

# **PHILIPPINE BIDDING DOCUMENTS**

## **SUPPLY AND DELIVERY OF LABORATORY SUPPLIES AND REAGENTS – FAILED ITEMS FOR CY 2025 – (REBID)**

Government of the Republic of the Philippines

**PB NO. VMC-2025-044**

**April 01, 2025  
10:00 am**

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# Preface

These Philippine Bidding Documents (PBDs) for the procurement of Goods through Competitive Bidding have been prepared by the Government of the Philippines for use by any branch, constitutional commission or office, agency, department, bureau, office, or instrumentality of the Government of the Philippines, National Government Agencies, including Government-Owned and/or Controlled Corporations, Government Financing Institutions, State Universities and Colleges, and Local Government Unit. The procedures and practices presented in this document have been developed through broad experience, and are for mandatory use in projects that are financed in whole or in part by the Government of the Philippines or any foreign government/foreign or international financing institution in accordance with the provisions of the 2016 revised Implementing Rules and Regulations of Republic Act No. 9184.

The Bidding Documents shall clearly and adequately define, among others: (i) the objectives, scope, and expected outputs and/or results of the proposed contract or Framework Agreement, as the case may be; (ii) the eligibility requirements of Bidders; (iii) the expected contract or Framework Agreement duration, the estimated quantity in the case of procurement of goods, delivery schedule and/or time frame; and (iv) the obligations, duties, and/or functions of the winning bidder.

Care should be taken to check the relevance of the provisions of the PBDs against the requirements of the specific Goods to be procured. If duplication of a subject is inevitable in other sections of the document prepared by the Procuring Entity, care must be exercised to avoid contradictions between clauses dealing with the same matter.

Moreover, each section is prepared with notes intended only as information for the Procuring Entity or the person drafting the Bidding Documents. They shall not be included in the final documents. The following general directions should be observed when using the documents:

- a. All the documents listed in the Table of Contents are normally required for the procurement of Goods. However, they should be adapted as necessary to the circumstances of the particular Procurement Project.
- b. Specific details, such as the “*name of the Procuring Entity*” and “*address for bid submission*,” should be furnished in the Instructions to Bidders, Bid Data Sheet, and Special Conditions of Contract. The final documents should contain neither blank spaces nor options.
- c. This Preface and the footnotes or notes in italics included in the Invitation to Bid, Bid Data Sheet, General Conditions of Contract, Special Conditions of Contract, Schedule of Requirements, and Specifications are not part of the text of the final document, although they contain instructions that the Procuring Entity should strictly follow.

- d. The cover should be modified as required to identify the Bidding Documents as to the Procurement Project, Project Identification Number, and Procuring Entity, in addition to the date of issue.
- e. Modifications for specific Procurement Project details should be provided in the Special Conditions of Contract as amendments to the Conditions of Contract. For easy completion, whenever reference has to be made to specific clauses in the Bid Data Sheet or Special Conditions of Contract, these terms shall be printed in bold typeface on Sections I (Instructions to Bidders) and III (General Conditions of Contract), respectively.
- f. For guidelines on the use of Bidding Forms and the procurement of Foreign-Assisted Projects, these will be covered by a separate issuance of the Government Procurement Policy Board.

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# ***Glossary of Acronyms, Terms, and Abbreviations***

**ABC** –Approved Budget for the Contract.

**BAC** – Bids and Awards Committee.

**Bid** – A signed offer or proposal to undertake a contract submitted by a bidder in response to and in consonance with the requirements of the bidding documents. Also referred to as *Proposal* and *Tender*. (2016 revised IRR, Section 5[c])

**Bidder** – Refers to a contractor, manufacturer, supplier, distributor and/or consultant who submits a bid in response to the requirements of the Bidding Documents. (2016 revised IRR, Section 5[d])

**Bidding Documents** – The documents issued by the Procuring Entity as the bases for bids, furnishing all information necessary for a prospective bidder to prepare a bid for the Goods, Infrastructure Projects, and/or Consulting Services required by the Procuring Entity. (2016 revised IRR, Section 5[e])

**BIR** – Bureau of Internal Revenue.

**BSP** – Bangko Sentral ng Pilipinas.

**Consulting Services** – Refer to services for Infrastructure Projects and other types of projects or activities of the GOP requiring adequate external technical and professional expertise that are beyond the capability and/or capacity of the GOP to undertake such as, but not limited to: (i) advisory and review services; (ii) pre-investment or feasibility studies; (iii) design; (iv) construction supervision; (v) management and related services; and (vi) other technical services or special studies. (2016 revised IRR, Section 5[i])

**CDA** - Cooperative Development Authority.

**Contract** – Refers to the agreement entered into between the Procuring Entity and the Supplier or Manufacturer or Distributor or Service Provider for procurement of Goods and Services; Contractor for Procurement of Infrastructure Projects; or Consultant or Consulting Firm for Procurement of Consulting Services; as the case may be, as recorded in the Contract Form signed by the parties, including all attachments and appendices thereto and all documents incorporated by reference therein.

**CIF** – Cost Insurance and Freight.

**CIP** – Carriage and Insurance Paid.

**CPI** – Consumer Price Index.

**DDP** – Refers to the quoted price of the Goods, which means “delivered duty paid.”

**DTI** – Department of Trade and Industry.

**EXW** – Ex works.

**FCA** – “Free Carrier” shipping point.

**FOB** – “Free on Board” shipping point.

**Foreign-funded Procurement or Foreign-Assisted Project**–Refers to procurement whose funding source is from a foreign government, foreign or international financing institution as specified in the Treaty or International or Executive Agreement. (2016 revised IRR, Section 5[b]).

**Framework Agreement** – Refers to a written agreement between a procuring entity and a supplier or service provider that identifies the terms and conditions, under which specific purchases, otherwise known as “Call-Offs,” are made for the duration of the agreement. It is in the nature of an option contract between the procuring entity and the bidder(s) granting the procuring entity the option to either place an order for any of the goods or services identified in the Framework Agreement List or not buy at all, within a minimum period of one (1) year to a maximum period of three (3) years. (GPPB Resolution No. 27-2019)

**GFI** – Government Financial Institution.

**GOCC** –Government-owned and/or –controlled corporation.

**Goods** – Refer to all items, supplies, materials and general support services, except Consulting Services and Infrastructure Projects, which may be needed in the transaction of public businesses or in the pursuit of any government undertaking, project or activity, whether in the nature of equipment, furniture, stationery, materials for construction, or personal property of any kind, including non-personal or contractual services such as the repair and maintenance of equipment and furniture, as well as trucking, hauling, janitorial, security, and related or analogous services, as well as procurement of materials and supplies provided by the Procuring Entity for such services. The term “related” or “analogous services” shall include, but is not limited to, lease or purchase of office space, media advertisements, health maintenance services, and other services essential to the operation of the Procuring Entity. (2016 revised IRR, Section 5[r])

**GOP** – Government of the Philippines.

**GPPB** –Government Procurement Policy Board.

**INCOTERMS** – International Commercial Terms.

**Infrastructure Projects** – Include the construction, improvement, rehabilitation, demolition, repair, restoration or maintenance of roads and bridges, railways, airports, seaports, communication facilities, civil works components of information technology projects, irrigation, flood control and drainage, water supply, sanitation, sewerage and solid waste management systems, shore protection, energy/power and electrification facilities, national

buildings, school buildings, hospital buildings, and other related construction projects of the government. Also referred to as *civil works or works*. (2016 revised IRR, Section 5[u])

**LGUs** – Local Government Units.

**NFCC** – Net Financial Contracting Capacity.

**NGA** – National Government Agency.

**PhilGEPS** - Philippine Government Electronic Procurement System.

**Procurement Project** – refers to a specific or identified procurement covering goods, infrastructure project or consulting services. A Procurement Project shall be described, detailed, and scheduled in the Project Procurement Management Plan prepared by the agency which shall be consolidated in the procuring entity's Annual Procurement Plan. (GPPB Circular No. 06-2019 dated 17 July 2019)

**PSA** – Philippine Statistics Authority.

**SEC** – Securities and Exchange Commission.

**SLCC** – Single Largest Completed Contract.

**Supplier** – refers to a citizen, or any corporate body or commercial company duly organized and registered under the laws where it is established, habitually established in business and engaged in the manufacture or sale of the merchandise or performance of the general services covered by his bid. (Item 3.8 of GPPB Resolution No. 13-2019, dated 23 May 2019). Supplier as used in these Bidding Documents may likewise refer to a distributor, manufacturer, contractor, or consultant.

**UN** – United Nations.

## ***Section I. Invitation to Bid***

### **Notes on the Invitation to Bid**

The Invitation to Bid (IB) provides information that enables potential Bidders to decide whether to participate in the procurement at hand. The IB shall be posted in accordance with Section 21.2 of the 2016 revised IRR of RA No. 9184.

Apart from the essential items listed in the Bidding Documents, the IB should also indicate the following:

- a. The date of availability of the Bidding Documents, which shall be from the time the IB is first advertised/posted until the deadline for the submission and receipt of bids;
- b. The place where the Bidding Documents may be acquired or the website where it may be downloaded;
- c. The deadline for the submission and receipt of bids; and
- d. Any important bid evaluation criteria (*e.g.*, the application of a margin of preference in bid evaluation).

The IB should be incorporated in the Bidding Documents. The information contained in the IB must conform to the Bidding Documents and in particular to the relevant information in the Bid Data Sheet.



Republic of the Philippines  
Department of Health  
Metro Manila Center for Health Development  
**VALENZUELA MEDICAL CENTER**



## INVITATION TO BID

### SUPPLY AND DELIVERY OF LABORATORY SUPPLIES AND REAGENTS – FAILED ITEMS FOR CY 2025 – (REBID)

**Public Bidding No. VMC 2025-044**

1. The *Valenzuela Medical Center (VMC)*, through the *General Appropriations Act/Income CY 2025* intends to apply the sum of **Philippine Currency: TEN MILLION THREE HUNDRED FORTY-SEVEN THOUSAND FIVE HUNDRED THREE PESOS AND 04/100 ONLY (P 10,347,503.04)** being the ABC to payments under the contract for Procurement of **SUPPLY AND DELIVERY OF LABORATORY SUPPLIES AND REAGENTS – FAILED ITEMS FOR CY 2025 – (REBID)**. Bids received in excess of the ABC shall be automatically rejected at bid opening.
2. The *Valenzuela Medical Center (VMC)* now invites bids for the above Procurement Project. Delivery of the Goods is required within the required period under Section IV Schedule of Requirements. Bidders should have completed at least three (3) years from the date of submission and receipt of bids, a contract similar to the Project. The description of an eligible bidder is contained in the Bidding Documents, particularly, in Section II (Instructions to Bidders).
3. Bidding will be conducted through open competitive bidding procedures using a non-discretionary “*pass/fail*” criterion as specified in the 2016 revised Implementing Rules and Regulations (IRR) of Republic Act (RA) No. 9184.
  - a. Bidding is restricted to Filipino citizens/sole proprietorships, partnerships, or organizations with at least sixty percent (60%) interest or outstanding capital stock belonging to citizens of the Philippines, and to citizens or organizations of a country the laws or regulations of which grant similar rights or privileges to Filipino citizens, pursuant to RA No. 5183.
4. Prospective Bidders may obtain further information *starting March 07, 2025* and inspect the Bidding Documents at the address given below during *9:00am – 11:00am and 2:00pm – 4:00pm*.
5. A complete set of Bidding Documents may be acquired by interested Bidders on *November 04, 2024* from the given address below and upon payment of a non-refundable fee in the amount of:

<b>ABC to be Bid</b>	<b>Maximum Cost of Bidding Documents (in Philippine Peso)</b>
500,000 and below	500.00
More than 500,000 up to 1 Million	1,000.00
More than 1 Million up to 5 Million	5,000.00
More than 5 Million up to 10 Million	10,000.00

The Procuring Entity shall allow the bidder to present its proof of payment for the fees *either in person, by facsimile, or through electronic means*.

