INVITATION TO BID

Procurement of medical products and reagents for diagnostics and treatment of children with oncological and oncohematological diseases

For the National Public Health Programme to the Ministry of Health (MoH) in Ukraine

ITB UKR-HP-2017-27

Ukraine



United Nations Development Programme
April 2017

Section 1: Letter of Invitation



ITB UKR-HP-2017-27

Procurement of medical products and reagents for diagnostics and treatment of children with oncological and oncohematological diseases

Dear Bidders,

The Government of Ukraine is in urgent need to secure medicines and essential health commodities at affordable prices and in sufficient quantities. Following recently adopted legislation allowing selected international organizations to provide procurement support services, the Ministry of Health of Ukraine has requested the United Nations Development Programme (UNDP) to support the procurement and distribution of a number of 2016 State Programme medicines and other medical products as an emergency measure.

Therefore, the United Nations Development Programme (UNDP) hereby invites you to submit a Bid in response to this Invitation to Bid (ITB) for the above-referenced subject.

This ITB includes the following documents:

Section 1 – This Letter of Invitation

Section 2 – Instructions to Bidders (including Data Sheet)

Section 3 – Schedule of Requirements and Technical Specifications

Section 4 – Criteria for award and checklist of documents required

Section 5 – Bid Submission Form

Section 6 - Documents Establishing the Eligibility and Qualifications of the Bidder

Section 7 – Technical Bid Form

Section 8 – Price Schedule Form

Section 9 - Form of Bid Security

Section 10 – Form for Performance Security (may be required from winning entity)

Section 11 – General Terms and Conditions for Goods

Your offer, comprising of the Technical and Financial Proposal in one archived file, should be submitted in accordance with Section 2.

You are kindly requested to submit an acknowledgment letter to UNDP to the following address:

United Nations Development Programme in Ukraine health.procurement.ua@undp.org
Attention: Procurement Unit

Mandatory subject of email: ITB- UKR-HP-2017-27 "Procurement of medical products and reagents for diagnostics and treatment of children with oncological and oncohematological diseases"

The letter should be received by UNDP preferably no later than 24.04.2017. The same letter should

advise whether your company intends to submit a Bid. If that is not the case, UNDP would appreciate your indicating the reason, for our records.

If you have received this ITB through a direct invitation by UNDP, transferring this invitation to another firm requires notifying UNDP accordingly.

Should you require any clarification, kindly communicate with the contact person identified in the attached Data Sheet as the focal point for queries on this ITB.

UNDP looks forward to receiving your Bid and thanks you in advance for your interest in UNDP procurement opportunities.

Yours sincerely,

Ms. Andra Brige, UNDP Deputy Country Director Operations

Section 2: Instruction to Bidders

Definitions

- a) "Bid" refers to the Bidder's response to the Invitation to Bid, including the Bid Submission Form, Technical Bid and Price Schedule and all other documentation attached thereto as required by the ITB.
- b) "Bidder" refers to any legal entity that may submit, or has submitted, a Bid for the supply of goods and provision of related services requested by UNDP.
- c) "Contract" refers to the legal instrument that will be signed by and between the UNDP and the successful Bidder, all the attached documents thereto, including the General Terms and Conditions (GTC) and the Appendices.
- d) "Country" refers to the country indicated in the Data Sheet.
- e) "Data Sheet" refers to such part of the Instructions to Bidders used to reflect conditions of the tendering process that are specific for the requirements of the ITB.
- f) "Day" refers to calendar day.
- g) "Goods" refer to any tangible product, commodity, article, material, wares, equipment, assets or merchandise that UNDP requires under this ITB.
- h) "Government" refers to the Government of the country where the goods and related services provided/rendered specified under the Contract will be delivered or undertaken.
- i) "Instructions to Bidders" refers to the complete set of documents which provides Bidders with all information needed and procedures to be followed in the course of preparing their Bid
- j) "ITB" refers to the Invitation to Bid consisting of instructions and references prepared by UNDP for purposes of selecting the best supplier or service provider to fulfil the requirement indicated in the Schedule of Requirements and Technical Specifications.
- k) "LOI" (Section 1 of the ITB) refers to the Letter of Invitation sent by UNDP to Bidders.
- "Material Deviation" refers to any contents or characteristics of the bid that is significantly different from an essential aspect or requirement of the ITB, and (i) substantially alters the scope and quality of the requirements; (ii) limits the rights of UNDP and/or the obligations of the offeror; and (iii) adversely impacts the fairness and principles of the procurement process, such as those that compromise the competitive position of other offerors.
- m) "Schedule of Requirements and Technical Specifications" refers to the document included in this ITB as Section 3 which lists the goods required by UNDP, their specifications, the related services, activities, tasks to be performed, and other information pertinent to UNDP's receipt and acceptance of the goods.
- n) "Services" refers to the entire scope of tasks related or ancillary to the completion or delivery of the goods required by UNDP under the ITB.

o) "Supplemental Information to the ITB" refers to a written communication issued by UNDP to prospective Bidders containing clarifications, responses to queries received from prospective Bidders, or changes to be made in the ITB, at any time after the release of the ITB but before the deadline for the submission of Bid.

A. GENERAL

- 1. UNDP hereby solicits Bids as a response to this Invitation to Bid (ITB). Bidders must strictly adhere to all the requirements of this ITB. No changes, substitutions or other alterations to the rules and provisions stipulated in this ITB may be made or assumed unless it is instructed or approved in writing by UNDP in the form of Supplemental Information to the ITB.
- 2. Submission of a Bid shall be deemed as an acknowledgement by the Bidder that all obligations stipulated by this ITB will be met and, unless specified otherwise, the Bidder has read, understood and agreed to all the instructions in this ITB.
- 3. Any Bid submitted will be regarded as an offer by the Bidder and does not constitute or imply the acceptance of any Bid by UNDP. UNDP is under no obligation to award a contract to any Bidder as a result of this ITB.
- 4. UNDP implements a policy of zero tolerance on proscribed practices, including fraud, corruption, collusion, unethical practices, and obstruction. UNDP is committed to preventing, identifying and addressing all acts of fraud and corrupt practices against UNDP as well as third parties involved in UNDP activities. (See http://www.undp.org/content/dam/undp/library/corporate/Procurement/english/Procurement%20Fraud%20Notice.pdf
 - http://www.undp.org/content/undp/en/home/operations/procurement/protestandsanctions/for full description of the policies)
- 5. In responding to this ITB, UNDP requires all Bidders to conduct themselves in a professional, objective and impartial manner, and they must at all times hold UNDP's interests paramount. Bidders must strictly avoid conflicts with other assignments or their own interests, and act without consideration for future work. All Bidders found to have a conflict of interest shall be disqualified. Without limitation on the generality of the above, Bidders, and any of their affiliates, shall be considered to have a conflict of interest with one or more parties in this solicitation process, if they:
 - 5.1 Are, or have been associated in the past, with a firm or any of its affiliates which have been engaged UNDP to provide services for the preparation of the design, Schedule of Requirements and Technical Specifications, cost analysis/estimation, and other documents to be used for the procurement of the goods and related services in this selection process;
 - 5.2 Were involved in the preparation and/or design of the programme/project related to the goods and related services requested under this ITB; or
 - 5.3 Are found to be in conflict for any other reason, as may be established by, or at the discretion of, UNDP.

In the event of any uncertainty in the interpretation of what is potentially a conflict of interest, Bidders must disclose the condition to UNDP and seek UNDP's confirmation on whether or not such conflict exists.

- 6. Similarly, the following must be disclosed in the Bid:
 - 6.1 Bidders who are owners, part-owners, officers, directors, controlling shareholders, or key personnel who are family of UNDP staff involved in the procurement functions and/or the Government of the country or any Implementing Partner receiving the goods and related services under this ITB; and
 - 6.4 Others that could potentially lead to actual or perceived conflict of interest, collusion or unfair competition practices.

Failure of such disclosure may result in the rejection of the Bid.

- 7. The eligibility of Bidders that are wholly or partly owned by the Government shall be subject to UNDP's further evaluation and review of various factors such as being registered as an independent entity, the extent of Government ownership/share, receipt of subsidies, mandate, access to information in relation to this ITB, and others that may lead to undue advantage against other Bidders, and the eventual rejection of the Bid.
- 8. All Bidders must adhere to the UNDP Supplier Code of Conduct, which may be found at this link: http://web.ng.undp.org/procurement/undp-supplier-code-of-conduct.pdf

B. CONTENTS OF BID

9. Sections of Bid

Bidders are required to complete, sign and submit the documents as per Data Sheet.

10. Clarification of Bid

- 10.1 Bidders may request clarification of any of the ITB documents no later than the number of days indicated in the **Data Sheet** (DS no. 16) prior to the Bid submission date. Any request for clarification must be sent in writing via courier or through electronic means to the UNDP address indicated in the **Data Sheet** (DS no. 17). UNDP will respond in writing, transmitted by electronic means and will transmit copies of the response (including an explanation of the query but without identifying the source of inquiry) to all Bidders who have provided confirmation of their intention to submit a Bid.
- 10.2 UNDP shall endeavor to provide such responses to clarifications in an expeditious manner, but any delay in such response shall not cause an obligation on the part of UNDP to extend the submission date of the Bid, unless UNDP deems that such an extension is justified and necessary.

11. Amendment of Bid

11.1 At any time prior to the deadline for submission of Bid, UNDP may for any reason, such as in response to a clarification requested by a Bidder, modify the ITB in the form of a Supplemental Information to the ITB. All prospective Bidders will be notified in writing of all changes/amendments and additional instructions through Supplemental Information to the ITB and through the method specified in the **Data Sheet** (DS No. 18).

11.2 In order to afford prospective Bidders reasonable time to consider the amendments in preparing their Bid, UNDP may, at its discretion, extend the deadline for submission of Bid, if the nature of the amendment to the ITB justifies such an extension.

C. PREPARATION OF BID

12. Cost

The Bidder shall bear any and all costs related to the preparation and/or submission of the Bid, regardless of whether its Bid was selected or not. UNDP shall in no case be responsible or liable for those costs, regardless of the conduct or outcome of the procurement process.

13. Language

The Bid, as well as any and all related correspondence exchanged by the Bidder and UNDP, shall be written in the language (s) specified in the **Data Sheet** (DS No. 4). Any printed literature furnished by the Bidder written in a language other than the language indicated in the **Data Sheet**, must be accompanied by a translation in the preferred language indicated in the **Data Sheet**. For purposes of interpretation of the Bid, and in the event of discrepancy or inconsistency in meaning, the version translated into the preferred language shall govern. Upon conclusion of a contract, the language of the contract shall govern the relationship between the contractor and UNDP.

