

STANDARD BIDDING DOCUMENT

Procurement of Therapeutic Goods

(Pharmaceuticals)

(Single Stage Two Envelop Procedure)

(National Competitive Bidding)

FOR

Procurement of Medicines (HIT Hospital)

Bid No. 5171/IT-3035/2024-25/Medicines/FOR/Med/Proc dated 11 October 2024



**PROCUREMENT DIRECTORATE HIT TAXILA
TAXILA CANTT**

Bid Submission Date / Time: 21 November 2024 at 10:30 AM

Bid Opening Date / Time: 21 November 2024 at 11:00 AM

***Feel free to contact for any query regarding details
of tendered items as well as Tender Clauses***

HEAVY INDUSTRIES TAXILA
TENDER NOTICE

Date: 25 October 2024

1. Sealed tenders are invited from reputed firms possessing NTN Certificates for supply of under mentioned items. Procurement will be carried out in the light of PPRA and MoDP rules: -

Tender No	Nomenclature	A/U	Qty	Delivery Period	Last date of Collection of Tender Documents	Date and time of Tender	
						Submission	Opening
IT-3035	Procurement of Medicines (HIT Hospital)	Details at Section E of this tender			<i>No physical collection involved (information available on PPRA and HIT website)</i>	21 November 2024 at 10:30 AM	21 November 2024 at 11:00 AM

2. Bidder must submit the following document along with check list at time of submission of tender: -

- a. Photocopy of Registration / Pre-qualification / Indexation letter issued by the HIT.
- b. Attested copy of Registration certificate issued by Sales Tax Department, copy of NTN certificate.
- c. Audit report of last 3 x FYs.
- d. Attested Bank Statement for last one year.
- e. Attested copy of CNIC of MD.
- f. Trade link between firm and OEM (in case of distributor / agent).
- g. Certificate on a judicial paper worth Rs 100 duly attested by Oath Commissioner that firm is neither defaulter nor blacklisted by any Govt Org directly or indirectly.
- h. Detail of list of contracts last 3 x FYs / works experience with Govt / Semi Govt Org.
- i. Copy of registration letter with Govt / Semi Govt organization (if any).
- j. Postal order of Rs 2,000/- should be enclosed with Technical Bid in favour of Director Procurement HIT Taxila

NOTE

In case of any query regarding Nomenclature, Specifications, firms must contact following officials well before Bid Opening date: -

- a. **Maj Aatif Ayub Malik (Procurement Directorate)**
Tel: (051) 9315333 Ext 63215
Fax: (051) 9315029
E-Mail: dirproccte@hit.gov.pk
- b. **HIT Hospital**
Tel: (051) 9315333 Ext 66180

TABLE OF CONTENT

Ser	Forms	Details	Page No	
1.	PART – I	Bidding Procedure & Requirements		
	a. Section – A	Invitation to Bids	5-6	
	b. Section – B	Instruction to Bidders (ITBs)		
		(1) Introduction		7-9
		(2) Bidding Documents		10-11
		(3) Preparation of Bids		12-18
		(4) Submission of Bids		19
		(5) Opening and Evaluation of Bids		20-28
		(6) Award of Contract		29-30
		(7) Grievance Redressal & Complaint Review Mechanism		31
	(8) Mechanism of Blacklisting		32-33	
	c. Section – C	Bid data sheet	34-37	
	d. Section – D	Eligible countries	38	
e. Section – E	Schedule of requirements, technical specifications	39		
f. Section – F	Evaluation and qualification criteria	40-42		
g. Section – G	Standard forms	43-50		
2.	PART – II	Conditions of the Contract		
	h. Section - H	General Conditions of the Contract (GCC)	51-62	
	i. Section – I	Special Conditions of the Contract (SCC)	63-66	
	j. Section - J	Contract Forms		
		(1) Form of Contract		67
		(2) Performance Guarantee Form		68
	(3) Integrity Pact		69	

PART-I

BIDDING PROCEDURE & REQUIREMENTS

HEAVY INDUSTRIES TAXILA



Bid No. 5171/IT-3035/2024-25/Medicines/FOR/Med/Proc dated 11 October 2024

Procurement of Medicines (HIT Hospital)

INVITATION TO TENDER / INVITATION TO BID

Date: 25 October 2024

1. This Invitation to Tender (IT) / Invitation to Bid (ITB) follows the Procurement Notice (PN) or Procurement Advertisement (PA) for the subject procurement which will also appear in 2 x newspaper (1 x English and 1 x URDU) on _____ 2024.

2. Heavy Industries Taxila invites sealed bids from eligible Suppliers for the provision of Medical Store items (Medicines) for. The complete original bid (technical & commercial) along with one additional photocopy of **technical bid, properly filled in**, and enclosed in sealed envelope(s) must be delivered as under: -

<u>Ser</u>	<u>Activity</u>	<u>Response</u>	<u>Remarks</u>
a.	Bid submission	10:30 AM, 21 November 2024	Tender Box available at Gate No 5
b.	Bid opening (Technical)	11 AM, 21 November 2024	Venue: Procurement Directorate HIT Taxila
c.	Pre-bid meeting	_____ 2024	Venue _____ Procurement Directorate HIT Taxila
d.	Bid submission address	Gate -5, Heavy Industries Taxila, Taxila Cantt	Complete Bid be dropped in Tender Box before deadline.

3. All bids must be accompanied by a **Bid Security / Earnest Money** in the form of Call Deposit Receipt (**CDR**), Bank Draft (**BD**) or Pay Order (**PO**) in favour of Director Procurement, Heavy Industries, Taxila as per the instructions provided in this Tender.

4. Firms shall nominate a Lead Member / authorized representative, on the firm's letterhead, with authority to conduct all business for and on behalf of the firms during the bidding process, and

in case of award of contract, during the execution of contract.

5. Appointment of the Lead Member shall be subsequently confirmed by submission of a valid Power of Attorney before signing of the contract by the firm winning the contract.

6. This IT consists of **76x pages** and comprises of following forms: -

a. **Part - I**

- (1) **Section – A**. Invitation letter and general instructions to the firms.
- (2) **Section – B**. Instruction to Bidders (ITBs)
- (3) **Section - C**. Bid Data Sheet
- (4) **Section - D**. Eligible Countries
- (5) **Section - E**. Schedule of Requirement, Technical Specification
- (6) **Section – F**. Evaluation and Qualification Criteria
- (7) **Section – G**. Standard Forms

b. **Part – II**. Conditions of the contract

- (1) **Section – H**. General Condition of the contract (GCC)
- (2) **Section – I**. Special Condition of the Contract (SCC)
- (3) **Section - J**. Contract Forms

7. Firms will fill and return, with their offers, the forms Section – C and Section G and the questionnaires duly stamped / signed by the authorized person / signatory

Yours faithfully,

**Deputy Assistant Director Procurement
Procurement Directorate
Heavy Industries Taxila
Taxila Cantt**

INSTRUCTION TO BIDDERS (ITBs)**INTRODUCTION**

a. Scope of Bid	1.1	The Procurement Directorate, as indicated in the Bid Data Sheet (BDS) invites Bids for the supply of Therapeutic Goods (Pharmaceuticals) and related services incidental thereto as specified in the BDS described in Section E - Technical Specifications & Schedule of Requirements. The Name, Identification and Number of Lots (contracts) of the procurement are specified in BDS.
	1.2	The successful Bidders will be expected to supply the Therapeutic Goods (Pharmaceuticals) within the specified period and timeline(s) as stated in the BDS.
b. Source of Funds	2.1	Source of funds is referred in Clause-2 of Invitation for Bids.
c. Eligible Bidders	3.1	A Bidder may be natural person, company or firm or public or semi-public agency of Pakistan or any foreign country, or any combination of them with a formal existing agreement (on Judicial Papers) in the form of a joint venture, consortium, or association. In the case of a joint venture, consortium, or association, all members shall be jointly and severally liable for the execution of the Contract in accordance with the terms and conditions of the Contract. The joint venture, consortium, or association shall nominate a Lead Member as nominated in the BDS, who shall have the authority to conduct all business for and on behalf of any and all the members of the joint venture, consortium, or association during the Bidding process, and in case of award of contract, during the execution of contract.
	3.2	The appointment of Lead Member in the joint venture, consortium, or association shall be confirmed by submission of a valid Power of Attorney to the Procurement Directorate.
	3.3	Verifiable copy of the agreement that forms a joint venture, consortium or association shall be required to be submitted as part of the Bid.
	3.4	Any bid submitted by the joint venture, consortium or association shall indicate the part of proposed contract to be performed by each party and each party shall be evaluated (or post qualified if required) with respect to its contribution only, and the responsibilities of each party shall not be substantially altered without prior written approval of the Procurement Directorate and in line with any instructions issued by the Authority.
	3.5	The invitation for Bids is open to all prospective supplier, manufacturers or authorized agents/dealers subject to any provisions of incorporation or licensing by the respective national incorporating agency or statutory body established for that particular trade or business.
	3.6 .	Foreign Bidders must satisfy all relevant licensing/or registration requirement with the appropriate national incorporating body or the statutory body, before participating in the national/international competitive Bidding with the exception of such procurements made by the foreign missions of Pakistan. For such purpose the bidder must have to initiate the registration process before the bid submission and the necessary evidence shall be submitted to the Procurement Directorate along with their bid,

	<p>however, the final award will be subject to the complete registration process.</p>
3.7	<p>A Bidder shall not have a conflict of interest. All Bidders found to have a conflict of interest shall be disqualified. A Bidders may be considered to have a conflict of interest with one or more parties in this Bidding process, if they:</p> <ul style="list-style-type: none"> (a) are associated or have been associated in the past, directly or indirectly with a firm or any of its affiliates which have been engaged by the Procurement Directorate to provide consulting services for the preparation of the design, specifications and other documents to be used for the procurement of the pharmaceuticals to be procured under this Invitation for Bids. (b) have controlling shareholders in common; or (c) receive or have received any direct or indirect subsidy from any of them; or (d) have the same legal representative for purposes of this Bid; or (e) have a relationship with each other, directly or through common third parties, that puts them in a position to have access to information about or influence on the Bid of another Bidder, or influence the decisions of the Procurement Directorate regarding this Bidding process; or (f) Submit more than one Bid in this Bidding process. (g) Participated as a consultant in the preparation of the design or technical specifications of the services that are the subject of the Bid.
3.8	<p>A Bidder may be ineligible if –</p> <ul style="list-style-type: none"> (a) he is declared bankrupt or, in the case of company or firm, insolvent; (b) payments in favor of the Bidder is suspended in accordance with the judgment of a court of law other than a judgment declaring bankruptcy and resulting (in accordance with the national laws) in the total or partial loss of the right to administer and dispose of its property; (c) legal proceedings are instituted against such Bidder involving an order suspending payments and which may result, in accordance with the national laws, in a declaration of bankruptcy or in any other situation entailing the total or partial loss of the right to administer and dispose of the property; (d) the Bidder is convicted, by a final judgment, of any offence involving professional conduct; (e) the Bidder is blacklisted and hence debarred due to involvement in corrupt and fraudulent practices, or performance failure or due to breach of bid securing declaration. (f) The firm, supplier and contractor is blacklisted or debarred by a foreign country, international organization, or other foreign institutions for the period defined by them.
3.9	<p>Bidders shall provide to the Procurement Directorate evidence of their eligibility, proof of compliance with the necessary legal, technical and financial requirements and their capability and, adequacy of resources to carry out the contract effectively.</p>

	3.10	Bidders shall provide such evidence of their continued eligibility to the satisfaction of the Procurement Directorate, as the Procurement Directorate shall reasonably request.
	3.11	Bidders shall submit proposals relating to the nature, conditions and modalities of sub-contracting wherever the sub-contracting of any elements of the contract amounting to the more than ten (10) percent of the Bid price is envisaged.
	3.12	Firms/companies/suppliers/dealers duly registered with relevant tax and other registration authorities required under Federal Government's rules, laws, statutes or relevant instructions;
	3.13	The Importer/Agent/distributor/supplier must possess valid authorization from the Manufacturer. The authorization certificate must be attested. However, in case of Manufacturer, they should have a documentary proof as prescribed in the Bid Form
d. Eligible Therapeutic Goods (Pharmaceuticals) and Related Services	4.1	All therapeutic goods (Pharmaceuticals) and related services to be supplied under the contract shall have their origin in eligible source countries, and all expenditures made under the contract will be limited to such Therapeutic goods and services. For purpose of this Bid, ineligible countries are stated in the section-D titled as "Eligible Countries".
	4.2	For purposes of this Clause, "origin" means the place where the goods are mined, grown, cultivated, produced, manufactured, or processed, or through manufacture, procession, or assembly, another commercially recognized article results that differs substantially in its basic characteristics from its imported components or the place from where the related services are/to be supplied.
	4.3	The nationality of the supplier that supplies, assembles, distributes, or sells the Therapeutic Goods and services shall not determine the origin of the goods.
	4.4	To establish the eligibility of the therapeutic Goods and the related services, Bidders shall fill the country of origin declarations included in the Form of Bid.
	4.5	If so required in the BDS, the Bidder shall demonstrate that it has been duly authorized by the manufacturer of the goods to deliver in Pakistan (or in respective country in case of procurement by the Pakistani Missions abroad), the therapeutic goods indicated in its Bid.
	4.6	All Therapeutic Goods and related services to be supplied under the contract shall conform to the policies of the Government of Pakistan in vogue. All expenditures made under the contract shall be limited to such Therapeutic Goods and services. For purposes of this clause, (a) the term.
	4.7	For the purposes of this Clause, the term "Therapeutic Goods" includes Drug or alternative medicine or medical devices or biological or other related product as may be notified by Drug Regulatory Authority of Pakistan. , and "related services" includes services such as insurance, Transportation port releases, installation, training and initial maintenance and after sales services.
e. One Bid per Bidder	5.1	A bidder shall submit only one Bid, in the same bidding process, either individually as a Bidder or as a member in a joint venture or any similar arrangement.
	5.2	No bidder can be a sub-contractor while submitting a Bid individually or as a member of a joint venture in the same Bidding process.

	5.3	A person or a firm cannot be a sub-contractor with more than one bidder in the same bidding process.
f. Cost of Bidding	6.1	The Bidder shall bear all costs associated with the preparation and submission of its Bid, and the Procurement Directorate shall in no case be responsible or liable for those costs, regardless of the conduct or outcome of the bidding process.
g. Bidding for Selective Items	7.1	A Bidder, if he so chooses, can bid for selective items from the list of Therapeutic Goods provided for in the Schedule of Requirements. A Bidder is also at a liberty to bid for all the Therapeutic Goods mentioned in the Schedule of Requirements provided he fulfills the requirements. However, a Bidder cannot bid for partial quantities of an item in the Schedule of requirement. The bid must be for the whole quantity of an item as required in the schedule of requirement.

BIDDING DOCUMENTS

h. Contents of Bidding Documents	8.1	The therapeutic goods required, bidding procedures, and terms and conditions of the contract are prescribed in the Bidding Documents. In addition to the Invitation to Bids, the Bidding Documents which should be read in conjunction with any addenda issued in accordance with ITB 10.2 include: Section A Invitation to Bids Section B Instructions to Bidders (ITBs) Section C Bid Data Sheet (BDS) Section D Eligible Countries Section E Technical Specifications, Schedule of Requirements Section F Evaluation and Qualification Criteria Section G Standard Forms Section H General Conditions of Contract (GCC) Section I Special Conditions of Contract (SCC) Section J Contract Forms
	8.2	The 02 x copies to be completed and returned with the Bid is specified in the BDS.
	8.3	The Procurement Directorate is not responsible for the completeness of the Bidding Documents and their addenda, if they were not obtained directly from the Procurement Directorate or the signed pdf version downloaded from the website of the Heavy Industries Taxila and PPRA website. However, Procurement Directorate shall place both the pdf and same editable version to facilitate the bidder for filling the forms.
	8.4.	The Bidder is expected to examine all instructions, forms, terms and specifications in the Bidding Documents. Failure to furnish all the information required in the Bidding Documents will be at the Bidder's risk and may result in the rejection of his Bid.
i. Clarification of Bidding Documents	9.1	A prospective Bidder requiring any clarification of the Bidding Documents may notify the Procurement Directorate in writing or in electronic form that provides record of the content of communication at the Procurement Directorate's address indicated in the BDS.
	9.2	The Procurement Directorate will within three (3) working days after receiving the request for clarification, respond in writing or in electronic form to any request for clarification provided that such request is received not later than three (03) days prior to the deadline for the submission of Bids as prescribed in ITB 23.1. However, this clause shall not apply in case of alternate methods of Procurement.

	9.3	<p>Copies of the Procurement Directorate's response will be forwarded to all identified Prospective Bidders through an identified source of communication, including a description of the inquiry, but without identifying its source.</p> <p>In case of downloading of the Bidding Documents from the website of PA, the response of all such queries will also be available on the same link available at the website.</p>
	9.4	<p>Should the Procurement Directorate deem it necessary to amend the Bidding Documents as a result of a clarification, it shall do so following the procedure under ITB 10.</p>
	9.5	<p>If indicated in the BDS, the Bidder's designated representative is invited at the Bidder's cost to attend a pre-Bid meeting at the place, date and time mentioned in the BDS. During this pre-Bid meeting, prospective Bidders may request clarification of the schedule of requirement, the Evaluation Criteria or any other aspects of the Bidding Documents.</p>
	9.6	<p>Minutes of the pre-Bid meeting, if applicable, including the text of the questions asked by Bidders, including those during the meeting (without identifying the source) and the responses given, together with any responses prepared after the meeting will be transmitted promptly to all prospective Bidders who have obtained the Bidding Documents. Any modification to the Bidding Documents that may become necessary as a result of the pre-Bid meeting shall be made by the Procurement Directorate exclusively through the use of an Addendum pursuant to ITB 10. Non-attendance at the pre-Bid meeting will not be a cause for disqualification of a Bidder.</p>
j. Amendment of Bidding Documents	10.1	<p>Before the deadline for submission of Bids, the Procurement Directorate for any reason, whether at its own initiative or in response to a clarification requested by a prospective Bidder or pre-Bid meeting may modify the Bidding Documents by issuing addenda.</p>
	10.2	<p>Any addendum issued including the notice of any extension of the deadline shall be part of the Bidding Documents pursuant to ITB 8.1 and shall be communicated in writing or in any identified electronic form that provide record of the content of communication to all the bidders who have obtained the Bidding Documents from the Procurement Directorate. The Procurement Directorate shall promptly publish the Addendum at the Procurement Directorate's web page identified in the BDS:</p> <p>Provided that the bidder who had either already submitted their bid or handed over the bid to the courier prior to the issuance of any such addendum shall have the right to withdraw his already filed bid and submit the revised bid prior to the original or extended bid submission deadline.</p>
	10.3	<p>To give prospective Bidders reasonable time in which to take an addendum/corrigendum into account in preparing their Bids, the Procurement Directorate may, at its discretion, extend the deadline for the submission of Bids:</p> <p>Provided that the Procurement Directorate shall extend the deadline for submission of Bid, if such an addendum is issued within last three (03) days of the Bid submission deadline.</p>

PREPARATION OF BIDS

k. Language of Bid	11.1	The Bid prepared by the Bidder, as well as all correspondence and documents relating to the Bid exchanged by the Bidder and the Procurement Directorate shall either be in Urdu or English or both.
	11.2	Where the use of local language is found essential, the original documentation shall be in Urdu or English, which shall be retained on record; for all other purposes their translations in local language shall be used: Provided that such use of local language ensures maximum economy and efficiency in the procurement.
	11.3	In case of Dispute reference shall be made to the original documentation retained on the record.
l. Documents and Sample(s) Constituting the Bid	12.1	<p>The Bid prepared by the Bidder shall constitute the following components: -</p> <ul style="list-style-type: none"> (a) Form of Bid and Bid Prices completed in accordance with ITB 15 and 16; (b) Details of the Sample(s) where applicable and requested in the BDS. (c) Documentary evidence established in accordance with ITB 14 that the Bidder is eligible and/or qualified for the subject bidding process; (d) Documentary evidence established in accordance with ITB 14.3(a) that the Bidder has been authorized by the manufacturer to deliver the therapeutic goods into Pakistan, where required and where the supplier is not the manufacturer of those therapeutic goods; (e) Documentary evidence established in accordance with ITB 13 that the goods and related services to be supplied by the Bidder are eligible therapeutic goods and services, and conform to the Bidding Documents; (f) Documentary evidence of manufacturing license and GMP certificates; (g) Quality Control Procedures; (h) Capacity of the Manufacturer; (i) Bid security or Bid Securing Declaration furnished in accordance with ITB 19; (j) Duly Notarized Power of Attorney authorizing the signatory of the Bidder to submit the bid; and (k) Any other document required in the BDS.
	12.2	<p>Where a sample(s) is required by a Procurement Directorate, the sample shall be:</p> <ul style="list-style-type: none"> (a) submitted as part of the bid, in the quantities, dimensions and other details requested in the BDS; (b) carriage paid; (c) received on, or before, the closing time and date for the submission of bids; and (d) evaluated to determine compliance with all characteristics listed in the BDS.
	12.3	<p>The Procurement Directorate shall retain the sample(s) of the successful Bidder. A Procurement Directorate shall reject the Bid if the sample(s)-</p> <ul style="list-style-type: none"> (a) do(es) not conform to all characteristics prescribed in the bidding documents; and (b) is/are not submitted within the specified time clearly mentioned in the