*[NOTE: For lot procurement, the maximum fee for the Bidding Documents for each lot shall be based on its ABC, in accordance with the Guidelines issued by the GPPB; provided that the total fees for the Bidding Documents of all lots shall not exceed the maximum fee prescribed in the Guidelines for the sum of the ABC of all lots.]*

6. The *Valenzuela Medical Center (VMC)* will hold a Pre-Bid Conference<sup>1</sup> on *March 19, 2025, 10:00 am* at *Valenzuela Medical Center, Annex 1 Building, Padrigal St., Karuhatan, Valenzuela*, which shall be open to prospective bidders.
7. Bids must be duly received by the BAC Secretariat through manual submission at the office address indicated below, online or electronic submission, or both on or before *April 01, 2025, 10:00 AM*. Late bids shall not be accepted.
8. All Bids must be accompanied by a bid security in any of the acceptable forms and in the amount stated in **ITB** Clause 14.
9. Bid opening shall be on *April 01, 2025, 10:00 AM*. at the given address below. Bids will be opened in the presence of the bidders' representatives who choose to attend the activity.
10. Prospective bidders shall provide use of a back-up data or cloud storage for large files uploaded for online bid submissions.
11. The *Valenzuela Medical Center (VMC)* reserves the right to reject any and all bids, declare a failure of bidding, or not award the contract at any time prior to contract award in accordance with Sections 35.6 and 41 of the 2016 revised IRR of RA No. 9184, without thereby incurring any liability to the affected bidder or bidders.
12. For further information, please refer to:

BAC Secretariats' Office  
Valenzuela Medical Center  
Annex Building, 2<sup>nd</sup> Floor,  
Padrigal St., Valenzuela City  
Telefax No. 294-4625  
Email: [ymc\\_bac@yahoo.com](mailto:ymc_bac@yahoo.com)

**SGD. SHIRLENE V. VIANZON**  
Chairperson, Bids and Awards Committee

## ***Section II. Instructions to Bidders***

### **Notes on the Instructions to Bidders**

This Section on the Instruction to Bidders (ITB) provides the information necessary for bidders to prepare responsive bids, in accordance with the requirements of the Procuring Entity. It also provides information on bid submission, eligibility check, opening and evaluation of bids, post-qualification, and on the award of contract.

## **1. Scope of Bid**

The *Valenzuela Medical Center (VMC)* wishes to receive Bids for the **SUPPLY AND DELIVERY OF LABORATORY SUPPLIES AND REAGENTS – FAILED ITEMS FOR CY 2025 – (REBID)**, with identification number *under Public Bidding VMC 2025-044*.

*[Note: The Project Identification Number is assigned by the Procuring Entity based on its own coding scheme and is not the same as the PhilGEPS reference number, which is generated after the posting of the bid opportunity on the PhilGEPS website.]*

The Procurement Project (referred to herein as “Project”) is composed of [indicate number of lots or items], the details of which are described in Section VII (Technical Specifications).

## **2. Funding Information**

**2.1.** The GOP through the source of funding as indicated below for **CY 2025** in the amount of **TEN MILLION THREE HUNDRED FORTY-SEVEN THOUSAND FIVE HUNDRED THREE PESOS AND 04/100 ONLY (P 10,347,503.04)**

2.2. The source of funding is:

- a. NGA, the General Appropriations Act or Special Appropriations.

## **3. Bidding Requirements**

The Bidding for the Project shall be governed by all the provisions of RA No. 9184 and its 2016 revised IRR, including its Generic Procurement Manuals and associated policies, rules and regulations as the primary source thereof, while the herein clauses shall serve as the secondary source thereof.

Any amendments made to the IRR and other GPPB issuances shall be applicable only to the ongoing posting, advertisement, or **IB** by the BAC through the issuance of a supplemental or bid bulletin.

The Bidder, by the act of submitting its Bid, shall be deemed to have verified and accepted the general requirements of this Project, including other factors that may affect the cost, duration and execution or implementation of the contract, project, or work and examine all instructions, forms, terms, and project requirements in the Bidding Documents.

## **4. Corrupt, Fraudulent, Collusive, and Coercive Practices**

The Procuring Entity, as well as the Bidders and Suppliers, shall observe the highest standard of ethics during the procurement and execution of the contract. They or through an agent shall not engage in corrupt, fraudulent, collusive, coercive, and

obstructive practices defined under Annex “I” of the 2016 revised IRR of RA No. 9184 or other integrity violations in competing for the Project.

## **5. Eligible Bidders**

- 5.1. Only Bids of Bidders found to be legally, technically, and financially capable will be evaluated.
- 5.2. Foreign ownership limited to those allowed under the rules may participate in this Project.
- 5.3. Pursuant to Section 23.4.1.3 of the 2016 revised IRR of RA No.9184, the Bidder shall have an SLCC that is at least one (1) contract similar to the Project the value of which, adjusted to current prices using the PSA’s CPI, must be at least equivalent to:
  - a. For the procurement of Expendable Supplies: The Bidder must have completed a single contract that is similar to this Project, equivalent to at least twenty-five percent (25%) of the ABC.
- 5.4. The Bidders shall comply with the eligibility criteria under Section 23.4.1 of the 2016 IRR of RA No. 9184.

## **6. Origin of Goods**

There is no restriction on the origin of goods other than those prohibited by a decision of the UN Security Council taken under Chapter VII of the Charter of the UN, subject to Domestic Preference requirements under **ITB** Clause 18.

## **7. Subcontracts**

- 7.1. The Bidder may subcontract portions of the Project to the extent allowed by the Procuring Entity as stated herein, but in no case more than twenty percent (20%) of the Project.

The Procuring Entity has prescribed that:

*[Select one, delete other/s]*

- a. Subcontracting is not allowed.

## **8. Pre-Bid Conference**

The Procuring Entity will hold a pre-bid conference for this Project on **March 19, 2025, 10:00 am at physical address, Valenzuela Medical Center, Annex 1 Building, Padrigal St. Karuhatan, Valenzuela City.**

## **9. Clarification and Amendment of Bidding Documents**

Prospective bidders may request for clarification on and/or interpretation of any part of the Bidding Documents. Such requests must be in writing and received by the Procuring Entity, either at its given address or through electronic mail indicated in the **IB**, at least ten (10) calendar days before the deadline set for the submission and receipt of Bids.

## **10. Documents comprising the Bid: Eligibility and Technical Components**

- 10.1. The first envelope shall contain the eligibility and technical documents of the Bid as specified in **Section VIII (Checklist of Technical and Financial Documents)**.
- 10.2. The Bidder's SLCC as indicated in **ITB Clause 5.3** should have been completed within *three (3) years period as provided in paragraph 2 of the **IB*** prior to the deadline for the submission and receipt of bids.
- 10.3. If the eligibility requirements or statements, the bids, and all other documents for submission to the BAC are in foreign language other than English, it must be accompanied by a translation in English, which shall be authenticated by the appropriate Philippine foreign service establishment, post, or the equivalent office having jurisdiction over the foreign bidder's affairs in the Philippines. Similar to the required authentication above, for Contracting Parties to the Apostille Convention, only the translated documents shall be authenticated through an apostille pursuant to GPPB Resolution No. 13-2019 dated 23 May 2019. The English translation shall govern, for purposes of interpretation of the bid.

## **11. Documents comprising the Bid: Financial Component**

- 11.1. The second bid envelope shall contain the financial documents for the Bid as specified in **Section VIII (Checklist of Technical and Financial Documents)**.
- 11.2. If the Bidder claims preference as a Domestic Bidder or Domestic Entity, a certification issued by DTI shall be provided by the Bidder in accordance with Section 43.1.3 of the 2016 revised IRR of RA No. 9184.
- 11.3. Any bid exceeding the ABC indicated in paragraph 1 of the **IB** shall not be accepted.
- 11.4. For Foreign-funded Procurement, a ceiling may be applied to bid prices provided the conditions are met under Section 31.2 of the 2016 revised IRR of RA No. 9184.
- 11.5. *[Include if Framework Agreement will be used:]* Financial proposals for single or multi-year Framework Agreement shall be submitted before the deadline of submission of bids as prescribed in the **IB**. For multi-year Framework Agreement, evaluation of the financial proposal during this stage is for purposes of determining eligibility and whether or not such financial proposal is within the ABC.

## **12. Bid Prices**

- 12.1. Prices indicated on the Price Schedule shall be entered separately in the following manner:
- a. For Goods offered from within the Procuring Entity's country:
    - i. The price of the Goods quoted EXW (ex-works, ex-factory, ex-warehouse, ex-showroom, or off-the-shelf, as applicable);
    - ii. The cost of all customs duties and sales and other taxes already paid or payable;
    - iii. The cost of transportation, insurance, and other costs incidental to delivery of the Goods to their final destination; and
    - iv. The price of other (incidental) services, if any, listed in e.
  - b. For Goods offered from abroad:
    - i. Unless otherwise stated in the **BDS**, the price of the Goods shall be quoted delivered duty paid (DDP) with the place of destination in the Philippines as specified in the **BDS**. In quoting the price, the Bidder shall be free to use transportation through carriers registered in any eligible country. Similarly, the Bidder may obtain insurance services from any eligible source country.
    - ii. The price of other (incidental) services, if any, as listed in **Section VII (Technical Specifications)**.

## **13. Bid and Payment Currencies**

- 13.1. For Goods that the Bidder will supply from outside the Philippines, the bid prices may be quoted in the local currency or tradeable currency accepted by the BSP at the discretion of the Bidder. However, for purposes of bid evaluation, Bids denominated in foreign currencies, shall be converted to Philippine currency based on the exchange rate as published in the BSP reference rate bulletin on the day of the bid opening.
- 13.2. Payment of the contract price shall be made in:

*[Select one, delete the other/s]*

- a. Philippine Pesos.