14. Bid Submission Form

The Bidder shall submit the Bid Submission Form using the form provided in Section 5 of this ITB.

15. Technical Bid Format and Content

Unless otherwise stated in the **Data Sheet** (DS no. 28), the Bidder shall structure the Technical Bid as follows:

- 15.1 Expertise of Firm/Organization this section should provide details regarding management structure of the organization, organizational capability/resources, and experience of organization/firm, the list of projects/contracts (both completed and ongoing, both domestic and international) which are related or similar in nature to the requirements of the ITB, manufacturing capacity of plant if Bidder is a manufacturer, authorization from the manufacturer of the goods if Bidder is not a manufacturer, and proof of financial stability and adequacy of resources to complete the delivery of goods and provision of related services required by the ITB (see ITB Clause 18 and DS No. 26 for further details). The same shall apply to any other entity participating in the ITB as a Joint Venture or Consortium.
- 15.2 Technical Specifications and Implementation Plan this section should demonstrate the Bidder's response to the Schedule of Requirements and Technical Specifications by identifying the specific components proposed; how each of the requirements shall be met point by point; providing a detailed specification and description of the goods required, plans and drawings where needed; the essential performance characteristics, identifying the works/portions of the work that will be subcontracted; a list of the major subcontractors, and demonstrating how the bid meets or exceeds the requirements,

while ensuring appropriateness of the bid to the local conditions and the rest of the project operating environment during the entire life of the goods provided. Details of technical bid must be laid out and supported by an Implementation Timetable, including Transportation and Delivery Schedule where needed, that is within the duration of the contract as specified in the **Data Sheet** (DSnoS.29 and 30).

Bidders must be fully aware that the goods and related services that UNDP require may be transferred, immediately or eventually, by UNDP to the Government partners, or to an entity nominated by the latter, in accordance with UNDP's policies and procedures. All bidders are therefore required to submit the following in their bids:

- a) A statement of whether any import or export licences are required in respect of the goods to be purchased or services to be rendered, including any restrictions in the country of origin, use or dual use nature of the goods or services, including any disposition to end users;
- Confirmation that the Bidder has obtained license of this nature in the past, and have an expectation of obtaining all the necessary licenses, should their bid be rendered the most responsive; and
- c) Complete documentation, information and declaration of any goods classified or may be classified as "Dangerous Goods".
- 15.3 Management Structure and Key Personnel This section should include the comprehensive curriculum vitae (CVs)of key personnel that will be assigned to support the implementation of the technical bid, clearly defining their roles and responsibilities. CVs should establish competence and demonstrate qualifications in areas relevant to the requirements of this ITB.

In complying with this section, the Bidder assures and confirms to UNDP that the personnel being nominated are available to fulfil the demands of the Contract during its stated full term. If any of the key personnel later becomes unavailable, except for unavoidable reasons such as death or medical incapacity, among other possibilities, UNDP reserves the right to render the Bid non-responsive. Any deliberate substitution of personnel arising from unavoidable reasons, including delay in the implementation of the project of programme through no fault of the Bidder, shall be made only with UNDP's acceptance of the justification for substitution, and UNDP's approval of the qualification of the replacement who shall be either of equal or superior credentials as the one being replaced.

- 15.4 Where the **Data Sheet** requires the submission of the Bid Security, the Bid Security shall be included along with the Technical Bid. The Bid Security may be forfeited by UNDP, and reject the Bidthe bid rejected, in the event of any or any combination of the following conditions:
 - a) If the Bidder withdraws its offer during the period of the Bid Validity specified in the **Data Sheet** (DS no. 11), or;
 - b) If the Bid Security amount is found to be less than what is required by UNDP as indicated in the **Data Sheet** (DS no. 9), or;
 - c) In the case the successful Bidder fails:
 - i. to sign the Contract after UNDP has awarded it;
 - ii. to comply with UNDP's variation of requirement, as per ITB Clause 35; or

iii. to furnish Performance Security, insurances, or other documents that UNDP may require as a condition to rendering effective the contract that may be awarded to the Bidder.

16. Price Schedule

The Price Schedule shall be prepared using the attached standard form (Section 8). It shall list all major cost components associated with the goods and related services, and the detailed breakdown of such costs. All goods and services described in the Technical Bid must be priced separately on a one-to-one correspondence. Any output and activities described in the Technical Bid but not priced in the Price Schedule, shall be assumed to be included in the prices of the items or activities, as well as in the final total price of the bid.

17. Currencies

All prices shall be quoted in the currency indicated in the **Data Sheet** (DS no. 15). However, where Bids are quoted in different currencies, for the purposes of comparison of all Bid:

- 17.1 UNDP will convert the currency quoted in the Bid into the UNDP preferred currency, in accordance with the prevailing UN operational rate of exchange on the last day of submission of Bid; and
- 17.2 In the event that the Bid found to be the most responsive to the ITB requirement is quoted in another currency different from the preferred currency as per **Data Sheet** (DS no. 15), then UNDP shall reserve the right to award the contract in the currency of UNDP's preference, using the conversion method specified above.

18. Documents Establishing the Eligibility and Qualifications of the Bidder

- 18.1 The Bidder shall furnish documentary evidence of its status as an eligible and qualified vendor, using the forms provided under Section 5, Bidder Information Forms. In order to award a contract to a Bidder, its qualifications must be documented to UNDP's satisfactions. These include, but are not limited to the following:
 - a) That, in the case of a Bidder offering to supply goods under the Contract which the Bidder did not manufacture or otherwise produce, the Bidder has been duly authorized by the goods' manufacturer or producer to supply the goods in the country of final destination;
 - b) That the Bidder has the financial, technical, and production capability necessary to perform the Contract; and
 - c) That, to the best of the Bidder's knowledge, it is not included in the UN 1267 List or the UN Ineligibility List, nor in any and all of UNDP's list of suspended and removed vendors.
- 18.2 Bids submitted by two (2) or more Bidders shall all be rejected by UNDP if they are found to have <u>any</u> of the following:
 - a) they have at least one controlling partner, director or shareholder in common; or
 - b) any one of them receive or have received any direct or indirect subsidy from the other/s; or
 - c) they have the same legal representative for purposes of this ITB; or

- d) they 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 regarding this ITB process;
- e) they are subcontractors to each other's bid, or a subcontractor to one bid also submits another Bid under its name as lead Bidder; or
- f) an expert proposed to be in the bid of one Bidder participates in more than one Bid received for this ITB process. This condition does not apply to subcontractors being included in more than one Bid.

19. Joint Venture, Consortium or Association

If the Bidder is a group of legal entities that will form or have formed a joint venture, consortium or association at the time of the submission of the Bid, they shall confirm in their Bid that: (i) they have designated one party to act as a lead entity, duly vested with authority to legally bind the members of the joint venture jointly and severally, and this shall be duly evidenced by a duly notarized Agreement among the legal entities, which shall be submitted along with the Bid; and (ii) if they are awarded the contract, the contract shall be entered into, by and between UNDP and the designated lead entity, who shall be acting for and on behalf of all entities that comprise the joint venture.

After the bid has been submitted to UNDP, the lead entity identified to represent the joint venture shall not be altered without the prior written consent of UNDP. Furthermore, neither the lead entity nor the member entities of the joint venture can:

- a) Submit another Bid, either in its own capacity; nor
- b) As a lead entity or a member entity for another joint venture submitting another Bid. The description of the organization of the joint venture/consortium/association must clearly define the expected role of each of the entity in the joint venture in delivering the requirements of the ITB, both in the bid and in the Joint Venture Agreement. All entities that comprise the joint venture shall be subject to the eligibility and qualification assessment by UNDP.

Where a joint venture is presenting its track record and experience in a similar undertaking as those required in the ITB, it should present such information in the following manner:

- a) Those that were undertaken together by the joint venture; and
- b) Those that were undertaken by the individual entities of the joint venture expected to be involved in the performance of the services defined in the ITB.

Previous contracts completed by individual experts working privately but who are permanently or were temporarily associated with any of the member firms cannot be claimed as the experience of the joint venture or those of its members, but should only be claimed by the individual experts themselves in their presentation of their individual credentials.

If the Bid of a joint venture is determined by UNDP as the most responsive Bid that offers the best value for money, UNDP shall award the contract to the joint venture, in the name of its designated lead entity, who shall sign the contract for and on behalf of all the member entities.

20. Alternative Bid

Unless otherwise specified in the Data Sheet (DS nos. 5 and 6), alternative bid shall not be

considered. Where the conditions for its acceptance are met, or justifications are clearly established, UNDP reserves the right to award a contract based on an alternative bid.

21. Validity Period

- 21.1 Bid shall remain valid for the period specified in the **Data Sheet** (DS no. 8), commencing on the submission deadline date also indicated in the **Data Sheet** (DS no. 21). A Bid valid for a shorter period shall be immediately rejected by UNDP and rendered non-responsive.
- 21.2 In exceptional circumstances, prior to the expiration of the Bid validity period, UNDP may request Bidders to extend the period of validity of their Bid. The request and the responses shall be made in writing, and shall be considered integral to the Bid.

22. Bidder's Conference

When appropriate, a Bidder's conference will be conducted at the date, time and location specified in the **Data Sheet** (DS no. 7). All Bidders are encouraged to attend. Non-attendance, however, shall <u>not</u> result in disqualification of an interested Bidder. Minutes of the Bidder's conference will be either posted on the UNDP website, or disseminated to the individual firms who have registered or expressed interest with the contract, whether or not they attended the conference. No verbal statement made during the conference shall modify the terms and conditions of the ITB unless such statement is specifically written in the Minutes of the Conference, or issued/posted as an amendment in the form of a Supplemental Information to the ITB.

D. SUBMISSION AND OPENING OF BID

23. Submission

- 23.1 The Technical Bid and the Price Schedule <u>must be submitted</u> by electronic method of transmission at the Bid submission address indicated in the data sheet.
- 23.2 Bidders must submit their Bid in the manner specified in the **Data Sheet** (DS nos. 22 and 23). When the Bid is expected to be in transit for more than 24 hours, the Bidder must ensure that sufficient lead time has been provided in order to comply with UNDP's deadline for submission. UNDP shall indicate for its record that the official date and time of receiving the Bid is the <u>actual</u> date and time when the said Bid has physically arrived at the UNDP premises indicated in the **Data Sheet** (DS no. 20).
- 23.3 Bidders submitting Bid by mail or by hand shall enclose the original and each copy of the Bid, in separate sealed envelopes, duly marking each of the envelopes as "Original Bid" and the others as "Copy of Bid". The two envelopes, consisting of original and copies, shall then be sealed in an outer envelope. The number of copies required shall be as specified in the **Data Sheet** (DS no. 19). In the event of any discrepancy between the contents of the "Original Bid" and the "Copy of Bid", the contents of the original shall govern. The original version of the Bid shall be signed or initialed by the Bidder or person(s) duly authorized to commit the Bidder on every page. The authorization shall be communicated through a document evidencing such authorization issued by the highest official of the firm, or a Power of Attorney, accompanying the Bid.