		Bid Data Sheet.
	12.4	Where it is not possible to avoid using a propriety article as a sample, a Bidder shall make it clear that the propriety article is displayed only as an example of the type or quality of the therapeutic goods being Bided for, and that competition shall not thereby be limited to the extent of that article only.
	12.5	Samples made up from materials supplied by a Procurement Directorate shall not be returned to a Bidder nor shall a Procurement Directorate be liable for the cost of making them.
	12.6	All samples produced from materials belonging to an unsuccessful Bidder shall be kept by the Procurement Directorate till thirty (30) days from the date of award of contract or exhaust of all the grievance forums (including those pending at Authority's Level or in some Court of Law). The sample shall be the property of the Procurement Directorate and shall dispose of such sample in such manner as described by the Drug Regulatory Authority of Pakistan.
m. Documents Establishing Eligibility of Goods and Related Services and Conformity to Bidding Documents	13.1	Pursuant to ITB 13, the Bidder shall furnish, as part of its Bid, all those documents establishing the eligibility in conformity to the terms and conditions specified in the Bidding Documents for all therapeutic goods and related services which the Bidder proposes to deliver.
	13.2	The documentary evidence of the eligibility of the therapeutic goods and related services shall consist of a statement in the Price Schedule of the country of origin of the goods and related services offered which shall be confirmed by a certificate of origin issued at the time of shipment.
	13.3	The documentary evidence of conformity of the therapeutic goods and related services to the Bidding Documents may be in the form of literature, drawings, and data, and shall consist of: <ul style="list-style-type: none"> (a) a detailed description of the essential technical specifications and performance characteristics of the Therapeutic Goods; (b) an item-by-item commentary on the Procurement Directorate's Technical Specifications demonstrating substantial responsiveness of the Therapeutic Goods and Services to those specifications, or a statement of deviations and exceptions to the provisions of the Technical Specifications; (c) any other procurement specific documentation requirement as stated in the BDS.
	13.4	The Bidder shall also furnish a list giving full particulars, including available sources and current prices of therapeutic goods, spare parts, special tools, etc., necessary for the proper and continuing functioning of the Goods during the period specified in the BDS following commencement of the use of the therapeutic goods by the Procurement Directorate.
	13.5	For purposes of the commentary to be furnished pursuant to ITB 13.3(c) above, the Bidder shall note that standards for workmanship, material, and equipment, as well as references to brand names or catalogue numbers designated by the Procurement Directorate in its Technical Specifications, are intended to be descriptive only and not restrictive. The Bidder may substitute alternative standards, brand names, and/or catalogue numbers in its Bid, provided that it demonstrates to the Procurement Directorate's

		satisfaction that the substitutions ensure substantial equivalence to those designated in the Technical Specifications.
	13.6	The required documents and other accompanying documents must be in English. In case any other language than English is used the pertinent translation into English shall be attached to the original version.
n. Documents Establishing Eligibility and Qualification of the Bidder	14.1	Pursuant to ITB 12, the Bidder shall furnish, as part of its Bid, all those documents establishing the Bidder's eligibility to participate in the bidding process and/or its qualification to perform the contract if its Bid is accepted.
	14.2	The documentary evidence of the Bidder's eligibility to Bid shall establish to the satisfaction of the Procurement Directorate that the Bidder, at the time of submission of its bid, is from an eligible country as defined in Section-4 titled as "Eligible Countries".
	14.3	The documentary evidence of the Bidder's qualifications to perform the contract if its Bid is accepted shall establish to the satisfaction of Procurement Directorate that: (a) in the case of a Bidder offering to deliver goods under the contract which the Bidder did not manufacture or otherwise produce, the Bidder has been duly authorized by the therapeutic goods' Manufacturer or producer to deliver the goods in Pakistan; (b) the Bidder has the financial, technical, and supply/production capability necessary to perform the Contract, meets the qualification criteria specified in BDS. (c) in the case of a Bidder not doing business within Pakistan, the Bidder is or will be (if awarded the contract) represented by an Agent in Pakistan equipped, and able to carry out the Supplier's obligations prescribed in the Conditions of Contract and/or Technical Specifications. (d) that the Bidder meets the qualification criteria listed in the Bid Data Sheet.
o. Form of Bid	15.1	The Bidder shall fill the Form of Bid furnished in the Bidding Documents. The Bid Form must be completed without any alterations to its format and no substitute shall be accepted.
p. Bid Prices	16.1	The Bid Prices quoted by the Bidder in the Form of Bid and in the Price Schedules shall conform to the requirements specified below in ITB Clause 16 or exclusively mentioned hereafter in the bidding documents.
	16.2	All items in the Statement of Work must be listed and priced separately in the Price Schedule(s). If a Price Schedule shows items listed but not priced, their prices shall be construed to be included in the prices of other items.
	16.3	Items not listed in the Price Schedule shall be assumed not to be included in the Bid, and provided that the Bid is still substantially responsive in their absence or due to their nominal nature, the corresponding average price of the respective item(s) of the remaining substantially responsive bidder(s) shall be construed to be the price of those missing item(s): Provided that: (a) where there is only one (substantially) responsive bidder, or (b) where there is provision for alternate proposals and the respective items are not listed in the other bids, the Procurement Directorate may fix the price of missing items in

	accordance with market survey, and the same shall be considered as final price.
16.4	The Bid price to be quoted in the Form of Bid in accordance with ITB 16.1 shall be the total price of the Bid.
16.5	The Bidder shall indicate on the appropriate Price Schedule, the unit prices (where applicable) and total Bid price of the therapeutic goods it proposes to deliver under the contract.
16.6	<p>Prices indicated on the Price Schedule shall be entered separately in the following manner:</p> <p>(a) For therapeutic goods manufactured from within Pakistan (or within the country where procurement is being done in case of foreign missions abroad):</p> <p>(1) The price of the therapeutic goods quoted EXW (ex-works, ex-factory, ex-warehouse, ex-showroom, or off-the-shelf, as applicable), including all customs duties and sales and other taxes already paid or payable:</p> <p style="padding-left: 40px;">i. on the components and raw material used in the manufacturing or assembly of goods quoted ex- works or ex-factory;</p> <p style="text-align: center;">or</p> <p style="padding-left: 40px;">ii. on the previously imported goods of foreign origin quoted ex-warehouse, ex-showroom, or off-the-shelf.</p> <p>(2) All applicable taxes which will be payable on the therapeutic goods if the contract is awarded.</p> <p>(3) The price for inland transportation, insurance, and other local costs incidental to delivery of the therapeutic goods to their final destination, if specified in the BDS.</p> <p>(4) The price of other (incidental or allied) services, if any, listed in the BDS.</p>
16.7	<p>Prices proposed on the Price Schedule for therapeutic goods and related services shall be disaggregated, where appropriate as indicated in this Clause. This desegregation shall be solely for the purpose of facilitating the comparison of Bids by the Procurement Directorate. This, shall not in any way limit the Procurement Directorate's right to contract on any of the terms and conditions offered: -</p> <p>(a) For Therapeutic Goods: -</p> <p style="padding-left: 40px;">(1) the price of the therapeutic Goods, quoted as per applicable INCOTERMS as specified in the BDS</p> <p style="padding-left: 40px;">(2) all customs duties, sales tax, and other taxes applicable on therapeutic goods or on the components and raw materials used in their manufacture or assembly, if the contract is awarded to the Bidder, and</p> <p>(b) For Related Services</p> <p style="padding-left: 40px;">(1) The price of the related services, and</p> <p style="padding-left: 40px;">(2) All customs duties, sales tax and other taxes applicable in Pakistan, paid or payable, on the related services, if the contract is awarded to the Bidder.</p>

	16.8	Prices quoted by the Bidder shall be fixed during the Bidder's performance of the contract and not subject to variation on any account. A Bid submitted with an adjustable price will be treated as non-responsive and shall be rejected, pursuant to ITB 28.
	16.9	If so indicated in the Invitation to Bids and Instructions to Bidders, that Bids are being invited for individual contracts (Lots) or for any combination of contracts (packages), Bidders wishing to offer any price reduction for the award of more than one contract shall specify in their Bid the price reductions applicable to each package, or alternatively, to individual contracts (Lots) within a package.
q. Bid Currencies	17.1	Prices shall be quoted in the following currencies: (a) For therapeutic goods and services that the Bidder will deliver from within Pakistan, the prices shall be quoted in Pakistani Rupees, unless otherwise specified in the BDS. (b) For therapeutic goods and related services that the Bidder will deliver from outside Pakistan, or for imported parts or components of goods and related services originating outside Pakistan, the Bid prices shall be quoted in any freely convertible currency of another country. If the Bidder wishes to be paid in a combination of amounts in different currencies, it may quote its price accordingly but use no more than three foreign currencies.
	17.2	For the purposes of comparison of bids quoted in different currencies, the price shall be converted into a single currency specified in the bidding documents. The rate of exchange shall be the selling rate, prevailing on the date of opening of (financial part of) bids specified in the bidding documents, as notified by the State Bank of Pakistan on that day. Currency of the contract shall be as described in the BDS.
	17.3	Bidders shall indicate details of their expected foreign currency requirements in the Bid.
	17.4	Bidders may be required by the Procurement Directorate to clarify their foreign currency requirements and to substantiate that the amounts included in Lump Sum and in the SCC are reasonable and responsive to ITB 17.1.
r. Bid Validity Period	18.1	Bids shall remain valid for the period specified in the BDS after the Bid submission deadline prescribed by the Procurement Directorate. A Bid valid for a shorter period shall be rejected by the Procurement Directorate as non-responsive. The period of Bid validity will be determined from the complementary bid securing instrument i.e. the expiry period of bid security or bid securing declaration as the case may be.
	18.2	Under exceptional circumstances, prior to the expiration of the initial Bid validity period, the Procurement Directorate may request the Bidders' consent to an extension of the period of validity of their Bids only once, for the period not more than the period of initial bid validity. The request and the Bidders responses shall be made in writing or in electronic forms that provide record of the content of communication. The Bid Security provided under ITB 19 shall also be suitably extended. A Bidder may refuse the request without forfeiting its Bid security or causing to be executed its Bid Securing Declaration. A Bidder agreeing to the request will not be required

		nor permitted to modify its Bid, but will be required to extend the validity of its Bid Security or Bid Securing Declaration for the period of the extension, and in compliance with ITB 19 in all respects.
	18.3	If the award is delayed by a period exceeding sixty (60) days beyond the expiry of the initial Bid validity period, the contract price may be adjusted by a factor specified in the request for extension. However, the Bid evaluation shall be based on the already quoted Bid Price without taking into consideration on the above correction.
s. Bid Security or Bid Securing Declaration	19.1	Pursuant to ITB 12, unless otherwise specified in the BDS, the Bidder shall furnish as part of its Bid, a Bid Security in form of fixed amount not exceeding five percent of the estimated value of procurement determined by the Procurement Directorate and in the amount and currency specified in the BDS or Bid Securing Declaration as specified in the BDS in the format provided in Section - G (Standard Forms).
	19.2	The Bid Security or Bid Securing Declaration is required to protect the Procurement Directorate against the risk of Bidder's conduct which would warrant the security's forfeiture, pursuant to ITB 19.9.
	19.3	The Bid Security shall be denominated in the local currency or in another freely convertible currency, and it shall be in the form specified in the BDS which shall be in any of the following: (a) A bank guarantee, an irrevocable letter of credit issued by a Scheduled bank in the form provided in the Bidding Documents or another form acceptable to the Procurement Directorate and valid for twenty-eight (28) days beyond the end of the validity of the Bid. This shall also apply if the period for Bid Validity is extended. In either case, the form must include the complete name of the Bidder; (b) A Call deposit receipt or cash deposit receipt (CDR) (c) Another security if indicated in the BDS
	19.4	The Bid Security or Bid Securing Declaration shall be in accordance with the Form of the Bid Security or Bid Securing Declaration included in Section - G (Standard Forms) or another form approved by the Procurement Directorate prior to the Bid submission.
	19.5	The Bid Security shall be payable promptly upon written demand by the Procurement Directorate in case any of the conditions listed in ITB 19.9 are invoked.
	19.6	Any Bid not accompanied by a Bid Security or Bid Securing Declaration in accordance with ITB 19.1 or 19.3 shall be rejected by the Procurement Directorate as non-responsive, pursuant to ITB 29.
	19.7	Unsuccessful Bidders' Bid Security will be discharged or returned as promptly as possible, however in no case later than thirty (30) days after the expiration of the period of Bid Validity prescribed by the Procurement Directorate pursuant to ITB-18. The Procurement Directorate shall make no claim to the amount of the Bid Security, and shall promptly return the Bid Security document, after whichever of the following that occurs earliest: (a) the expiry of the Bid Security; (b) the entry into force of a procurement contract and the provision of a performance security (or guarantee), for the performance of the contract if such a security (or guarantee), is required by the Biding

		<p>documents;</p> <p>(c) the rejection by the Procurement Directorate of all Bids;</p> <p>(d) the withdrawal of the Bid prior to the deadline for the submission of Bids, unless the Bidding documents stipulate that no such withdrawal is permitted.</p>
	19.8	The successful Bidder's Bid Security will be discharged upon the Bidder signing the contract pursuant to ITB 42, or furnishing the performance security (or guarantee), pursuant to ITB 43.
	19.9	<p>The Bid Security may be forfeited or the Bid Securing Declaration executed:</p> <p>(a) if a Bidder:</p> <p>(1) Withdraws its Bid during the period of Bid Validity as specified by the Procurement Directorate, and referred by the bidder on the Form of Bid except as provided for in ITB 18.2; or</p> <p>(2) does not accept the correction of errors pursuant to ITB 31.3; or</p> <p>(b) In the case of a successful Bidder, if the Bidder fails:</p> <p>(1) To sign the contract in accordance with ITB 42; or</p> <p>(2) To furnish performance security (or guarantee) in accordance with ITB 43.</p>
t. Alternative Bids by Bidders	20.1	Bidders shall submit offers that comply with the requirements of the Bidding Documents, including the basic Bidder's technical design as indicated in the specifications and Schedule of Requirements. Alternatives will not be considered, unless specifically allowed for in the BDS. If so allowed, ITB 20.2 shall prevail.
	20.2	When alternative schedule for delivery of goods is explicitly invited, a statement of that effect will be included in the BDS as will the method for evaluating different schedule for delivery of goods.
	20.3	If so allowed in the BDS, Bidders wishing to offer technical alternatives to the requirements of the Bidding Documents must also submit a Bid that complies with the requirements of the Bidding Documents, including the basic technical design as indicated in the specifications. In addition to submitting the basic Bid, the Bidder shall provide all information necessary for a complete evaluation of the alternative by the Procurement Directorate, including technical specifications, breakdown of prices, and other relevant details. Only the technical alternatives, if any, of the Most Advantageous Bidder conforming to the basic technical requirements (without altering the bid price) shall be considered by the Procurement Directorate.
u. Bid Security Validity	21.1	Bid security submitted shall be valid for a period specified in the BDS
v. Format and Signing of Bid	22.1	The Bidder shall prepare an original and the 2 x copies of the Bid as indicated in the BDS, clearly marking each "ORIGINAL" and "COPY," as appropriate. In the event of any discrepancy between them, the original shall prevail:
	22.2	The original and the copy or copies of the Bid shall be typed or written in indelible ink and shall be signed by the Bidder or a person or persons duly authorized to sign on behalf of the Bidder. This authorization shall consist of a written confirmation as specified in the BDS and shall be attached to the Bid. The name and position held by each person signing the

		authorization must be typed or printed below the signature. All pages of the Bid, except for un-amended printed literature, shall be initialed by the person or persons signing the Bid.
	22.3	Any interlineations, erasures, or overwriting shall be valid only if they are signed by the person or persons signing the Bidder.

SUBMISSION OF BIDS

w. Sealing and Marking of Bids	23.1	In case of Single Stage Two Envelope Procedure, The Bid shall comprise two envelopes submitted simultaneously, one called the Technical Proposal and the other Financial Proposal. Both envelopes to be enclosed together in an outer single envelope called the Bid. Each Bidder shall submit his bid as under: (a) Bidder shall submit his TECHNICAL PROPOSAL and FINANCIAL PROPOSAL in separate inner envelopes and enclosed in a single outer envelope. (b) ORIGINAL and each copy of the Bid shall be separately sealed and put in separate envelopes and marked as such. (c) The envelopes containing the ORIGINAL and copies will be put in one sealed envelope and addressed / identified as given in Sub- Clause 22.2.
	23.2	The inner and outer envelopes shall: (a) Be addressed to the Gate no 5 Heavy Industries Taxila at the address provided in the Bidding Data; (b) Bear the name and identification number of the contract as defined in the Bidding Data; and provide a warning not to open before the time and date for bid opening, as specified in the Bidding Data pursuant to ITB 24.1. (c) In addition to the identification required in Sub- Clause 22.2 hereof, the inner envelope shall indicate the name and address of the bidder to enable the bid to be returned unopened in case it is declared "late" pursuant to Clause IB.25
	23.3	If all envelopes are not sealed and marked as required by ITB 23.2, ITB 23.3 and ITB 23.4 or incorrectly marked, the Procurement Directorate will assume no responsibility for the misplacement or premature opening of Bid.
x. Deadline for Submission of Bids	24.1	Bids shall be received by the Procurement Directorate no later than the date and time specified in the BDS.
	24.2	The Procurement Directorate may, in exceptional circumstances and at its discretion, extend the deadline for the submission of Bids by amending the Bidding Documents in accordance with ITB 10, in which case all rights and obligations of the Procurement Directorate and Bidders previously subject to the deadline will thereafter be subject to the new deadline.
y. Late Bids	25.1	The Procurement Directorate shall not consider for evaluation any Bid that arrives after the deadline for submission of Bids, in accordance with ITB 24.
	25.2	Any Bid received by the Procurement Directorate after the deadline for submission of Bids shall be declared late, recorded, rejected and returned unopened to the Bidder.
z. Modification,	26.1	A Bidder may modify or substitute or withdraw its Bid after it has been

Substitution and Withdrawal of Bids		submitted, provided that written notice of the modification, substitution or withdrawal of the Bid, is received by the Procurement Directorate prior to the deadline for submission of Bids.
	26.2	The Bidder modification, substitution or withdrawal notice shall be prepared, sealed, marked, and dispatched with the outer and inner envelopes additionally marked "MODIFICATION" "SUBSTITUTION" or "WITHDRAWAL" as appropriate. The notice may also be sent postmarked no later than the deadline for submission of Bids.
	26.3	Bids may only be modified by withdrawal of the original Bids and submission of a replacement Bids. Modifications submitted in any other way shall not be taken into account in the evaluation of bids.
	26.4	No Bids may be withdrawn, replaced or modified in the interval between the deadline for submission of Bids and the expiration of the period of Bid validity specified by the Bidder on the Form of Bid. Withdrawal of a Bid during this interval shall result in the Bidder forfeiture of its Bid security or execution of the Bid Securing Declaration