## **14. Bid Security**

- 14.1. The Bidder shall submit a Bid Securing Declaration<sup>2</sup> or any form of Bid Security in the amount indicated in the **BDS**, which shall be not less than the percentage of the ABC in accordance with the schedule in the **BDS**.
- 14.2. The Bid and bid security shall be valid until *[indicate date]*. Any Bid not accompanied by an acceptable bid security shall be rejected by the Procuring Entity as non-responsive.
- 14.3. *[Include if Framework Agreement will be used:]* In the case of Framework Agreement, other than the grounds for forfeiture under the 2016 revised IRR, the bid security may also be forfeited if the successful bidder fails to sign the Framework Agreement, or fails to furnish the performance security or performance securing declaration. Without prejudice on its forfeiture, bid securities shall be returned only after the posting of performance security or performance securing declaration, as the case may be, by the winning Bidder or compliant Bidders and the signing of the Framework Agreement.

## 15. Sealing and Marking of Bids

Each Bidder shall submit one copy of the first and second components of its Bid.

The Procuring Entity may request additional hard copies and/or electronic copies of the Bid. However, failure of the Bidders to comply with the said request shall not be a ground for disqualification.

If the Procuring Entity allows the submission of bids through online submission or any other electronic means, the Bidder shall submit an electronic copy of its Bid, which must be digitally signed. An electronic copy that cannot be opened or is corrupted shall be considered non-responsive and, thus, automatically disqualified.

## 16. Deadline for Submission of Bids

- 16.1. The Bidders shall submit on the specified date and time (**April 01, 2025 @ 10:00 am**) and either at its physical address, **Valenzuela Medical Center, Annex 1 Building, Padrigal St. Karuhatan, Valenzuela City** as indicated in paragraph 7 of the **IB**.

## 17. Opening and Preliminary Examination of Bids

- 17.1. The BAC shall open the Bids in public at the time, on the date, and at the place specified in paragraph 9 of the **IB**. The Bidders' representatives who are present shall sign a register evidencing their attendance. In case videoconferencing, webcasting or other similar technologies will be used, attendance of participants shall likewise be recorded by the BAC Secretariat.

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<sup>2</sup> In the case of Framework Agreement, the undertaking shall refer to entering into contract with the Procuring Entity and furnishing of the performance security or the performance securing declaration within ten (10) calendar days from receipt of Notice to Execute Framework Agreement.

In case the Bids cannot be opened as scheduled due to justifiable reasons, the rescheduling requirements under Section 29 of the 2016 revised IRR of RA No. 9184 shall prevail.

- 17.2. The preliminary examination of bids shall be governed by Section 30 of the 2016 revised IRR of RA No. 9184.

## **18. Domestic Preference**

- 18.1. The Procuring Entity will grant a margin of preference for the purpose of comparison of Bids in accordance with Section 43.1.2 of the 2016 revised IRR of RA No. 9184.

## **19. Detailed Evaluation and Comparison of Bids**

- 19.1. The Procuring BAC shall immediately conduct a detailed evaluation of all Bids rated “*passed*,” using non-discretionary pass/fail criteria. The BAC shall consider the conditions in the evaluation of Bids under Section 32.2 of the 2016 revised IRR of RA No. 9184.

*[Include the following options if Framework Agreement will be used:]*

- a. In the case of single-year Framework Agreement, the Lowest Calculated Bid shall be determined outright after the detailed evaluation;
  - b. For multi-year Framework Agreement, the determination of the eligibility and the compliance of bidders with the technical and financial aspects of the projects shall be initially made by the BAC, in accordance with Item 7.4.2 of the Guidelines on the Use of Framework Agreement.
- 19.2. If the Project allows partial bids, bidders may submit a proposal on any of the lots or items, and evaluation will be undertaken on a per lot or item basis, as the case maybe. In this case, the Bid Security as required by **ITB** Clause 15 shall be submitted for each lot or item separately.
  - 19.3. The descriptions of the lots or items shall be indicated in **Section VII (Technical Specifications)**, although the ABCs of these lots or items are indicated in the **BDS** for purposes of the NFCC computation pursuant to Section 23.4.2.6 of the 2016 revised IRR of RA No. 9184. The NFCC must be sufficient for the total of the ABCs for all the lots or items participated in by the prospective Bidder.
  - 19.4. The Project shall be awarded as follows:

*[Select one, delete the other/s]*

Option 1 – One Project having several items that shall be awarded as one contract.

Option 2 – One Project having several items grouped into several lots, which shall be awarded as separate contracts per lot.

Option 3 - One Project having several items, which shall be awarded as separate contracts per item.

*[Delete Options 2 and 3 if Framework Agreement will be used.]*

- 19.5. Except for bidders submitting a committed Line of Credit from a Universal or Commercial Bank in lieu of its NFCC computation, all Bids must include the NFCC computation pursuant to Section 23.4.1.4 of the 2016 revised IRR of RA No. 9184, which must be sufficient for the total of the ABCs for all the lots or items participated in by the prospective Bidder. For bidders submitting the committed Line of Credit, it must be at least equal to ten percent (10%) of the ABCs for all the lots or items participated in by the prospective Bidder.

## **20. Post-Qualification**

- 20.1. *[Include if Framework Agreement will be used:]* For multi-year Framework Agreement, all bidders initially determined to be eligible and financially compliant shall be subject to initial post-qualification. The BAC shall then recommend the execution of a Framework Agreement among all eligible, technically and financially compliant bidders and the Procuring Entity and shall be issued by HoPE a Notice to Execute Framework Agreement. The determination of the Lowest Calculated Bid (LCB) shall not be performed by the BAC until a Mini-Competition is conducted among the bidders who executed a Framework Agreement. When a Call for Mini-Competition is made, the BAC shall allow the bidders to submit their best financial proposals on such pre-scheduled date, time and place to determine the bidder with the LCB.
- 20.2. Within a non-extendible period of five (5) calendar days from receipt by the Bidder of the notice from the BAC that it submitted the Lowest Calculated Bid, *{[Include if Framework Agreement will be used:]}* or in the case of multi-year Framework Agreement, that it is one of the eligible bidders who have submitted bids that are found to be technically and financially compliant,}the Bidder shall submit its latest income and business tax returns filed and paid through the BIR Electronic Filing and Payment System (eFPS) and other appropriate licenses and permits required by law and stated in the **BDS**. *{[Include if Framework Agreement will be used:]}* For every mini-competition in Framework Agreement, the LCB shall likewise submit the required documents for final Post Qualification.}

## **21. Signing of the Contract**

- 21.1. The documents required in Section 37.2 of the 2016 revised IRR of RA No. 9184 shall form part of the Contract. Additional Contract documents are indicated in the **BDS**.

*[Include the following clauses if Framework Agreement will be used:]*

- 21.2. At the same time as the Procuring Entity notifies the successful Bidder that its bid has been accepted, the Procuring Entity shall send the Framework Agreement Form to the Bidder, which contract has been provided in the Bidding Documents, incorporating therein all agreements between the parties.

- 21.3. Within ten (10) calendar days from receipt of the Notice to Execute Framework Agreement with the Procuring Entity, the successful Bidder or its duly authorized representative shall formally enter into a Framework Agreement with the procuring entity for an amount of One Peso to be paid to the procuring entity as a consideration for the option granted by the procuring entity to procure the items in the Framework Agreement List when the need arises.
- 21.4. The Procuring Entity shall enter into a Framework Agreement with the successful Bidder within the same ten (10) calendar day period provided that all the documentary requirements are complied with.
- 21.5. The following documents shall form part of the Framework Agreement:
  - a. Framework Agreement Form;
  - b. Bidding Documents;
  - c. Call-offs;
  - d. Winning bidder's bid, including the Technical and Financial Proposals, and all other documents/statements submitted (*e.g.*, bidder's response to request for clarifications on the bid), including corrections to the bid, if any, resulting from the Procuring Entity's bid evaluation;
  - e. Performance Security or Performance Securing Declaration, as the case may be;
  - f. Notice to Execute Framework Agreement; and
  - g. Other contract documents that may be required by existing laws and/or specified in the **BDS**.

## ***Section III. Bid Data Sheet***

### **Notes on the Bid Data Sheet**

The Bid Data Sheet (BDS) consists of provisions that supplement, amend, or specify in detail, information, or requirements included in the ITB found in Section II, which are specific to each procurement.

This Section is intended to assist the Procuring Entity in providing the specific information in relation to corresponding clauses in the ITB and has to be prepared for each specific procurement.

The Procuring Entity should specify in the BDS information and requirements specific to the circumstances of the Procuring Entity, the processing of the procurement, and the bid evaluation criteria that will apply to the Bids. In preparing the BDS, the following aspects should be checked:

- a. Information that specifies and complements provisions of the ITB must be incorporated.
- b. Amendments and/or supplements, if any, to provisions of the ITB as necessitated by the circumstances of the specific procurement, must also be incorporated.

# Bid Data Sheet

ITB Clause	
5.3	For this purpose, contracts similar to the Project shall be:  a. Similar to Laboratory Supplies and Reagents b. At least three (3) years
7.1	[Specify the portions of Goods to be subcontracted, which shall not be a significant or material component of the Project as determined by the Procuring Entity.] N/A
12	The price of the Goods shall be quoted DDP[state place of destination] or the applicable International Commercial Terms (INCOTERMS) for this Project.
14.1	The bid security shall be in the form of a Bid Securing Declaration, or any of the following forms and amounts:  a. The amount of not less than <u>206,950.06</u> [Indicate the amount equivalent to two percent (2%) of ABC], if bid security is in cash, cashier's/manager's check, bank draft/guarantee or irrevocable letter of credit; or  b. The amount of not less than <u>517,375.15</u> [Indicate the amount equivalent to five percent (5%) of ABC] if bid security is in Surety Bond.
19.3	[In case the Project will be awarded by lot, list the grouping of lots by specifying the group title, items, and the quantity for every identified lot, and the corresponding ABC for each lot.] ABC  [In case the project will be awarded by item, list each item indicating its quantity and ABC.] N/A
20.2	[List here any licenses and permits relevant to the Project and the corresponding law requiring it.]  <i>Refer to Checklist</i>
21.2	[List here any additional contract documents relevant to the Project that may be required by existing laws and/or the Procuring Entity.)  <i>Refer to Checklist</i>

## ***Section IV. General Conditions of Contract***

### **Notes on the General Conditions of Contract**

The General Conditions of Contract (GCC) in this Section, read in conjunction with the Special Conditions of Contract in Section V and other documents listed therein, should be a complete document expressing all the rights and obligations of the parties.

Matters governing performance of the Supplier, payments under the contract, or matters affecting the risks, rights, and obligations of the parties under the contract are included in the GCC and Special Conditions of Contract.

Any complementary information, which may be needed, shall be introduced only through the Special Conditions of Contract.

## **1. Scope of Contract**

This Contract shall include all such items, although not specifically mentioned, that can be reasonably inferred as being required for its completion as if such items were expressly mentioned herein. All the provisions of RA No. 9184 and its 2016 revised IRR, including the Generic Procurement Manual, and associated issuances, constitute the primary source for the terms and conditions of the Contract, and thus, applicable in contract implementation. Herein clauses shall serve as the secondary source for the terms and conditions of the Contract.

This is without prejudice to Sections 74.1 and 74.2 of the 2016 revised IRR of RA No. 9184 allowing the GPPB to amend the IRR, which shall be applied to all procurement activities, the advertisement, posting, or invitation of which were issued after the effectivity of the said amendment.