23.4 Bidders must be aware that the mere act of submission of a Bid, in and of itself, implies that the Bidder accepts the General Contract Terms and Conditions of UNDP as attached hereto as Section 11.

24. Deadline for Submission of Bid and Late Bids

Bid must be received by UNDP at the address and no later than the date and time specified in the **Data Sheet** (DSno.20 and 21).

UNDP shall not consider any Bid that arrives after the deadline for submission of Bid. Any Bid received by UNDP after the deadline for submission of Bid shall be declared late, rejected, and returned unopened to the Bidder.

25. Withdrawal, Substitution, and Modification of Bid

- 25.1 Bidders are expected to have sole responsibility for taking steps to carefully examine in detail the full consistency of its Bid to the requirements of the ITB, keeping in mind that material deficiencies in providing information requested by UNDP, or lack clarity in the description of goods and related services to be provided, may result in the rejection of the Bid. The Bidder shall assume any responsibility regarding erroneous interpretations or conclusions made by the Bidder in the course of understanding the ITB out of the set of information furnished by UNDP.
- 25.2 A Bidder may withdraw, substitute or modify its Bid after it has been submitted by sending a written notice in accordance with ITB Clause 23, duly signed by an authorized representative, and shall include a copy of the authorization (or a Power of Attorney). The corresponding substitution or modification of the Bid must accompany the respective written notice. All notices must be received by UNDP prior to the deadline for submission and submitted in accordance with ITB Clause 23 (except that withdrawal notices do not require copies). The respective envelopes shall be clearly marked "WITHDRAWAL," "SUBSTITUTION," or MODIFICATION".
- 25.3 Bid requested to be withdrawn shall be returned unopened to the Bidders.
- 25.4 No Bid may be withdrawn, substituted, or modified in the interval between the deadline for submission of Bid and the expiration of the period of Bid validity specified by the Bidder on the Bid Submission Form or any extension thereof.

26. Bid Opening

UNDP will open the Bid in the presence of an ad-hoc committee formed by UNDP of at least two (2) members. If electronic submission is permitted, any specific electronic Bid opening procedures shall be as specified in the **Data Sheet** (DS no. 23).

The Bidders' names, modifications, withdrawals, the condition of the envelope labels/seals, the number of folders/files and all other such other details as UNDP may consider appropriate, will be announced at the opening. No Bid shall be rejected at the opening stage, except for late submission, for which the Bid shall be returned unopened to the Bidder.

27. Confidentiality

Information relating to the examination, evaluation, and comparison of Bid, and the recommendation of contract award, shall not be disclosed to Bidders or any other persons not officially concerned with such process, even after publication of the contract award.

Any effort by a Bidder to influence UNDP in the examination, evaluation and comparison of the Bid or contract award decisions may, at UNDP's decision, result in the rejection of its Bid.

If a Bidder is unsuccessful, the Bidder may seek a meeting with UNDP for a debriefing. The purpose of the debriefing is discussing the strengths and weaknesses of the Bidder's submission, in order to assist the Bidder in improving the bid presented to UNDP. The content of other bid and how they compare to the Bidder's submission shall not be discussed.

E. EVALUATION OF BID

28. Preliminary Examination of Bid

UNDP shall examine the Bid to determine whether they are complete with respect to minimum documentary requirements, whether the documents have been properly signed, whether or not the Bidder is in the UN Security Council 1267/1989 Committee's list of terrorists and terrorist financiers, and in UNDP's list of suspended and removed vendors, and whether the Bid are generally in order, among other indicators that may be used at this stage. UNDP may reject any Bid at this stage.

29. Evaluation of Bid

- 29.1 UNDP shall examine the Bid to confirm that all terms and conditions under the UNDP General Terms and Conditions and Special Conditions have been accepted by the Bidder without any deviation or reservation.
- 29.2 The evaluation team shall review and evaluate the Bids on the basis of their responsiveness to the Schedule of Requirements and Technical Specifications and other documentation provided, applying the procedure indicated in the **Data Sheet** (DS No. 25). Absolutely no changes may be made by UNDP in the criteria after all Bids have been received.
- 29.1 UNDP reserves the right to undertake a post-qualification exercise, aimed at determining, to its satisfaction the validity of the information provided by the Bidder. Such post-qualification shall be fully documented and, among those that may be listed in the **Data Sheet** (DS No.33), may include, but need not be limited to, all or any combination of the following:
 - a) Verification of accuracy, correctness and authenticity of the information provided by the bidder on the legal, technical and financial documents submitted;
 - b) Validation of extent of compliance to the ITB requirements and evaluation criteria based on what has so far been found by the evaluation team;
 - c) Inquiry and reference checking with Government entities with jurisdiction on the bidder, or any other entity that may have done business with the bidder;
 - d) Inquiry and reference checking with other previous clients on the quality of performance on on-going or previous contracts completed;

- e) Physical inspection of the bidder's plant, factory, branches or other places where business transpires, with or without notice to the bidder;
- f) Testing and sampling of completed goods similar to the requirements of UNDP, where available; and
- g) Other means that UNDP may deem appropriate, at any stage within the selection process, prior to awarding the contract.

30. Clarification of Bid

To assist in the examination, evaluation and comparison of bids, UNDP may, at its discretion, ask any Bidder to clarify its Bid.

UNDP's request for clarification and the Bidder's response shall be in writing. Notwithstanding the written communication, no change in the prices or substance of the Bid shall be sought, offered, or permitted, except to provide clarification, and confirm the correction of any arithmetic errors discovered by UNDP in the evaluation of the Bid, in accordance with ITB Clause 35.

Any unsolicited clarification submitted by a Bidder in respect to its Bid, which is not a response to a request by UNDP, shall not be considered during the review and evaluation of the Bid.

31. Responsiveness of Bid

UNDP's determination of a Bid's responsiveness will be based on the contents of the Bid itself.

A substantially responsive Bid is one that conforms to all the terms, conditions, and specifications of the ITB without material deviation, reservation, or omission.

If a Bid is not substantially responsive, it shall be rejected by UNDP and may not subsequently be made responsive by the Bidder by correction of the material deviation, reservation, or omission.

32. Nonconformities, Reparable Errors and Omissions

- 32.3 Provided that a Bid is substantially responsive, UNDP may waive any non-conformities or omissions in the Bid that, in the opinion of UNDP, do not constitute a material deviation.
- 32.4 Provided that a Bid is substantially responsive, UNDP 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 Bid related to documentation requirements. Such omission shall not be related to any aspect of the price of the Bid. Failure of the Bidder to comply with the request may result in the rejection of its Bid.
- 32.5 Provided that the Bid is substantially responsive, UNDP shall correct arithmetical errors as follows:
 - a) if there is a discrepancy between the unit price and the line item total that is obtained by multiplying the unit price by the quantity, the unit price shall prevail and the line item total shall be corrected, unless in the opinion of UNDP there is an obvious misplacement of the decimal point in the unit price, in which case the line item total 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 subtotals, the subtotals shall prevail and the total shall be corrected; and
- c) if there is a discrepancy between words and figures, the amount in words shall prevail, unless the amount expressed in words is related to an arithmetic error, in which case the amount in figures shall prevail subject to the above.
- 32.6 If the Bidder does not accept the correction of errors made by UNDP, its Bid shall be rejected.

F. AWARD OF CONTRACT

33. Right to Accept, Reject, or Render Non-Responsive Any or All Bid

- 33.1 UNDP reserves the right to accept or reject any Bid, to render any or all of the Bids as non-responsive, and to reject all Bids at any time prior to award of contract, without incurring any liability, or obligation to inform the affected Bidder(s) of the grounds for UNDP's action. Furthermore, UNDP is not obligated to award the contract to the lowest price offer.
- 33.2 UNDP shall also verify, and immediately reject their respective Bid, if the Bidders are found to appear in the UN's Consolidated List of Individuals and Entities with Association to Terrorist Organizations, in the List of Vendors Suspended or Removed from the UN Secretariat Procurement Division Vendor Roster, the UN Ineligibility List, and other such lists that as may be established or recognized by UNDP policy on Vendor Sanctions. (See http://www.undp.org/content/undp/en/home/operations/procurement/

34. Award Criteria

nt protest/

Prior to expiration of the period of Bid validity, UNDP shall award the contract to the qualified and eligible Bidder that is found to be responsive to the requirements of the Schedule of Requirements and Technical Specification, and has offered the lowest price (See DS No. 32).

35. Right to Vary Requirements at the Time of Award

At the time of award of Contract, UNDP reserves the right to vary the quantity of the goods and/or related services, by up to a maximum twenty five per cent (25%) of the total offer, without any change in the unit price or other terms and conditions.

36. Contract Signature

Within fifteen (15) days from the date of receipt of the Contract, the successful Bidder shall sign and date the Contract and return it to UNDP.

Failure of the successful Bidder to comply with the requirement of ITB Section F.3 and this provision shall constitute sufficient grounds for the annulment of the award, and forfeiture of the Bid Security if any, and on which event, UNDP may award the Contract to the Bidder with the second highest rated Bid, or call for new Bid.

37. Performance Security

A performance security, if required, shall be provided in the amount and form provided in Section 9 and by the deadline indicated in the **Data Sheet** (DS no. 14), as applicable. Where a Performance Security will be required, the submission of the said document, and the confirmation of its acceptance by UNDP, shall be a condition for the effectivity of the Contract that will be signed by and between the successful Bidder and UNDP.

38. Bank Guarantee for Advanced Payment

Except when the interests of UNDP so require, it is the UNDP's preference to make no advanced payment(s) on contracts (i.e., payments without having received any outputs). In the event that the Bidder requires an advanced payment upon contract signature, and if such request is duly accepted by UNDP, and the said advanced payment exceeds 20% of the total Bid price, or exceed the amount of USD 30,000, UNDP shall require the Bidder to submit a Bank Guarantee in the same amount as the advanced payment.