OPENING AND EVALUATION OF BIDS

aa. Opening of Bids	27.1	The Procurement Directorate will open all Bids, in public, in the presence of Bidders' or their representatives who choose to attend, and other parties with a legitimate interest in the Bid proceedings at the place, on the date and at the time, specified in the BDS. The Bidders' representatives present shall sign a Index Sheet as proof of their attendance.
	27.2	First, envelopes marked "WITHDRAWAL" shall be opened and read out and the envelope with the corresponding bid shall not be opened, but returned to the Bidder. No bid withdrawal shall be permitted unless the corresponding Withdrawal Notice contains a valid authorization to request the withdrawal and is read out at bid opening.
	27.3	Second, outer envelopes marked "SUBSTITUTION" shall be opened. The inner envelopes containing the Substitution Bid shall be exchanged for the corresponding Original Bid being substituted, which is to be returned to the Bidder unopened. No envelope shall be substituted unless the corresponding Substitution Notice contains a valid authorization to request the substitution and is read out and recorded at bid opening.
	27.4	Next, outer envelopes marked "MODIFICATION" shall be opened. No Technical Proposal and/or Financial Proposal shall be modified unless the corresponding Modification Notice contains a valid authorization to request the modification and is read out and recorded at the opening of the Bids. Any Modification shall be read out along with the Original Bid except in case of Single Stage Two Envelope Procedure where only the Technical Proposal, both Original as well as Modification, are to be opened, read out, and recorded at the opening. Financial Proposal, both Original and Modification, will remain unopened till the prescribed financial bid opening date.
	27.5	In case of Single Stage Two Envelope Procedure, the Board of Officers will open the Technical Proposals in public at the address, date and time specified in the BDS in the presence of Bidders' designated representatives who choose to attend and other parties with a legitimate interest in the Bid proceedings. The Financial Proposals will remain

		unopened and will be held in custody of the Procurement Directorate until the specified time of their opening.
	27.6	The envelopes holding the Technical Proposals shall be opened one at a time, and the following read out and recorded: (a) the name of the Bidder; (b) whether there is a modification or substitution; (c) the presence of a Bid Security, if required; and (d) Any other details as the Procurement Directorate may consider appropriate.
	27.7	Bids not opened and not read out at the Bid opening shall not be considered further for evaluation, irrespective of the circumstances. In particular, any discount offered by a Bidder which is not read out at Bid opening shall not be considered further.
	27.8	Bidders are advised to send in a representative with the knowledge of the content of the Bid who shall verify the information read out from the submitted documents. Failure to send a representative or to point out any un-read information by the sent Bidder's representative shall indemnify the Procurement Directorate against any claim or failure to read out the correct information contained in the Bidder's Bid.
	27.9	No Bid will be rejected at the time of Bid opening except for late Bids which will be returned unopened to the Bidder, pursuant to ITB 25.
	27.10	The Procurement Directorate shall prepare minutes of the Bid opening. The record of the Bid opening shall include, as a minimum: the name of the Bidder and whether or not there is a withdrawal, substitution or modification, the Bid price if applicable, including any discounts and alternative offers and the presence or absence of a Bid Security or Bid Securing Declaration.
	27.11	The Bidders' representatives who are present shall be requested to sign on the attendance sheet. The omission of a Bidder's signature on the record shall not invalidate the contents and affect the record. A copy of the record shall be distributed to all the Bidders.
	27.12	A copy of the minutes of the Bid opening shall be furnished to individual Bidders upon request.
	27.13	In case of Single Stage Two Envelop Bidding Procedure, after the evaluation and approval of technical proposal the Procurement Directorate, shall at a time within the bid validity period, publically open the financial proposals of the technically accepted bids only. The financial proposal of bids found technically non-responsive shall be returned unopened to the respective bidders subject to redress of the grievances from all tiers of grievances.
bb. Confidentiality	28.1	Information relating to the examination, clarification, evaluation and comparison of Bids and recommendation of contract award shall not be disclosed to Bidders or any other persons not officially concerned with such process until the time of the announcement of the respective evaluation report.
	28.2	Any effort by a Bidder to influence the Procurement Directorate processing of Bids or award decisions may result in the rejection of its Bid.
	28.3	Notwithstanding ITB 28.2 from the time of Bid opening to the time of contract award, if any Bidder wishes to contact the Procurement Directorate on any matter related to the Bidding process, it should do so in

		writing or in electronic forms that provides record of the content of communication.
cc. Clarification of Bids	29.1	To assist in the examination, evaluation and comparison of Bids of the Bidders, the Procurement Directorate may, ask any Bidder for a clarification of its Bid including breakdown of prices. Any clarification submitted by a Bidder that is not in response to a request by the Procurement Directorate shall not be considered.
	29.2	The request for clarification and the response shall be in writing or in electronic forms that provide record of the content of communication. In case of Single Stage Two Envelope Procedure, no change in the prices or substance of the Bid shall be sought, offered, or permitted, whereas in case of Single Stage One Envelope Procedure, only the correction of arithmetic errors discovered by the Procurement Directorate in the evaluation of Bids should be sought in accordance with ITB 31.
	29.3	The alteration or modification in the bid which in any affect the following parameters will be considered as a change in the substance of a bid: (a) Evaluation & qualification criteria; (b) Required scope of work or specifications; (c) All securities requirements; (d) Tax requirements; (e) Terms and conditions of bidding documents. (f) Change in the ranking of the bidder
	29.4	From the time of Bid opening to the time of Contract award if any Bidder wishes to contact the Procurement Directorate on any matter related to the Bid it should do so in writing or in electronic forms that provide record of the content of communication.
dd. Preliminary Examination of Bids	30.1	Prior to the detailed evaluation of Bids, the Procurement Directorate will determine whether each Bid: (a) Meets the eligibility criteria defined in ITB 3 and ITB 4; (b) Has been prepared as per the format and contents defined by the Procurement Directorate in the Bidding Documents; (c) Has been properly signed; (d) Is accompanied by the required securities; and (e) Is substantially responsive to the requirements of the Bidding Documents. The Procurement Directorate's determination of a Bid's responsiveness will be based on the contents of the Bid itself.
	30.2	A substantially responsive Bid is one which conforms to all the terms, conditions, and specifications of the Bidding Documents, without material deviation or reservation. A material deviation or reservation is one that: - (a) Affects in any substantial way the scope, quality, or performance of the Services; (b) Limits in any substantial way, inconsistent with the Bidding Documents, the Procurement Directorate's rights or the Bidders obligations under the Contract; or (c) If rectified, would affect unfairly the competitive position of other Bidders presenting substantially responsive Bids. (d) Failure to sign the bid form and price schedules by the authorized

	<p>person or persons;</p> <p>(e) Failure to satisfy eligibility requirements;</p> <p>(f) Failure to submit a bid security as specified in the bidding documents;</p> <p>(g) Failure to satisfy the bid validity period;</p> <p>(h) Inability to meet the critical delivery schedule or work schedule clearly specified in the bidding documents, where such schedule is a crucial condition with which bidders must comply;</p> <p>(i) Failure to comply with minimum experience criteria as specified in the bidding documents;</p> <p>(j) Conditional Bids such as conditions in a bid which limit the bidder's responsibility to accept an award;</p> <p>(k) Stipulating price adjustment when fixed price Bids were invited;</p> <p>(l) Subcontracting in a substantially different amount or manner than that permitted;</p> <p>(m) Failure to submit major supporting documents required by the bidding documents to determine substantial responsiveness of a bid</p>
30.3	<p>The Procurement Directorate will confirm that the documents and information specified under ITB 12, 13 and 14 have been provided in the Bid. If any of these documents or information is missing, or is not provided in accordance with the Instructions to Bidders, the Bid shall be rejected.</p>
30.4	<p>The Procurement Directorate may waive off any minor informality, nonconformity, or irregularity in a Bid which does not constitute a material deviation, provided such waiver does not prejudice or affect the relative ranking of any Bidder.</p> <p>Explanation: A minor informality, non-conformity or irregularity is one that is merely a matter of form and not of substance. It also pertains to some immaterial defect in a Bid or variation of a bid from the exact requirements of the invitation that can be corrected or waived without being prejudicial to other bidders. The defect or variation is immaterial when the effect on quantity, quality, or delivery is negligible when contrasted with the total cost or scope of the supplies or services being acquired. The Procurement Directorate either shall give the bidder an opportunity to cure any deficiency resulting from a minor informality or irregularity in a bid or waive the deficiency, whichever is advantageous to the Procurement Directorate. Examples of minor informalities or irregularities include failure of a bidder to –</p> <p>(a) Submit the number of copies of signed bids required by the invitation;</p> <p>(b) Furnish required information concerning the number of its employees;</p> <p>(c) The firm submitting a bid has formally adopted or authorized, before the date set for opening of bids, the execution of documents by typewritten, printed, or stamped signature and submits evidence of such authorization and the bid carries such a signature.</p>
30.5	<p>Provided that a Technical Bid is substantially responsive, the Procurement Directorate may request the Bidder to submit the necessary information or documentation, within a reasonable period of time, to rectify nonmaterial nonconformities or omissions in the Technical Bid related to documentation requirements. Requesting information or documentation on such nonconformities shall not be related to any such aspect of the technical Proposal linked with the ranking of the bidders. Failure of the Bidder to comply with the request may result in the rejection of its Bid.</p>

	30.6	Provided that a Technical Bid is substantially responsive, the Procurement Directorate shall rectify quantifiable nonmaterial nonconformities or omissions related to the Financial Proposal. To this effect, the Bid Price shall be adjusted, for comparison purposes only, to reflect the price of the missing or nonconforming item or component.
	30.7	If a Bid is not substantially responsive, it will be rejected by the Procurement Directorate and may not subsequently be evaluated for complete technical responsiveness.
ee. Examination of Terms and Conditions; Technical Evaluation	31.1	The Procurement Directorate shall examine the Bid to confirm that all terms and conditions specified in the GCC and the SCC have been accepted by the Bidder without any material deviation or reservation.
	31.2	The Procurement Directorate shall evaluate the technical aspects of the Bid submitted in accordance with ITB 23, to confirm that all requirements specified in Section E – Schedule of Requirements, Technical Specifications of the Bidding Documents have been met without material deviation or reservation.
	31.3	If after the examination of the terms and conditions and the technical evaluation, the Procurement Directorate determines that the Bid is not substantially responsive in accordance with ITB 31, it shall reject the Bid.
ff. Correction of Errors	32.1	Bids determined to be substantially responsive will be checked for any arithmetic errors. Errors will be corrected as follows: - (a) if there is a discrepancy between unit prices and the total price that is obtained by multiplying the unit price and quantity, the unit price shall prevail, and the total price shall be corrected, unless in the opinion of the Procurement Directorate there is an obvious misplacement of the decimal point in the unit price, in which the total price as quoted shall govern and the unit price shall be corrected; (b) if there is an error in a total corresponding to the addition or subtraction of sub-totals, the sub-totals shall prevail and the total shall be corrected; and (c) Where there is a discrepancy between the amounts in figures and in words, the amount in words will govern. (d) Where there is discrepancy between grand total of price schedule and amount mentioned on the Form of Bid, the amount referred in Price Schedule shall be treated as correct subject to elimination of other errors.
	32.2	The amount stated in the Bid will, be adjusted by the Procurement Directorate in accordance with the above procedure for the correction of errors and, with, the concurrence of the Bidder, shall be considered as binding upon the Bidder. If the Bidder does not accept the corrected amount, its Bid will then be rejected, and the Bid Security may be forfeited or the Bid Securing Declaration may be executed in accordance with ITB 19.9.
gg. Conversion to Single Currency	33.1	To facilitate evaluation and comparison, the Procurement Directorate will convert all Bid prices expressed in the amounts in various currencies in which the Bid prices are payable. For the purposes of comparison of bids quoted in different currencies, the price shall be converted into a single currency specified in the bidding documents. The rate of exchange shall

		be the selling rate, prevailing on the date of opening of (financial part of) bids specified in the bidding documents, as notified by the State Bank of Pakistan on that day.
	33.2	The currency selected for converting Bid prices to a common base for the purpose of evaluation, along with the source and date of the exchange rate, are specified in the BDS .
hh.	Evaluation of Bids	
	34.1	The Concerned Project and Procurement Directorate shall evaluate and compare only the Bids determined to be substantially responsive, pursuant to ITB 30 .
	34.2	In evaluating the Technical Proposal of each Bid, the Concerned Project shall use the criteria and methodologies listed in the BDS and in terms of Statement of Requirements and Technical Specifications. No other evaluation criteria or methodologies shall be permitted.
	34.3	The Procurement Directorate's evaluation of a Bid will take into account: (a) In the case of therapeutic goods manufactured in Pakistan or therapeutic goods of foreign origin already imported in Pakistan, Income Tax, General Sales Tax and other similar/applicable taxes, which will be payable on the goods if a contract is awarded to the Bidder; (b) In the case of therapeutic goods of foreign origin offered from abroad, customs duties and other similar import taxes which will be payable on the goods if the contract is awarded to the Bidder; and
	34.4	The comparison shall be between the EXW price of the therapeutic goods offered from within Pakistan, such price to include all costs, as well as duties and taxes paid or payable on components and raw material incorporated or to be incorporated in the therapeutic goods, and named port of destination, border point, or named place of destination} in accordance with applicable INCOTERM in the price of the therapeutic goods offered from outside Pakistan.
	34.5	In evaluating the Bidders, the evaluation committee will, in addition to the Bid price quoted in accordance with ITB 16.1, take account of one or more of the following factors as specified in the BDS , and quantified in ITB 33.5: (a) Cost of inland transportation, insurance, and other costs within the Pakistan incidental to delivery of the therapeutic goods to their final destination. (b) Delivery schedule offered in the Bid; (c) Deviations in payment schedule from that specified in the Special
		Conditions of Contract; (d) The cost of components, mandatory spare parts, and service; (e) The availability (in Pakistan) of spare parts and after-sales services for the equipment offered in the Bid; (f) The projected operating and maintenance costs during the life of the equipment; (g) The performance and productivity of the equipment offered; and/or (h) Other specific criteria indicated in the TBS and/or in the Technical Specifications.
	34.6	For factors retained in BDS , pursuant to ITB 34.4 one or more of the following quantification methods will be applied, as detailed in the BDS : (a) Inland transportation from EXW/port of entry/border point, Insurance and incidentals.

(b) Inland transportation, insurance, and other incidental costs for delivery of the therapeutic goods from EXW/port of entry/border point to Project Site named in the **BDS** will be computed for each Bid by the PA on the basis of published tariffs by the rail or road transport agencies, insurance companies, and/or other appropriate sources. To facilitate such computation, Bidder shall furnish in its Bid the estimated dimensions and shipping weight and the approximate EXW or as per applicable INCOTERM value of each package. The above cost will be added by the Procurement Directorate to EXW or as per applicable INCOTERM price.

(c) *Delivery schedule.*

i) The Procurement Directorate requires that the therapeutic goods under the Invitation for Bids shall be delivered (shipped) at the time specified in the Schedule of Requirements. The estimated time of arrival of the therapeutic goods at the Project Site will be calculated for each Bid after allowing for reasonable international and inland transportation time. Treating the Bid resulting in such time of arrival as the base, a delivery “adjustment” will be calculated for other Bids by applying a percentage, specified in the **BDS**, of the EXW or as per applicable INCOTERM price for each week of delay beyond the base, and this will be added to the Bid price for evaluation. No credit shall be given to early delivery.

Or

ii) The therapeutic goods covered under this invitation are required to be delivered (shipped) within an acceptable range of weeks specified in the Schedule of Requirement. No credit will be given to earlier deliveries, and Bids offering delivery beyond this range will be treated as non-responsive. Within this acceptable range, an adjustment per week, as specified in the **BDS**, will be added for evaluation to the Bid price of Bids offering deliveries later than the earliest delivery period specified in the Schedule of Requirements.

Or

(iii) The therapeutic goods covered under this invitation are required to be delivered (shipped) in partial shipments, as specified in the Schedule of Requirements. Bids offering deliveries earlier or later than the specified deliveries will be adjusted in the evaluation by adding to the Bid price a factor equal to a percentage, specified in the **BDS**, of EXW or as per applicable INCOTERM price per week of variation from the specified delivery schedule.

(d) *Deviation in payment schedule.*

i) Bidders shall state their Bid price for the payment schedule outlined in the **SCC**. Bids will be evaluated on the basis of this base price. Bidders are, however, permitted to state an alternative payment schedule and indicate the reduction in Bid price they wish to offer for such alternative payment schedule. The Procurement Directorate may consider the alternative payment schedule offered by the selected Bidder.

Or

		<p>ii) The SCC stipulates the payment schedule offered by the Procurement Directorate. If a Bid deviates from the schedule and if such deviation is considered acceptable to the Procurement Directorate, the Bid will be evaluated by calculating interest earned for any earlier payments involved in the terms outlined in the Bid as compared with those stipulated in this invitation, at the rate per annum specified in the BDS.</p> <p>(e) <i>Specific additional criteria.</i> Other specific additional criteria to be considered in the evaluation and the evaluation method shall be detailed in the BDS and/or the Technical Specifications.</p>
	34.7	If these Bidding Documents allow Bidders to quote separate prices for different Lots, and the award to a single Bidder of multiple Lots, the methodology of evaluation to determine the lowest evaluated Lot combinations, including any discounts offered in the Form of Bid, is specified in the BDS .
ii. Domestic Preference	35.1	If the BDS so specifies, the Procurement Directorate will grant a margin of preference to certain goods in line with the rules, regulations, regulatory guides or instructions issued by the Authority from time to time.
jj. Determination of Most Advantageous Bid	36.1	In case where the Procurement Directorate adopts the Cost Based Evaluation Technique and, the Bid with the lowest evaluated price from amongst those which are eligible, compliant and substantially responsive shall be the Most Advantageous Bid.
	36.2	<p>The Procurement Directorate may adopt the Quality & Cost Based Selection Technique due to the following reasons:</p> <p>i. In addition to the mandatory requirements and mandatory technical specifications, requires parameters specified in Evaluation Criteria to be evaluated while determining the quality of the therapeutic goods:</p> <p>ii. In such cases, the Concerned Project may allocate certain weightage to these factors as a part of Evaluation Criteria, and may determine the ranking of the bidders on the basis of combined evaluation in accordance with provisions of Rule 2(1)(h) of PPR-2004.</p>
kk. Post-qualification of Bidder and/or Abnormally Low Financial Proposal	37.1	<p>Where the Bid price is considered to be abnormally low, the Concerned Project shall perform price analysis either during determination of Most Advantageous Bid. The following process shall apply:</p> <p>(a) The Concerned Project may reject a Bid if the Concerned Project has determined that the price in combination with other constituent elements of the Bid is abnormally low in relation to the subject matter of the procurement (i.e. scope of the procurement) and raises concerns as to the capability and capacity of the respective Bidder to perform that contract;</p> <p>(b) Before rejecting an abnormally low Bid the Concerned Project shall request the Bidder an explanation of the Bid or of those parts which it considers contribute to the Bid being abnormally low; take account of the evidence provided in response to a request in writing; and subsequently verify the Bid or parts of the Bid being abnormally low;</p> <p>(c) The decision of the Concerned Project to reject a Bid and reasons for the decision shall be recorded in the procurement proceedings and</p>

	<p>promptly communicated to the Bidder concerned;</p> <p>(d) The Concerned Project shall not incur any liability solely by rejecting abnormally Bid; and</p> <p>(e) An abnormally low Bid means, in the light of the Concerned Project estimate and of all the Bids submitted, the Bid appears to be abnormally low by not providing a margin for normal levels of profit.</p> <p>Guidance for Concerned Project:</p> <p>In order to identify the Abnormally Low Bid (ALB) following approaches can be considered to minimize the scope of subjectivity:</p> <p>(i) Comparing the bid price with the cost estimate;</p> <p>(ii) Comparing the bid price with the bids offered by other bidders submitting substantially responsive bids; and</p> <p>(iii) Comparing the bid price with prices paid in similar contracts in the recent past either government- or development partner-funded.</p>
37.2	The Concerned Project will determine to its satisfaction whether the Bidder that is selected as having submitted the most advantageous Bid is qualified to perform the contract satisfactorily, in accordance with the criteria listed in ITB 14.3.
37.3	The determination will take into account the Bidder's financial, technical, and production capabilities. It will be based upon an examination of the documentary evidence of the Bidder's qualifications submitted by the Bidder, pursuant to ITB 14.3, as well as such other information as the Concerned Project deems necessary and appropriate. Factors not included in these Bidding Documents shall not be used in the evaluation of the Bidders' qualifications.
37.4	<p>Procurement Directorate may seek "Certificate for Independent Price Determination" from the Bidder and the results of reference checks may be used in determining award of contract.</p> <p>Explanation: The Certificate shall be furnished by the bidder. The bidder shall certify that the price is determined keeping in view of all the essential aspects such as raw material, its processing, value addition, optimization of resources due to economy of scale, transportation, insurance and margin of profit etc.</p>
37.5	An affirmative determination will be a prerequisite for award of the contract to the Bidder. A negative determination will result in rejection of the Bidder's Bid, in which event the Concerned Project will proceed to the next ranked bidder to make a similar determination of that Bidder's capabilities to perform satisfactorily.

AWARD OF CONTRACT

<p>ll. Criteria of Award</p>	<p>38.1</p>	<p>Subject to ITB 37 and 39, the Procurement Directorate will award the Contract to the Bidder whose Bid has been determined to be substantially responsive to the Bidding Documents and who has been declared as Most Advantageous Bidder, provided that such Bidder has been determined to be:</p> <p>(a) Eligible in accordance with the provisions of ITB 3;</p> <p>(b) Is determined to be qualified to perform the Contract satisfactorily; and</p> <p>(c) Successful negotiations have been concluded, if any.</p>
<p>mm. Negotiations</p>	<p>39.1</p>	<p>Negotiations may be undertaken with the Most Advantageous Bid relating to the following areas:</p> <p>(a) A minor alteration to the technical details of the statement of requirements;</p> <p>(b) Reduction of quantities for budgetary reasons, where the reduction is in excess of any provided for in the Bidding documents;</p> <p>(c) A minor amendment to the special conditions of Contract;</p> <p>(d) Finalizing payment arrangements;</p> <p>(e) Delivery arrangements;</p> <p>(f) The methodology for provision of related services; or</p> <p>(g) Clarifying details that were not apparent or could not be finalized at the time of Bidding;</p>
	<p>39.2</p>	<p>Where negotiation fails to result into an agreement, the Procurement Directorate may invite the next ranked Bidder for negotiations. Where negotiations are commenced with the next ranked Bidder, the Procurement Directorate shall not reopen earlier negotiations.</p>
<p>nn. Procurement Directorate's Right to reject All Bids</p>	<p>40.1</p>	<p>Notwithstanding ITB 38, the Procurement Directorate reserves the right to reject all the bids, and to annul the Bidding process at any time prior to award of contract, without thereby incurring any liability to the affected Bidder or Bidders. However, the Authority (i.e. PPR) may call from the Procurement Directorate the justification of those grounds.</p>
	<p>40.2</p>	<p>Notice of the rejection of all Bids shall be given promptly to all Bidders that have submitted Bids.</p>
	<p>40.3</p>	<p>The Procurement Directorate shall upon request communicate to any Bidder the grounds for its rejection of its Bids, but is not required to justify those grounds.</p>
<p>oo. Procurement Directorate's Right to Vary Quantities at the Time of Award</p>	<p>41.1</p>	<p>The Procurement Directorate reserves the right at the time of contract award to increase or decrease the quantity of goods or related services originally specified in these Bidding Documents (schedule of requirements) provided this does not exceed by the percentage indicated in the BDS, without any change in unit price or other terms and conditions of the Bid and Bidding Documents.</p>
<p>pp. Notification of Award</p>	<p>42.1</p>	<p>Prior to the award of contract, the Procurement Directorate shall issue a Final Evaluation Report giving justification for acceptance or rejection of the bids.</p>
	<p>42.2</p>	<p>Where no grievance have been lodged, the Bidder whose Bid has been</p>

		accepted will be notified of the award by the Procurement Directorate prior to expiration of the Bid Validity period in writing or electronic forms that provide record of the content of communication. The Letter of Acceptance / contract signing will state the sum that the Procurement Directorate will pay the successful Bidder in consideration for the execution of the scope of works as prescribed by the Contract (hereinafter and in the Contract called the "Contract Price).
qq. Signing of Contract	43.1	Promptly after notification of award, Procurement Directorate shall send the successful Bidder the draft agreement, incorporating all terms and conditions as agreed by the parties to the contract.
	43.2	Immediately after the Redressal of grievance by the GRC, and after fulfillment of all condition's precedent of the Contract Form, the successful Bidder and the Procurement Directorate shall sign the contract.
	43.3	Where no formal signing of a contract is required, purchase order issued to the bidder shall be construed to be the contract.
rr. Performance Guarantee	44.1	After the receipt of the Letter of Acceptance / Contract, the successful Bidder, within the specified time, shall deliver to the Procurement Directorate a Performance Guarantee in the form stipulated in the BDS and SCC , denominated in the type and proportions of currencies in the Letter of Acceptance / contracts and in accordance with the Conditions of Contract.
	44.2	If the Performance Guarantee is provided by the successful Bidder and it shall be in the form specified in the BDS which shall be in any of the following: (a) Call deposit receipt or Cash deposit receipt or Bank Guarantee; (b) Irrevocable letter of credit issued by a Scheduled bank or in the case of an irrevocable letter of credit issued by a foreign bank, the letter shall be confirmed or authenticated by a Scheduled bank; (c) Bank guarantee confirmed by a reputable local bank or, in the case of a successful foreign Bidder, bonded by a foreign bank; or Any Performance Security (or guarantee) submitted shall be enforceable in Pakistan.
	44.3	Failure of the successful Bidder to comply with the requirement of ITB 44.1 shall constitute sufficient grounds for the annulment of the award and forfeiture of the Bid Security, in which event the Procurement Directorate may make the award to the next ranked Bidder or call for new Bids.
ss. Advance Payment	45.1	The advance payment will not be provided in normal circumstances. However, in case where international incoterms are involved, the same will be dealt with standard international practices and in the manner as prescribed in ITB 45.2.
	45.2	The Procurement Directorate will provide an Advance Payment as stipulated in the Conditions of Contract, subject to a maximum amount, as stated in the BDS . The Advance Payment request shall be accompanied by an Advance Payment Security (Guarantee) in the form provided during conclusion of contract. For the purpose of receiving the Advance Payment, the Bidder shall make and estimate of, and include in its Bid, the expenses that will be incurred in order to commence Delivery of Goods. These expenses will relate to the purchase of equipment, machinery,

		materials, and on the engagement of labor during the first month beginning with the date of the Procurement Directorate's "Notice to Commence" as specified in the SCC .
tt. Arbitrator	46.1	The Arbitrator shall be appointed by mutual consent of the both parties as per the provisions specified in the SCC.
uu. Corrupt & Fraudulent Practices	47.1	Procuring Agencies (including beneficiaries of Government funded projects and procurement) as well as Bidders/Suppliers/Contractors under Government financed contracts, observe the highest standard of ethics during the procurement and execution of such contracts, and will avoid to engage in any corrupt and fraudulent practices.