Additional requirements for the completion of this Contract shall be provided in the **Special Conditions of Contract (SCC)**.

## **2. Advance Payment and Terms of Payment**

- 2.1. Advance payment of the contract amount is provided under Annex “D” of the revised 2016 IRR of RA No. 9184.
- 2.2. The Procuring Entity is allowed to determine the terms of payment on the partial or staggered delivery of the Goods procured, provided such partial payment shall correspond to the value of the goods delivered and accepted in accordance with prevailing accounting and auditing rules and regulations. The terms of payment are indicated in the **SCC**.

*[Include the following clauses if Framework Agreement will be used:]*

- 2.3. For a single-year Framework Agreement, prices charged by the Supplier for Goods delivered and/or services performed under a Call-Off shall not vary from the prices quoted by the Supplier in its bid.
- 2.4. For multi-year Framework Agreement, prices charged by the Supplier for Goods delivered and/or services performed under a Call-Off shall not vary from the prices quoted by the Supplier during conduct of Mini-Competition.

## **3. Performance Security**

Within ten (10) calendar days from receipt of the Notice of Award by the Bidder from the Procuring Entity but in no case later than prior to the signing of the Contract by both parties, the successful Bidder shall furnish the performance security in any of the forms prescribed in Section 39 of the 2016 revised IRR of RA No. 9184.*{[Include if Framework Agreement will be used:] In the case of Framework Agreement, the Bidder may opt to furnish the performance security or a Performance Securing Declaration as defined under the Guidelines on the Use of Framework Agreement.}*

## **4. Inspection and Tests**

The Procuring Entity or its representative shall have the right to inspect and/or to test the Goods to confirm their conformity to the Project {[Include if Framework Agreement will be used:] or Framework Agreement} specifications at no extra cost to the Procuring Entity in accordance with the Generic Procurement Manual. In addition to tests in the SCC, **Section IV (Technical Specifications)** shall specify what inspections and/or tests the Procuring Entity requires, and where they are to be conducted. The Procuring Entity shall notify the Supplier in writing, in a timely manner, of the identity of any representatives retained for these purposes.

All reasonable facilities and assistance for the inspection and testing of Goods, including access to drawings and production data, shall be provided by the Supplier to the authorized inspectors at no charge to the Procuring Entity.

## **5. Warranty**

- 5.1 In order to assure that manufacturing defects shall be corrected by the Supplier, a warranty shall be required from the Supplier as provided under Section 62.1 of the 2016 revised IRR of RA No. 9184.
- 5.2 The Procuring Entity shall promptly notify the Supplier in writing of any claims arising under this warranty. Upon receipt of such notice, the Supplier shall, repair or replace the defective Goods or parts thereof without cost to the Procuring Entity, pursuant to the Generic Procurement Manual.

## **6. Liability of the Supplier**

The Supplier's liability under this Contract shall be as provided by the laws of the Republic of the Philippines.

If the Supplier is a joint venture, all partners to the joint venture shall be jointly and severally liable to the Procuring Entity.

## ***Section V. Special Conditions of Contract***

### **Notes on the Special Conditions of Contract**

Similar to the BDS, the clauses in this Section are intended to assist the Procuring Entity in providing contract-specific information in relation to corresponding clauses in the GCC found in Section IV.

The Special Conditions of Contract (SCC) complement the GCC, specifying contractual requirements linked to the special circumstances of the Procuring Entity, the Procuring Entity's country, the sector, and the Goods purchased. In preparing this Section, the following aspects should be checked:

- a. Information that complements provisions of the GCC must be incorporated.
- b. Amendments and/or supplements to provisions of the GCC as necessitated by the circumstances of the specific purchase must also be incorporated.

However, no special condition which defeats or negates the general intent and purpose of the provisions of the GCC should be incorporated herein.

# Special Conditions of Contract

GCC Clause	
1	<p style="color: red;"><i>[List here any additional requirements for the completion of this Contract. The following requirements and the corresponding provisions may be deleted, amended, or retained depending on its applicability to this Contract:]</i></p> <p><b>Delivery and Documents –</b></p> <p>For purposes of the Contract, “EXW,” “FOB,” “FCA,” “CIF,” “CIP,” “DDP” and other trade terms used to describe the obligations of the parties shall have the meanings assigned to them by the current edition of INCOTERMS published by the International Chamber of Commerce, Paris. The Delivery terms of this Contract shall be as follows:</p> <p><i>[For Goods supplied from abroad, state:]</i> “The delivery terms applicable to the Contract are DDP delivered [indicate place of destination]. In accordance with INCOTERMS.”</p> <p><i>[For Goods supplied from within the Philippines, state:]</i> “The delivery terms applicable to this Contract are delivered [indicate place of destination]. Risk and title will pass from the Supplier to the Procuring Entity upon receipt and final acceptance of the Goods at their final destination.”</p> <p>Delivery of the Goods shall be made by the Supplier in accordance with the terms specified in Section VI (Schedule of Requirements).</p> <p>For purposes of this Clause the Procuring Entity’s Representative at the Project Site is [indicate name(s)].</p> <p><b>Incidental Services –</b></p> <p>The Supplier is required to provide all of the following services, including additional services, if any, specified in Section VI. Schedule of Requirements: <i>Select appropriate requirements and delete the rest.</i></p> <ul style="list-style-type: none"> <li>a. performance or supervision of on-site assembly and/or start-up of the supplied Goods;</li> <li>b. furnishing of tools required for assembly and/or maintenance of the supplied Goods;</li> <li>c. furnishing of a detailed operations and maintenance manual for each appropriate unit of the supplied Goods;</li> <li>d. performance or supervision or maintenance and/or repair of the supplied Goods, for a period of time agreed by the parties, provided that this service shall not relieve the Supplier of any warranty obligations under this Contract; and</li> </ul>

	<p>e. training of the Procuring Entity's personnel, at the Supplier's plant and/or on-site, in assembly, start-up, operation, maintenance, and/or repair of the supplied Goods.</p> <p>f. <i>[Specify additional incidental service requirements, as needed.]</i></p> <p>The Contract price for the Goods shall include the prices charged by the Supplier for incidental services and shall not exceed the prevailing rates charged to other parties by the Supplier for similar services.</p> <p><b>Spare Parts –</b></p> <p>The Supplier is required to provide all of the following materials, notifications, and information pertaining to spare parts manufactured or distributed by the Supplier:</p> <p><i>Select appropriate requirements and delete the rest.</i></p> <ul style="list-style-type: none"> <li>a. such spare parts as the Procuring Entity may elect to purchase from the Supplier, provided that this election shall not relieve the Supplier of any warranty obligations under this Contract; and</li> <li>b. in the event of termination of production of the spare parts: <ul style="list-style-type: none"> <li>i. advance notification to the Procuring Entity of the pending termination, in sufficient time to permit the Procuring Entity to procure needed requirements; and</li> <li>ii. following such termination, furnishing at no cost to the Procuring Entity, the blueprints, drawings, and specifications of the spare parts, if requested.</li> </ul> </li> </ul> <p>The spare parts and other components required are listed in <b>Section VI (Schedule of Requirements)</b> and the cost thereof are included in the contract price.</p> <p>The Supplier shall carry sufficient inventories to assure ex-stock supply of consumable spare parts or components for the Goods for a period of <i>[indicate here the time period specified. If not used indicate a time period of three times the warranty period]</i>.</p> <p>Spare parts or components shall be supplied as promptly as possible, but in any case, within <i>[insert appropriate time period]</i> months of placing the order.</p>
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	<p><b>Packaging –</b></p> <p>The Supplier shall provide such packaging of the Goods as is required to prevent their damage or deterioration during transit to their final destination, as indicated in this Contract. The packaging shall be sufficient to withstand, without limitation, rough handling during transit and exposure to extreme temperatures, salt and precipitation during transit, and open storage. Packaging case size and weights shall take into consideration, where appropriate, the remoteness of the Goods' final destination and the absence of heavy handling facilities at all points in transit.</p> <p>The packaging, marking, and documentation within and outside the packages shall comply strictly with such special requirements as shall be expressly provided for in the Contract, including additional requirements, if any, specified below, and in any subsequent instructions ordered by the Procuring Entity.</p> <p>The outer packaging must be clearly marked on at least four (4) sides as follows:</p> <p>Name of the Procuring Entity  Name of the Supplier  Contract Description  Final Destination  Gross weight  Any special lifting instructions  Any special handling instructions  Any relevant HAZCHEM classifications</p>
	<p>A packaging list identifying the contents and quantities of the package is to be placed on an accessible point of the outer packaging if practical. If not practical the packaging list is to be placed inside the outer packaging but outside the secondary packaging.</p> <p><b>Transportation –</b></p> <p>Where the Supplier is required under Contract to deliver the Goods CIF, CIP, or DDP, transport of the Goods to the port of destination or such other named place of destination in the Philippines, as shall be specified in this Contract, shall be arranged and paid for by the Supplier, and the cost thereof shall be included in the Contract Price.</p> <p>Where the Supplier is required under this Contract to transport the Goods to a specified place of destination within the Philippines, defined as the Project Site, transport to such place of destination in the Philippines, including insurance and storage, as shall be specified in this Contract, shall be arranged by the Supplier, and related costs shall be included in the contract price.</p>

	<p>Where the Supplier is required under Contract to deliver the Goods CIF, CIP or DDP, Goods are to be transported on carriers of Philippine registry. In the event that no carrier of Philippine registry is available, Goods may be shipped by a carrier which is not of Philippine registry provided that the Supplier obtains and presents to the Procuring Entity certification to this effect from the nearest Philippine consulate to the port of dispatch. In the event that carriers of Philippine registry are available but their schedule delays the Supplier in its performance of this Contract the period from when the Goods were first ready for shipment and the actual date of shipment the period of delay will be considered force majeure.</p> <p>The Procuring Entity accepts no liability for the damage of Goods during transit other than those prescribed by INCOTERMS for DDP deliveries. In the case of Goods supplied from within the Philippines or supplied by domestic Suppliers risk and title will not be deemed to have passed to the Procuring Entity until their receipt and final acceptance at the final destination.</p> <p><b>Intellectual Property Rights –</b></p> <p>The Supplier shall indemnify the Procuring Entity against all third-party claims of infringement of patent, trademark, or industrial design rights arising from use of the Goods or any part thereof.</p>
2.2	<p><i>[If partial payment is allowed, state]</i> “The terms of payment shall be as follows: _____.”</p>
4	<p>The inspections and tests that will be conducted are: <i>[Indicate the applicable inspections and tests]</i></p>