39. Vendor Protest

UNDP's vendor protest procedure provides an opportunity for appeal to those persons or firms not awarded a purchase order or contract through a competitive procurement process. In the event that a Bidder believes that it was not treated fairly, the following link provides further details regarding UNDP vendor protest procedures:

http://www.undp.org/procurement/protest.shtml

Instructions to Bidders

DATA SHEET

The following data for the supply of goods and related services shall complement / supplement the provisions in the Instruction to Bidders. In the case of a conflict between the Instruction to Bidders and the Data Sheet, the provisions in the **Data Sheet shall prevail.**

DS No.	Cross Ref. to Instructions	Data	Specific Instructions / Requirements
1		Project Title:	Procurement support to the Ministry of Health of Ukraine
2		Title of Goods/Services/Work Required:	Procurement of medical products and reagents for diagnostics and treatment of children with oncological and oncohematological diseases in 160 lots In accordance with the Technical Specifications as
			per Section 3.
3		Country:	Ukraine
4	C.13		☑ English
		Language of the Bid:	As this particular procurement case is subject to review and approval by UNDP HQ Advisory Committee on Procurement, Members of which are English speakers, the Bidders are requested to submit their Bids in English.
			☑ Ukrainian/Russian
5	C.20	Conditions for Submitting Bid for Parts or sub-parts of the Total Requirements	⊠ Not allowed
6	C.20	Conditions for Submitting Alternative Bid	☑ Shall not be considered
7	C.22	A pre-Bid conference will be held on:	Time: 16:00 hrs local time Date: April 25, 2017 Venue: Alexanian conference hall (Operations building); UN Office in Ukraine; 1 Klovskyi Street Companies can participate at pre-bid conference
			through skype conference as well. Interested companies should send confirmations by email.

8	C.21.1	Period of Bid Validity	,
		commencing on the submission date	□ 90 days☑ 120 days
9	B.9.5 C.15.4 b)	Bid Security	□ Required Bid security is required in the amount of:
			USD 10,000 for a bid for one or more lots cumulatively exceeding 200,000.00 USD up to USD 399,999.99
			OR USD 20,000 for a bid for one or more lots cumulatively exceeding 400,000.00 USD
10	B.9.5	Acceptable forms of Bid Security	 ☑ Bank Guarantee Bid Security shall be submitted in the form of Bank Guarantee as per template provided in the Section 9
11	B.9.5 C.15.4 a)	Validity of Bid Security	150 days
12		Advanced Payment upon signing of contract	⊠ Not allowed
13		Liquidated Damages	If the Supplier fails to supply the specified Goods within the time period(s) stipulated in the individual contract (Purchase Orders), the UNDP may without prejudice to its other remedies under the contract, deduct 0.5% of the complete consignment for each day of delay until actual delivery, up to maximum deduction of 10% of the value of the Purchase Order. Once the maximum is reached, UNDP may consider termination of the PO.

14	F.37	Performance Security	✓ May be required from winning entity for all
			contracts (Purchase Orders) exceeding 300,000 USD as per template provided in the Section 10 Amount: 10 % of the contract amount Form: Bank guarantee.
15	C.17 C.17.2	Preferred Currency of Bid and Method for Currency conversion	☑ United States Dollars (USD) - strongly advised to use as a risk mitigation measure against the impact of the local currency devaluation.
			UNDP will execute payments in USD to international suppliers.
			Payments to local (Ukrainian) suppliers will be executed either in USD or UAH based on UN Operational Exchange Rate effective at the date of payment (please refer to treasury.un.org).
			☑ Local Currency (UAH)
			Prices submitted by Bidders will be evaluated versus each other based on UN Operational exchange rate effective at the closure day of the bid submission (please refer to treasury.un.org)
16	B.10.1	Deadline for submitting requests for clarifications/ questions	5 calendar days before the submission date.
17	B.10.1	Contact Details for submitting clarifications/questions	Focal Person in UNDP: Procurement Unit Tel. No. :+38 044 253 59 84
			E-mail address dedicated for this purpose: health.procurement.ua@undp.org
18	B.11.1	Manner of Disseminating Supplemental Information to the ITB and responses/clarifications to queries	☑ Direct communication to prospective Bidders by email, and Posting on the website http://procurement-notices.undp.org
19	D.23.3	No. of copies of Bid that must be submitted	1 (one)
20	D.23.1 b) D.23.2	Bid submission address	tenders.ua@undp.org
	D.24		Please note that bids received through any other address will not be considered.
			Any bid sent to the private email addresses of any UNDP staff will not be accepted.

21	C.21.1 D.24	Deadline of Bid Submission	Date and Time:May 19, 2017 10:00 AM, Kyiv time (UTC +2:00) Attention, the date is changed!
22	D.23.2	Manner of Submitting Bid	☑ Electronic submission of Bid for technical and financial offers
23	D.23.2 D.26	Conditions and Procedures for electronic submission and opening, if allowed	 ☑ Official Address for e-submission: tenders.ua@undp.org ☑ Format: PDF files preferred in ZIP archives only. ☑ Max. File Size per transmission: [5 MB] ☑ Max. No. of transmission: [10] No. of copies to be transmitted: [1] ☑ Mandatory subject of email: ITB-UKR-HP-2017-27 "Procurement of medical products and reagents for diagnostics and treatment of children with oncological and oncohematological diseases" ☑ Virus Scanning Software to be Used prior to transmission: [Files should not contain any viruses or malware software.] ☑ Time Zone to be Recognized: [UTC+2, Kyiv time] ☑ Other conditions: N/A PLEASE make all efforts to provide your proposal in 1 archived PDF file not exceeding 5 MB size. Bidders are solely responsible for ensuring that any and all files sent to UNDP are readable, that is, uncorrupted, in the indicated electronic format, and free from viruses and malware. Failure to provide readable files will result in the Bid being rejected. Please take into consideration the fact that emails are delivered within 5-10 mins, therefore avoid last minute submission, which might lead to late submission.
24	D.23.1 c)	Date, time and venue for opening of Bid	Date and Time: May 19, 2017 2:00 PM, Kyiv time (UTC +2:00) Attention, the date is changed! Any bidder that intends to participate in the public bid opening shall notify UNDP by address health.procurement.ua@undp.org at least 24 hours in advance.
			Companies can participate at the Bid Opening procedure through skype conference as well.

			Interested companies should send confirmations by email.
			Venue: Alexanian conference hall (Operations building); UN Office in Ukraine; 1 Klovskyi Street
25		Evaluation method to be used in selecting the most responsive Bid	As per DS # 32
26	C.15.1	Required Documents that must be Submitted to Establish Qualification of Bidders	 ☑ Duly filled-in, signed and stamped Sections 4-8. ☑ Copy of properly furnished Bid Security (as per DS# 9). Please use template provided in the Section 9.
			Original should be provided to UNDP within 1 week after the Deadline of Bid Submission (DS #21) to below address, otherwise the bid will be rejected: 1 Klovskyi Uzviz, Kyiv, Ukraine 01021 To: UNDP Procurement Unit
			☑ Copies of required documents to establish conformity of Bidder to the qualifications requirements and products quoted to product standards and requirements as per Section 4 "Criteria for award and checklist of documents required" and Section 3 "Schedule of Requirements and Technical Specifications"
27		Other documents that may be Submitted to Establish Eligibility	N/A
28	C.15	Structure of the Technical Bid and List of Documents to be Submitted	As per DS # 26.
29	C.15.2	Latest Expected date for commencement of Contract	June 12, 2017
30	C.15.2	Maximum Expected duration of contract	As per Deadlines described in the Section 3
31		UNDP will award the contract to:	☑ One or more Bidders, depending on the following factors: Lowest-priced technically responsive offer per Lot
			*) the discount factor will be considered if such proposed by Bidder for awarding of more than one lot only at the stage of contracting and will not be considered for evaluation purposes.

32	F.34	Criteria for the Award and Evaluation of Bid	Award Criteria ☑ Non-Discretionary "Pass/Fail" Qualifying Criteria on the requirements listed in the Section 4 "Criteria for award and checklist of documents required" AND ☑ Lowest price offer of technically qualified/responsive Bid
33	E.29	Post qualification Actions	 ☑ Verification of accuracy, correctness and authenticity of the information provided by the bidder on the legal, technical and financial documents submitted; ☑ Validation of extent of compliance to the ITB requirements and evaluation criteria based on what has so far been found by the evaluation team; ☑ Inquiry and reference checking with other previous clients on the quality of performance on ongoing or previous contracts completed;
34		Conditions for Determining Contract Effectivity	☑ Provision of Performance Security, if required
35		Other Information Related to the ITB	Administrative Requirements: Prior to technical evaluation, submitted offers will be reviewed on a "Pass" or "Fail" basis to determine compliance with the below formal criteria/requirements: Bids must be submitted within the stipulated deadline; Bids must meet required Bid Validity; Bids must include copy of properly furnished Bid Security (as per DS 9). Original should be provided within 1 week after the Deadline of Bid Submission (as per DS #21), otherwise the Bid will be rejected. Bids have been signed by the proper authority Full compliance and agreement with UNDP General terms and conditions available by the link: http://www.undp.org/content/dam/undp/documents/procurement/documents/genconditionpurchaseorders.pdf. Other information is available on http://www.ua.undp.org/content/ukraine/en/home/operations/procurement.html For information, please contact procurement.ua@undp.org

Section 3:

Schedule of Requirements and Technical Specifications

1. EXECUTIVE SUMMARY

In April 2015, the Ministry of Health of Ukraine approached the UN System in Ukraine to support the procurement and distribution of medicines and other medical products in scope of health state programs as an emergency measure. This new approach to procurement in the public health sector was aimed to prevent corruption and protect the rights of patients in Ukraine to access affordable and quality medicines.

Under the budget of 2015, UNDP successfully procured medicines and medical products for eight state health programs, managing to achieve significant savings and deliver additional quantities. In 2016 UNDP Ukraine continues to support the Ministry of Health of Ukraine with procurement of essential and vital medicines for 23 programs.

UNDP in Ukraine is fully committed to play its role in resolving the immediate crisis and to support the Ministry of Health of Ukraine in its efforts to reform the procurement and supply management system for it to correspond to the highest standards of transparency, accountability, cost-efficiency, equity and sustainability.

The main objective of this ITB is to source high quality medical supplies from reliable suppliers and in accordance with the value-for-money principle needed to meet the current health crisis. This ITB targets to source medical products and reagents for diagnostics and treatment of children with oncological and oncohematological diseases.

2. PRODUCT STANDARDS

In view of the specific emergency experienced by the country, and the urgency with which UNDP has been requested to procure these medical products, these standards below are specific for this procurement action and in no way, constitute an obligation from UNDP to use any of these standards in future procurement actions.

<u>UNDP</u> will procure the medical product only under the following product standards quality criteria (Option 1 or Option 2):

OPTION 1 (1.1.+1.2.)

