1 uL							
Sample T	ypes: Capillary/Venous whole blood							
Packaging: 25 strips per canister/box/manufacturer's standard								
With one	glucometer control solution for every 500 strips							
Assay Ra	nge: 0.50-33.30mmol/L (9-600mg/DL)							
	Period: Thirty (30) to Sixty (60) calendar days after f the Notice to Proceed							
Delivery	Place: DOH MMCHD Pasig Warehouse							
B. Upon	delivery, the following shall be complied with:							
Shelf life	at least 18 months upon delivery							
On each canister/box, the following should be legibly imprinted or stickered with non-removable or permanent sticker/label that is binding and with residue and tearing, if removed:								
the Tech	tional Requirements to be attached to nical Specifications form arranged, numbered ed as enumerated below:							
(CPR) or	and current Certificate Product Registration Valid Extension issued by the Philippine Food Administration (PFDA);							
award. If	must be valid for the entire period of the the CPR is about to expire, the supplier must mitted a copy of an application of renewal to							

the FDA at least 3 months before the expiry date copy of the expiring CPR which is stamped with "Extension of Validity" shall be submitted as pro [AO 2019-0041]	an
2. Valid and current License to Operate (LTO) for consuppliers, distributors and traders issued by Philippe Food and Drugs Administration (PFDA). Provided, to the application for renewal was made timely as DOH AO No. 2016-003: In case of expired LTO, following copies may be submitted: (i) expired LTO, application for renewal with FDA document track number; and, (iii) Official Receipt as proof of payme of renewal of LTO	pine that per the ; (ii) king
3. Product Insert/Product Information or download from the internet and other manufacture unamended sales literature, unconditional statement of specification and compliance issued by manufacturer, samples, independent test data etc. appropriate for cross-referencing statement compliance to the technical specification in accordate to what is indicated in Technical Specifications;	er's ents the , as of
4. Certification from Manufacturer/Distributor/Importer/Wholesaler reflected in the Certificate of Product Registration the product/s to be bid) that the Bidder is authorized dealer or distributor of the product	
5. Certificate of Compliance to the Electronic Drug P Monitoring System (EDPMS) issued by either Pharmaceutical Division (PD) of the DOH or D Regional Health Office/Centers for Health Developm pursuant to DOH Administrative Order No. 2018-0 and RA 9502 and its IRR;	the DOH nent
In case of an expired Certificate of Compliance EDPMS, refer to DOH Department Circular (DC) No 0001, "Interim Guidelines on the Certificate of Com	o.2023-

D. Additional requirement by the Lowest/Single Calculated Bid (L/SCB) as part of post qualification:

- 1. You are requested to submit within (5) five days upon receipt of this notice three (3) copies of all documents needed for Post Qualification of the following documents:
 - a. Eligibility Documents
 - i. (Mayor's Permit (latest annual and quarterly)
 - ii. SEC/DTI Registration,

Procurement Activities for Drugs and Medicines."

to Electronic Drug Price Monitoring System for Government

- iii. Tax Clearance)
- b. Certificate of Registration from BIR
- c. Income Tax Returns latest payment
- d. Bid Bulletin
- e. Product Sample /Brochure

- f. Authority from the Manufacturer to Distribute the Product
- g. License to Operate
- h. And other documents stated in BDS
- 2. One (1) original sample of the manufacturer's product to be submitted and returned after evaluation. The sample submitted and approved during the evaluation shall be the same item to be delivered upon contract award. The prototype of the labeling instruction must be part of the sample submitted; however, the technical specifications of the labeling instruction of the product must be complied with upon delivery.
- E. Product Recall & Disposal:

1. The Supplier must ensure the quality of products and if there will be problems in the quality, the Supplier will recall and replace the products distributed in the regions/hospitals/treatment hubs/RHU/HC/BHSS based on Guidelines on Product Recall, FDA Circular No. 2016-012;

2. In instances of product recalls due to failures of suppliers and manufacturers to comply with standards of safety and quality, the cost associated with proper disposal/ destruction, handling or pull out from health facilities where these products have already been distributed shall be borne by the supplier (subject to the latest policy for disposal) (DOH Administrative Order (AO) No. 2019-0041)

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

ltem Number	Description	Quantity	Total ABC (Php)	Delivery Site	Delivered, Weeks/Months
	Glucometer Strips	228,250 strips	P 1,369,500.00	DOH-MMCHD Pasig Warehouse	30 -60 calendar days after receipt of approved P.O./NTP

Signature over Printed Name

[date of signing]

In the capacity of:

Duly authorized to sign bid for and on behalf of:

[title or other appropriate designation] (Name of Company) [Complete office address] [Contact No.] [Fax No.] [Email Address]