## ***Section VI. Schedule of Requirements***

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

Item Number	Description	Quantity		Unit Price	Total Amount	Delivered, Weeks/Months
1	Lugol's Iodine, 500ml (Expiration upon delivery: at least 2 years)	bot	4	2,010.00	8,040.00	a. 1 <sup>st</sup> Delivery 20 CD upon the receipt of NTP
2	Drug test liquid control sample (Positive and negative) 5ml	pc	20	4,000.00	80,000.00	
3	Differential Stain for Blood Smear Specification: Composition: Includes 4-step staining set (Solution 1 - 250 ml colorless; Solution 2 - 250 ml red solution; Solution 3 - 250 ml blue solution; Solution 4 250 ml colorless solution) Functionality: Quick dry stain for peripheral blood and bone marrow smear. Packaging: Approximately 500 tests, Stability: Must be stable for a minimum of 12 months at room temperature. Other Inclusions: Stained smear must appear pinkish gray. Microscopically, erythrocytes stain must appear pink to orange. Eosinophils must appear bright orange. Platelets must appear purple granules. Neutrophils cytoplasm must appear light pink. And lymphocytes cytoplasm must appear light to medium blue. Expiration upon delivery: at least 2 years	set	2	3,992.00	7,984.00	b. Succeeding delivery with schedules
4	Evacuated Tube - Na Citrate, (Specification: 3.2%, blue top, 100's, 1.8 ml max bld vol., disposable, sterile, plastic, for single use, with label indicating additive content, approximate blood draw, expiration and lot #, field for patient name, age and ward) Expiration upon delivery: at least 2 years	pack	40	466.00	18,640.00	
5	Needle, multisample, flash back blood collection, Specification: 100's/box, sterile, gauge 21, multiple sample draw, transparent/clear hub, with 1 adaptor per box. Expiration upon delivery should be at least 2 years	box	38	1,172.50	44,555.00	
	For HBA1c Machine Analyzer reagent free use of machine with complete set of computer and printer; Result printout customizable to end-user's format; Can generate printout of summary of reports run daily as specified by end-user; LIS ready, the winning bidder should					

	shoulder any expenses related to LIS connectivity. Fully automated; User-friendly; flexible (different sample tubes can be loaded in any order in any rack); with sample loader and STAT position for emergency assay; compact size, Bench top Lot bid 6				
6	HBA1C (Specification: Complete Set of Reagents, controls, calibrators and supplies; High Performance Liquid Chromatography (HPLC) Method; NGSP and IFCC Certified; at least 400 tests per set) (Expiration upon delivery: at least 2 years)	test	4800	141.00	676,800.00
7	Standard kit, Densi Check plus, 3 standards, 1 blank	kit	1	15,000.00	15,000.00
8	Taxo V (Expiration upon delivery: at least 18 months)	cart	1	1,550.00	1,550.00
9	Taxo X (Expiration upon delivery: at least 18 months)	cart	1	1,550.00	1,550.00
10	Antibiotic Discs, 50's/cartridge - Amikacin, (30 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	2	518.00	1,036.00
11	Antibiotic Discs, 50's/cartridge - Ampicillin (10 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	3	518.00	1,554.00
12	Antibiotic Discs, 50's/cartridge - Azithromycin, (15 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	2	518.00	1,036.00
13	Antibiotic Discs, 50's/cartridge - Aztreonam, (30 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	3	518.00	1,554.00
14	Antibiotic Discs, 50's/cartridge - Cefazolin, (30 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	2	518.00	1,036.00
15	Antibiotic Discs, 50's/cartridge - Cefepime, (30 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	2	518.00	1,036.00
16	Antibiotic Discs, 50's/cartridge - Cefotaxime, (30 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	2	518.00	1,036.00
17	Antibiotic Discs, 50's/cartridge - Cefoxitin, (30 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	2	518.00	1,036.00
18	Antibiotic Discs, 50's/cartridge - Chloramphenicol, (30mcg or ug) (Expiration upon delivery: at least 18 months)	cart	2	518.00	1,036.00

19	Antibiotic Discs, 50's/cartridge - Co-Amoxiclav (Amoxicillin /Clavulanic Acid 30 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	4	518.00	2,072.00	
20	Antibiotic Discs, 50's/cartridge - Cefuroxime (30 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	4	518.00	2,072.00	
21	Antibiotic Discs, 50's/cartridge - Ceftriaxone (30 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	4	518.00	2,072.00	
22	Taxo X + V (Expiration upon delivery: at least 18 months)	cart	1	1,600.00	1,600.00	
23	Antibiotic Discs, 50's/cartridge - Co-trimoxazole (Sulfamethoxazole 23.75ug/Trimethoprim 1.25ug) (Expiration upon delivery: at least 18 months)	cart	4	525.00	2,100.00	
24	Antibiotic Discs - Levofloxacin, 50 Discs/ cart (5mcg or ug) (Expiration upon delivery: at least 18 months)	cart	2	518.00	1,036.00	
25	Antibiotic Discs, 50's/cartridge - Meropenem (10 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	4	518.00	2,072.00	
26	Antibiotic Discs, 50's/cartridge - Nalidixic Acid, (30 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	1	518.00	518	
27	Antibiotic Discs, 50's/cartridge - Norfloxacin, (10 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	1	518.00	518	
28	Antibiotic Discs, 50's/cartridge - Novobiocin, (5 ug) (Expiration upon delivery: at least 18 months)	cart	1	943.00	943	
29	Antibiotic Discs, 50's/cartridge - Ertapenem (10 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	4	593.00	2,372.00	
30	Antibiotic Discs, 50's/cartridge - Imipenem (10 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	4	518.00	2,072.00	
31	Antibiotic Discs, 50's/cartridge - Ceftazidime (30 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	4	518.00	2,072.00	
32	Antibiotic Discs, 50's/cartridge - Ciprofloxacin (5 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	4	518.00	2,072.00	

33	Antibiotic Discs, 50's/cartridge - Erythromycin (15 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	3	518.00	1,554.00	
34	Antibiotic Discs, 50's/cartridge - Gentamicin (10 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	3	518.00	1,554.00	
35	Antibiotic Discs, 50's/cartridge - Piperacillin 10ug/Tazobactam100ug (Expiration upon delivery: at least 18 months)	cart	3	525.00	1,575.00	
36	Antibiotic Discs, 50's/cartridge - Tetracycline (30 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	2	518.00	1,036.00	
37	Antibiotic Discs, 4150's/cartridge - Vancomycin (30 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	2	518.00	1,036.00	
38	Antibiotic Discs, 50's/cartridge - Penicillin (10 UI) (Expiration upon delivery: at least 18 months)	cart	3	518.00	1,554.00	
39	Antibiotic Discs, 50's/cartridge – Ofloxacin (5 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	1	518.00	518	
40	Antibiotic Discs, 50's/cartridge - Oxacillin (1 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	3	518.00	1,554.00	
41	Antibiotic Discs, 50's/cartridge - Clindamycin (2 mcg or ug) (Expiration upon delivery: at least 18 months)	cart	3	518.00	1,554.00	
42	Coagulase test (Rabbit Plasma / Fetal Bovine) (Expiration upon delivery: at least 18 months)	bot	1	3,500.00	3,500.00	
43	Chocolate Agar Plate +300 mg/l Bacitracin, commercially prepared (10 plates/pack) (Expiration upon delivery: at least 1 month)	plate	500	95.00	47,500.00	
44	Additive/supplement to Thayer Martin Medium for Cultivation of nutritionally fastidious microorganism (Isovitalex or Vitox) (Expiration upon delivery: at least 18 months)	bot	1	7,500.00	7,500.00	
45	Dehydrated Culture Media, 500 grams- Brain Heart Infusion Brot (Expiration upon delivery: at least 18 months)	bot	1	5,500.00	5,500.00	
46	Dehydrated Culture Media, 500 grams- Citrate Media (Expiration upon delivery: at least 18 months)	bot	1	5,700.00	5,700.00	

47	Dehydrated Culture Media, 500 grams-Haemophilus Test Medium (Expiration upon delivery: at least 18 months)	bot	1	10,000.00	10,000.00	
48	Salmonella Typing Sera (POLYVALENT)3 ml/kit, for serologic identification of Salmonella (Expiration upon delivery: at least 18 months)	bot	1	10,000.00	10,000.00	
49	Cryogenic vial 4ml, (Specification: round bottom, sterile, internal thread, stable self-standing base, compatible with 81 placer cryogenic box used in the laboratory, can withstand -80°C)	pc	50	19.50	975	
	Reagent free use of machine for Fully Automated Immunology Analyzer Floor Type for Serology section, with controls and other consumables of the machine, with current and valid certificates of product registration (FDA) Result printout customizable to end-user's format; Can generate printout summary of reports run daily as specified by end-user; LIS ready and the winning bidder should shoulder any expenses related to LIS connectivity. with complete set of computers and printer. (LOT BID): 50-68					
	1. Shall provide for FREE latest model of fully automated equipment necessary FLOOR TYPE for the testing, data processing and analysis, generation of results, and efficient system operation with the following features:					
	a. Batch and random-access analyzer using Enhanced Chemiluminescence Assay or Chemiluminescent Microparticle Immunoassay (CMIA) as test methodology, using samples such as: serum, plasma or whole blood, using Levy –Jennings, and Westgard rules for quality controls;					
	b. Quantitative detection of Transfusion Transmissible Infection (TTI) markers for Hepatitis B, HIV, HCV, and Syphilis (e.g. HBsAg, HCV antibody and Antigen, HIV 1 and 2 antibodies, and anti-TP) in serum or plasma samples;					
	c. Preferred reagents shall have specificity and sensitivity levels of not less than 99.5%.					
	d. Must be capable of sample and reagent identification by barcode scanner					
	e. All assays shall have a current and valid certificate of product registration from the Food and Drug Administration (FDA).					

	f. All assays shall have a current or latest proof of Kit Evaluation from STD AIDS Cooperative Central Laboratory (SACCL) or from the National Reference Laboratory for Transfusion Transmissible Infections (Research Institute for Tropical Medicine);				
	g. Uses e-connectivity technology;				
	h. Uses single-use tips and single use cuvettes or microwells;				
	i. Must be able to detect for clots, bubbles and short samples;				
	j. No preparation, no mixing or reconstitution required for reagents;				
	k. On board stability of reagents should be two months or more;				
	l. WITH OR WITHOUT manual primes, purges, washes, or tubing maintenance;				
	m. WITH OR WITHOUT water or drain required and with self-contained onboard waste management;				
	n. Capable of running small sample volume (not more than 100 uL per assay)				
	o. Dimension of analyzer shall fit in the allotted space in the designated area in the SEROLOGY section				
	p. throughput of up to 100-130 test/hour				
	2. Shall continuously provide and replenish consumables including controls and calibrators;				
	3. Shall assure replacement of unused reagents and consumables (delivered within six months of expiration date) before expiration and will provide a guarantee letter of replacement;				
	4. Shall replace reagents and consumables with three-months remaining shelf life before the expiration date;				
	5. Shall provide FREE transport, installation (including minor civil works) and regular maintenance and calibration of all equipment and replacement of parts or units duly coordinated with concerned laboratory staff and EFMS personnel;				
	6. Supplier shall issue a commitment statement that it shall provide FREE training and seminar and/or re-training of laboratory staff and				