1.1. Medical products must be produced and controlled in accordance with product standards and quality system standards recommended by the World Health Organization (WHO) AND/OR

the International Medical Devise Regulators Forum (IMDRF) (former Global Harmonization Task Force (GHTF). For more information see http://www.imdrf.org/. The GHTF founding members are Australia, Canada, the European Union (EU), Japan and the United States of America (USA). To be compliant with this criterion bidders are requested to provide at least one of

the following pre-market approval(s) / market clearance(s)/registration(s):

- Canada Medical Device license, OR;
- European Union EC Full Quality Assurance Certificate or EC Production Quality Assurance
 Certificate cate or EC Type-Examination Certificate (CE/ Conformité Européenne mark) or /
 Conformité Européenne 92/42 or CE/ Conformité Européenne 98/79, OR;
- Australia TGA Production Quality Assurance Certificate or TGA Type-Examination Certificate or TGA Full Quality Assurance Certificate issued by Therauptic Goods Administration, OR;
- Japan PMDA (Pharmaceuticals and Medical Devices Agency) approval or JMHLW (Japan Ministry of Health, Labour and Welfare) Minister's approval, OR;
- USA PMA (pre-market approval) letter or BLA license (Biologics License Application) or 510k device letter issued by US Food and Drug Administration.

NB: In case product is considered RUO "for research use only", UNDP will conduct additional verification of product's compliance to the products standards. If Bidder proposes RUO product, evidence of at least on successful supply within countries-founding members of GHTF shall be provided from the medical institution/diagnostic center/laboratory center (since March 2014).

In case within one lot UNDP will obtain proposals for RUO "for research use only", ASR "analyte-specific reagent" and IVD 'in-vitro diagnostics", preference will be given to ASR and IVD product(s).

AND

- 1.2.a. Suppliers and manufacturers must provide an evidence of conformity* to at least one of the following conformity with the following Quality Management System standards as recognized by GHTF standards:
 - ISO13485/ISO 13488, or
 - ISO 9001, or
 - United States QS 21 CFR part 820, or
 - Japan QS Standard for medical devices.
- *) The evidence(s) of conformity shall indicate, as applicable:
 - a) Manufacturer's certified quality management system standard(s)
 - b) Assessment body (name, country)
 - c) Last audit date and
 - d) Expiration date
 - e) Certificate number

OR

1.2.b. Suppliers/manufacturers shall provide a copy of valid GMP Certificate issued by The Pharmaceutical Inspection Co-operation Scheme (PIC/S) authorities for the manufacturing site(s) of the proposed product(s).

OPTION 2 (2.1.+2.2.)

- 2.1. The proposed product(s) must have registered/certified for the use in Ukraine:
 - Registration Certificate issued by the State Administration on Pharmaceutical Products and Drugs Control Service of Ukraine and at least one successfully completed supply of this product in the similar volume in/to Ukraine within the past three years confirmed by medical institution/diagnostic center/laboratory center (since March 2014), OR;

 Declaration of Conformity with the requirements of technical regulations issued for the use in Ukraine (Resolution of the Cabinet of Ministers of Ukraine #753, #754, #755¹) and at least one successfully completed supply of this product in the similar volume in/to Ukraine within the past three years confirmed by medical institution/diagnostic center/laboratory center (since March 2014)

AND

- 2.2.a. Suppliers and manufacturers must provide an evidence of conformity* to at least one of the following conformity with the following Quality Management System standards as recognized by GHTF standards:
 - ISO13485/ISO 13488, or
 - ISO 9001, or
 - United States QS 21 CFR part 820, or
 - Japan QS Standard for medical devices.
 - -
- *) The evidence(s) of conformity shall indicate, as applicable:
 - a) Manufacturer's certified quality management system standard(s)
 - b) Assessment body (name, country)
 - c) Last audit date and
 - d) Expiration date
 - e) Certificate number

OR

2.2.b. Suppliers/manufacturers shall provide a copy of valid GMP Certificate issued by the Pharmaceutical Inspection Co-operation Scheme (PIC/S) authorities for the manufacturing site(s) of the proposed product(s).

Bidders shall demonstrate their compliane in the Annex 4 – Compliance of product/s to the requirements.

NB: If branded product is requested and equivalent product is allowed to be proposed as per section "Product List and Technical Specification", the Bidder must provide technical specification of item quoted and statement of deviations from branded product.

3. **REGISTRATION / AUTHORIZATION FOR USE IN UKRAINE**

Where a medical product has not yet been registered in Ukraine the suppliers of the Goods who wish to provide to, or within Ukraine, must make sure that the Goods comply with one and most relevant of the following regulations::

- a) Registration Certificate issued by the State Administration on Pharmaceutical Products and Drugs Control Service of Ukraine;
- b) Declaration of Conformity with the requirements of technical regulations (Resolution of the

¹ 1. Technical regulations on medical devices approved by the Cabinet of Ministers of Ukraine on 02/10/2013 № 753 (http://zakon3.rada.gov.ua/laws/show/753-2013-%D0%BF);

^{2.} Technical regulations on medical products for diagnostics in vitro, approved by the Cabinet of Ministers of Ukraine on 02/10/2013 № 754 (http://zakon1.rada.gov.ua/laws/show/754-2013-%D0%BF);

^{3.}Technical regulations on implanted active medical devices approved by the Cabinet of Ministers of Ukraine on 02/10/2013 № 755 (http://zakon1.rada.gov.ua/laws/show/755-2013-%D0%BF).

Cabinet of Ministers of Ukraine #753, #754, #755 dd. 02.10.2013).

UNDP will evaluate offers for both registered and non-registered medical products. Non-registered products must meet **quality standards as per OPTION1.** Bidders offering non-registered products that are compliant with quality standards, must start the registration/certification process preferably before, but not later than 5 days after, signing a conditional contract for the supply of product(s). Failure to obtain registration/certification and submit the required documents to UNDP will serve, at no claim to UNDP, as a ground for contract termination, liquidating Bid Security or Performance Security amount and either awarding the next qualified Bidder or initiating a new bidding process. The decision to transfer the award or initiate a new ITB will be at the discretion of UNDP.

4. PRODUCTS LIST AND TECHNICAL SPECIFICATION:

	4. PRODUCTS LIST AND TECHNICAL S			
Lot	Medical Device Name	Назва медичного виробу	иом	Quantity
	Medical devices for treatment or	f children with oncological oncohema	tologicals disea	se
1	Leukocyte Reduction Filter or device for Platelet Concentrate (for bedside application)	Фільтри або пристрій для видалення лейкоцитів з тромбоконцентрату (для приліжкового використання)	unit	936
2	Infusion filter for 96-hour use	Фільтри для інфузій (96-годинні)	unit	6,146
3	Leukocyte Reduction Filter or device for Red Blood Cells (for bedside application)	Фільтри або пристрій для видалення лейкоцитів з еритроцитарної маси (для приліжкового використання)	unit	2,026
4	SAG-M conservation medium, bag with 100 ml solution	Стабілізатор SAG-M, пакети з розчином 100 мл	unit	551
5	Dual container PLASMAFLEX/BLUEFLEX for Macotronic device or its equivalent	Подвійний контейнер PLASMAFLEX/BLUEFLEX до апарата Macotronic або еквівалент	unit	100
6	Set for collection of Red Blood Cells with filter for Automated Complex Apheresis device MCS+ (for platelet concentrate collection)	Набір для збору клітин крові з фільтром до апарата для автоматичного комплексного аферезу MCS+ (для заготівлі тромбоконцентрату)	set	220
7	Cryo Freezing Container 60—100 ml	Контейнери для кріозаморожування 60-100 мл	unit	350
8	Container with ACD-A Anticoagulant solution for apheresis devices	Контейнер з розчином антикоагулянту АЦД-А для апарата аферезу	unit	8,189
9	Container for collection of bone marrow	Контейнер для збору кісткового мозку	unit	50
10	C5L Platelet Set (5-day storage)	Комплект C5L тромбоцитаферезу (зберігання 5 діб)	kit	740

Lot	Medical Device Name	Назва медичного виробу	иом	Quantity
11	C4L Platelet Set (5-day storage)	Комплект С4L тромбоцитаферезу (зберігання 5 діб)	kit	120
12	Optia Collection Set for Spectra Optia Apheresis System (or equivalent)	Комплект для збору Optia до системи аферезу SpectraOptia (або еквівалент)	kit	100
13	AMICUS Mononuclear Cell (MNC) Apheresis Kit, Double Needle or equivalent	Комплект для аферезу "Amicus" МНК з двоголковим доступом або еквівалент	kit	140
14	Amicus Apheresis Kit, Single needle for Automated Blood Cell Separator or equivalent	Комплект для автоматичного цитаферезу клітин крові до клітинного сепаратора Amicus одноголоковий або еквівалент	kit	1,309
15	Amicus Apheresis Kit, Double needle for Automated Blood Cell Separator or equivalent	Комплект для автоматичного цитаферезу клітин крові до клітинного сепаратора Amicus двоголоковий або еквівалент	kit	1,184
16	Dual Needle Extended Life Platelet Set with LRS Chamber for COBE Spectra Apheresis System (or equivalent)	Комплект двоголковий для збору довгоживучих тромбоцитів з камерою LRS COBE Spectra до системи для аферезу COBE Spectra (або еквівалент)	kit	70
17	Trima Accel LRS PLT/RBC/Plasma Set for Trima Accel Automated Blood Collection System (or equivalent)	Комплект TrimaAccel для тромбоцитів LRS, плазми та еритроцитів до системи автоматичного збору компонентів крові TrimaAccel (або еквівалент)	kit	3,084
18	S5L single needle platelet apheresis set or equivalent	Комплект S5L тромбоцитаферезу, одноголкове підключення або еквівалент	kit	50
19	PIR Plasma Treatment Set for blood cell separator COM TEC or equivalent	Комплект PIR до сепаратора клітин крові COM.TEC або еквівалент	kit	5
20	TSCD Wafers, plate for TSCD or equivalent	TSCD Wafers, пластини до апарата TSCD або еквівалент	unit	51

Laboratory reagents for the Reference Laboratory for diagnosis of oncohematological diseases of the National Children's Specialized Hospital "OKHMATDYT" Laboratory reagents for immunocytological diagnosis of leukemias and lymphomas, detection of minimal residual disease, enumeration of haematopoietic stem sell collections using a flow cytofluorometer CytoMix FC-500