GRIEVANCE REDRESSAL & COMPLAINT REVIEW MECHANISM

vv. Constitution of Grievance Redressal	48.1	Technical Directorate, Heavy Industries Taxila shall constitute a Grievance Redressal Committee (GRC) comprising of odd number of person with proper power and authorization to address the complaint. The GRC shall not have any of the members of Procurement Evaluation Committee. The committee must have one subject specialist depending the nature of the procurement.
ww. GRC Procedure	49.1	Any party can file its written complaint against the eligibility parameters or any other terms and conditions prescribed in the bidding documents found contrary to provision of Procurement Regulatory Framework, and the same shall be addressed by the GRC well before the bid submission deadline.
	49.2	Any Bidder feeling aggrieved by any act of the Procurement Directorate after the submission of his bid may lodge a written complaint concerning his grievances not later than seven days of the announcement of technical evaluation report and five days after issuance of final evaluation report.
	49.3	In case, the complaint is filed against the technical evaluation report, the Procurement Directorate shall halt the procurement proceedings till final decision.
	49.4	In case, the complaint is filed after the issuance of the final evaluation report, the complainant cannot raise any objection on technical evaluation of the report: Provided that the complainant may raise the objection on any part of the final evaluation report in case where single stage one envelopes bidding procedure is adopted.
	49.5	The GRC, in both the cases shall investigate and decide upon the complaint within ten days of its receipt.
	49.6	Any bidder or the Procurement Directorate not satisfied with the decision of the GRC may file Appeal before the Appellate Committee of the Authority on prescribed format after depositing the Prescribed fee.
	49.7	The Committee, upon receipt of the Appeal against the decision of the GRC complete in all respect shall serve notices in writing upon all the parties to appeal.

	49.8	The committee shall call the record from the concerned Procurement Directorate or the GRC as the case may be, and the same shall be provided within prescribed time.
	49.9	The committee may after examination of the relevant record and hearing all the concerned parties, shall decide the complaint within fifteen (15) days of receipt of the Appeal.
	49.10	The decision of the Committee shall be in writing and shall be signed by the Head and each Member of the Committee. The decision of the committee shall be final.

MECHANISM OF BLACKLISTING

xx.Mechanism of Blacklisting	50.1	The Procurement Directorate shall bar for not more than the time prescribed in Rule-19 of the Public Procurement Rules, 2004, from participating in their respective procurement proceedings, bidder or contractor who either: (1) Involved in corrupt and fraudulent practices as defined in Rule-2 of Public Procurement Rules; (2) Fails to perform his contractual obligations; and (3) Fails to abide by the id securing declaration;
	50.2	The show cause notice shall contain: (a) precise allegation, against the bidder or contractor; (b) the maximum period for which the Procurement Directorate proposes to debar the bidder or contractor from participating in any public procurement of the Procurement Directorate; and (c) the statement, if needed, about the intention of the Procurement Directorate to make a request to the Authority for debarring the bidder or contractor from participating in public procurements of all the procuring agencies.
	50.3	The Procurement Directorate shall give minimum of seven days to the bidder or contractor for submission of written reply of the show cause notice
	50.4	In case, the bidder or contractor fails to submit written reply within the requisite time, the Procurement Directorate may issue notice for personal hearing to the bidder or contractor/ authorize representative of the bidder or contractor and the Procurement Directorate shall decide the matter on the basis of available record and personal hearing, if availed.
	50.5	In case the bidder or contractor submits written reply of the show cause notice, the Procurement Directorate may decide to file the matter or direct issuance of a notice to the bidder or contractor for personal hearing.
	50.6	The Procurement Directorate shall give minimum of seven days to the bidder or contractor for appearance before the specified officer of the Procurement Directorate for personal hearing. The specified officer shall decide the matter on the basis of the available record and personal hearing of the bidder or contractor, if availed
	50.7	The Procurement Directorate shall decide the matter within fifteen days from the date of personal hearing unless the personal hearing is adjourned to a next date and in such an eventuality, the period of personal hearing shall be reckoned from the last date of personal hearing.

	50.8	The Procurement Directorate shall communicate to the bidder or contractor the order of debarring the bidder or contractor from participating in any public procurement with a statement that the bidder or contractor may, within thirty days, prefer a representation against the order before the Authority.
	50.9	Such blacklisting or barring action shall be communicated by the Procurement Directorate to the Authority and respective bidder or bidders in the form of decision containing the grounds for such action. The same shall be publicized by the Authority after examining the record whether the procedure defined in blacklisting and debarment mechanism has been adhered to by the Procurement Directorate.
	50.10	The bidder may file the review petition before the Review Petition Committee Authority within thirty days of communication of such blacklisting or barring action after depositing the prescribed fee and in accordance with "Procedure of filing and disposal of review petition under Rule-19(3) Regulations, 2021". The Committee shall evaluate the case and decide within ninety days of filing of review petition.
	50.11	The committee shall serve a notice in writing upon all respondent of the review petition. The notices shall be accompanied by the copies of review petition and all attached documents of the review petition including the decision of the Procurement Directorate. The parties may file written statements along with essential documents in support of their contentions. The Committee may pass such order on the representation may deem fit.
	50.12	The Authority on the basis of decision made by the committee either may debar a bidder or contractor from participating in any public procurement process of all or some of the procuring agencies for such period as the deemed appropriate or acquit the bidder from the allegations. The decision of the Authority shall be final.

BID DATA SHEET
Bid Data Sheet (BDS)

1. The following specific data for the therapeutic goods to be procured shall complement, supplement, or amend the provisions in the Instructions to Bidders (ITBs). Whenever there is a conflict, the provisions herein shall prevail over those in ITBs.

BDS Clause Number	ITB Number	Amendments of, and Supplements to, Clauses in the Instruction to Bidders
Introduction		
a.	1.1	Procurement Directorate The subject of procurement is: Procurement of Medicine (HIT Hospital) Commencement date for delivery of Therapeutic Goods (DP): 01 Month
b.	2.1	Financial year for the operations of the Procurement Directorate: <i>2024/25</i> Name of Project HIT Hospital Name of financing institution: MoDP (Budget Dte HIT)
c.	3.1	Maximum number of members in the joint venture, consortium or association shall be: N/A.
d.	4.1	Ineligible Source country(s) are India and Israel
e.	4.5	Demonstration of authorization by manufacturer: DRAP
Bidding Documents		
f.	7.1	The bid shall be quoted complete Package (Package Deal), The bidder should not bid a single item or single quantity.
g.	9.1	The address for clarification of Bidding Documents is Assistant Director Procurement, Procurement Directorate Heavy Industries Taxila – Email address dirprocde@hit.gov.pk Fax No. 0519315029
	9.5	Pre-bid meeting will not be held
Preparation of Bids		
h.	11.1	The Language of all correspondences and documents related to the Bid is: <i>English</i>
i.	12.1(b)	Advance samples to be provided to HIT hospital after technical opening within one x week
j.	12.1 (k)	In addition to the documents stated in ITB 12 , the following documents must be included with the Bid (1) Photocopy of Registration / Pre-qualification / Indexation letter issued by the HIT. (2) Attested copy of Registration certificate issued by Sales Tax Department, copy of NTN certificate. (3) Audit report of last 3 x FYs. (4) Attested Bank Statement for last one year. (5) Attested copy of CNIC of MD. (6) Trade link between firm and OEM (in case of distributor / agent). (7) Certificate on a judicial paper worth Rs 100 duly attested by Oath Commissioner that firm is neither defaulter nor blacklisted by any Govt Organization directly or indirectly. (8) Detail of list of contracts last 3 x FYs / works experience with Govt / Semi Govt Organization. (9) Copy of registration letter with Govt / Semi Govt organization (if any). (10) Postal order of Rs 2,000/- should be enclosed with Technical Bid in favour of Director Procurement HIT, Taxila

		(11) DRAP valid license.																																				
k.	12.2 (d)	N/A																																				
l.	13.3 (c)	The Bidder is required to include with its Bid, documentation from the manufacturer of the therapeutic goods, that it has been duly authorized to deliver, in Pakistan, the therapeutic goods indicated in its Bid. "OEM Authorization letter / Trade link with OEM / Agency Agreement"																																				
m.	14.3 (b)	The qualification criteria required from Bidders in ITB 14.3(b) is modified as follows: <table border="1"> <thead> <tr> <th>Ser</th> <th>Description</th> <th>Maximum Points</th> <th>Passing Marks</th> </tr> </thead> <tbody> <tr> <td colspan="4">a. Performance Evaluation</td> </tr> <tr> <td>(1)</td> <td>Financial Soundness</td> <td>15</td> <td>7.5</td> </tr> <tr> <td>(2)</td> <td>Past Experience / Record</td> <td>15</td> <td>7.5</td> </tr> <tr> <td>(3)</td> <td>Past Performance</td> <td>20</td> <td>10</td> </tr> <tr> <td colspan="2" style="text-align: right;">Sub Total</td> <td>50</td> <td>25</td> </tr> <tr> <td colspan="4">b. Technical Evaluation</td> </tr> <tr> <td>(1)</td> <td>Technical Evaluation Parameters</td> <td>50</td> <td>35</td> </tr> <tr> <td colspan="2" style="text-align: right;">Total</td> <td>100</td> <td>60</td> </tr> </tbody> </table>	Ser	Description	Maximum Points	Passing Marks	a. Performance Evaluation				(1)	Financial Soundness	15	7.5	(2)	Past Experience / Record	15	7.5	(3)	Past Performance	20	10	Sub Total		50	25	b. Technical Evaluation				(1)	Technical Evaluation Parameters	50	35	Total		100	60
Ser	Description	Maximum Points	Passing Marks																																			
a. Performance Evaluation																																						
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(1)	Technical Evaluation Parameters	50	35																																			
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n.	16.7 (a) (iii), (iv) (optional)	For therapeutic goods manufactured from within Pakistan the price quoted shall be consignee address including all transportation charges etc.																																				
o.	16.8	The price shall be fixed.																																				
p.	17.1 (a)	For therapeutic goods and related services originating in Pakistan the currency of the Bid shall be <i>Pakistani Rupees</i> ;																																				
q.	17.2	N/A																																				
r.	18.1	The Bid Validity period shall be 180 days .																																				
s.	19.1	<p>(1) The Bid Security / EM will be in the form of a CDR / Bank Draft / Pay Order in the favour of Director Procurement, Heavy Industries Taxila, for the amount in Pakistan Rupees. Bid Security in the form of crossed cheque / cheques shall be liable for rejection. Rates are as under: -</p> <p>(a) <u>Prequalified/Registered/ Indexed Firms/ Pre-qualified Firms.</u> 2% of the quoted value subject to the maximum ceiling of Rs. 0.5 million.</p> <p>(b) <u>Pre-qualified/Registered/ Pre-qualified but Un-indexed Firms.</u> 3% of the quoted value subject to the maximum ceiling of Rs. 0.75 million.</p> <p>(c) <u>Unregistered/Un-indexed Firms.</u> 5% of the quoted value subject to the maximum ceiling of Rs. 1 million.</p> <p>(2) The actual bid security / Earnest Money showing the amount will be placed inside the envelope of the "Financial Proposal".</p> <p>(3) Copy of the muted bid security / Earnest Money envelope (hiding the actual amount) will be placed inside the "Technical Proposal" envelope.</p>																																				
t.	19.3	The Bid Security shall be in the form of: <i>CDR or Bank Guarantee</i>																																				
u.	19.3 (c)	N/A																																				
v.	20.1	N/A																																				

w.	21.1	The Bid Security shall be Valid for <i>One Year</i>
x.	22.1	The number of copies of the Bid to be completed and returned shall be <i>02 x copies (1 x Original and 1 x Copy)</i> .
y.	22.2	Written confirmation of authorization <i>CEO / MD</i>
Submission of Bids		
z.	23.2 (a)	Submission of Bid: - Lieutenant Colonel Muhammad Shuja Chaudhry Assistant Director Procurement Gate No 5 (Tender Document) Heavy Industries Taxila, Taxila Cantt Tel: (051) 9315333 Ext 63215 Fax: (051) 9315029 E-Mail: dirproccte@hit.gov.pk ITB title and No: Bid No. 5171/IT-3035/2024-25/Medicines/FOR/Med/Proc dated 25 October 2024 Time and date for submission: 21 November 2024 before 1030 hours
aa.	23.2 (b)	Procurement of Medicines (HIT Hospital) ITB title and No: Bid No. 5171/IT-3035/2024-25/Medicines/FOR/Med/Proc dated 25 October 2024 Time and date for submission: 21 November 2024 before 1030 hours
bb.	24.1	The deadline for Bid submission is (1) Day: <i>Tuesday</i> (2) Date: 21 November 2024 (3) Time: 1030 Hours
Opening and Evaluation of Bids		
cc.	27.1	The Bid opening shall take place at: Conference Room of Procurement Directorate Heavy Industries Taxila Cantt Floor/Room No: <i>2nd Floor</i> City/Town: <i>Taxila</i> Country: <i>Pakistan</i> Day: <i>Tuesday</i> Date: 21 November 2024 Time: 1030 Hours
dd.	34.1 (h)	N/A
ee.	34.2 (b)	Delivery schedule. 01 Month Option (i) Option (ii) Option (iii)
ff.	34.5	This tender is on package deal. Lowest bidder will be awarded the contract against packages.
gg.	35.1	N/A
hh.	36	Evaluation Techniques Least Cost Based Selection (LCBS)

		After meeting the requirements of eligibility, qualification and substantial responsiveness, the bid in compliance with all the mandatory (technical) specifications/requirements and/or requisite quality threshold (if any), and having lowest evaluated cost (or financial proposal) shall be considered highest ranked bid.
Award of Contract		
ii.	41.1	Percentage for quantity increase or decrease is 15%.
jj.	44.1	The Performance guarantee shall be 5% of <i>the Contract Price</i>
kk.	44.2	The Performance guarantee shall be in the form of <i>CDR or Performance Bank Guarantee</i>
ll.	45.1	The Advance Payment if essential shall be limited to “ N/A ”
mm.	45.2	Maximum amount of Advance payment shall be “ N/A ”
nn.	46.1	Arbitrator shall be appointed by mutual consent of the both parties.
Review of GRC Decisions		
oo.	50.1	<u>Lodging Complaints</u> Brigadier Muhammad Khalid Hayat Director Procurement Heavy Industries Taxila, Taxila Cantt Tel: (051) 9315333 Ext 63211 Fax: (051) 9315029 E-Mail: dirproccte@hit.gov.pk
pp.		The Joint Appellant Committee (JAC) to submit a copy of grievance: Ministry of Defence Production (MoDP) Tel : 051-9270989 The Address of PPRA to submit a copy of grievance: Grievance Redressal Appellate Committee, Public Procurement Regulatory Authority 1 st Floor, G-5/2, Islamabad, Pakistan Tel: +92-51-9202254

ELIGIBLE COUNTRIES

2. All the bidders are allowed to participate in the subject procurement without regard to nationality, except bidders of some nationality, prohibited in accordance with policy of the Federal Government.

3. Following countries are ineligible to participate in the procurement process:

- a. India
- b. Israel

4. Ministry of Interior, Government of Pakistan has notified List of Business-Friendly Countries (BVL), information can be accessed through following link:

<http://www.dgip.gov.pk/Files/Visa%20Categories.aspx#L>

SCHEDULE OF REQUIREMENTS

1. Procurement will be as per packages mentioned below. Firms having technically qualified will be selected on package wise least cost method: -

Ser	Packages	No of Items	Annexures
a.	Package 1: Anesthesia Drugs	10	Appendix I to Section E
b.	Package 2: Anti – Asthmatic	13	Appendix II to Section E
c.	Package 3: Anti – Biotic	61	Appendix III to Section E
d.	Package 4: Anti – Diabetic	21	Appendix IV to Section E
e.	Package 5: Anti – Histamine	14	Appendix V to Section E
f.	Package 6: Cardiovascular	41	Appendix VI to Section E
g.	Package 7: Immune Sera & Immunoglobulin	5	Appendix VII to Section E
h.	Package 8: IV Fluids	8	Appendix VIII to Section E
i.	Package 9: Miscellaneous	28	Appendix IX to Section E
j.	Package 10: Moisturizing Agent	4	Appendix X to Section E
k.	Package 11: Multivitamins, Supplements & Minerals	22	Appendix XI to Section E
l.	Package 12: Analgesic	31	Appendix XII to Section E
m.	Package 13: Gastroenterology	22	Appendix XIII to Section E
n.	Package 14: Anti-Psychotic	19	Appendix XIV to Section E
Total		299	

- Note:
- Failure to complete the intended supplies even after 60 days will result in the forfeiture of performance security and the company/firm/supplier may be blacklisted.
 - The disposable should be delivered in packing with clear marking **“for HIT Hosp use only”**.
 - Short Dated Stocks (expiry of within 6 months) would be replaced within one month of being notified.