	hospital biomedical technicians to make them competent in the use of the equipment; shall issue certificates of training to the personnel concerned;				
	7. Shall perform verification/validation of equipment in conjunction with laboratory staff, and provision of FREE reagents and consumables for that purpose;				
	8. Has the capacity to upgrade the equipment provided when necessary or when the need arises upon request by end user without additional costs to procured reagents;				
	9. Shall provide connection for Laboratory Information System (LIS) of choice of end user and/or Hospital Information System (HIS) - shall be shouldered by the winning bidder.				
	10. Analyzer and accessory equipment must be brand new (equipment not more than three years old are acceptable);				
	11. Shall provide for FREE printer with scanner, paper and ink including maintenance and replacement if malfunctioning within forty-eight hours from notice;				
	12. Shall provide for FREE compatible uninterruptible power supply (UPS) and automatic voltage regulator (AVR) for the analyzer as well as corresponding computer hardware and software integrated with the analyzer;				
	13. Supplier or distributor shall be responsible for securing the equipment and its accessories from rodents and other pests and shall immediately repair or replace the equipment and its accessories should damage occur due to these pests within forty-eight (48) hours from notice;				
	14. Shall provide FREE regular or scheduled preventive maintenance or calibration and ASAP repair or replacement of parts or units of the equipment for FREE duly coordinated with concerned laboratory staff and EFMS Biomedical personnel;				
	15. Shall ensure prompt response by designate engineer/s for correction of equipment failure or malfunction;				
	16. Shall provide a back-up machine for FREE, including transportation and installation, in case				

	of machine breakdown which cannot be resolved within 48 hours from notice;				
	17. Shall shoulder the cost or reimburse the expenses for examinations sent out related to equipment failure or defects in the reagents or consumables which cannot be resolved within forty-eight hours from notice;				
	18. Shall replenish consumed reagents and consumables incidental to repeated test runs due to equipment failure or malfunction including that of the UPS and AVR, defects in reagent or consumables; or due to trial runs during trainings or retraining; or during scheduled preventive maintenance or calibration of equipment;				
	19. Shall provide updated Material Safety Data Sheet (MSDS) for chemical reagents upon delivery of the item;				
	20. Manufacturing or reagents and equipment are covered by relevant ISO certification or quality certification of similar nature. (FOR POST-QUAL DOCUMENTS)				
	21. Products carried are covered by current Certificate of Product Registration or in the absence of the former, proof of renewal or updated Certificate of Exemption issued by the Philippine FDA;				
	22. The supplier shall have installations of the same equipment and its accessories and using the same reagents in the Philippines. A certification from at least one institution having such installation shall be provided. (FOR POST-QUAL DOCUMENTS);				
	23. The supplier shall submit a certificate of distributorship from the principal manufacturer of the equipment and reagents (FOR POST-QUAL DOCUMENTS);				
	24. Shall allow delivery on staggered or as per need basis;				
	25. Supplier /distributor is preferably, but not necessarily, ISO 9001:2015 certified (FOR POST-QUAL DOCUMENTS);				
	26. Should there be persistence of erroneous, invalid, or inconsistent results that may jeopardize the safety of the patients or the quality of services provided despite attempts to repair, the contract shall be terminated at the discretion of the department. The supplier shall refund whatever is paid equivalent to the value				

	of unused reagents or wasted reagents due to repeat analyses or repairs; or if not yet paid, shall not be paid for the said reagents. The supplier shall be compelled to immediately remove the equipment from the section.				
50	Anti-HBc (CLIA)	test	700	356.33	249,431.00
51	HBc IgM(CLIA)	test	600	436.50	261,900.00
52	Anti-Hbe (CLIA)	test	500	410.00	205,000.00
53	Hbe Ag (CLIA)	test	800	417.33	333,864.00
54	HAV IgM (CLIA)	test	200	295.50	59,100.00
55	Anti-HAV IgG (CLIA)	test	200	325.00	65,000.00
56	Anti-Hbs(CLIA)	test	1000	480.00	480,000.00
57	T3, (CLIA)	test	200	223.50	44,700.00
58	T4, (CLIA)	test	200	243.50	48,700.00
59	TSH, (CLIA)	test	1600	257.67	412,272.00
60	FT3, (CLIA)	test	1500	234.33	351,495.00
61	FT4, (CLIA)	test	1500	244.33	366,495.00
62	PSA, (CLIA)	test	700	393.00	275,100.00
63	CA 125, (CLIA)	test	400	614.50	245,800.00
64	CEA, (CLIA)	test	500	429.50	214,750.00
65	Troponin I HIGHLY SENSITIVE, (CLIA)	test	3000	775.50	2,326,500.00
66	Beta-HCG, (CLIA)	test	400	611.00	244,400.00
67	Procalcitonin, (CLIA)	test	500	1,119.50	559,750.00
68	Ferritin, (CLIA)	test	500	414.50	207,250.00
69	Ammonia Water/Scott's Tap 378.5 ml	bot	1	2,250.00	2,250.00
70	Clear Frozen Section Compound Mounting Medium, 118ml dipper bottle	bot	1	1,350.00	1,350.00
	Antigen decloaker: for free of use, bench top with pressure chamber unit, built in slide rack (able to hold at least 70 slides per run), able to retrieve antigen within 60minutes or faster, digital display, working temperature of 60C-110C (temperature increment of 5C or lower), 220-230V (50-60Hz, 1000W). Expiry of				

	reagents should be at least 2 years upon delivery (Lot bid: 71-78)				
71	Polymer Detection System (contain pre-diluted reagents: Peroxidase Block, Protein Block, Post Primary Block, Polymer, DAB Chromogen, DAB Substrate Buffer (Polymer) and Hematoxylin) 50 tests	kit	4	20,096.03	80,384.12
72	Epitope Retrieval Solution pH6, 1L, 1600 tests	bot	2	35,741.57	71,483.14
73	Epitope Retrieval Solution pH9, 1L, 1600 tests	bot	2	35,741.57	71,483.14
74	Wash Solution 10x conc., 1000ml, 640 tests	bot	6	28,752.50	172,515.00
75	Primary Antibodies for ER, 7ml, 46 tests	bot	4	34,717.75	138,871.00
76	Primary Antibodies for PR, 7ml, 46 tests	bot	4	34,717.75	138,871.00
77	Primary Antibodies for HER2, 6ml, 46 tests	bot	4	35,944.41	143,777.64
78	Pen, light blue hydrophobic reagent soluble in commonly used clearing agents; designed to minimize wastage of reagents by allowing the user to ring the tissue(s) or cells to be stained thereby localizing the staining reagents, 1000 tests	pc	1	18,331.00	18,331.00
	Supply and Delivery of the following inclusive of consumables, controls, extraction kits, VTM, and swabs LOT BID # 79 (compatible with the existing machine SAN SURE):				
79	COVID-19 RT PCR nucleic acid and Diagnostic Kit	test	4320	350.00	1,512,000.00
80	Respirator Mask (Specification: NIOSH Approved for at least 95 percent filtration efficiency; Cushioning nose foam, collapsed resistant cup shape, two strap design with welded dual point attachment, Fit tested to molecular laboratory staff, Medium Size)	pc	720	75.00	54,000.00
					<b>10,347,503.04</b>

## **Section VII. Technical Specifications**

### **Notes for Preparing the Technical Specifications**

A set of precise and clear specifications is a prerequisite for Bidders to respond realistically and competitively to the requirements of the Procuring Entity without qualifying their Bids. In the context of Competitive Bidding, the specifications (*e.g.* production/delivery schedule, manpower requirements, and after-sales service/parts, descriptions of the lots or items) must be prepared to permit the widest possible competition and, at the same time, present a clear statement of the required standards of workmanship, materials, and performance of the goods and services to be procured. Only if this is done will the objectives of transparency, equity, efficiency, fairness, and economy in procurement be realized, responsiveness of bids be ensured, and the subsequent task of bid evaluation and post-qualification facilitated. The specifications should require that all items, materials and accessories to be included or incorporated in the goods be new, unused, and of the most recent or current models, and that they include or incorporate all recent improvements in design and materials unless otherwise provided in the Contract.

Samples of specifications from previous similar procurements are useful in this respect. The use of metric units is encouraged. Depending on the complexity of the goods and the repetitiveness of the type of procurement, it may be advantageous to standardize the General Technical Specifications and incorporate them in a separate subsection. The General Technical Specifications should cover all classes of workmanship, materials, and equipment commonly involved in manufacturing similar goods. Deletions or addenda should then adapt the General Technical Specifications to the particular procurement.

Care must be taken in drafting specifications to ensure that they are not restrictive. In the specification of standards for equipment, materials, and workmanship, recognized Philippine and international standards should be used as much as possible. Where other particular standards are used, whether national standards or other standards, the specifications should state that equipment, materials, and workmanship that meet other authoritative standards, and which ensure at least a substantially equal quality than the standards mentioned, will also be acceptable. The following clause may be inserted in the Special Conditions of Contract or the Technical Specifications.

#### **Sample Clause: Equivalency of Standards and Codes**

Wherever reference is made in the Technical Specifications to specific standards and codes to be met by the goods and materials to be furnished or tested, the provisions of the latest edition or revision of the relevant standards and codes shall apply, unless otherwise expressly stated in the Contract. Where such standards and codes are national or relate to a particular country or region, other authoritative standards that ensure substantial equivalence to the standards and codes specified will be acceptable.

Reference to brand name and catalogue number should be avoided as far as possible; where unavoidable they should always be followed by the words "*or at least equivalent.*" References to brand names cannot be used when the funding source is the GOP.

Where appropriate, drawings, including site plans as required, may be furnished by the Procuring Entity with the Bidding Documents. Similarly, the Supplier may be requested to provide drawings or samples either with its Bid or for prior review by the Procuring Entity during contract execution.

Bidders are also required, as part of the technical specifications, to complete their statement of compliance demonstrating how the items comply with the specification.