Lot	Medical Device Name	Назва медичного виробу	иом	Quantity
21	Monoclonal Antibody to CD1a, Fluorochrome PE, 100 tests	Моноклональне антитіло CD1a, мічене флюоресцентним барвником PE, 100 тестів	vial	1
22	Monoclonal Antibody to CD2, Fluorochrome PC5, 100 tests	Моноклональне антитіло CD2, мічене флюоресцентним барвником PC5, 100 тестів	vial	2
23	Monoclonal Antibody to CD3, Fluorochrome FITC, 100 tests	Моноклональне антитіло CD3, мічене флюоресцентним барвником FITC, 100 тестів	vial	3
24	Monoclonal Antibody to CD4, Fluorochrome PC5, 100 tests	Моноклональне антитіло CD4, мічене флюоресцентним барвником PC5, 100 тестів	vial	2
25	Monoclonal Antibody to CD5, Fluorochrome PC5, 100 tests	Моноклональне антитіло CD5, мічене флюоресцентним барвником PC5, 100 тестів	vial	1
26	Monoclonal Antibody to CD7, Fluorochrome FITC, 100 tests	Моноклональне антитіло CD7, мічене флюоресцентним барвником FITC, 100 тестів	vial	3
27	Monoclonal Antibody to CD8, Fluorochrome FITC, 100 tests	Моноклональне антитіло CD8, мічене флюоресцентним барвником FITC, 100 тестів	vial	1
28	Monoclonal Antibody to CD8, Fluorochrome PE, 100 tests	Моноклональне антитіло CD8, мічене флюоресцентним барвником PE, 100 тестів	vial	3
29	Monoclonal Antibody to CD8, Fluorochrome PC5, 100 tests	Моноклональне антитіло CD8, мічене флюоресцентним барвником PC5, 100 тестів	vial	1
30	Monoclonal Antibody to CD10, Fluorochrome PC5, 100 tests	Моноклональне антитіло CD10, мічене флюоресцентним барвником PC5, 100 тестів	vial	1
31	Monoclonal Antibody to CD13, Fluorochrome PE, 100 tests	Моноклональне антитіло CD13, мічене флюоресцентним барвником PE, 100 тестів	vial	1
32	Monoclonal Antibody to CD14, Fluorochrome PE, 100 tests	Моноклональне антитіло CD14, мічене флюоресцентним барвником PE, 100 тестів	vial	1

Lot	Medical Device Name	Назва медичного виробу	иом	Quantity
33	Monoclonal Antibody to CD15, Fluorochrome PE, 100 tests	Моноклональне антитіло CD15, мічене флюоресцентним барвником PE, 100 тестів	vial	1
34	Monoclonal Antibody to CD16, Fluorochrome PE, 100 tests	Моноклональне антитіло CD16, мічене флюоресцентним барвником PE, 100 тестів	vial	1
35	Monoclonal Antibody to CD19, Fluorochrome ECD, 100 tests	Моноклональне антитіло CD19, мічене флюоресцентним барвником ECD, 100 тестів	vial	1
36	Monoclonal Antibody to CD19, Fluorochrome PC5, 100 tests	Моноклональне антитіло CD19, мічене флюоресцентним барвником PC5, 100 тестів	vial	1
37	Monoclonal Antibody to CD20, Fluorochrome FITC, 100 tests	Моноклональне антитіло CD20, мічене флюоресцентним барвником FITC, 100 тестів	vial	1
38	Monoclonal Antibody to CD22, Fluorochrome PE, 100 tests	Моноклональне антитіло CD22, мічене флюоресцентним барвником PE, 100 тестів	vial	1
39	Monoclonal Antibody to CD25, Fluorochrome PC5, 100 tests	Моноклональне антитіло CD25, мічене флюоресцентним барвником PC5, 100 тестів	vial	1
40	Monoclonal Antibody to CD33, Fluorochrome PC5, 100 tests	Моноклональне антитіло CD33, мічене флюоресцентним барвником PC5, 100 тестів	vial	1
41	Monoclonal Antibody to CD45, Fluorochrome ECD, 100 tests	Моноклональне антитіло CD45, мічене флюоресцентним барвником ECD, 100 тестів	vial	6
42	Monoclonal Antibody to CD45, Fluorochrome PC5, 100 tests	Моноклональне антитіло CD45, мічене флюоресцентним барвником PC5, 100 тестів	vial	2
43	Monoclonal Antibody to CD117, Fluorochrome PE, 100 tests	Моноклональне антитіло CD117, мічене флюоресцентним барвником PE, 100 тестів	vial	1
44	Monoclonal Antibody to CD64, Fluorochrome PE, 2 ml (tests??)	Моноклональне антитіло CD64, мічене флюоресцентним барвником PE, 2 мл	vial	1

Lot	Medical Device Name	Назва медичного виробу	иом	Quantity
45	Monoclonal Antibody to CD65, Fluorochrome FITC, 100 tests	Моноклональне антитіло CD65, мічене флюоресцентним барвником FITC, 100 тестів	vial	1
46	Monoclonal Antibody to CD79a, Fluorochrome PE, 50 tests	Моноклональне антитіло CD79a, мічене флюоресцентним барвником PE, 50 тестів	vial	1
47	Monoclonal Antibody to HLA-DR, Fluorochrome FITC, 100 tests	Моноклональне антитіло HLA- DR, мічене флюоресцентним барвником FITC, 100 тестів	vial	3
48	Monoclonal Antibody to HLA-DR, Fluorochrome PC5, 100 tests	Моноклональне антитіло HLA- DR, мічене флюоресцентним барвником PC5, 100 тестів	vial	1
49	Monoclonal Antibody to CD3- FITC/CD(16+56) PE, 50 tests	Моноклональне антитіло CD3- FITC/CD(16+56)PE, 50 тестів	vial	3
50	Monoclonal Antibody to Anti-TdT, Fluorochrome FITC, 50 tests	Моноклональне антитіло Anti- TdT, мічене флюоресцентним барвником FITC, 50 тестів	vial	1
51	Monoclonal Antibody to kappa/lambda/CD19, Fluorochrome FITC/PE/PC5, 25 tests	Моноклональне антитіло kappa/lambda/CD19, мічене флюоресцентним барвником FITC/PE/PC5, 25 тестів	vial	3
52	IsoFlow Sheath Fluid,10 L or equivalent	Обжимна рідина IsoFlow,10 л або еквівалент	pack	5
53	Lysing solution for intracellular antigen analysis IntraPrep, 150 tests or equivalent	Лізуючий розчин для внутрішньоклітинного дослідження IntraPrep, 150 тестів або еквівалент	pack	1
54	Lysing solution Optilyse, 200 tests or equivalent	Лізуючий розчин Optilyse, 200 тестів або еквівалент	vial	2
55	Cleaning solution Cleans (5 L) or equivalent	Миючий розчин Cleans (5 л) або еквівалент	pack	1
56	Flow-Count Fluorospheres, 200 tests	FlowCountFluorospheres, 200 тестів флюоросфери Flow-Count	pack	1

Lot	Medical Device Name	Назва медичного виробу	иом	Quantity	
57	Test Tube, 12X75MM, Blue (250/PK) Analysis tubes, 12 x 75 mm, blue (250 units in a pack)	Test Tube, 12X75MM, Blue (250/PK) пробірки для аналізу, 12 х 75 мм, блакитні (250 штук в упаковці)	pack	4	
	Reagents for HLA	-typing and selection of marrow done	ors		
58	Micro SSP HLA Class I and II ABDR DNA typing tray — Class I & II (10 typings), One Lambda Inc., or equivalent	Micro SSP HLA Клас I та II ABDR планшет для ДНК типування - Клас I & II (10 типувань), One Lambda Inc., або еквівалент	set	50	
59	Micro SSP HLA Class I C Locus Specific Typing Tray (16 typings), One Lambda Inc., or equivalent	Micro SSP HLA Клас I С Локус специфічний планшет для ДНК типування (16 типувань), One Lambda Inc., або еквівалент	set	10	
60	Micro SSP general class II DNA Typing Tray — DQB1 (24 typings), One Lambda Inc., or equivalent	Micro SSP загальний клас II планшет для ДНК типування - DQB1 (24 типування), One Lambda Inc., або еквівалент	set	7	
61	Kit for DNA extraction in blood and body fluids Purelink Genomic DNA kit, (50 selections), Life Technologies Corp. / Thermo Fisher Scientific, or equivalent	Набір для виділення ДНК з крові та біологічних рідин Perelink Genomic DNA kit (50 виділень), Life Technologies Corp. / ThermoFisher Scientific, або еквівалент	set	16	
62	Taq Polymerase, 50 μl, One Lambda Inc., or equivalent	Таq Полімераза, 50 мкл, One Lambda Inc., або еквівалент	pack	100	
63	5XTB buffer with Et Br, 100 ml, One Lambda Inc., or equivalent	5XTB буфер з Et Br, 100 мл, One Lambda Inc., або еквівалент	pack	20	
Labo	Laboratory reagents for the molecular-genetic and molecular-cytogenetic diagnostics of leukemia in children				
64	Purelink Total RNA blood kit for RNA isolation from blood (50 preps), Life Technologies Corp. / ThermoFisher Scientific, or equivalent	Набір для виділення РНК з крові Purelink Total RNA blood kit (50 виділень), Life Technologies Corp. / ThermoFisher Scientific, або еквівалент	pack	11	
65	CEP X SpectrumOrange/CEP Y (satellite III) SpectrunGreen Chromosome Enumeration DNA Probe Kit (20 µl in a pack), Abbott Molecular, Inc., or equivalent	СЕР X Спектрум оранж/СЕР Y (сателіт III) спектрум грін набір ДНК проб для підрахунку хромосом (20 мкл в упаковці),	pack	2	