TECHNICAL QUOTATION FORMAT

Ser	IT No	Nomenclature	A/U	Packing	Coy /Make	Remarks
			As per required Product List			

EVALUATION CRITERIA

1. Evaluation will be carried out on **package wise Least Cost method** as under: -

Ser	Evaluation Criteria	Total Marks	Minimum Passing Marks
a.	Performance Evaluation (Procurement Directorate)	50	25
b.	Technical Evaluation (User Factory)	50	35
Total		100	60
c.	Price Evaluation	100	1 st lowest

2. **Performance Evaluation (Procurement Directorate) – 50 Marks**

- a. **Financial Soundness.** Following parameters will be used:-

Ser	Description	Max Marks	Calculation Procedure	Remarks
(1)	Annual turnover of last 3 x Financial Years	5	Marks will be calculated as per the formula: - $\text{Score} = \frac{(Y1+Y2+Y3)}{(3 * X)} * 5$ <ul style="list-style-type: none"> Y1, Y2, Y3 respective annual turnovers of last three years X= Last purchased rate / estimated value of the quoted items available with HIT. 	<ul style="list-style-type: none"> Third Party generated verifiable audit reports for last three financial years to be provided for minimum of upto Rs 5 Mn (in FOR Case) and US\$ (0.05) Mn in FOB case Else Income Tax returns for last 3 x financial years, fully verified by ITO of the circle.
(2)	Working Capital of last three years	10	Marks will be calculated as per the formula: - $\text{Score} = \frac{(Y1+Y2+Y3)}{(3 * X)} * 10$ <ul style="list-style-type: none"> Y1, Y2 and Y3 being respective working capitals of last three years. X= Last Purchase Rate / Estimated value of the quoted items Available with HIT. 	
(3)	Litigation history	-	One mark will be deducted for each litigation history (if any), where decision went against the firm	Affidavit on judicial stamp paper
Total		15		

- b. **Past Experience / Past Record**

Ser	Description	Max Marks	Calculation Procedure
-----	-------------	-----------	-----------------------

(1)	Projects of similar nature and complexity	10	<ul style="list-style-type: none"> • 3 years (1.3 mark per contract Max 3.3 marks per year) • New Firms will be awarded 2.5 gratis marks
(2)	Status of enlistment with Govt Organization (Attested copies of Registration certificate to be enclosed)	5	<ul style="list-style-type: none"> • Full marks will be given on provision of at least 1 x Registration certificate • Non-registered firms will be awarded 2.5 gratis Marks
Total		15	

c. **Past Performance.** Marks for past performance shall be awarded on the basis of following criteria (New firms will be awarded pass marks):-

Ser	Description	Max Marks	Calculation Procedure
(1)	Contracted store supplied beyond DP in last 3 years	2.5	<ul style="list-style-type: none"> • X1 = Total value of last 3 years' contracts. • X2 = Total value of stores delivered within Delivery Period (last 3 years' contracts). <p>Formula: Score= $\frac{X2}{X1} * 2.5$</p>
(2)	Quantum of rejections of items in the last 3 years contracts	2.5	<ul style="list-style-type: none"> • X1 = Total value of last 3 years contracts. • X2 = Total value of items accepted in first go (last 3 years' contracts). <p>Formula: Score= $\frac{X2}{X1} * 2.5$</p>
(3)	Timely provision of documents/ bank guarantees / bid security money	2.5	<ul style="list-style-type: none"> • X1 = Total no of contracts concluded in last 3 years. • X2 = Total number of timely provided bank guarantees/ bid securities against the total no of contracts in last 3 years. <p>Formula: Score= $\frac{X2}{X1} * 2.5$</p>
(4)	No of contracts / items still pending beyond DP	5	One mark would be deducted for each contract in hand, whose deliveries are over due for more than 2 months
(5)	Risk and Expense action against firm approved	2.5	Half mark will be deducted against each Risk and Expense action approved / done within last 3 years
(6)	Response to HIT Procurement queries /problems	5	<ul style="list-style-type: none"> • Half mark will be deducted for each advice letter issued to firm • One mark will be deducted for each

	(last 3 years)		warning letter issued to the firm
Total		20	

d. **Project Technical Evaluation Parameters**

Ser	Description	Max Points	Remarks
(1)	Source of Items (Local / Imported), trade link with OEM (agency agreement if OEM not participation directly).	10	Yes/No
(2)	Reliability of Similar Items Provided Previously in HIT hosp / Civil / Military hosp (att supply order).	No of supply order 8-10=10 05-7=05 03-5=03 01-2=01	Yes/No
(3)	Valid Drug Manufacturing License (For Manufactures) Valid Sales License and Establishment Registration Certificate (For Sole Agents).	05	Yes/No
(4)	Valid GMP Certificate issued by DRAP (For Local Agents)	05	Yes/No
(5)	Undertaking by Firm on Notarized Stamp Paper of Rs. 100/- Binding the Firm to Supply the Stock in Compliance to SRO 470(1)/2017 Subject to Department Requirement.	05	Yes/No
(6)	Undertaking by Firm on Notarized Stamp Paper of Rs. 100/- Binding the Firm if required the supplied stocks would be sent for drug testing and in this regard firm will bear all expenses.	05	Yes/No
(7)	Provision of Two Packs of Samples for Technical Committee Evaluation.	10	Yes/No
Total		50	
Note:-			
➤ All points mentioned in d (1) to (7) are mandatory. Non-compliance with any will render company ineligible.			

LETTER OF BID

Date of this Bid submission: [_____ Tuesday, ___ November, _____ Year]

RFB No.: _____ [insert number of Bidding process]

Request for Bid No.: [_____] Bid No

Alternative No.: [_____] insert identification No if this is a Bid for an alternative]

To: **Assistant Director Procurement Directorate**

Gate No 5, Heavy industries taxila

We, the undersigned Bidder, hereby submit our Bid, in two parts, namely: M/s _____

- (a) the Technical Proposal, and
- (b) the Financial Proposal.

In submitting our Bid we make the following declarations:

1. **No reservations:** We have examined and have no reservations to the bidding document, including addenda issued in accordance with Instructions to Bidders (ITB 9);
2. **Eligibility:** We meet the eligibility requirements and have no conflict of interest in accordance with ITB 3;
3. **Bid/Proposal-Securing Declaration:** We have not been suspended nor declared ineligible by the Procurement Directorate based on execution of a Bid Securing Declaration or Proposal Securing Declaration in the Procurement Directorate's country in accordance with ITB 4;
4. **Conformity:** We offer to supply in conformity with the bidding document and in accordance with the Delivery Schedules specified in the Schedule of Requirements the following Goods: [insert a brief description of the Goods and Related Services];
5. **Total Price:** The total price of our Bid, excluding any discounts offered in item (c) below is:
In case of only one lot, the total price of the Bid is [insert the total price of the bid in words and figures, indicating the various amounts and the respective currencies];
In case of multiple lots, the total price of each lot is [insert the total price of each lot in words and figures, indicating the various amounts and the respective currencies];
In case of multiple lots, total price of all lots (sum of all lots) [insert the total price of all lots in words and figures, indicating the various amounts and the respective currencies];
6. **Discounts:** The discounts offered and the methodology for their application are:
 - a. The discounts offered are: [Specify in detail each discount offered]

b. The exact method of calculations to determine the net price after application of discounts is shown below: *[Specify in detail the method that shall be used to apply the discounts];*

7. **Bid Validity Period:** Our Bid shall be valid for the period specified in BDS 17.1 (as amended, if applicable) from the date fixed for the Bid submission deadline specified in BDS 23.1 (as amended, if applicable), and it shall remain binding upon us, and may be accepted at any time before the expiration of that period;
8. **Performance Security:** If our Bid is accepted, we commit to obtain a performance security in accordance with the bidding document;
9. **One Bid per Bidder:** We are not submitting any other Bid(s) as an individual Bidder, and we are not participating in any other bid(s) as a Joint Venture member or as a subcontractor, and meet the requirements, other than Alternative Bids submitted in accordance with ITB 19;
10. **Suspension and Debarment:** We, along with any of our subcontractors, suppliers, consultants, manufacturers, or service providers for any part of the contract, are not subject to, and not controlled by any entity or individual that is subject to, a temporary suspension or a debarment imposed by the Procurement Directorate. Further, we are not ineligible under Pakistan laws;
11. **State-owned enterprise or institution:** *[select the appropriate option and delete the other] [We are not a state-owned enterprise or institution] / [We are a state-owned enterprise or institution but meet the requirements of];*
12. **Binding Contract:** We understand that this Bid, together with your written acceptance thereof included in your Letter of Acceptance, shall constitute a binding contract between us, until a formal contract is prepared and executed;
13. **Not Bound to Accept:** We understand that you are not bound to accept the the Most Advantageous Bid or any other Bid that you may receive; and
14. **Fraud and Corruption:** We hereby certify that we have taken steps to ensure that no person acting for us, or on our behalf, engages in any type of Fraud and Corruption.

Name of the Bidder: **[insert complete name of Bidder]*

Name of the person duly authorized to sign the Bid on behalf of the Bidder: *** [insert complete name of person duly authorized to sign the Bid]*

Title of the person signing the Bid: *[insert complete title of the person signing the Bid]*

Signature of the person named above: *[insert signature of person whose name and capacity are shown above]*

Date signed *[insert date of signing]* **day of** *[insert month], [insert year]*

Bidder Information Form

Date: *[insert date (as day, month and year) of Bid submission]*

No.: *[insert number of Bidding process]*

Alternative No.: *[insert identification No if this is a Bid for an alternative]*

Page _____ of _____ pages

1.	Bidder's Name <i>[insert Bidder's legal name]</i>
2.	In case of JV, legal name of each member : <i>[insert legal name of each member in JV]</i>
3.	Bidder's actual or intended country of registration: <i>[insert actual or intended country of registration]</i>
4.	Bidder's year of registration: <i>[insert Bidder's year of registration]</i>
5.	Bidder's Address in country of registration: <i>[insert Bidder's legal address in country of registration]</i>
6.	Bidder's Authorized Representative Information Name: <i>[insert Authorized Representative's name]</i> Address: <i>[insert Authorized Representative's Address]</i> Telephone/Fax numbers: <i>[insert Authorized Representative's telephone/fax numbers]</i> Email Address: <i>[insert Authorized Representative's email address]</i>
7.	Attached are copies of original documents of <i>[check the box(es) of the attached original documents]</i> <input type="checkbox"/> Articles of Incorporation (or equivalent documents of constitution or association), and/or documents of registration of the legal entity named above. <input type="checkbox"/> In case of JV, letter of intent to form JV or JV agreement, in accordance with ITB 3.4. <input type="checkbox"/> Establishing that the Bidder is not under the supervision of the Procurement Directorate
8.	Included are the organizational chart, a list of Board of Directors, and the beneficial ownership.

COMMERCIAL OFFER
Price Schedule
Therapeutic Goods Manufactured Pakistan, or to be Imported

Ser	IT Number	Ser Number	Part Number	Nomenclature	Description / Specifications	A/U	Quantity	Price Per Unit (Rs)	Total Price (Rs)
1.									
2.									
3.									
Total Price without Taxes:									Rs. -----
							Details of applicable Taxes:		
							GST	@ ----- %	Rs. -----
							Income Tax	@ ----- %	Rs. -----
							-----Tax	@ ----- %	Rs. -----
							-----Tax	@ ----- %	Rs. -----
							-----Tax	@ ----- %	Rs. -----
Total of all Applicable Taxes									Rs -----
Total Price with Taxes (all applicable):									Rs. -----

Form of Bid Security

[The bank shall fill in this Bank Guarantee Form in accordance with the instructions indicated.]

[Guarantor letterhead or SWIFT identifier code]

Beneficiary: *[Purchaser to insert its name and address]*

No.: *[Purchaser to insert reference number for the Request for Bids]*

Alternative No.: *[Insert identification No if this is a Bid for an alternative]*

Date: *[Insert date of issue]*

BID GUARANTEE No.: *[Insert guarantee reference number]*

Guarantor: *[Insert name and address of place of issue, unless indicated in the letterhead]*

We have been informed that _____ *[insert name of the Bidder, which in the case of a joint venture shall be the name of the joint venture (whether legally constituted or prospective) or the names of all members thereof]* (hereinafter called "the Applicant") has submitted or will submit to the Beneficiary its Bid (hereinafter called "the Bid") for the execution of _____ under Request for Bids No. _____ ("the RFB").

Furthermore, we understand that, according to the Beneficiary's conditions, Bids must be supported by a Bid guarantee.

At the request of the Applicant, we, as Guarantor, hereby irrevocably undertake to pay the Beneficiary any sum or sums not exceeding in total an amount of _____ (_____) upon receipt by us of the Beneficiary's complying demand, supported by the Beneficiary's statement, whether in the demand itself or a separate signed document accompanying or identifying the demand, stating that either the Applicant:

- (a) has withdrawn its Bid during the period of Bid validity set forth in the Applicant's Letter of Bid ("the Bid Validity Period"), or any extension thereto provided by the Applicant; or
- (b) Having been notified of the acceptance of its Bid by the Beneficiary during the Bid Validity Period or any extension thereto provided by the Applicant, (i) has failed to sign the contract agreement, or (ii) has failed to furnish the performance security, in accordance with the Instructions to Bidders ("ITB") of the Beneficiary's bidding document.

This guarantee will expire: (a) if the Applicant is the successful Bidder, upon our receipt of copies of the Contract agreement signed by the Applicant and the performance security issued to the Beneficiary in relation to such Contract agreement; or (b) if the Applicant is not the successful Bidder, upon the earlier of (i) our receipt of a copy of the Beneficiary's notification to the Applicant of the results of the Bidding process; or (ii) twenty-eight days after the end of the Bid Validity Period.

Consequently, any demand for payment under this guarantee must be received by us at the office indicated above on or before that date.

[Signature(s)]

Note: All italicized text is for use in preparing this form and shall be deleted from the final product.

Form of Bid-Securing Declaration

[The Bidder shall fill in this Form in accordance with the instructions indicated.]

Date: *[date (as day, month and year)]*

No.: *[number of Bidding process]*

Alternative No.: *[insert identification No if this is a Bid for an alternative]*

To: *[complete name of Procurement Directorate]*

We, the undersigned, declare that:

We understand that, according to your conditions, Bids must be supported by a Bid-Securing Declaration.

We accept that we will be blacklisted and henceforth cross debarred for participating in respective category of public procurement proceedings for a period of (not more than) six months, if fail to abide with a bid securing declaration, however without indulging in corrupt and fraudulent practices, if we are in breach of our obligation(s) under the Bid conditions, because we:

- (a) have withdrawn our Bid during the period of Bid validity specified in the Letter of Bid; or
- (b) having been notified of the acceptance of our Bid by the Procurement Directorate during the period of Bid validity, (i) fail or refuse to sign the Contract; or (ii) fail or refuse to furnish the Performance Security (or guarantee), if required, in accordance with the ITB.

We understand this Bid Securing Declaration shall expire if we are not the successful Bidder, upon the earlier of (i) our receipt of your notification to us of the name of the successful Bidder; or (ii) twenty-eight days after the expiration of our Bid.

Name of the Bidder* _____

Name of the person duly authorized to sign the Bid on behalf of the Bidder** _____

Title of the person signing the Bid _____

Signature of the person named above _____

Date signed _____ day of _____, _____

*: In the case of the Bid submitted by joint venture specify the name of the Joint Venture as Bidder

**: Person signing the Bid shall have the power of attorney given by the Bidder attached to the Bid

[Note: In case of a Joint Venture, the Bid-Securing Declaration must be in the name of all members to the Joint Venture that submits the Bid.]

Letter of Acceptance

[Letter head paper of the Procurement Directorate]

[date]

To: *[name and address of the Supplier]*

This is to notify you that your Bid dated *[date]* for execution of the *[name of the Contract and identification number, as given in the Special Conditions of Contract]* for the Contract Price of the equivalent of *[amount in numbers and words] [name of currency]*, as corrected and modified in accordance with the Instructions to Bidders is hereby accepted by us.

We hereby confirm *[insert the name of the Appointing Authority]*, to be the Appointing Authority, to appoint the Arbitrator in case of any arisen disputes.

You are hereby informed that after you have read and return the attached draft Contract the parties to the contract shall sign the vetted contract within fourteen (14) working days.

You are hereby required to furnish the Performance Guarantee/Security in the form and the amount stipulated in the Special Conditions of the Contract within a period of fourteen (14) days after the receipt of Letter of Acceptance.

Authorized Signature:

Name and Title of Signatory:

Name of Agency:

Attachment: Contract

Copy: Appointing Authority and Supplier

Manufacturer's Authorization

*[The Bidder shall require the Manufacturer to fill in this Form in accordance with the instructions indicated. This letter of authorization should be on the letterhead of the Manufacturer and should be signed by a person with the proper authority to sign documents that are binding on the Manufacturer. The Bidder shall include it in its Bid, if so indicated in the **BDS.**]*

Date: *[insert date (as day, month and year) of Bid submission]*

No.: *[insert number of Bidding process]*

Alternative No.: *[insert identification No if this is a Bid for an alternative]*

To: *[insert complete name of Procurement Directorate]*

WHEREAS

We *[insert complete name of Manufacturer]*, who are official manufacturers of *[insert type of goods manufactured]*, having factories at *[insert full address of Manufacturer's factories]*, do hereby authorize *[insert complete name of Bidder]* to submit a Bid the purpose of which is to provide the following Therapeutic Goods, manufactured by us *[insert name and or brief description of the Therapeutic Goods]*, and to subsequently negotiate and sign the Contract.

We hereby extend our full guarantee and warranty in accordance with Clause 29 of the General Conditions of Contract, with respect to the Therapeutic Goods offered by the above firm.

Signed: *[insert signature(s) of authorized representative(s) of the Manufacturer]*

Name: *[insert complete name(s) of authorized representative(s) of the Manufacturer]*

Title: *[insert title]*

Dated on _____ day of _____, _____ *[insert date of signing]*

PART II
CONDITIONS OF THE CONTRACT

GENERAL CONDITIONS OF THE CONT (GCC)

1.	Definitions	1.1	The following words and expressions shall have the meanings hereby assigned to them:
			a. “Authority” means Public Procurement Regulatory Authority.
			b. The “Arbitrator” is the person appointed with mutual consent of both the parties, to resolve contractual disputes as provided for in the General Conditions of the Contract GCC Clause 34 hereunder.
			c. The “Contract” means the agreement entered into between the Procurement Directorate and the Supplier, as recorded in the Contract Form signed by the parties, including all attachments and appendices thereto and all documents incorporated by reference therein.
			d. The “Commencement Date” is the date when the Supplier shall commence execution of the contract as specified in the SCC .
			e. “Completion” means the fulfillment of the related services by the Supplier in accordance with the terms and conditions set forth in the contract.
			f. “Country of Origin” means the countries and territories eligible under the PPRA Rules 2004 and its corresponding Regulations as further elaborated in the SCC .
			g. The “Contract Price” is the price stated in the Letter of Acceptance and thereafter as adjusted in accordance with the provisions of the Contract.
			h. “Defective Goods” are those goods which are below standards, requirements or specifications stated by the Contract.
			i. “Delivery” means the transfer of the goods from the supplier equipment, machinery, and /or other materials which the Supplier is required to supply to the Consignee address under Contract.
			j. “Effective Contract date” is the date shown in the Certificate of Contract Commencement issued by the Procurement Directorate upon fulfillment of the conditions precedent stipulated in GCC Clause 3 .
			k. “Related Services” means those services ancillary to the delivery of the Goods, such as transportation and insurance, and any other incidental services, such as installation, commissioning, provision of technical assistance, training, initial maintenance and other such obligations of the Supplier covered under the Contract.
			l. “GCC” means the General Conditions of Contract contained in this section.
			m. “Intended Delivery Date” is the date on which it is intended that the Supplier shall effect delivery as specified in the SCC .
			n. “SCC” means the Special Conditions of Contract.
			o. “Supplier” means the individual private or government entity or a combination of the above whose Bid to perform the contract has been accepted by the Procurement Directorate and is named as such in the Contract Agreement, and includes the legal successors or permitted assigns of the supplier and shall be named in the SCC .
			p. “Project Name” means the name of the project stated in SCC .
			q. “Day” means calendar day.
			r. “Eligible Country” means the countries and territories eligible for

			participation in accordance with the policies of the Federal Government.
		s.	“End User” means the organization(s) where the goods will be used, as named in the SCC .
		t.	“Origin” means the place where the Goods were mined, grown, or produced or from which the Services are supplied. Goods are produced when, through manufacturing, processing, or substantial and major assembly of components, a commercially recognized new produce results that is substantially different in basic characteristics or in purpose or utility from its components.
		u.	“Force Majeure” means an unforeseeable event which is beyond reasonable control of either Party and which makes a Party’s performance of its obligations under the Contract impossible or so impractical as to be considered impossible under the circumstances. For the purposes of this Contract, “Force Majeure” means an event which is beyond the reasonable control of a Party, is not foreseeable, is unavoidable, and its origin is not due to negligence or lack of care on the part of a Party, and which makes a Party’s performance of its obligations hereunder impossible or so impractical as reasonably to be considered impossible in the circumstances. and includes, but is not limited to, war, riots, civil disorder, earthquake, fire, explosion, storm, flood, epidemics, or other adverse weather conditions, strikes, lockouts or other industrial action (except where such strikes, lockouts or other industrial action are within the power of the Party invoking Force Majeure to prevent), confiscation or any other action by Government agencies.
		v.	“Specification” means the Specification of the Goods and performance of incidental services in accordance with the relevant standards included in the Contract and any modification or addition made or approved by the Hospital.
		w.	The Supplier's Bid is the completed Bid document submitted by the Supplier to the Procurement Directorate.
2.	Application and interpretation	2.1	These General Conditions shall apply to the extent that they are not superseded by provisions of other parts of the Contract.
		2.2	In interpreting these Conditions of Contract headings and marginal notes are used for convenience only and shall not affect their interpretations unless specifically stated; references to singular include the plural and vice versa; and masculine include the feminine. Words have their ordinary meaning under the language of the Contract unless specifically defined.
		2.3	The documents forming the Contract shall be interpreted in the following order of priority: a. Form of Contract, b. Special Conditions of Contract, c. General Conditions of Contract, d. Letter of Acceptance, e. Certificate of Contract Commencement f. Specifications g. Contractor's Bid, and h. Any other document listed in the Special Conditions of Contract as forming part of the Contract.