# Technical Specifications

Item	Specification	Statement of Compliance
		<p><i>[Bidders must state here either “Comply” or “Not Comply” against each of the individual parameters of each Specification stating the corresponding performance parameter of the equipment offered. Statements of “Comply” or “Not Comply” must be supported by evidence in a Bidders Bid and cross-referenced to that evidence. Evidence shall be in the form of manufacturer’s un-amended sales literature, unconditional statements of specification and compliance issued by the manufacturer, samples, independent test data etc., as appropriate. A statement that is not supported by evidence or is subsequently found to be contradicted by the evidence presented will render the Bid under evaluation liable for rejection. A statement either in the Bidder’s statement of compliance or the supporting evidence that is found to be false either during Bid evaluation, post-qualification or the execution of the Contract may be regarded as fraudulent and render the Bidder or supplier liable for prosecution subject to the applicable laws and issuances.]</i></p>
1	Lugol's Iodine, 500ml (Expiration upon delivery: at least 2 years)	
2	Drug test liquid control sample (Positive and negative) 5ml	
3	Differential Stain for Blood Smear Specification: Composition: Includes 4-step staining set (Solution 1 - 250 ml colorless; Solution 2 - 250 ml red solution; Solution 3 - 250 ml blue solution; Solution 4 250 ml colorless solution) Functionality: Quick dry stain for peripheral	

	blood and bone marrow smear. Packaging: Approximately 500 tests, Stability: Must be stable for a minimum of 12 months at room temperature. Other Inclusions: Stained smear must appear pinkish gray. Microscopically, erythrocytes stain must appear pink to orange. Eosinophils must appear bright orange. Platelets must appear purple granules. Neutrophils cytoplasm must appear light pink. And lymphocytes cytoplasm must appear light to medium blue. Expiration upon delivery: at least 2 years	
4	Evacuated Tube - Na Citrate, (Specification: 3.2%, blue top, 100's, 1.8 ml max bld vol., disposable, sterile, plastic, for single use, with label indicating additive content, approximate blood draw, expiration and lot #, field for patient name, age and ward) Expiration upon delivery: at least 2 years	
5	Needle, multisample, flash back blood collection, Specification: 100's/box, sterile, gauge 21, multiple sample draw, transparent/clear hub, with 1 adaptor per box. Expiration upon delivery should be at least 2 years	
	For HBA1c Machine Analyzer reagent free use of machine with complete set of computer and printer; Result printout customizable to end-user's format; Can generate printout of summary of reports run daily as specified by end-user; LIS ready, the winning bidder should shoulder any expenses related to LIS connectivity. Fully automated; User-friendly; flexible (different sample tubes can be loaded in any order in any rack); with sample loader and STAT position for emergency assay; compact size, Bench top Lot bid 6	
6	HBA1C (Specification: Complete Set of Reagents, controls, calibrators and supplies; High Performance Liquid Chromatography (HPLC) Method; NGSP and IFCC Certified; at least 400 tests per set) (Expiration upon delivery: at least 2 years)	
7	Standard kit, Densi Check plus, 3 standards, 1 blank	
8	Taxo V (Expiration upon delivery: at least 18 months)	
9	Taxo X (Expiration upon delivery: at least 18 months)	

10	Antibiotic Discs, 50's/cartridge - Amikacin, (30 mcg or ug) (Expiration upon delivery: at least 18 months)	
11	Antibiotic Discs, 50's/cartridge - Ampicillin (10 mcg or ug) (Expiration upon delivery: at least 18 months)	
12	Antibiotic Discs, 50's/cartridge - Azithromycin, (15 mcg or ug) (Expiration upon delivery: at least 18 months)	
13	Antibiotic Discs, 50's/cartridge - Aztreonam, (30 mcg or ug) (Expiration upon delivery: at least 18 months)	
14	Antibiotic Discs, 50's/cartridge - Cefazolin, (30 mcg or ug) (Expiration upon delivery: at least 18 months)	
15	Antibiotic Discs, 50's/cartridge - Cefepime, (30 mcg or ug) (Expiration upon delivery: at least 18 months)	
16	Antibiotic Discs, 50's/cartridge - Cefotaxime, (30 mcg or ug) (Expiration upon delivery: at least 18 months)	
17	Antibiotic Discs, 50's/cartridge - Cefoxitin, (30 mcg or ug) (Expiration upon delivery: at least 18 months)	
18	Antibiotic Discs, 50's/cartridge - Chloramphenicol, (30mcg or ug) (Expiration upon delivery: at least 18 months)	
19	Antibiotic Discs, 50's/cartridge - Co-Amoxiclav (Amoxicillin /Clavulanic Acid 30 mcg or ug) (Expiration upon delivery: at least 18 months)	
20	Antibiotic Discs, 50's/cartridge - Cefuroxime (30 mcg or ug) (Expiration upon delivery: at least 18 months)	
21	Antibiotic Discs, 50's/cartridge - Ceftriaxone (30 mcg or ug) (Expiration upon delivery: at least 18 months)	
22	Taxo X + V (Expiration upon delivery: at least 18 months)	
23	Antibiotic Discs, 50's/cartridge - Co-trimoxazole (Sulfamethoxazole 23.75ug/Trimethoprim 1.25ug) (Expiration upon delivery: at least 18 months)	

24	Antibiotic Discs - Levofloxacin, 50 Discs/ cart (5mcg or ug) (Expiration upon delivery: at least 18 months)	
25	Antibiotic Discs, 50's/cartridge - Meropenem (10 mcg or ug) (Expiration upon delivery: at least 18 months)	
26	Antibiotic Discs, 50's/cartridge - Nalidixic Acid, (30 mcg or ug) (Expiration upon delivery: at least 18 months)	
27	Antibiotic Discs, 50's/cartridge - Norfloxacin, (10 mcg or ug) (Expiration upon delivery: at least 18 months)	
28	Antibiotic Discs, 50's/cartridge - Novobiocin, (5 ug) (Expiration upon delivery: at least 18 months)	
29	Antibiotic Discs, 50's/cartridge - Ertapenem (10 mcg or ug) (Expiration upon delivery: at least 18 months)	
30	Antibiotic Discs, 50's/cartridge - Imipenem (10 mcg or ug) (Expiration upon delivery: at least 18 months)	
31	Antibiotic Discs, 50's/cartridge - Ceftazidime (30 mcg or ug) (Expiration upon delivery: at least 18 months)	
32	Antibiotic Discs, 50's/cartridge - Ciprofloxacin (5 mcg or ug) (Expiration upon delivery: at least 18 months)	
33	Antibiotic Discs, 50's/cartridge - Erythromycin (15 mcg or ug) (Expiration upon delivery: at least 18 months)	
34	Antibiotic Discs, 50's/cartridge - Gentamicin (10 mcg or ug) (Expiration upon delivery: at least 18 months)	
35	Antibiotic Discs, 50's/cartridge - Piperacillin 10ug/Tazobactam100ug (Expiration upon delivery: at least 18 months)	
36	Antibiotic Discs, 50's/cartridge - Tetracycline (30 mcg or ug) (Expiration upon delivery: at least 18 months)	
37	Antibiotic Discs, 4150's/cartridge - Vancomycin (30 mcg or ug) (Expiration upon delivery: at least 18 months)	
38	Antibiotic Discs, 50's/cartridge - Penicillin (10 UI) (Expiration upon delivery: at least 18 months)	

39	Antibiotic Discs, 50's/cartridge – Ofloxacin (5 mcg or ug) (Expiration upon delivery: at least 18 months)	
40	Antibiotic Discs, 50's/cartridge - Oxacillin (1 mcg or ug) (Expiration upon delivery: at least 18 months)	
41	Antibiotic Discs, 50's/cartridge - Clindamycin (2 mcg or ug) (Expiration upon delivery: at least 18 months)	
42	Coagulase test (Rabbit Plasma / Fetal Bovine) (Expiration upon delivery: at least 18 months)	
43	Chocolate Agar Plate +300 mg/l Bacitracin, commercially prepared (10 plates/pack) (Expiration upon delivery: at least 1 month)	
44	Additive/supplement to Thayer Martin Medium for Cultivation of nutritionally fastidious microorganism (Isovitalex or Vitox) (Expiration upon delivery: at least 18 months)	
45	Dehydrated Culture Media, 500 grams- Brain Heart Infusion Brot (Expiration upon delivery: at least 18 months)	
46	Dehydrated Culture Media, 500 grams- Citrate Media (Expiration upon delivery: at least 18 months)	
47	Dehydrated Culture Media, 500 grams- Haemophilus Test Medium (Expiration upon delivery: at least 18 months)	
48	Salmonella Typing Sera (POLYVALENT)3 ml/kit, for serologic identification of Salmonella (Expiration upon delivery: at least 18 months)	
49	Cryogenic vial 4ml, (Specification: round bottom, sterile, internal thread, stable self-standing base, compatible with 81 placer cryogenic box used in the laboratory, can withstand -80°C)	
	Reagent free use of machine for Fully Automated Immunology Analyzer Floor Type for Serology section, with controls and other consumables of the machine, with current and valid certificates of product registration (FDA) Result printout customizable to end-user's format; Can generate printout summary of reports run daily as specified by end-user; LIS ready and the winning bidder should shoulder any expenses related to LIS connectivity. with complete set of computers and printer. (LOT BID): 50-68	

	1. Shall provide for FREE latest model of fully automated equipment necessary FLOOR TYPE for the testing, data processing and analysis, generation of results, and efficient system operation with the following features:	
	a. Batch and random-access analyzer using Enhanced Chemiluminescence Assay or Chemiluminescent Microparticle Immunoassay (CMIA) as test methodology, using samples such as: serum, plasma or whole blood, using Levy – Jennings, and Westgard rules for quality controls;	
	b. Quantitative detection of Transfusion Transmissible Infection (TTI) markers for Hepatitis B, HIV, HCV, and Syphilis (e.g. HBsAg, HCV antibody and Antigen, HIV 1 and 2 antibodies, and anti-TP) in serum or plasma samples;	
	c. Preferred reagents shall have specificity and sensitivity levels of not less than 99.5%.	
	d. Must be capable of sample and reagent identification by barcode scanner	
	e. All assays shall have a current and valid certificate of product registration from the Food and Drug Administration (FDA).	
	f. All assays shall have a current or latest proof of Kit Evaluation from STD AIDS Cooperative Central Laboratory (SACCL) or from the National Reference Laboratory for Transfusion Transmissible Infections (Research Institute for Tropical Medicine);	
	g. Uses e-connectivity technology:	
	h. Uses single-use tips and single use cuvettes or microwells;	
	i. Must be able to detect for clots, bubbles and short samples;	
	j. No preparation, no mixing or reconstitution required for reagents;	
	k. On board stability of reagents should be two months or more;	
	l. WITH OR WITHOUT manual primes, purges, washes, or tubing maintenance;	
	m. WITH OR WITHOUT water or drain required and with self-contained onboard waste management;	

	n. Capable of running small sample volume (not more than 100 uL per assay)	
	o. Dimension of analyzer shall fit in the allotted space in the designated area in the SEROLOGY section	
	p. throughput of up to 100-130 test/hour	
	2. Shall continuously provide and replenish consumables including controls and calibrators;	
	3. Shall assure replacement of unused reagents and consumables (delivered within six months of expiration date) before expiration and will provide a guarantee letter of replacement;	
	4. Shall replace reagents and consumables with three-months remaining shelf life before the expiration date;	
	5. Shall provide FREE transport, installation (including minor civil works) and regular maintenance and calibration of all equipment and replacement of parts or units duly coordinated with concerned laboratory staff and EFMS personnel;	
	6. Supplier shall issue a commitment statement that it shall provide FREE training and seminar and/or re-training of laboratory staff and hospital biomedical technicians to make them competent in the use of the equipment; shall issue certificates of training to the personnel concerned;	
	7. Shall perform verification/validation of equipment in conjunction with laboratory staff, and provision of FREE reagents and consumables for that purpose;	
	8. Has the capacity to upgrade the equipment provided when necessary or when the need arises upon request by end user without additional costs to procured reagents;	
	9. Shall provide connection for Laboratory Information System (LIS) of choice of end user and/or Hospital Information System (HIS) - shall be shouldered by the winning bidder.	
	10. Analyzer and accessory equipment must be brand new (equipment not more than three years old are acceptable);	
	11. Shall provide for FREE printer with scanner, paper and ink including maintenance and	