Lot	Medical Device Name	Назва медичного виробу	иом	Quantity
		Abbott Molecular, Inc., або еквівалент		
66	NP-40 (octylphenoxypolyethoxyethanol), (2 x 1000 μl in a pack), Abbott Molecular, Inc., or equivalent	НП-40 (октилфеноксиполі- етоксіетанол), (2 х 1000 мкл в упаковці), Abbott Molecular, Inc., або еквівалент	pack	6
67	DAPI II contrasting dye (2 x 500 μl in a pack), Abbott Molecular, Inc., or equivalent	DAPI II контрастуючий барвник, (2 x 500 мкл в упаковці), Abbott Molecular, Inc., або еквівалент	pack	44
68	CEP7 (D7Z1) Alpha Satellite DNA Probe (20 μl in a pack), Abbott Molecular, Inc., or equivalent	CEP7 (D7Z1) Альфа Сателіт ДНК проба (20 мкл в упаковці), Abbott Molecular, Inc., або еквівалент	set	1
69	LSI BCR/ABL Dual Color, Dual Fusion Translocation DNA Probe (20 µl in a pack), Abbott Molecular, Inc., or equivalent	LSI BCR/ABL двокольорова, подвійного злиття транслокаційна ДНК проба (20 мкл в упаковці), Abbott Molecular, Inc., або еквівалент	set	2
70	LSI MLL Dual Color Break Apart Rearrangement DNA Probe (20 µl in a pack), Abbott Molecular, Inc., or equivalent	LSI MLL двокольорова на точки розриву ДНК проба (20 мкл в упаковці), Abbott Molecular, Inc., або еквівалент	set	3
71	LSI LSI PML/RARa Dual Color, Translocation DNA Probe (20 µl in a pack), Abbott Molecular, Inc., or equivalent	LSI LSI PML/RARa двокольорова транслокаційна ДНК проба (20 мкл в упаковці), Abbott Molecular, Inc., або еквівалент	set	2
72	LSI CBFB Dual Color Break Apart Rearrangement DNA Probe (20 µl in a pack), Abbott Molecular, Inc., or equivalent	LSI CBFB двокольорова, на точки розриву ДНК проба (20 мкл в упаковці), Abbott Molecular, Inc., або еквівалент	set	2
73	LSI ETV6(TEL)/RUNX1(AML1)ES Dual Color Translocation Probe Set (20 µl in a pack), Abbott Molecular, Inc., or equivalent	LSITV6 (TEL)/RUNX1(AML)ES двокольоровий набір транслокаційних проб (20 мкл в упаковці), Abbott Molecular, Inc., або еквівалент	set	2
74	LSI RUNX1/ RUNX1T1DFFISH DNA Probe kit, Dual Color Translocation Probe Set (20 µl in a pack), Abbott Molecular, Inc., or equivalent	LSI RUNX1/ RUNX1T1DFFISH ДНК проби набір двокольоровий набір транслокаційних проб (20 мкл в упаковці), Abbott Molecular, Inc., або еквівалент	set	2
75	TelVysion Probe 21q Spectrum Orange (5 μl in a pack), Abbott Molecular, Inc., or equivalent	TelVysion проба 21q спектрум оранж (5 мкл в упаковці), Abbott Molecular, Inc., або еквівалент	set	4

Lot	Medical Device Name	Назва медичного виробу	UOM	Quantity
76	TelVysion Probe 22q Spectrum Green (5 μl in a pack), Abbott Molecular, Inc., or equivalent	TelVysion проба 22q спектрум грін (5 мкл в упаковці), Abbott Molecular, Inc., або еквівалент	set	1
77	TelVysion Probe Xq/Yq Spectrum Orange (5 μl in a pack), Abbott Molecular, Inc., or equivalent	TelVysion проба Хq/Yq спектрум оран (5 мкл в упаковці), Abbott Molecular, Inc., або еквівалент	set	1
78	TaqMan Universal PCR Master Mix (200 reactions), Life Technologies Corp. / ThermoFisher Scientific, or equivalent	ПЛР мастер-мікс TaqMan Universal (200 реакцій), Life Technologies Corp. / ThermoFisher Scientific, або еквівалент	vial	10
79	High Capacity cDNA Reverse Transcription Mix (200 reactions), Life Technologies Corp. / ThermoFisher Scientific, or equivalent	Суміш для зворотної транскрипції High Capacity cDNA (200 реакцій), Life Technologies Corp. / ThermoFisher Scientific, або еквівалент	vial	9
80	RPMI 1640 Medium(100 ml), Life Technologies Corp. / ThermoFisher Scientific, or equivalent	Середовище RPMI 1640 (100 мл), Life Technologies Corp. / ThermoFisher Scientific, або еквівалент	vial	27
81	Fetal Bovine Serum (100 ml), Life Technologies Corp. / ThermoFisher Scientific, or equivalent	Сироватка Fetal Bovine (100 мл), Life Technologies Corp. / ThermoFisher Scientific, або еквівалент	vial	7
82	MicroAmp Optical 96-Well Reaction Plate (10 plates in a pack), Life Technologies Corp. / ThermoFisher Scientific, or equivalent	96-лункові оптичні плашки MicroAmp (10 штук в упаковці), Life Technologies Corp. / ThermoFisher Scientific, або еквівалент	pack	5
83	MicroAmp Optical Adhesive Film (25 covers in a pack), Life Technologies Corp. / ThermoFisher Scientific, or equivalent	Плівка MicroAmp оптична (25 штук в упаковці), Life Technologies Corp. / ThermoFisher Scientific, або еквівалент	pack	2
84	Set of genes for genotyping, KIR Genotyping SSP Kit, dry format (10 test), One Lambda Inc., or equivalent	Набір для генотипування генів системи KIR genotyping SSP dry kit (10 тестів), One Lambda Inc., або еквівалент	pack	3
85	QuantideX qPCR BCR-ABL1 IS Kit for quantitative detection of BCR/ABL1 by real time PCR (60 reactions), Asuragen Inc., or equivalent	Набір для кількісного визначення мутації BCR/ABL1 методом ПЛР у реальному часі QuantideX qPCR BCR-ABL1 IS Kit (60 реакцій), Asuragen Inc., або еквівалент	set	1
86	Multiplex PCR qualitative detection of BCR/ABL1 mutations Kit with detection in agarose gel, Seeplex	Набір для мультиплексного якісного визначення мутації BCR/ABL1 методом ПЛР з детекцією в агарозному гелі	set	15

Lot	Medical Device Name	Назва медичного виробу	иом	Quantity			
	Leukemia BCR/ABL, 25 reactions, SeeGene Inc., or equivalent	Seeplex Leukemia BCR/ABL (25 реакцій), SeeGene, Inc., або еквівалент					
87	LeukoStrat™ FLT3 Mutation Assay Set for PCR-based detection of FLT3 mutations in agarose gel (33 reactions), Invivoscribe Technologies, Inc., or equivalent	Набір для визначення мутації гена FLT3 методом ПЛР з детекцією в агарозному гелі LeukoStrat [™] FLT3 Mutation Assay (33 реакції), Invivoscribe Technologies, Inc., або еквівалент	set	4			
88	ETV6(TEL)/RUNX1(AML) Kit for detecting mutations by real time PCR (24 reactions), Qiagen GmbH, or equivalent	Набір для визначення мутації ETV6(TEL)/RUNX1(AML) Kit методом ПЛР у реальному часі (24 реакції), Qiagen GmbH, або еквівалент	set	1			
89	Real time PCR detection of PML/RARAbcr1 mutations Kit (24 reactions), Qiagen GmbH, or equivalent	Набір для визначення мутації PML-RARA bcr1 Kit методом ПЛР у реальному часі (24 реакції), Qiagen GmbH, або еквівалент	set	1			
90	Real time PCR detection of PML/RARAbcr2 mutations Kit (24 reactions), Qiagen GmbH, or equivalent	Набір для визначення мутації PML-RARA bcr2 Kit методом ПЛР у реальному часі (24 реакції), Qiagen GmbH, або еквівалент	set	1			
91	Real time PCR detection of PML/RARAbcr2 mutations Kit (24 reactions), Qiagen GmbH, or equivalent	Набір для визначення мутації PML-RARA bcr3 Kit методом ПЛР у реальному часі (24 реакції), Qiagen GmbH, або еквівалент	set	1			
Laboratory reagents for the diagnosis and treatment of children with oncological diseases at the National Cancer Institute							
Laboratory reagents for immunocytological studies to enumerate haematopoietic stem sell collections, diagnose the immune system in children with solid malignant tumors using a flow cytofluorometer							
92	Monoclonal Antibody to CD34, Fluorochrome PE, 100 tests or analogue	Моноклональне антитіло CD34, мічене флюоресцентним барвником PE, 100 тестів або аналог	vial	2			
93	Monoclonal Antibody to HLA-DR, Fluorochrome PE, 100 tests or analogue	Моноклональне антитіло HLA- DR, мічене флюоресцентним барвником PE, 100 тестів або аналог	vial	1			
94	Sheath Fluid IsoFlow, 10 l or equivalent	Обжимна рідина IsoFlow, 10 л або еквівалент	pack	20			

Lot	Medical Device Name	Назва медичного виробу	иом	Quantity
95	Cleaning solution Cleans (5 I) or equivalent	Миючий розчин Cleans (5 л) або еквівалент	pack	10
96	Monoclonal Antibody to CD11c, Fluorochrome PE, 100 tests or analogue	Моноклональне антитіло CD11c, мічене флюоресцентним барвником PE, 100 тестів або аналог	vial	1
97	Monoclonal Antibody to CD69, Fluorochrome PC5, 100 tests or analogue	Моноклональне антитіло CD69, мічене флюоресцентним барвником PC5, 100 тестів або аналог	vial	1
98	Monoclonal Antibody to CD127, Fluorochrome PE, 100 tests or analogue	Моноклональне антитіло CD127, мічене флюоресцентним барвником РЕ, 100 тестів або аналог	vial	1
solic	Laboratory reagents for molecular genetic studies to confirm the diagnosis and ch solid malignant tumors in children, detecting micrometastases, minimal residual d effectiveness of polymerase chain reaction therapy on devices Applied Biosystems		l disease, and m	onitoring the
99	Sequence Detection Primer	Праймер Sequence Detection	vial	10
100	TaqMan fluorescent probe	Флуоресцентний зонд TaqMan	vial	5
101	TaqMan Universal PCR Master Mix	ПЛР мастер-мікс TaqMan Universal	vial	5
102	High Capacity cDNA Reverse Transcription Kit	Суміш для зворотної транскрипції High Capacity cDNA	set	1
103	MicroAmp Film	Плівка MicroAmp	pack	3
104	MicroAmp Optical 96-Well Reaction Plate	96-лункові плашки MicroAmp	pack	10
105	TaqMan SNP Genotyping Assays	Єсейна збірка TaqMan SNP Genotyping/	set	3

Lot	Medical Device Name	Назва медичного виробу	иом	Quantity
106	Set or extraction of RNA and DNA from paraffin-embedded samples NucleoSpin FFPE RNA/DNA	Набір для виділення РНК та ДНК з парафінізованих зразків NucleoSpin FFPE RNA/DNA	set	2
107	Set for extraction of microRNA and RNA from cells and tissues NucleoSpin miRNA	Набір для виділення мікро РНК та РНК з клітин та тканин NucleoSpin miRNA	set	2
108	Set for extraction of RNA from blood NucleoSpin RNA Blood	Набір для виділення РНК з крові NucleoSpin RNA Blood	set	1
109	Set for extraction of DNA from tissues NucleoSpin Tissue	Набір для виділення ДНК з тканин NucleoSpin Tissue	set	2
110	Set for extraction of RNA from cells and tissues NucleoSpin RNA II	Набір для виділення РНК з клітин та тканин NucleoSpin RNA II	set	2
111	RNase Reagent	Реагент РНКаза	vial	3
112	Reagent Proteinase K	Реагент Протеїназа К	vial	2
113	RNAlater Solution	Розчин RNAlater	vial	1
114	Fetal Bovine Serum	Сироватка Fetal Bovine	vial	2
115	RPMI 1640 Medium	Середовище RPMI 1640	vial	10
116	Versene Solution	Розчин Versene	vial	1