3.	Conditions Precedent	3.1	Having signed the Contract, it shall come into effect on the date on which the following conditions have been satisfied: - a. Submission of performance Security (or guarantee) in the form specified in the SCC ; b. Furnishing of Advance Payment Unconditional Guarantee.
		3.2	If the Condition precedent stipulated on GCC Clause 3.1 is not met by the date specified in the SCC this contract shall not come into effect;
		3.3	If the Procurement Directorate is satisfied that each of the conditions precedent in this contract has been satisfied (except to the extent waved by him, but subject to such conditions as he shall impose in respect of such waiver) he shall promptly issue to the supplier a certificate of Contract commencement, which shall confirm the start date.
4.	Governing Language	4.1	The Contract as all correspondence and documents relating to the contract exchanged by the Supplier and the Procurement Directorate shall be written in the language specified in SCC . The version of the Contract written in the specified language shall govern its interpretation.
5.	Applicable Law	5.1	The contract shall be governed and interpreted in accordance with the laws of Pakistan, unless otherwise specified in SCC .
6.	Country of Origin	6.1	The origin of Therapeutic Goods (Pharmaceuticals) and Services may be distinct from the nationality of the Supplier.
7.	Standards	7.1	The Therapeutic Goods supplied under this Contract shall conform to the standards mentioned in the Technical Specifications.
8.	Joint Venture Consortium or Association	8.1	If the Supplier is a joint venture, consortium, or association, all of the parties shall be jointly and severally liable to the Purchaser for the fulfillment of the provisions of the Contract and shall designate one party to act as a leader with authority to bind the joint venture, consortium, or association. The composition or the constitution of the joint venture, consortium, or association shall not be altered without the prior consent of the Purchaser
9.	Contract Documents	9.1	Subject to the order of precedence set forth in the Contract Agreement, all documents forming the Contract (and all parts thereof) are intended to be correlative, complementary, and mutually explanatory. The Contract Agreement shall be read as a whole.
10.	Use of Contract Documents and Information; Inspection and Audit by the Government of Pakistan	10.1	The Supplier shall not, without the Procurement Directorate's prior written consent, disclose the Contract, or any provision thereof, or any specification, plan, drawing, pattern, sample, or information furnished by or on behalf of the Procurement Directorate in connection therewith, to any person other than a person employed by the Supplier in the performance of the Contract. Disclosure to any such employed person shall be made in confidence and shall extend only as far as may be necessary for purposes of such performance.
		10.2	The Supplier shall not, without the Procurement Directorate's prior written consent, make use of any document or information enumerated in GCC Clause 10.1 except for purposes of performing the Contract.
		10.3	Any document, other than the Contract itself, enumerated in GCC Clause 10.1 shall remain the property of the Procurement Directorate and shall be returned (all copies) to the Procurement Directorate on completion of the Supplier's performance under the Contract if so required by the Procurement Directorate.

		10.4	The Supplier shall permit the Government of Pakistan or / and donor agencies involved in financing the project to inspect the Supplier's accounts and records relating to the performance of the Supplier and to have them audited by auditors appointed by the Government of Pakistan or / and the appropriate donor agencies, if so required by the Government of Pakistan or / and the appropriate donor agencies.
11.	Patent and Copy Rights	11.1	The Supplier shall indemnify the Procurement Directorate against all third-party claims of infringement of patent, trademark, or industrial design rights arising from use of the Therapeutic Goods or any part thereof in Pakistan.
		11.2	The patent right in all drawings, documents, and other materials containing data and information furnished to the Procurement Directorate by the Supplier herein shall remain vested in the supplier, or, if they are furnished to the Procurement Directorate directly, or through the Supplier by any third party, including suppliers of materials, the patent right in such materials shall remain vested in such third party.
12.	Scope of Supply	12.1	The therapeutic Goods and Related Services to be supplied shall be as specified in the Schedule of Requirements
13.	Performance Guarantee	13.1	The Performance Guarantee shall be provided to the Procurement Directorate no later than the date specified in the Letter of Acceptance and shall be issued in an amount and form and by a bank or surety acceptable to the Procurement Directorate, and denominated in the types and proportions of the currencies in which the Contract Price is payable as specified in the SCC .
		13.2	The proceeds of the Performance Guarantee shall be payable to the Procurement Directorate as compensation for any loss resulting from the Supplier's failure to complete its obligations under the Contract.
		13.3	The Performance Guarantee shall be in one of the following forms:
		a.	A bank guarantee, an irrevocable letter of credit issued by a reputable bank, or in the form provided in the Bidding Documents or another form acceptable to the Procurement Directorate; or
		b.	A cashier's or certified check.
13.4	The performance guarantee will be discharged by the Procurement Directorate and returned to the Supplier not later than thirty (30) days following the date of completion of the Supplier's performance obligations under the Contract, including any warranty obligations, unless otherwise specified in SCC .		
14.	Supplier's Responsibility	14.1	The Supplier shall supply all the Therapeutic Goods and Related Services included in the Scope of Supply in accordance with GCC Clause 12, and the Delivery and Completion Schedule, as per GCC Clause 17
15.	Inspections and Test	15.1	The Procurement Directorate or its representative shall have the right to inspect and /or to test the therapeutic Goods to confirm their conformity to the Contract specifications at no extra cost to the Procurement Directorate. SCC and the Technical Specifications shall specify what inspections and tests the Procurement Directorate shall notify the Supplier in writing or in electronic forms that provide record of the content of communication, in a timely manner, of the identity of any representatives retained for these purposes.

		<p>15.2 The inspections and tests may be conducted on the premises of the Supplier or its subcontractor(s), at point of delivery, and/or at the Therapeutic Goods' final destination. If conducted on the premises of the Supplier or its subcontractor(s), all reasonable facilities and assistance, including access to drawings and production data, shall be furnished to the inspectors at no charge to the Procurement Directorate.</p> <p>15.3 Should any inspected or tested Therapeutic Goods fail to conform to the Specifications, the Procurement Directorate may reject the Therapeutic Goods, and the Supplier shall replace the rejected Therapeutic Goods to meet specification requirements free of cost to the Procurement Directorate.</p> <p>15.4 The Procurement Directorate's right to inspect, test and, where necessary, reject Therapeutic Goods after its arrival in the Procurement Directorate's country shall in no way be limited or eared by reason of the Therapeutic Goods having previously been inspected, tested, and passed by the Procurement Directorate or its representative prior to the Therapeutic Goods' shipment from the country of origin.</p> <p>15.5 Nothing in GCC Clause 13 shall in any way release the supplier from any warranty or other obligations under this Contract.</p>
16.	Packing	<p>16.1 The supplier shall provide such packing of the Therapeutic Goods as required in the Technical Specification of the Bidding Documents to prevent their damage or deterioration during transit to their final destination, as indicated in the Contract. The packing shall be sufficient to withstand, without limitation, rough handling during transit and exposure to extreme temperatures, salt and precipitation during transit, and open storage. Packing case size and weights shall take into consideration, where appropriate, the remoteness of the Therapeutic Goods final destination and the absence of heavy handling facilities at all points in transit.</p> <p>16.2 The packing, 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 in SCC, and in any subsequent instructions ordered by the Procurement Directorate.</p>
17.	Delivery and Documents	<p>17.1 Documents to be submitted by the Supplier are specified in SCC.</p>
20.	Related Services	<p>20.1 The Supplier may be required to provide any or all of the following services, including additional services, if any, specified in SCC:</p> <p>a. Performance or supervision of on-site assembly, Installation Commissioning and/or start-up of the supplied Therapeutic Goods;</p> <p>b. Furnishing of tools required for assembly and/or maintenance of the supplied Therapeutic Goods;</p> <p>c. Furnishing of a detailed operations and maintenance manual for each appropriate unit of the supplied Therapeutic Goods;</p> <p>d. Performance or supervision or maintenance and/or repair of the supplied Therapeutic 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</p> <p>e. Training of the Procurement Directorate's personnel, at the Supplier's</p>

			plant and/or on-site, in assembly, start-up, operation, maintenance, and/or repair of the supplied Therapeutic Goods.
		20.2	Prices charged by the Supplier for related services, if not included in the Contract Price for the Therapeutic Goods, shall be agreed upon in advance by the parties and shall not exceed the prevailing rates charged to other parties by the Supplier for similar services.
21.	Warranty/ Defect Liability Period	21.1	The Supplier warrants that the Therapeutic Goods supplied under the Contract are new, unused, of the most recent or current models and that they incorporate all recent improvements in design and materials unless provided otherwise in the Contract. The Supplier further warrants that all Therapeutic Goods supplied under this Contract shall have no defect, arising from design, materials, or workmanship (except when the design and/or material is required by the Procurement Directorate, specifications) or from any act or omission of the Supplier, that may develop under normal use of the supplied Therapeutic Goods in the conditions prevailing in Pakistan.
		21.2	This warranty shall remain valid for a period specified in the SCC after the Therapeutic Goods, or any portion thereof as the case may be, have been delivered to and accepted at the final destination indicated in the Contract, or for a period specified in the SCC after the date of shipment from the port or place of loading in the source country, whichever period concludes earlier, unless specified otherwise in SCC .
		21.3	The Procurement Directorate shall promptly notify the Supplier in writing or in electronic forms that provide record of the content of communication of any claims arising under this warranty.
22.	Payment	22.1	The method and conditions of payment to be made to the Supplier under this Contract shall be specified in SCC .
		22.2	The Supplier's request(s) for payment shall be made to the Procurement Directorate in writing or in electronic forms that provide record of the content of communication, accompanied by an invoice describing, as appropriate, the Goods delivered and Services performed, and by documents submitted pursuant to GCC Clause 13 , and upon fulfillment of other obligations stipulated in the Contract.
		22.3	Payments shall be made promptly by the Procurement Directorate, within sixty (60) days after submission of an invoice or claim by the Supplier. If the Procurement Directorate makes a late payment, the Supplier shall be paid interest on the late payment. Interest shall be calculated from the date by which the payment should have been made up to the date when the late payment is made at the rate as specified in the SCC .
		22.4	The currency or currencies in which payment is made to the Supplier under this Contract shall be specified in SCC subject to the following general principle: payment will be made in the currency or currencies in which the payment has been requested in the Supplier's Bid.
		22.5	All payments shall be made in the currency or currencies specified in the SCC pursuant to GCC Clause 22.4
23.	Prices	23.1	The contract price shall be as specified in the Contract Agreement Subject to any additions and adjustments thereto or deductions there from, as may be made pursuant to the Contract.

		23.2	Prices charged by the Supplier for Therapeutic Goods delivered and Services performed under the Contract shall not vary from the prices quoted by the Supplier in its Bid, with the exception of any price adjustments authorized in SCC or in the Procurement Directorate's request for Bid Validity extension, as the case may be.
24.	Change Orders	24.1	The Procurement Directorate may at any time, by a written order given to the Supplier, make changes within the general scope of the Contract in any one or more of the following:
		a.	Drawings, designs, or specifications, where Therapeutic Goods to be furnished under the Contract are to be specifically manufactured for the Procurement Directorate;
		b.	The method of shipment or packing;
		c.	The place of delivery; and/or
		d.	The Services to be provided by the Supplier.
24.2	If any such change causes an increase or decrease in the cost of, or the time required for, the Supplier's performance of any provisions under the Contract an equitable adjustment shall be made in the Contract Price or delivery schedule, or both, and the Contract shall accordingly be amended. Any claims by the Supplier for adjustment under this clause must be asserted within thirty (30) days from the date of the Supplier's receipt of the Procurement Directorate change order.		
24.3	Prices to be charged by the supplier for any related services that might be needed but which were not included in the Contract shall be agreed upon in advance by the Parties and shall not exceed the prevailing rates charged to other parties by the Supplier for similar services.		
25.	Contract Amendments	25.1	Subject to GCC Clause 24 , no variation in or modification of the terms of the Contract shall be made except by written amendment signed by the parties.
26.	Assignment	26.1	Neither the Procurement Directorate nor the Supplier shall assign, in whole or in part, obligations under this Contract, except with the prior written consent of the other party.
27.	Sub-contracts	27.1	The Supplier shall consult the Procurement Directorate in the event of subcontracting under this contract if not already specified in the Bid. Subcontracting shall not alter the Supplier's obligations.
		27.2	Subcontracts must comply with the provision of GCC Clause 5 .
28.	Delays in the Supplier's Performance	28.1	Delivery of the Therapeutic Goods and performance of Services shall be made by the Supplier in accordance with the time schedule prescribed by the Procurement Directorate in the Schedule of Requirements.
		28.2	If at any time during performance of the Contract, the Supplier or its subcontractor(s) should encounter conditions impeding timely delivery of the Therapeutic Goods and performance of Services, the Supplier shall promptly notify the Procurement Directorate in writing or in electronic forms that provide record of the content of communication of the fact of the delay, its likely duration and its cause(s). As soon as practicable after receipt of the Supplier's notice, the Procurement Directorate shall evaluate the situation and may at its discretion extend the Supplier's time for performance, with or without liquidated damages, in which case the extension shall be ratified by the parties by amendment of Contract.

		28.3	Except as provided under GCC Clause 31 , a delay by the Supplier in the performance of its delivery obligations shall render the Supplier liable to the imposition of liquidated damages pursuant to GCC Clause 29 , unless an extension of time is agreed upon without the application of liquidated damages.
29.	Liquidated Damages	29.1	Subject to GCC Clause 31 , if the Supplier fails to deliver any or all of the Therapeutic Goods or to perform the Services within the period(s) specified in the Contract, the Procurement Directorate shall, without prejudice to its other remedies under the Contract, deduct from the Contract Price, as liquidated damages, a sum equivalent to the percentage specified in SCC of the delivered price of the delayed Therapeutic Goods or unperformed Services for each week or part thereof of delay until actual delivery or performance, up to a maximum deduction of the performance security (or guarantee) specified in SCC . Once the said maximum is reached, the Procurement Directorate may consider termination of the Contract.
30.	Termination for Default	30.1	The Procurement Directorate or the Supplier, without prejudice to any other remedy for breach of Contract, by written notice of default sent to the concerned party may terminate the Contract if the other party causes a fundamental breach of the Contract.
		30.2	Fundamental breaches of Contract shall include, but shall not be limited to the following:
		a.	the Supplier fails to deliver any or all of the Therapeutic Goods within the period(s) specified in the Contract, or within any extension thereof granted by the Procurement Directorate; or
		b.	the Supplier fails to perform any other obligation(s) under the Contract;
		c.	Supplier's failure to submit performance security (or guarantee) within the time stipulated in the SCC ;
		d.	the supplier has abandoned or repudiated the contract.
		e.	the Procurement Directorate or the Supplier is declared bankrupt or goes into liquidation other than for a reconstruction or amalgamation;
		f.	a payment is not paid by the Procurement Directorate to the Supplier after Number of days Specified in SCC from the due date for payment;
		g.	the Procurement Directorate gives Notice that Therapeutic goods delivered with a defect is a fundamental breach of Contract and the Supplier fails to correct it within a reasonable period of time determined by the Procurement Directorate; and
		h.	if the Procurement Directorate determines, based on the reasonable evidence, that the Supplier has engaged in corrupt, coercive, collusive, obstructive or fraudulent practices, in competing for or in executing the Contract.
		30.3	For the purpose of this clause:
			Corrupt and Fraudulent Practice " in respect of procurement process, shall be either one or any combination of the practices including,-
		a.	"coercive practices" which means any impairing or harming or threatening to impair or harm, directly or indirectly, any party or the property of the party to influence the actions of a party to achieve a wrongful gain or to cause a wrongful loss to another party;

			<p>b. "collusive practices" which means any arrangement between two or more parties to the procurement process designed to stifle open competition for any wrongful gain, and to establish prices at artificial, non-competitive levels;</p>
			<p>c. "corrupt practices" which means the offering, giving, receiving or soliciting, directly or indirectly, of anything of value to influence the acts of another party for wrongful gain;</p> <p>d. "fraudulent practices" which means any act or omission, including a misrepresentation, that knowingly or recklessly misleads, or attempts to mislead, a party to obtain a financial or other benefit or to avoid an obligation; and</p> <p>e. "obstructive practices" which means harming or threatening to harm, directly or indirectly, persons to influence their participation in a procurement process, or affect the execution of a contract..</p>
		30.4	<p>In the event the Procurement Directorate terminates the Contract in whole or in part, the Procurement Directorate may procure, upon such terms and in such manner as it deems appropriate, Therapeutic Goods or Services similar to those undelivered, and the Supplier shall be liable to the Procurement Directorate for any excess costs for such similar Therapeutic Goods or Services. However, the Supplier shall continue performance of the Contract to the extent not terminated.</p>
31.	Termination for Force Majeure	31.1	<p>Notwithstanding the provisions of GCC Clauses 29, and, neither Party shall have any liability or be deemed to be in breach of the Contract for any delay nor is other failure in performance of its obligations under the Contract, if such delay or failure is a result of an event of Force Majeure.</p> <p>For purpose of this clause, "Force Majeure" means an event which is beyond the reasonable control of a Party, is not foreseeable, is unavoidable, and its origin is not due to negligence or lack of care on the part of a Party, and which makes a Party's performance of its obligations hereunder impossible or so impractical as reasonably to be considered impossible in the circumstances, and includes, but is not limited to, war, riots, civil disorder, earthquake, fire, explosion, storm, flood, epidemics, or other adverse weather conditions, strikes, lockouts or other industrial action (except where such strikes, lockouts or other industrial action are within the power of the Party invoking Force Majeure to prevent</p>
		31.2	<p>If a Party (hereinafter referred to as "the Affected Party") is or will be prevented from performing its substantial obligation under the contract by Force Majeure, it shall give a Notice to the other Party giving full particulars of the event and circumstance of Force Majeure in writing or in electronic forms that provide record of the content of communication of such condition and the cause thereof. Unless otherwise directed by the Procurement Directorate in writing or in electronic forms that provide record of the content of communication, the Supplier shall continue to perform its obligations under the Contract as far as is reasonably practical, and shall seek all reasonable alternative means for performance not prevented by the Force Majeure event.</p>

32.	Termination for Insolvency	32.1	The Procurement Directorate may at any time terminate the Contract by giving written notice to the Supplier if the Supplier becomes bankrupt or otherwise insolvent. In this event, termination will be without compensation to the Supplier, provided that such termination will not prejudice or affect any right of action or remedy which has accrued or will accrue thereafter to the Procurement Directorate.
33.	Termination for Convenience	33.1	The Procurement Directorate, by written notice sent to the Supplier, may terminate the contract, in whole or in part, at any time for its convenience. The notice of termination shall specify that termination is for the Procurement Directorate's convenience, the Contract is terminated, and the date upon which such termination becomes effective.
		33.2	The Therapeutic Goods that are complete and ready for shipment within thirty (30) days after the Supplier's receipt of notice of termination shall be accepted by the Procurement Directorate at the Contract terms and price.
34.	Disputes Resolution	34.1	In the event of any dispute arising out of this contract, either party shall issue a notice of dispute to settle the dispute amicably. The parties hereto shall, within twenty-eight (28) days from the notice date, use their best efforts to settle the dispute amicably through mutual consultations and negotiation. Any unsolved dispute may be referred by either party to Arbitrator who will be Chairman HIT (sole Arbitrator).
		34.2	After the dispute has been referred to the arbitrator, within 30 days, or within such other period as may be proposed by the Parties, the Arbitrator shall give its decision. The rendered decision shall be binding to the Parties and not liable to be challenged in any Court of Law.
35.	Procedure for Disputes Resolution	35.1	The arbitration shall be conducted in accordance with the arbitration procedure published by the Institution named and in the place shown in the SCC.
		35.2	The rate of the Arbitrator's fee and administrative costs of arbitration shall be borne equally by the Parties. The rates and costs shall be in accordance with the rules of the Appointing Authority. In conducting arbitration to its finality each party shall bear its incurred costs and expenses.
		35.3	The arbitration shall be conducted in accordance with the arbitration procedure published by the institution named and in the place shown in the SCC.
36.	Replacement of Arbitrator	36.1	Should the Arbitrator resign or die, or should the Procurement Directorate and the Supplier agree that the Arbitrator is not functioning in accordance with the provisions of the contract, a new Arbitrator shall be appointed by mutual consent of the both parties.
37.	Limitation of Liability	37.1	Except in cases of criminal negligence or willful conduct, and in the case of infringement,
		a.	The supplier shall not be liable to the Procurement Directorate, whether in contract, tort, or otherwise, for any indirect or consequential loss or damage, loss of use, loss of production, or loss of profits or interest costs, provided that this exclusion shall not apply to any obligation of the Supplier to pay liquidated damages to the Procurement Directorate; and
		b.	The aggregate liability of the Supplier to the Procurement Directorate, whether under the Contract, in tort or otherwise, shall not exceed the total Contract Price, provided that this limitation shall not apply to the cost of repairing or replacing defective equipment or to any obligation of the

			Supplier to indemnify the Procurement Directorate with respect to patent infringement.
38. Notices	38.1	Any notice given by one party to the other pursuant to this Contract shall be sent to the other party in writing or in electronic forms that provide record of the content of communication and confirmed in writing or in electronic forms that provide record of the content of communication to the other party's address specified in SCC .	
		38.2	A notice shall be effective when delivered or on the notice's effective date, whichever is later.
39. Effectiveness of the Contract	39.1	Unless otherwise specified in the SCC, the Contract shall become effective on the date ("the Effective Date") that the Supplier receives written notification from the relevant authority in the Purchaser's Country that the Therapeutic Goods have been registered for use in the Procurement Directorate's Country.	
	39.2	If thirty (30) days, or such longer period specified in the SCC, elapse from the date of Contract signing and the Contract has not become effective pursuant to Sub-Clause 39.1 above, then either party may, by not less than seven (7) days' written notice to the other party, declare this Contract null and void. In such event, the Supplier's Performance Security shall be promptly returned.	
40. Extension	40.1	If at any time during performance of the Contract, the Supplier or its subcontractors should encounter conditions impeding timely delivery of the Therapeutic Goods or completion of Related Services, the Supplier shall promptly notify the Purchaser in writing of the delay, its likely duration, and its cause. As soon as practicable after receipt of the Supplier's notice, the Purchaser shall evaluate the situation and may at its discretion extend the Supplier's time for performance, in which case the extension shall be ratified by the parties by amendment of the Contract.	
	40.2	Except in case of Force Majeure, as provided under GCC Clause 31, a delay by the Supplier in the performance of its Delivery and Completion obligations shall render the Supplier liable to the imposition of liquidated damages pursuant to GCC Clause 29, unless an extension of time is agreed upon, pursuant to GCC Sub-Clause 39.1	
41. Taxes and Duties	41.1	A foreign Supplier shall be entirely responsible for all taxes, stamp duties, license fees, and other such levies imposed outside Pakistan.	
	41.2	If any tax exemptions, reductions, allowances or privileges may be available to the Supplier in Pakistan the Procurement Directorate shall use its best efforts to enable the Supplier to benefit from any such tax savings to the maximum allowable extent.	
	41.3	A local Supplier shall be entirely responsible for all taxes, duties, license fees, etc., incurred until delivery of the contracted Goods to the Procurement Directorate.	

