



Republic of the Philippines  
Department of Health

## METRO MANILA CENTER FOR HEALTH DEVELOPMENT

### SUPPLEMENTAL/ BID BULLETIN NO. 1

IB 2023 – 024

#### PROCUREMENT OF ETONOGESTREL SUBDERMAL IMPLANT (SHORT OF AWARD)

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

Revision and clarification to provisions/specifications in the Bidding Documents:	
FROM	TO
<b>Bid Data Sheet</b> <b>ITB Clause 5.3</b> For this purpose, contracts similar to the Project shall be: a. Medical Supplies, Medical Instruments b. Completed within three (3) years prior to the deadline for the submission and receipt of bids.	<b>Bid Data Sheet</b> <b>ITB Clause 5.3</b> For this purpose, contracts similar to the Project shall be: a. Medical Supplies/ Drugs and Medicine b. Completed within three (3) years prior to the deadline for the submission and receipt of bids.

Bidders are advised to use the following attached forms and submit together with all required documents for the submission of bids on 28<sup>th</sup> day of November 2022, 9:30 AM:

This Supplemental/Bid Bulletin No. 1 shall form an integral part of the Bidding Documents. All other provisions indicated in the bidding documents which are not affected by this Supplemental/Bid Bulletin No. 1 shall remain in effect.

For guidance and information of all concerned.

Issued this 18<sup>th</sup> day of November 2022 in MMCHD

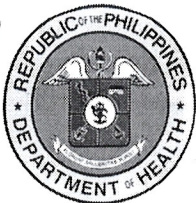
Approved by:

**ALELI ANNIE GRACE P. SUDIACAL, MD, MPH**  
Director III / BAC Chairperson



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

Republic of the Philippines Department of Health Metro Manila Center for Health Development  <b>TECHNICAL SPECIFICATIONS</b>			
Item No. 1	<b>PROCUREMENT OF ETONOGESTREL SUBDERMAL IMPLANT (SHORT OF AWARD)</b>	Qty./Unit	17,114 ROD
Name of Manufacturer:		Country of Origin	
Brand:		Model: (if applicable)	
<b>ABC: P 25,671,000.00</b>			
PURCHASER'S SPECIFICATION		STATEMENT OF COMPLIANCE	
<p><b><u>SPECIFICATIONS:</u></b></p> <p>&gt; Form and Strength: 68 mg subdermal implant</p> <p>&gt; <b>Primary Packaging-</b> each box contains 1 unit of singled-rod contraceptive implant preloaded in a sterile disposable applicator, 1 patient card, 1 physician card and 1 product information insert</p> <p>&gt; <b>Secondary Packaging</b> - standard packaging of the manufacturer as approved by PFDA including product insert or encryption/imprint inside the box</p> <p><b>Shelf Life</b></p> <ol style="list-style-type: none"> <li>1. Must be fresh commercial stock with a total shelf life of Sixty (60) months upon manufacture but not less than forty-two (42) months from the date of delivery.</li> </ol> <p><b>Labelling Instructions:</b></p> <p>&gt;Standard labeling instruction as approved by PFDA pursuant to Administrative Order No. 2016-0008</p> <p>In addition to the labelling requirements of FDA</p> <ol style="list-style-type: none"> <li>a. Each box the following should be imprinted or stickered with non-removable or permanent sticker or label that is binding, and with residue and tearing, If removed;</li> </ol> <p>Philippine Government Property – Department of Health NOT FOR SALE</p>			



## METRO MANILA CENTER FOR HEALTH DEVELOPMENT

Date of Manufacture:

Date of Expiry:

Batch/Lot No.:

**Delivery Schedule:**

>60-90 calendar days (CD)

**Delivery Site:**

>DOH-MMCHD Pasig Warehouse

**Additional Technical Documents:**

- a. Valid PFDA Certificate of Product Registration (CPR) or Valid Extension
- b. PFDA License to Operate (LTO) for Drugs Distributors and Traders
- c. Product insert/ Product Information
- d. Certificate from the manufacturer/main distributor that Bidder is an authorized dealer/exclusive distributor of the product

**FDA Test Analysis:**

>The minimum number of the sample units required for each test analysis of delivered medicines shall be based on the PFDA Circular 2014-014

**ADDITIONAL REQUIREMENT TO BE SUBMITTED BY THE SINGLE/LOWEST CALCULATED BIDDER (SCB/LCB) AS PART OF POST QUALIFICATION:**

1. One (1) original sample of 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 award of contract. Prototype of the labelling instruction must be part of the sample submitted however, the technical specifications of the labelling instruction of the product must be complied upon delivery.