	replacement if malfunctioning within forty-eight hours from notice;	
	12. Shall provide for FREE compatible uninterruptible power supply (UPS) and automatic voltage regulator (AVR) for the analyzer as well as corresponding computer hardware and software integrated with the analyzer;	
	13. Supplier or distributor shall be responsible for securing the equipment and its accessories from rodents and other pests and shall immediately repair or replace the equipment and its accessories should damage occur due to these pests within forty-eight (48) hours from notice;	
	14. Shall provide FREE regular or scheduled preventive maintenance or calibration and ASAP repair or replacement of parts or units of the equipment for FREE duly coordinated with concerned laboratory staff and EFMS Biomedical personnel;	
	15. Shall ensure prompt response by designate engineer/s for correction of equipment failure or malfunction;	
	16. Shall provide a back-up machine for FREE, including transportation and installation, in case of machine breakdown which cannot be resolved within 48 hours from notice;	
	17. Shall shoulder the cost or reimburse the expenses for examinations sent out related to equipment failure or defects in the reagents or consumables which cannot be resolved within forty-eight hours from notice;	
	18. Shall replenish consumed reagents and consumables incidental to repeated test runs due to equipment failure or malfunction including that of the UPS and AVR, defects in reagent or consumables; or due to trial runs during trainings or retraining; or during scheduled preventive maintenance or calibration of equipment;	
	19. Shall provide updated Material Safety Data Sheet (MSDS) for chemical reagents upon delivery of the item;	
	20. Manufacturing or reagents and equipment are covered by relevant ISO certification or quality certification of similar nature. (FOR POST-QUAL DOCUMENTS)	
	21. Products carried are covered by current Certificate of Product Registration or in the absence of the former, proof of renewal or	

	updated Certificate of Exemption issued by the Philippine FDA;	
	22. The supplier shall have installations of the same equipment and its accessories and using the same reagents in the Philippines. A certification from at least one institution having such installation shall be provided. (FOR POST-QUAL DOCUMENTS);	
	23. The supplier shall submit a certificate of distributorship from the principal manufacturer of the equipment and reagents (FOR POST-QUAL DOCUMENTS);	
	24. Shall allow delivery on staggered or as per need basis;	
	25. Supplier /distributor is preferably, but not necessarily, ISO 9001:2015 certified (FOR POST-QUAL DOCUMENTS);	
	26. Should there be persistence of erroneous, invalid, or inconsistent results that may jeopardize the safety of the patients or the quality of services provided despite attempts to repair, the contract shall be terminated at the discretion of the department. The supplier shall refund whatever is paid equivalent to the value of unused reagents or wasted reagents due to repeat analyses or repairs; or if not yet paid, shall not be paid for the said reagents. The supplier shall be compelled to immediately remove the equipment from the section.	
50	Anti-HBc (CLIA)	
51	HBc IgM(CLIA)	
52	Anti-Hbe (CLIA)	
53	Hbe Ag (CLIA)	
54	HAV IgM (CLIA)	
55	Anti-HAV IgG (CLIA)	
56	Anti-Hbs(CLIA)	
57	T3, (CLIA)	
58	T4, (CLIA)	
59	TSH, (CLIA)	

60	FT3, (CLIA)	
61	FT4, (CLIA)	
62	PSA, (CLIA)	
63	CA 125, (CLIA)	
64	CEA, (CLIA)	
65	Troponin I HIGHLY SENSITIVE, (CLIA)	
66	Beta-HCG, (CLIA)	
67	Procalcitonin, (CLIA)	
68	Ferritin, (CLIA)	
69	Ammonia Water/Scott's Tap 378.5 ml	
70	Clear Frozen Section Compound Mounting Medium, 118ml dipper bottle	
	Antigen decloaker: for free of use, bench top with pressure chamber unit, built in slide rack (able to hold at least 70 slides per run), able to retrieve antigen within 60minutes or faster, digital display, working temperature of 60C-110C (temperature increment of 5C or lower), 220-230V (50-60Hz, 1000W). Expiry of reagents should be at least 2 years upon delivery (Lot bid: 71-78)	
71	Polymer Detection System (contain pre-diluted reagents: Peroxidase Block, Protein Block, Post Primary Block, Polymer, DAB Chromogen, DAB Substrate Buffer (Polymer) and Hematoxylin) 50 tests	
72	Epitope Retrieval Solution pH6, 1L, 1600 tests	
73	Epitope Retrieval Solution pH9, 1L, 1600 tests	
74	Wash Solution 10x conc., 1000ml, 640 tests	
75	Primary Antibodies for ER, 7ml, 46 tests	
76	Primary Antibodies for PR, 7ml, 46 tests	
77	Primary Antibodies for HER2, 6ml, 46 tests	
78	Pen, light blue hydrophobic reagent soluble in commonly used clearing agents; designed to minimize wastage of reagents by allowing the user to ring the tissue(s) or cells to be stained	

	thereby localizing the staining reagents, 1000 tests	
	Supply and Delivery of the following inclusive of consumables, controls, extraction kits, VTM, and swabs LOT BID # 79 (compatible with the existing machine SAN SURE):	
79	COVID-19 RT PCR nucleic acid and Diagnostic Kit	
80	Respirator Mask (Specification: NIOSH Approved for at least 95 percent filtration efficiency; Cushioning nose foam, collapsed resistant cup shape, two strap design with welded dual point attachment, Fit tested to molecular laboratory staff, Medium Size)	

## ***Section VIII. Checklist of Technical and Financial Documents***

### **Notes on the Checklist of Technical and Financial Documents**

The prescribed documents in the checklist are mandatory to be submitted in the Bid, but shall be subject to the following:

- a. GPPB Resolution No. 09-2020 on the efficient procurement measures during a State of Calamity or other similar issuances that shall allow the use of alternate documents in lieu of the mandated requirements; or
- b. Any subsequent GPPB issuances adjusting the documentary requirements after the effectivity of the adoption of the PBDs.

The BAC shall be checking the submitted documents of each Bidder against this checklist to ascertain if they are all present, using a non-discretionary “pass/fail” criterion pursuant to Section 30 of the 2016 revised IRR of RA No. 9184.

# Checklist of Technical and Financial Documents

## VALENZUELA MEDICAL CENTER

PUBLIC BIDDING NO. VMC-2025-044

PROJECT : **SUPPLY AND DELIVERY OF LABORATORY SUPPLIES AND REAGENTS – FAILED ITEMS FOR CY 2025 – (REBID)**

BIDDER :

### I. TECHNICAL COMPONENT ENVELOPE

#### Class "A" Documents

##### Legal Documents

<input type="checkbox"/>	(a) Valid PhilGEPS Registration Certificate Platinum Membership) (all pages);
<input type="checkbox"/>	(b) Statement of the prospective bidder of <b>all</b> its ongoing government and private contracts, including contracts awarded but not yet started, if any, whether similar or not similar in nature and complexity to the contract to be bid; <b>and</b>
<input type="checkbox"/>	(c) Statement of the bidder's Single Largest Completed Contract (SLCC) similar to the contract to be bid, except under conditions provided for in Sections 23.4.1.3 and 23.4.2.4 of the 2016 revised IRR of RA No. 9184, within the last three (3) years as provided in the Bidding Documents; <b>and</b>
<input type="checkbox"/>	(d) Original copy of Bid Security. If in the form of a Surety Bond, submit also a certification issued by the Insurance Commission; <b>or</b> Original copy of Notarized Bid Securing Declaration
<input type="checkbox"/>	(e) Conformity with the Technical Specifications, which may include production/delivery schedule, manpower requirements, and/or after-sales/parts, if applicable; <b>and</b>
<input type="checkbox"/>	(f) Original duly signed Omnibus Sworn Statement (OSS); ➤ <b>For corporation/Partnership/Cooperative – attach</b> Original Notarized Secretary's Certificate ➤ For JVA - attach Original Special Power of Attorney of all members of the joint venture giving full power and authority to its officer to sign the OSS and do acts to represent the Bidder.

##### Financial Documents

<input type="checkbox"/>	(g) The prospective bidder's computation of Net Financial Contracting Capacity (NFCC); <b>or</b> A committed Line of Credit from a Universal or Commercial Bank in lieuof its NFCC computation.
<input type="checkbox"/>	(h) If applicable, a duly signed joint venture agreement (JVA) in case the joint venture is already in existence <b>or</b> duly notarized statements from all the potential joint venture partners stating that they will enter into and abide by the provisions of the JVA in the instance that the bid is successful.

### II. FINANCIAL COMPONENT ENVELOPE

<input type="checkbox"/>	(i)Original of duly signed and accomplished Financial Bid Form
<input type="checkbox"/>	(j) Original of duly signed and accomplished Price Schedule(s).
<input type="checkbox"/>	(k) Certificate of Product Registration (CPR) or Certificate of Exemption Arranged chronologically per Item or Certificate of Exemption
<input type="checkbox"/>	<i>Other documentary requirements under RA No. 9184 (as applicable)</i>
<input type="checkbox"/>	(l) [For foreign bidders claiming by reason of their country's extension of reciprocal rights to Filipinos] Certification from the relevant government office of their country stating that Filipinos are allowed to participate in government procurement activities for the same item or product.
<input type="checkbox"/>	(m) Certification from the DTI if the Bidder claims preference as a Domestic Bidder or Domestic Entity.

### **ADDITIONAL REQUIREMENTS BY VMC (POST-QUALIFICATION)**

<input type="checkbox"/>	CTC copy of Official Receipt as proof of payment of bidding documents.
<input type="checkbox"/>	Registration certificate from Securities and Exchange Commission (SEC), Department of Trade and Industry (DTI) for sole proprietorship, or Cooperative Development Authority (CDA) for cooperatives or its equivalent document.
<input type="checkbox"/>	Mayor's or Business permit issued by the city or municipality where the principal place of business of the prospective bidder is located, or the equivalent document for Exclusive Economic Zones or Areas.
<input type="checkbox"/>	Tax clearance per E.O. No. 398, s. 2005, as finally reviewed and approved by the Bureau of Internal Revenue (BIR).
<input type="checkbox"/>	The Supplier's audited financial statements, showing, among others, the Supplier's total and current assets and liabilities, stamped "received" by the BIR
<input type="checkbox"/>	2023 ITR or its duly accredited and authorized institutions, for the preceding calendar year which should not be earlier than two (2) years from the date of bid submission
<input type="checkbox"/>	Certificate of Good Performance from at least one (1) Government or Private Hospital / Agency except from VMC (CY 2024-present)
<input type="checkbox"/>	Certificate of Good Performance with Satisfactory rating from laboratory section for existing suppliers for CY 2023-2024.
<input type="checkbox"/>	Special Power of Attorney (SPA) for authorized representative if OSS is Sole Proprietorship
<input type="checkbox"/>	Proof of evidence for Single Largest Completed Contract (SLCC) – Purchase Order or Notice of Award or Contract Agreement equivalent to 25% of amount to be bid
<input type="checkbox"/>	Certificate from the manufacturer to distribute their products or Exclusive Distributorship or any equivalent document
<input type="checkbox"/>	License to Operate (LTO)
<input type="checkbox"/>	Certificate of Stocks Availability (Notarized)