PART II
CONDITIONS OF THE CONTRACT

SPECIAL CONDITIONS OF THE CONTRACT
(SCC)

1. The following Special Conditions of Contract (SCC) shall supplement the GCC. Whenever there is a conflict, the provisions herein shall prevail over those in the GCC. The corresponding clause number of the GCC is indicated in parentheses.

SCC Clause Number	GCC Clause Number	Amendments of, and Supplements to, Clauses in the GCC
Definitions (GCC 1)		
1.	1.1	The Procurement Directorate is: <i>[Name and address]</i>
2.	1.1(j)	The Supplier is: <i>[Name and address]</i>
3.	1.1(q)	The title of the subject procurement or The Project is: <i>[write the name of title or project]</i>
Governing Language (GCC 4)		
4.	4.1	The Governing Language shall be:
Applicable Law (GCC 5)		
5.	5.1	The Applicable Law shall be: Laws of the
Country of Origin (GCC 6)		
6.	6.1	Country of Origin is
Performance Security (or guarantee) (GCC 13)		
7.	13.1	The amount of performance security (or guarantee), as a percentage of the Contract Price, shall be: <i>[below the ten (10) percent of the Contract Price]</i>
8.	13.4	After delivery and acceptance of the Therapeutic Goods, 5% percent of the Performance Security (or guarantee) shall be withheld to cover the Supplier's warranty obligations in accordance with GCC Clause 21.2 .
Inspections and Tests (GCC 15)		
9.	15.1	Inspection and tests prior to shipment of Therapeutic Goods and at final acceptance are as follows: Quality and quantity inspection shall be carried out prior to shipment of Therapeutic Goods by the manufacturer(s) at the supplier's own expense and responsibility in terms of the items specified in the specifications. The supplier shall submit the inspection certificate issued by himself which should be attached with the certificate(s) of the manufacturer(s) to the Procurement Directorate in order to ensure that the Therapeutic goods are manufactured in compliance with the contract.
Packing (GCC Clause 16)		
10.	16.2	The following SCC shall supplement GCC Clause 16.2 : The Therapeutic Goods shall be packed properly in accordance with standard export packing specified by the Procurement Directorate in the Technical Specification.
Delivery and Documents (GCC Clause 17)		
11.	17.1	For Therapeutic Goods from within Pakistan: Upon delivery of the Therapeutic Goods to the transporter, the Supplier shall notify the Procurement Directorate and mail the following documents to the Procurement Directorate: a. one original plus four copies of the Supplier's invoice showing Therapeutic Goods' description, quantity, unit price, and total amount; b. delivery note, railway receipt, or truck receipt; c. Manufacturer's or Supplier's warranty certificate; d. inspection certificate issued by the nominated inspection agency, and the Supplier's factory inspection report; and e. Certificate of country of origin issued by Pakistan Chamber of Commerce and

		Industry or equivalent authority in the country of origin in duplicate. The above documents shall be received by the Procurement Directorate before arrival of the Therapeutic Goods and, if not received, the Supplier will be responsible for any consequent expenses.
Related Services (GCC Clause 20)		
13.	20.1	Related services to be provided are: <i>[Selected services covered under GCC Clause 16 and/or other should be specified with the desired features. The price quoted in the Bid price or agreed with the selected Supplier shall be included in the Contract Price.]</i>
Warranty (GCC Clause 21)		
14.	21.2	In partial modification of the provisions, the warranty period shall be ___-__ hours of operation or _____ months from date of acceptance of the Goods or (_____) months from the date of shipment, whichever occurs earlier. The Supplier shall, in addition, comply with the performance and/or consumption guarantees specified under the Contract. If, for reasons attributable to the Supplier, these guarantees are not attained in whole or in part, the Supplier shall, at its discretion, either: a. make such changes, modifications, and/or additions to the Goods or any part thereof as may be necessary in order to attain the contractual guarantees specified in the Contract at its own cost and expense and to carry out further performance tests in accordance with SCC 4, or b. pay liquidated damages to the Procurement Directorate with respect to the failure to meet the contractual guarantees. The rate of these liquidated damages shall be 0.20 percent per day of undelivered materials/goods value up to the sum equivalent to the amount of ten percent of the contract value.
15.	21.4 & 21.5	The period for correction of defects in the warranty period is:
Payment (GCC Clause 22)		
16.	22.1	The method and conditions of payment to be made to the Supplier under this Contract shall be as follows: Payment for Therapeutic Goods and Services supplied from within Pakistan: Payment for Therapeutic Goods and Services supplied from within Pakistan shall be made in Pakistani Rupees, as follows: a. Advance Payment: percent of the Contract Price shall be paid within thirty (30) days of signing of the Contract against a simple receipt and a bank guarantee for the equivalent amount and in the form provided in the Bidding Documents or another form acceptable to the Procurement Directorate. b. On Delivery: percent of the Contract Price shall be paid on receipt of the Therapeutic Goods and upon submission of the documents specified in GCC Clause 11 . c. On Acceptance: The remaining 100% percent of the Contract Price shall be paid to the Supplier within thirty (30) days after the date of the acceptance certificate for the respective delivery issued by the Procurement Directorate.
	22.2	Rate to be used for paying the Supplier's interest on the late payment made by Procurement Directorate shall be <i>[insert: rate]</i> .
Prices (GCC 23)		
17	23.1	Prices shall be adjusted in accordance with provisions in the Attachment to SCC. <i>[To be inserted only if price is subject to adjustment.]</i>
Liquidated Damages (GCC Clause 29)		
18.	29.1	Applicable rate: <i>[insert rate]</i>

		Maximum deduction: is equal to the performance security. Note: 0.1 to 0.2 per cent per day of undelivered materials/good's value.
	Procedure for Dispute Resolution (GCC Clause 34)	
19.	34.3	<p>Dispute Resolution</p> <p>(a) For Contracts to be entered with foreign Contractor/ Service Provider: All disputes arising in connection with the present Contract shall be finally settled under the Rules of Conciliation and Arbitration of the International Chamber of Commerce by one or more arbitrators appointed in accordance with said Rules.</p> <p>(b) For Contracts to be entered with nationals of Pakistan:</p> <ol style="list-style-type: none"> 1. If any dispute of any kind whatsoever shall arise between the Procurement Directorate and the Supplier in connection with or arising out of the Contract, including without prejudice to the generality of foregoing, any question regarding its existence, validity, termination and the execution of the Contract– whether during developing phase or after their completion and whether before or after the termination, abandonment or breach of the Contract – the parties shall seek to resolve any such dispute or difference by mutual diligent negotiations in good faith within 7 (seven) days following a notice sent by one Party to the other Party in this regard. 2. At future of negotiation the dispute shall be resolved through mediation and mediator shall be appointed with the mutual consent of the both parties. 3. At the event of failure of mediation to resolve the dispute relating to this contract such dispute shall finally be resolved through binding Arbitration by sole arbitrator in accordance with Arbitration Act 1940. The arbitrator shall be appointed by mutual consent of the both parties. The Arbitration shall take place in [<i>Insert name of the city</i>] and proceedings will be conducted in –[<i>Specify language</i>] language. 4. The cost of the mediation and arbitration shall be shared by the parties in equal proportion however the both parties shall bear their own costs and lawyer's fees regarding their own participation in the mediation and arbitration. However, the Arbitrator may make an award of costs upon the conclusion of the arbitration making any party to the dispute liable to pay the costs of another party to the dispute. 5. Arbitration proceedings as mentioned in the above clause regarding resolution of disputes may be commenced prior to, during or after delivery of goods. 6. Notwithstanding any reference to the arbitration herein, the parties shall continue to perform their respective obligations under the Contract unless they otherwise agree that the Procurement Directorate shall pay the Supplier any monies due to the Supplier.
	Notices (GCC Clause 38)	
20.	38.1	<p>— Procurement Directorate's address for notice purposes:</p> <p>—Supplier's address for notice purposes:</p>
	Effectiveness of the Contract (GCC Clause 39)	
21.	39.1	The Effective Date of the Contract is [<i>insert: date of Contract signing if EITHER: (i) the Therapeutic Goods have already been registered at the time of Contracting signing OR (ii) registration of the Therapeutic Goods is not a requirement under the Applicable Law. Otherwise, delete and insert "NOT USED."</i>]
	39.2	The time period shall be [<i>insert: a number greater than 30] days. [If not used, delete and insert "NOT USED."]</i>

CONTRACT FORMS

Form of Contract

THIS AGREEMENT made the _____ day of _____ 20____ between *[name and address of Procurement Directorate]* of Pakistan (hereinafter called “the Procurement Directorate”) of the one part and *[name of Supplier]* of *[city and country of Supplier]* (hereinafter called “the Supplier”) of the other part:

WHEREAS the Procurement Directorate invited Bids for certain Therapeutic goods (Pharmaceuticals) and related-services, viz., *[brief description of therapeutic goods and services]* and has accepted a Bid by the Supplier for the supply of those therapeutic goods and related services in the sum of *[contract price in words and figures]* (hereinafter called “the Contract Price”).

NOW THIS CONTRACT WITNESSETH AS FOLLOWS:

1. In this Contract words and expressions shall have the same meanings as are respectively assigned to them in the Conditions of Contract referred to.
2. The following documents shall be deemed to form and be read and construed as part of this Contract, In the event of any ambiguity or conflict between the Contract Documents listed below, the order of precedence shall be the order in which the Contract Documents are listed below:-
 - (a) This form of Contract;
 - (b) the Form of Bid and the Price Schedule submitted by the Bidder;
 - (c) the Schedule of Requirements;
 - (d) the Technical Specifications;
 - (e) the Special Conditions of Contract;
 - (f) the General Conditions of the Contract;
 - (g) the Procurement Directorate's Letter of Acceptance; and
 - (h) *[add here: any other documents]*
3. In consideration of the payments to be made by the Procurement Directorate to the Supplier as hereinafter mentioned, the Supplier hereby covenants with the Procurement Directorate to provide the goods and related services and to remedy defects therein in conformity in all respects with the provisions of the Contract.
4. The Procurement Directorate hereby covenants to pay the Supplier in consideration of the provision of the goods and related services and the remedying of defects therein, the Contract Price or such other sum as may become payable under the provisions of the contract at the times and in the manner prescribed by the contract.

IN WITNESS whereof the parties hereto have caused this Contract to be executed in accordance with their respective laws the day and year first above written.

Signed, sealed, delivered by _____ the _____ (for the Procurement Directorate)

Witness to the signatures of the Procurement Directorate:

.....

Signed, sealed, delivered by _____ the _____ (for the Procurement Directorate)

Witness to the signatures of the Supplier:

Performance Guarantee Form

To: *[name of Procurement Directorate]*

WHEREAS *[name of Supplier]* (hereinafter called "the Supplier") has undertaken, in pursuance of Contract No. *[reference number of the contract]* dated *[insert date]* to delivery *[description of therapeutic goods and services]* (hereinafter called "the Contract").

AND WHEREAS it has been stipulated by you in the said Contract that the Supplier shall furnish you with a Bank Guarantee by a reputable bank for the sum specified therein as security for compliance with the Supplier's performance obligations in accordance with the Contract.

AND WHEREAS we have agreed to give the Supplier a guarantee:

THEREFORE, WE hereby affirm that we are Guarantors and responsible to you, on behalf of the Supplier, up to a total of *[amount of the guarantee in words and figures]*, and we undertake to pay you, upon your first written demand declaring the Supplier to be in default under the Contract and without cavil or argument, any sum or sums within the limits of *[amount of guarantee]* as aforesaid, without your needing to prove or to show grounds or reasons for your demand or the sum specified therein.

This guarantee is valid until the: *[insert date]*

Signature and seal of the Guarantors

[name of bank or financial institution]

[address]

[date]

Integrity Pact

DECLARATION OF FEES, COMMISSION AND BROKERAGE ETC. PAYABLE BY THE SUPPLIERS OF GOODS, SERVICES & WORKS IN CONTRACTS WORTH RS.10.00 MILLION OR MORE

Contract Number: _____

Dated: _____

Contract Value: _____

Contract Title: _____

[Name of Supplier] hereby declares that it has not obtained or induced the procurement of any contract, right, interest, privilege or other obligation or benefit from Government of Pakistan or any administrative subdivision or agency thereof or any other entity owned or controlled by it (GoP) through any corrupt business practice.

Without limiting the generality of the foregoing [Name of Supplier] represents and warrants that it has fully declared the brokerage, commission, fee etc. paid or payable to anyone and not given or agreed to give and shall not give or agree to give to anyone within or outside Pakistan either directly or indirectly through any natural or juridical person, including its affiliate, agent, associate, broker, consultant, director, promoter, shareholder, sponsor or subsidiary, any commission, gratification, bribe, finder's fee or kickback, whether described as consultations fee or otherwise, with the object of obtaining or inducing the procurement of a contract, right, interest, privilege or other obligation or benefit in whatsoever form from GoP, except that which has been expressly declared pursuant hereto.

[Name of Supplier] certifies that it has made and will make full disclosure of all agreements and arrangements with all persons in respect of or related to the transaction with GoP and has not taken any action or will not take any action to circumvent the above declaration, representative or warranty.

[Name of Supplier] accepts full responsibility and strict liability for making and false declaration, not making full disclosure, misrepresenting fact or taking any action likely to defeat the purpose of this declaration, representation and warranty. It agrees that any contract, right interest, privilege or other obligation or benefit obtained or procured as aforesaid shall, without prejudice to any other right and remedies available to GoP under any law, contract or other instrument, be voidable at the option of GoP.

Notwithstanding any rights and remedies exercised by GoP in this regard, [Name of Supplier] agrees to indemnify GoP for any loss or damage incurred by it on account of its corrupt business practices and further pay compensation to GoP in an amount equivalent to ten time the sum of any commission, gratification, bribe, finder's fee or kickback given by [Name of Supplier] as aforesaid for the purpose of obtaining or inducing the procurement of any contract, right, interest, privilege or other obligation or benefit in whatsoever form from GoP.

[Buyer]

[Seller/Supplier]

Tender No. 5171/IT-3035/2024-25/Medicines/FOR//Med/Proc dated 11 October 2024
Procurement of Medicines for HIT Hospital

Appendix I to Section E

Ser	APP Ser	Nomenclature	A/U	Qty Demanded	First Batch	Second Batch
Package 1: Anaesthesia Drugs						
1.	204	Injection - Atracurium Besylate 5mg/ml	Amp	3360	2000	1360
2.	206	Injection - Bupivacaine HCl 10ml	Amp	600	300	300
3.	207	Injection - Bupivacaine HCl 15mg+ Glucose Anhydrous 165 mg Spinal	Amp	1200	800	400
4.	243	Injection - Ketamine 50mg/ml	Amp	480	280	200
5.	247	Injection - Lignocain with Adernalin 10ML	Amp	600	420	180
6.	248	Injection - Lignocaine HCL 10mg	Amp	1800	1200	600
7.	256	Injection - Midazolam 5mg/5ml	Amp	600	450	150
8.	258	Injection - Neopyrolate 0.5mg/ml+Neostigmine 2.5mg	Amp	720	420	300
9.	268	Injection - Propofol 200mg/10ml	Amp	720	320	400
10.	285	Tube - Lignocain Gel 1% 15gm	Tube	1440	840	600

Appendix II to Section E

Ser	APP Ser	Nomenclature	A/U	Qty Demanded	First Batch	Second Batch
Package 1: Anti-Asthmatic						
1.	23	Capsule - Budesonide+Formoterol Fumarate 400/6mcg	No	18960	8960	10000
2.	105	Tablet - Montelukast 10mg	No	19800	14000	5800
3.	106	Tablet - Montelukast 5mg	No	3720	1700	2020
4.	142	Capsule - Tiotropium 18mcg	No	20160	5060	15100
5.	151	Syrup - Acephylline Piperazine and Diphenhydramine 120ml/Cough Prep Syp 120ml	Bott	6240	3240	3000
6.	175	Ipratropium Solution for nebulization bott of 20ml	Bott	780	500	280
7.	176	Beclomethasone Nebules For Nebulization 2ml	Amp	17160	8100	9060
8.	177	Ipratropium Solution for nebulization 2ml	Amp	23400	13000	10400
9.	178	Salbutamol Respiratory Solution 20ml	Bott	504	300	204
10.	184	Sachet - Acetylcysteine	No	1380	580	800
11.	281	Formoterol Fumarate Beclomethasone Dipropionate Inhaler 6/100mcg Inhaler	Amp	720	320	400
12.	282	Salbutamol Inhaler	No	300	180	120
13.	288	Beclomethasone (Dipropionate) 50mcg/actu Nasal Spray	Bott	432	232	200

Appendix III to Section E

Ser	APP Ser	Nomenclature	A/U	Qty Demanded	First Batch	Second Batch
Package 3: Anti-Biotic						
1.	6	Capsule - Amoxicillin 500mg	No	10200	5200	5000
2.	7	Tablet - Amoxicillin+Clavulanic Acid 625mg	No	30720	18000	12720
3.	8	Tablet - Amoxicilline+Clavulanic Acid 1gm	No	2520	1020	1500
4.	9	Tablet - Artemether+Lumefantrine 40/240mg	No	264	0	264
5.	10	Tablet - Artemether+Lumefantrine 80/480mg	No	2640	640	2000
6.	17	Tablet - Azithromycin 250mg	No	39600	9600	30000
7.	24	Tablet - Canesteen V 500mg	No	336	200	136
8.	28	Capsule - Cefaclor 500mg	No	1440	920	520

9.	29	Capsule - Cefixime 400mg	No	7680	2600	5080
10.	30	Capsule - Cefuroxime 250mg	No	2160	1160	1000
11.	33	Capsule - Cephadrine 500mg	No	11760	4560	7200
12.	36	Tablet - Ciprofloxacin 500mg	No	21600	13000	8600
13.	37	Tablet - Clarithromycin 500mg	No	6240	2200	4040
14.	52	Capsule - Doxycycline 100mg	No	9360	3360	6000
15.	66	Capsule - Fluconazole 150mg	No	1920	800	1120
16.	85	Tablet - Levofloxacin 250mg	No	4320	2300	2020
17.	86	Tablet - Levofloxacin 500mg	No	13440	8040	5400
18.	88	Tablet - Linezolid 600mg	No	2640	1640	1000
19.	102	Tablet - Metronidazole 400mg	No	38400	28000	10400
20.	107	Tablet - Moxifloxacin 400mg	No	2160	800	1360
21.	163	Amoxicillin+Clavulanic Acid 156.25/31mg Susp	Bott	4200	2200	2000
22.	164	Syrup - Amoxicillin 125mg/5ml	Bott	360	200	160
23.	165	Amoxicillin+Clavulanic Acid 312.50mg Suspension of 90ml	Bott	360	360	0
24.	166	Syrup - Artemether+Lumefantrine 15/90mg/5ml	Bott	168	100	68
25.	167	Syrup - Azithromycin 200mg/5ml Susp 15ml	Bott	360	200	160
26.	168	Syrup - Cefixime 100mg/5ml 30ml	Bott	3720	2000	1720
27.	169	Syrup - Ciprofloxacin 125mg/5ml	Bott	360	200	160
28.	171	Metronidazole 200mg/5ml Susp of 90ml	Bott	960	560	400
29.	188	Polymyxin b sulphate+Neomycin+Lignocain Skin Ointment 10gm	Tube	3180	2080	1100
30.	189	Polymyxin b sulphate+Neomycin Eye Ointment	Tube	2160	660	1500
31.	191	Povidone-Iodine Scrub 450ml	Bott	192	92	100
32.	192	Povidone-Iodine Solution 60 ml	Bott	1020	620	400
33.	193	Ketoconazole Lotion 60ml	Bott	1020	620	400
34.	195	Povidone-Iodine Solution 450ml	Bott	744	254	490
35.	201	Injection - Amikacin 100mg	Vial	780	500	280
36.	202	Injection - Amikacin 500mg	Vial	2640	2000	640
37.	203	Injection - Amoxicillin+Clavulanic Potassium 1000 / 200gm	Vial	3360	2000	1360
38.	209	Injection -Cefoprazone+Sulbactam 2gm	Vial	2640	1800	840
39.	210	Injection - Cefotaxime 1gm	Vial	2640	0	2640
40.	211	Injection - Cefotaxime 500mg	Vial	360	360	0
41.	212	Injection - Ceftazidime 1gm	Vial	4200	2000	2200
42.	213	Injection - Ceftazidime 500mg	Vial	600	400	200
43.	214	Injection - Ceftriaxone 1gm	Vial	19800	10800	9000
44.	215	Injection - Ceftriaxone 500mg	Vial	1440	840	600
45.	216	Injection - Cefoprazone+Sulbactam 1gm	Vial	1440	840	600
46.	218	Injection - Ciprofloxacin 200mg	Vial	1440	800	640
47.	229	Injection - Gentamycin 80mg	Amp	1140	700	440
48.	246	Injection - Levofloxacin 500mg	Vial	120	80	40
49.	249	Injection - Linezolid 600mg	Bott	720	360	360
50.	252	Injection - Meropenem 1gm	Vial	2160	1400	760
51.	253	Injection - Meropenem 500mg	Vial	1176	600	576
52.	255	Injection - Metronidazole 500mg	Bott	8040	5040	3000
53.	273	Injection - Tazobactam+Pipracillin 0.5gm+4gm	Vial	2160	1400	760
54.	278	Injection - Vancomycin 1gm	Vial	1020	600	420
55.	284	Lignocain+Alcohol Oral Gel	Tube	1440	840	600
56.	286	Gentamicin 0.3% Eye/Ear Drops 10ml	Bott	240	140	100
57.	287	Ciprofloxacin 0.3% w/v Ear Drops	Bott	456	150	306
58.	292	Fusidic Acid+Hydrocortisone 10gm/15 gm	Tube	840	540	300

