



SUPPLEMENTAL/ BID BULLETIN NO. 1

IB#2024-130
Procurement of Protein Purified Derived Solution

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

1. Query during Pre-bidding Conference:		
Technical Specifications	Query	Response of the End User Unit
Shelf Life: Must have a minimum shelf life of eighteen (18) months remaining at the time of delivery	1.Shelf Life: Must have a minimum shelf life of twelve (12) months remaining at the time of delivery.	1.Shelf Life: Must have a minimum shelf life of twelve (12) months remaining at the time of delivery. GRANTED

Bidders are advised to use the **following attached forms and submit them together with all required documents for the submission of bids on the 22nd day of August 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 13th day of August 2024 in MMCHD

Approved by:


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

Republic of the Philippines
 Department of Health
 Metro Manila Center for Health Development
TECHNICAL SPECIFICATIONS

Item No. 1	Protein Purified Derived Solution	Qty./Unit	1,000 vials
Name of Manufacturer:		Country of Origin (if applicable)	
Brand:		Model: (if applicable)	
ABC: P 1,000,000.00			
PURCHASER'S SPECIFICATION		STATEMENT OF COMPLIANCE	
<p>Specifications:</p> <p>Tuberculin Purified Protein Derivative Solution for Injection, Intra dermal</p> <p>Dosage Strength and form: 5- TU/O 1ml, 5ml</p> <p>Multi Dose Vial</p> <p>Storage 2 to 8 degrees Celsius (35 to 46 degrees)</p> <p>Transport at 2 to 8 degrees Celsius (35 to 46 degrees)</p> <p>Do not Freeze</p> <p>Enough antigen for 50 test per vial</p> <p>Delivery Period: 60 calendar days from the receipt of approved NTP</p>			
<p>B. Upon delivery, the following shall be complied with:</p> <p>1.Shelf Life:</p> <p><i>Must have a minimum shelf life of twelve (12) months remaining at the time of delivery.</i></p> <p>2. Packaging Instructions:</p> <p>Standard packaging of the manufacturer as approved by PFDA, including product insert or encryption/imprint inside the box</p> <p>3. Labeling instructions:</p> <p style="padding-left: 40px;">In each box, the following should be legibly imprinted or stickered:</p> <p style="padding-left: 80px;">“Philippine Government Property-Department of Health”</p> <p style="text-align: center; padding-left: 40px;">NOT FOR SALE</p> <p style="padding-left: 40px;">Date of Manufacture: _____</p> <p style="padding-left: 40px;">Date of Expiry: _____</p> <p style="padding-left: 40px;">Batch/Lot No.: _____</p>			
<p>C. Additional Requirements to be attached to the Technical Specifications form arranged, numbered, and tabbed as enumerated below:</p> <p>1. Valid and current Certificate Product Registration (CPR) , Certificate of Medical Device Registration (CMDR), or Certificate of Medical Device Notification (CMDN) issued by the Philippine Food and Drug Administration (PFDA);</p>			

The CPR must be valid for the entire period of the award. If the CPR/CMDN/CMDR is about to expire, the supplier must have submitted a copy of an application of renewal to the FDA at least 3 months before the expiry date (a copy of the expiring CPR, which is stamped with an "Extension of Validity" shall be submitted as proof); [AO 2019-0041]

2. Valid and current License to Operate (LTO) for Medical Device Exporter / Trader /Importer / Distributor / Wholesaler issued by Philippine Food and Drugs Administration (PFDA)

3. Product Insert/Product Information or downloaded from the internet and other manufacturer's unamended sales literature, unconditional statements of specification and compliance issued by the manufacturer, samples, independent test data, etc., as appropriate for a cross-referencing statement of compliance to the technical specification in accordance to what is indicated in Technical Specifications;

4. The bidder shall submit any of the following whichever is applicable:

a. If the bidder is a manufacturer, certificate that the bidder manufactures the products/item or

b. If the bidder is an Exclusive/Authorized Distributor or Dealer of the products/items, a Certificate or Contract from the manufacturer must be provided as proof that the bidder is an Exclusive/Authorized Distributor or Dealer of the products/items or

c. If the bidder is an agent of the exclusive distributor or dealer, the following must be provided:

i. Certificate or Distributor/Dealership Agreement by the Manufacturer with the distributor or dealer; and

ii. Certificate/Contract between the distributor/dealer and the bidder.