59.	293	Fusidic Acid Cream 10 gm Tube	Tube	2640	2000	640
60.	294	Terbinafine 10gm Cream	Tube	948	700	248
61.	297	Clotrimazole V 1% Cream of 10/15gm	Tube	672	300	372

Appendix IV to Section E

Ser	APP Ser	Nomenclature	A/U	Qty Demanded	First Batch	Second Batch
Package 4: Anti-Diabetic						
1.	41	Tablet - Dapagliflozin 05mg	No	2160	860	1300
2.	42	Tablet - Dapagliflozin 10mg	No	4320	2300	2020
3.	57	Tablet - Empagliflozin+Linagliptin 25/5mg	No	4320	3320	1000
4.	71	Tablet - Gliclazide MR 30mg Tablet	No	1680	960	720
5.	72	Gliclazide MR 60mg	No	4920	2900	2020
6.	73	Tablet - Glimepride 1mg	No	6720	5000	1720
7.	74	Tablet - Glimepride 2mg	No	34320	20320	14000
8.	75	Tablet - Glimepride 3mg	No	6720	4020	2700
9.	76	Tablet - Glimepride 4mg	No	26520	15500	11020
10.	99	Tablet - Metformin 500mg	No	27000	20000	7000
11.	132	Tablet - Sitagliptin 100mg	No	3120	700	2420
12.	133	Tablet - Sitagliptin 50mg	No	4320	1320	3000
13.	134	Tablet - Sitagliptin+Metformin 50/1000mg	No	22320	12020	10300
14.	135	Tablet - Sitagliptin+Metformin 50/500mg	No	43920	28020	15900
15.	136	Tablet - Sitagliptin+Metformin XR 100/1000mg	No	4320	1300	3020
16.	137	Tablet - Sitagliptin+Metformin XR 50/1000mg	No	5040	2000	3040
17.	138	Tablet - Sitagliptin+Metformin XR 50/500mg	No	4200	2200	2000
18.	149	Tablet - Vildagliptin 50mg	No	10200	6000	4200
19.	150	Tablet - Vildagliptin+Metformin 50/1000mg	No	4200	2000	2200
20.	238	Injection - Human Insulin Regular 10ml	Vial	144	80	64
21.	240	Injection - Humulin 70/30 10ml Vial	Vial	2640	2000	640

Appendix V to Section E

Ser	APP Ser	Nomenclature	A/U	Qty Demanded	First Batch	Second Batch
Package 5: Anti-Histamine						
1.	34	Tablet - Cetrizine 10mg	No	22200	5000	17200
2.	56	Tablet - Ebastine 10mg	No	23400	13400	10000
3.	63	Tablet - Fexofenadine 120mg	No	3840	1840	2000
4.	64	Tablet - Fexofenadine 60mg	No	3360	1300	2060
5.	65	Tablet - Fexofenadine+Pseudoephedrine 120mg/60mg	No	15120	9120	6000
6.	90	Tablet - Loratadin 10mg	No	28320	10300	18020
7.	152	Syrup - Cetrizine 5mg/5ml 120/60ml	Bott	4560	1560	3000
8.	153	Syrup - Chlorpherniramine 5mg/5ml of 120ml	Bott	960	460	500
9.	154	Syrup - Dextromethorphan 10gm+Pseudoephedrine 30mg+Chlorpheniramine Meeate 2mg 120ml	Bott	3360	1360	2000
10.	159	Syrup - Loratadin 5mg/5ml	Bott	720	720	0
11.	220	Injection - Dexamethasone 4mg	Amp	9720	6000	3720
12.	241	Injection - Hydrocortisone 100mg	Vial	17040	12000	5040
13.	291	Cream - Hydrocortisone 1% w/v 10gm	No	504	304	200
14.	296	Cream - Clotrimazole+Hydrocortisone 1% 10gm Tube	Tube	780	280	500

Appendix VI to Section E

Ser	APP Ser	Nomenclature	A/U	Qty Demanded	First Batch	Second Batch
Package 6: Cardiovascular						

1.	3	Tablet - Amlodipine 10mg	No	11760	6760	5000
2.	4	Tablet - Amlodipine 5mg	No	15360	7360	8000
3.	5	Tablet - Amlodipine+Valsartan+Hydrochlorothiazide 5/160/12.5mg	No	2520	1520	1000
4.	11	Tablet - Aspirin 300mg Soluable	No	6600	5000	1600
5.	12	Tablet - Aspirin Enteric Coated 75mg	No	38400	20400	18000
6.	13	Tablet - Atenolol 50mg	No	4320	1320	3000
7.	14	Tablet - Atenolol 50mg+Hydrochlorothiazide 12.5mg	No	3360	1360	2000
8.	15	Tablet - Atorvastatin 10mg	No	21000	11000	10000
9.	16	Tablet - Atorvastatin 20mg	No	19800	9800	10000
10.	20	Tablet - Bisoprolol 2.5mg	No	12240	7000	5240
11.	21	Tablet - Bisoprolol 5mg	No	15000	8000	7000
12.	25	Tablet - Captopril 25mg	No	9000	6000	3000
13.	27	Tablet - Carvedilol 6.25	No	8640	3640	5000
14.	39	Tablet - Clopidogrel 75mg	No	9600	4600	5000
15.	40	Tablet - Clopidogrel+Aspirin 75/75	No	11760	5060	6700
16.	48	Capsule - Diltiazem 90mg SR	No	1560	560	1000
17.	70	Tablet - Furosemide 40mg	No	3120	1800	1320
18.	77	Tablet - Glyceryl Trinitrate 0.5mg	No	1440	440	1000
19.	78	Glyceryl Trinitrate 2.6mg	No	19800	7000	12800
20.	89	Tablet - Lisinopril 5mg	No	2640	640	2000
21.	91	Tablet - Losaratan Potassium 50mg	No	21600	13000	8600
22.	92	Tablet - Losartan Potassium 25 mg	No	5040	4040	1000
23.	93	Tablet - Losartan Potassium+Hydrochlorothiazide 50/12.5mg	No	3720	1700	2020
24.	100	Tablet - Methyldopa 250mg	No	7800	2800	5000
25.	109	Tablet - Nebivilol 2.5mg	No	5040	3000	2040
26.	110	Tablet - Nebivilol 5mg	No	4920	2900	2020
27.	123	Tablet - Propranolol 10mg	No	19800	13000	6800
28.	124	Tablet - Propranolol 40mg	No	5040	2040	3000
29.	128	Tablet - Rosuvastatin 10mg	No	18600	6600	12000
30.	129	Tablet - Rosuvastatin 20mg	No	8160	5100	3060
31.	146	Tablet - Valsartan+Amlodipine 10/160	No	9720	6020	3700
32.	147	Tablet - Valsartan+Amlodipine 5/160mg	No	3360	1360	2000
33.	148	Tablet - Valsartan+Amlodipine 5/80mg	No	26580	18080	8500
34.	200	Injection - Adrenaline 1mg /ml	Amp	600	300	300
35.	205	Injection - Atropine 1mg/ml	Amp	300	300	0
36.	223	Injection - Dopamine 200mg /5ml	Amp	300	220	80
37.	225	Injection - Enoxaparin 40mg	PFS	2016	1500	516
38.	226	Injection - Enoxaparin 60mg	PFS	1440	800	640
39.	228	Injection - Furosemide 20mg	Amp	7320	4020	3300
40.	232	Injection - Glyceryl Trinitrate 10mg/10ml	Amp	480	280	200
41.	233	Injection - Heparin Sodium 5000IU/ml 5ml	Vial	624	324	300

Appendix VII to Section E

Ser	APP Ser	Nomenclature	A/U	Qty Demanded	First Batch	Second Batch
Package 7: Immune Sera & Immunoglonulin						
1.	234	Injection - Human Albumin 100ml	Bott	96	60	36
2.	235	Injection - Human Albumin 50ml	Bott	240	140	100
3.	237	Injection - Human Immunoglobuline 10ml	Bott	144	80	64
4.	239	Injection - Human Rabbits Immunoglobuline 300 IU	PFs	168	90	78
5.	269	Injection - Rho Immunoglobuline 300 mcg	PFs	96	50	46

Appendix VIII to Section E

Ser	APP Ser	Nomenclature	A/U	Qty Demanded	First Batch	Second Batch
Package 8: IV Fluids						
1.	230	Injection - Glucose 5% + Sodium Chloride 0.9%1000ml	Bag	1440	440	1000
2.	231	Injection - Glucose 5% 1000ml	Bag	960	360	600
3.	259	Injection - Normal Saline 1/2 500mg	Bag	360	160	200
4.	265	Injection - Peads Saline 500ml	Bag	480	300	180
5.	266	Injection - Polygeline 500ml	Bag	120	80	40
6.	270	Injection - Ringer Lactate 1000ml	Bag	19800	9800	10000
7.	271	Injection - Sodium Chloride 0.9% 1000ml	Bag	2640	10000	-7360
8.	272	Injection - Sodium Chloride 0.9% 100ml	Bag	50520	29020	21500

Appendix IX to Section E

Ser	APP Ser	Nomenclature	A/U	Qty Demanded	First Batch	Second Batch
Package 9: Miscalleneous						
1.	18	Tablet - Betahistine 16mg	No	5520	1520	4000
2.	19	Tablet - Betahistine 8mg	No	4320	1820	2500
3.	38	Tablet - Clomiphene Citrate 50mg	No	960	460	500
4.	49	Tablet - Dinoprostone 10mg Vaginal Pessary	No	360	200	160
5.	50	Diocahedral Smecitite 3gm Sachet	No	5040	3000	2040
6.	139	Capsule - Tamsulosin Hcl 0.4mg	No	23400	8000	15400
7.	140	Tablet - Tamsulosin Hcl 0.4mg+Dutasteride 0.5mg	No	3360	1360	2000
8.	141	Thyroxin Sodium 50mcg	No	25200	15000	10200
9.	145	Capsule - Tranexamic Acid 500mg	No	3720	2000	1720
10.	157	Syrup - Lactulose 3.35mg/5ml 120ml	Bott	3720	1720	2000
11.	162	Syrup - Aluminium Hydroxide+Magnesium Hydroxide+Simethicone+Carbexymethylcellulose of 120ml	Bott	3120	1120	2000
12.	174	Sodium Alginate 500mg+Calcium Carbonate 160mg+Sodium Bicarbonate 267mg/10ml Suspension	Bott	3120	1800	1320
13.	179	Saccharomyces Boulardii 250mg Sachet	No	5040	3040	2000
14.	185	Polythylen Glycol+Sodium Bicarbonate+Sodium Chloride Sachet	No	2640	1640	1000
15.	190	Cream - Clobederm 10gm Tube	Tube	4320	2300	2020
16.	194	Calamin Lotion 100ml Topical	Bott	624	424	200
17.	197	Methylated Spirit bott of one ltr	Bott	456	156	300
18.	198	Sodium biphosphate 19.2gm+Sodium Phosphate 7.2gm 120ml Rectal Liquid Solution	Bott	744	344	400
19.	236	Injection - Human Diploid Cell Vaccine Rabbies 0.5mg	Vial	780	400	380
20.	262	Injection - Oxytocin 5IU/ml	Amp	10200	4000	6200
21.	274	Injection - Tetanus Toxide 0.5ml /10ml vial	amp	8520	4020	4500
22.	276	Injection - Tranexamic Acid 500mg	Amp	3840	1800	2040
23.	277	Injection - Triamcinolone 40mg/ml	Vial	480	300	180
24.	289	Sodium Bicarbonate+Glycerien Ear Drops 10ml Bott	Bott	576	300	276
25.	290	Cream - Adapeline 0.1% 10gm Tube	No	780	480	300
26.	295	Cream - Silver Sulphadiazine 1% w/v 20gm	Tube	624	224	400
27.	298	Cream - Crotamiton+Sulphur 10% Topical 10gm	Tube	360	360	0
28.	299	Cream - Permethrine 5% w/v 20gm/15gm Tube	Tube	1056	680	376

Appendix X to Section E

Ser	APP Ser	Nomenclature	A/U	Qty Demanded	First Batch	Second Batch
Package 10: Moisturizing Agent						
1.	186	White Soft Parrafine Jar of 100gm	Bott	2640	1640	1000
2.	187	White soft Parafine Oint	Kg	264	64	200
3.	196	Glycerine Pure 50ml Bott	Bott	900	600	300
4.	199	Liquid Parrafine 500ml/100ml Bottle	bott	744	500	244

Appendix XI to Section E

Ser	APP Ser	Nomenclature	A/U	Qty Demanded	First Batch	Second Batch
Package 11: Multivitamins, Supplements & Minerals						
1.	69	Tablet - Folic Acid 5mg Tab bott of 100	No	24000	8000	16000
2.	80	Capsule - Iron III Hydroxide Polymaltose+Vitamin B Complex	No	50760	30060	20700
3.	94	Tablet - M/Vit With Minerals	No	41400	30400	11000
4.	95	Tablet - Mecobalamin 500mcg	No	51000	31000	20000
5.	114	Oral Rehydration Salt Sachet (Flavoured)	No	10200	8200	2000
6.	116	Tablet - Ossein Mineral Complex+ Vit D	No	57840	37040	20800
7.	156	Syrup - Iron Hydroxide+Polymaltose Complex	Bott	780	480	300
8.	160	Syrup - Vitamin B Complex 120ml	Bott	1800	1000	800
9.	161	Syrup - Zinc Sulphate 10mg/5ml of 60ml	Bott	1800	1050	750
10.	172	Ossein Mineral Complex+ Vit D Susp Of 60ml	Bott	600	400	200
11.	180	Myo-Inositol 2000mg+Folic Acid 400mcg Sachet	No	1440	440	1000
12.	181	Cranberry Sachet	No	18240	6200	12040
13.	183	Calcium+Vitamin C Sachet	No	34200	14000	20200
14.	208	Injection - Carboxymaltos 50mg/ml 10ml	Vial	144	64	80
15.	217	Injection - Cholecalciferol 60000IU	Amp	5040	3000	2040
16.	227	Injection - Erythropoietin 10000IU	PFs	60	60	0
17.	242	Injection - Iron sucrose 100mg/5ml	Amp	3120	1420	1700
18.	250	Injection - Magnesium Sulphate 500mg/ml 10ml	Amp	216	100	116
19.	251	Injection - mecobalamin 500mcg	Amp	2640	1040	1600
20.	267	Injection - Potassium Chloride 20ml Amp	Amp	180	180	0
21.	279	Injection - Vitamin B Complex	Amp	2160	1000	1160
22.	280	Injection - Vitamin K 1 ml	Amp	960	560	400

Appendix XII to Section E

Ser	APP Ser	Nomenclature	A/U	Qty Demanded	First Batch	Second Batch
Package 12: Analgesic						
1.	1	Tablet - Aceclofenac 100mg	No	9720	6000	3720
2.	31	Capsule - Celecoxib 100mg	No	7800	2800	5000
3.	32	Capsule - Celecoxib 200mg	No	2640	1240	1400
4.	35	Tablet - Chymotrypsin+Trypsin	No	4560	1560	3000
5.	44	Tablet - Diclofenac Potassium 50mg	No	17520	7520	10000
6.	45	Tablet - Diclofenac Sodium 50mg	No	78000	4800	73200
7.	46	Tablet - Diclofenac Sodium+Misoprostol 50mg	No	9840	2840	7000
8.	47	Tablet - Diclofenac Sodium+Misoprostol 75mg	No	1440	640	800
9.	68	Tablet - Flurbiprofen 100mg	No	6480	4480	2000
10.	79	Tablet - Iburprofen 400mg	No	18600	12000	6600
11.	96	Tablet - Mefenemic Acid 250/500mg	No	25380	5300	20080
12.	97	Tablet - Meloxicam 15mg	No	4320	1320	3000

13.	98	Tablet - Meloxicam 7.5mg	No	2640	1200	1440
14.	108	Tablet - Naproxin Sodium 500mg	No	14640	8040	6600
15.	111	Tablet - Nimsulide 100mg	No	22200	15000	7200
16.	115	Tablet - Orphenadine+Paracetamol 50/650mg	No	100800	62000	38800
17.	118	Tablet - Paracetamol 500mg	No	234000	180000	54000
18.	119	Paracetamol+Tramadol 325/37.5mg	No	26400	16000	10400
19.	120	Tablet - Piroxicam 20mg	No	20160	15060	5100
20.	121	Capsule - Pregablin 50mg	No	5760	2700	3060
21.	122	Capsule - Pregablin 75mg	No	7800	3800	4000
22.	130	Tablet - Seratiopeptidase 10mg	No	12960	8060	4900
23.	143	Tablet - Tizanidine 2mg	No	38400	25400	13000
24.	170	Ibuprofen 200mg/5ml Susp	Bott	6600	3600	3000
25.	173	Syrup - Paracetamol 125mg/5ml	Bott	5160	2160	3000
26.	221	Injection - Diclofenac Sodium 75mg /3ml	Amp	18600	12000	6600
27.	244	Injection - Ketorolac 30mg	Amp	15360	9300	6060
28.	257	Injection - Nalbuphine 10mg Inj	Amp	1320	520	800
29.	264	Paracetamol 1000mg Infusion	Bott	14640	8040	6600
30.	275	Injection - Tramadol 100mg	Amp	1440	900	540
31.	283	Diclofenac Diethylammonium Gel	Tube	2640	1600	1040

Appendix XIII to Section E

Ser	APP Ser	Nomenclature	A/U	Qty Demanded	First Batch	Second Batch
Package 13: Gastroenterology						
1.	43	Capsule - Dexlansoprazole 60mg	No	10320	4020	6300
2.	51	Tablet - Domperidone 10mg	No	39600	19000	20600
3.	53	Tablet - Doxylamine Succinate 10mg+Pyridoxine 10mg	No	7320	3320	4000
4.	54	Tablet - Drotaverine 40mg	No	12600	7000	5600
5.	60	Capsule - Esomeprazole 20mg	No	14640	5040	9600
6.	61	Capsule - Esomeprazole 40mg	No	18720	12020	6700
7.	62	Tablet - Famotidine 40mg	No	10200	4000	6200
8.	81	Tablet - Itopride 50mg	No	31200	20000	11200
9.	82	Capsule - Lansoprazole 30mg	No	2160	1160	1000
10.	87	Tablet - Levosulpride 25mg	No	45600	25000	20600
11.	101	Tablet - Metocopramide 10mg	No	3360	1360	2000
12.	112	Capsule - Omeprazole 20mg	No	69600	39600	30000
13.	113	Capsule - Omeprazole 40mg	No	73800	33000	40800
14.	117	Tablet - Pantoprazole 40mg	No	22320	12300	10020
15.	155	Syrup - Domperidone 5mg/5ml of 60ml	Bott	1260	860	400
16.	182	Omeprazole Insta+Sodium Bicarbonate 20mg Sachet	No	5760	2760	3000
17.	222	Injection - Dimenhydrinate 50mg /1ml	Amp	7800	5000	2800
18.	224	Injection - Drotaverin 40mg	Amp	6720	3720	3000
19.	254	Injection - Metoclopramide 10mg/2ml	Amp	8160	5060	3100
20.	260	Injection - Omeprazole 40mg	Vial	16200	8000	8200
21.	261	Injection - Ondanseteron 8mg/4ml	Amp	6960	3900	3060
22.	263	Injection - Pantoprazole 40mg	Vial	1704	900	804

Appendix XIV to Section E

Ser	APP Ser	Nomenclature	A/U	Qty Demanded	First Batch	Second Batch
Package 14: Anti-Psycotic						
1.	2	Tablet - Alprazolam 0.5mg	No	19440	9240	10200
2.	22	Tablet - Bromazepam 3mg	No	17400	7400	10000

3.	26	Tablet - Carbamazapine 200mg	No	22200	12000	10200
4.	55	Capsule - Duloxetine 30mg	No	4560	2500	2060
5.	58	Tablet - Escitalopram 5mg	No	1800	800	1000
6.	59	Tablet - Esitalopram 10mg	No	6720	3720	3000
7.	67	Capsule - Fluoxetine 20mg	No	19440	12040	7400
8.	83	Tablet - Levetiracetam 250mg	No	1320	720	600
9.	84	Tablet - Levetiracetam 500mg	No	8160	5100	3060
10.	103	Tablet - Mirtazepine 15mg	No	1440	640	800
11.	104	Tablet - Mirtazepine 30mg	No	3720	1500	2220
12.	125	Tablet - Quetiapine 25mg	No	4920	2920	2000
13.	126	Tablet - Risperidone 1mg	No	2520	1500	1020
14.	127	Tablet - Risperidone 2mg	No	2640	1400	1240
15.	131	Tablet - Sertraline 50mg	No	21840	15040	6800
16.	144	Tablet - Topiramate 25mg	No	2640	1640	1000
17.	158	Syrup - Levetiracetam 100mg/5ml of 30ml	Bott	120	100	20
18.	219	Injection - Citocoline 1gm	Amp	360	200	160
19.	245	Injection - Levetiracetam 500mg	Amp	1440	1000	440