Laboratory reagents for molecular cytogenetic studies to confirm diagnosis and select treatment strategy, detect micrometastases, minimal residual disease and monitor the effectiveness of therapy of pediatric malignant solid tumors using fluorescence in situ hybridization (FISH) applying fluorescent microscope

Lot	Medical Device Name	Назва медичного виробу	иом	Quantity
117	LSI 1p36 / LSI 1q25 and LSI 19q13 / LSI 19p13 Dual Color DNA Probe Set	LSI 1p36/LSI 1q25 та LSI 19q13/LSI 19p13 двокольоровий набір ДНК проб	20 μl in a pack	1
118	LSI FOXO1 Dual Color, DNA Probe Set Break Apart Rearrangement FISH	LSI FOXO1 двокольоровий набір ДНК проб на точки розриву FISH	20 μl in a pack	1
119	LSI N-MYC SG(2q24)/CEP 2 SO Dual Color DNA Probe Set	LSI N-MYC SG(2q24)/CEP 2 SO двокольоровий набір ДНК проб	20 μl/pack	2
120	LSI EWSR1 Dual Color, DNA Probe Set Break Apart Rearrangement	LSI EWSR1 двокольоровий набір ДНК проб на точки розриву	20 mcl in a pack	1
121	LSI/WCP Hybridization Buffer	LSI/WCP гібридизаційний буфер	2 x 150 μl/pack	2
122	Mixture of sodium chloride and sodium citrate	Суміш хлориду та цитрату соди	500 g in a pack	2
	Laboratory reagents for immunohist	ochemical diagnostics of solid malign	ant tumors in cl	nildren
123	FLEX Monoclonal Mouse Anti-Human Cytokeratin, Clone AE1/AE3, Ready- to-Use, DAKO AS/AS+	FLEX Моноклональне антитіло "миша проти людини", цитокератин, AE1/AE3, готове до використання, DAKO AS/AS+	vial	3
124	FLEX Polyclonal Rabbit Anti-Human Alpha-1-Fetoprotein, Ready-to-Use	FLEX Поліклональне антитіло "кролик проти людини", Альфа- 1-фетопротеїн, готове до використання	vial	2
125	FLEX Monoclonal Mouse Anti-Human Calretinin, DAK-Calret 1, Ready-to- Use	FLEX Моноклональне антитіло "миша проти людини", калретінин, DAK-Calret 1, готове до використання	vial	1
126	FLEX Monoclonal Mouse Anti-Human β-Catenin, β-Catenin-1, Ready-to-Use	FLEX Моноклональне антитіло "миша проти людини", β- катенін, β-Catenin-1, готове до використання	vial	3
127	FLEX Monoclonal Mouse Anti- Human, CD10, Clone 56C6, Ready-to- Use	FLEX Моноклональне антитіло "миша проти людини", КД10, 56C6, готове до використання	vial	1

Lot	Medical Device Name	Назва медичного виробу	иом	Quantity
128	FLEX Monoclonal Mouse Anti- Human, CD1a, Clone 010, Ready-to- Use, DAKO AS/AS+	FLEX Моноклональне антитіло "миша проти людини", КД1а, 010, готове до використання, DAKO AS/AS+	vial	1
129	FLEX Monoclonal Mouse Anti- Human, CD31, Endothelial Cell, Clone JC70A, Ready-to-Use	FLEX Моноклональне антитіло "миша проти людини", КД31, ендотеліальні клітини, JC70A, готове до використання	vial	1
130	FLEX Monoclonal Mouse Anti-Human CD34 Class II Clone QBEnd 10, Ready- to-Use	FLEX Моноклональне антитіло "миша проти людини", КД34 клас II, QBEnd 10, готове до використання	vial	1
131	FLEX Monoclonal Mouse Anti-Human CD68, Clone KP1, Ready-to-Use	FLEX Моноклональне антитіло "миша проти людини", КД68, KP1, готове до використання	vial	1
132	FLEX Monoclonal Mouse Anti- Human, CD99, MIC2 Gene Products, Ewing's Sarcoma Marker Clone 12E7, Ready-to-Use, AS/AS+	FLEX Моноклональне антитіло "миша проти людини", КД99 MIC2 генні продукти, маркер саркоми Евінга 12Е7 готове до використання, AS/AS+	vial	1
133	FLEX Polyclonal Rabbit Anti-Human Carcinoembryonic Antigen, Ready- to-Use	FLEX Поліклональне антитіло "кролик проти людини", раково- ембріональний антиген, готове до використання	vial	1
134	Monoclonal Mouse Anti-Human, collagen IV, CIV 22	Моноклональне антитіло "миша проти людини", колаген IV, CIV 22	vial	1
135	FLEX Monoclonal Mouse Anti- Human, Desmin Clone D33, Ready- to-Use	FLEX Моноклональне антитіло "миша проти людини", десмін, D33, готове до використання	vial	1
136	FLEX Monoclonal Mouse Anti- Human, E-Cadherin, Clone NCH-38, Ready-to-Use (Dako AS/AS+)	FLEX Моноклональне антитіло "миша проти людини", Е- кадхерін, NCH-38, готове до використання (Dako AS/AS+)	vial	1
137	FLEX Monoclonal Mouse Anti- Human, Epithelial Membrane Antigen, Clone E29, Ready-to-Use	FLEX Моноклональне антитіло "миша проти людини", епітеліально-мембранний антиген, E29, готове до використання	vial	1
138	FLEX Polyclonal Rabbit Anti-Glial Fibrillary Acidic Protein, Ready-to- Use	FLEX Поліклональне антитіло "кролик проти гліального фібрилярного кислого протеїну", готове до використання	vial	1

Lot	Medical Device Name	Назва медичного виробу	иом	Quantity
139	FLEX Monoclonal Mouse Anti-Human Hepatocyte Clone OCH1E5, Ready- to-Use	FLEX Моноклональне антитіло "миша проти людини", гепатоцити, ОСН1Е5, готове до використання	vial	1
140	FLEX Monoclonal Mouse Anti- Human, Ki-67 Antigen, Clone MIB-1, Ready-to-Use	FLEX Моноклональне антитіло "миша проти людини", Кі-67 антиген, МІВ-1, готове до використання	vial	1
141	FLEX Monoclonal Mouse Anti-Human Actin (Muscle) Clone HHF35, Ready- to-Use	FLEX Моноклональне антитіло "миша проти людини", м'язевий актин, HHF35, готове до використання	vial	1
142	Monoclonal Mouse Anti-Human, MyoD1, 5.8A	Моноклональне антитіло "миша проти людини", MyoD1, 5.8A	vial	1
143	FLEX Monoclonal Mouse Anti- Myogenin, Clone F5D, Ready-to-Use, DAKO AS/AS+	FLEX Моноклональне антитіло "миша проти міогеніну", F5D, готове до використання, DAKO AS/AS+	vial	1
144	Monoclonal Mouse Anti-Human Mesothelial Cell Clone HBME-1	Моноклональне антитіло "миша проти людини", мезотеліальні клітини, HBME-1	vial	2
145	FLEX Monoclonal Mouse Anti-Human Neuron-Specific Enolase (NSE), Clone BBS/NC/VI-H14, Ready-to-Use	FLEX Моноклональне антитіло "миша проти людини", НСЄ, BBS/NC/VI-H14, готове до використання	vial	1
146	FLEX Monoclonal Mouse Anti-Human Progesterone Receptor, Clone PgR 636, Ready-to-Use, DAKO AS/AS+	FLEX Моноклональне антитіло "миша проти людини", прогестероновий рецептор, PgR 636, готове до використання, DAKO AS/AS+	vial	2
147	FLEX Polyclonal Rabbit Anti S100, Ready-to-Use	FLEX Поліклональне антитіло "кролик проти S100", готове до використання	vial	1
148	FLEX Monoclonal Mouse Anti-Human Actin (Smooth Muscle), Clone 1A4, Ready-to-Use	FLEX Моноклональне антитіло "миша проти людини", гладенько-м'язевий актин, 1A4, готове до використання	vial	1
149	Monoclonal Mouse Anti-Human Vascular Endothelial Growth Factor (VEGF), Clone VG1	Моноклональне антитіло "миша проти людини", васкулярний ендотеліальний фактор росту (VEGF), VG1	vial	1

Lot	Medical Device Name	Назва медичного виробу	UOM	Quantity
150	Monoclonal Mouse Anti-Human Von Willebrand Factor, Clone F8/86	Моноклональне антитіло "миша проти людини", фактор Вон Вілєбранта , F8/86	vial	3
151	FLEX Monoclonal Mouse Anti-Human WT1 Protein, Clone 6F-H2, DAKO AS/AS+	FLEX Моноклональне антитіло "миша проти людини", WT1 протеїн, 6F-H2, DAKO AS/AS+	vial	9
152	Monoclonal Mouse Anti-Human, epidermal growth factor receptor (EGFR), E30	Моноклональне антитіло "миша проти людини", рецептор до епідермального ростового фактору (EGFR), E30	vial	1
153	FLEX Monoclonal Mouse Anti-Human CD45, Leucocyte Common Antigen, Clones 2B11 + PD7/26, Ready-to-Use	FLEX Моноклональне антитіло "миша проти людини", КД45, LCA, 2B11+PD7/26, готове до використання	vial	1
154	EnVision FLEX+, High pH (Dako Autostainer Instruments)	EnVision FLEX+, Високий pH (Dako Autostainer Instruments)	pack	1
155	Mounting Medium	Заключне середовище	vial	1
156	Rabbit Monoclonal Anti- Synaptophysin antibody	Моноклональне антитіло "кролик проти сінаптофізину"	vial	1
157	Rabbit Monoclonal Anti- Chromogranin A antibody	Моноклональне антитіло "кролик проти хромограніну А"	vial	1
158	Elite PAP Pen	Elite PAP імуномаркер	pack	1
159	Disposable Microtome Blades Patho Cutter R22	Змінні леза до мікротому Patho Cutter R22	pack	1
160	HistoBond® + S slides	Скельця S HistoBond® +	pack	25

NB. UNDP reserves the right to vary the quantity of the goods by up to a maximum twenty-five per cent (25%) of the total offer, without any change in the unit price or other terms and conditions.

4. DELIVERY TIMEFRAMES

Early delivery of medical products to Ukraine is critical therefore we encourage shortest delivery periods, preferably within 4 months at the latest after signing the contract.

Further to the Schedule of Requirements in the preceding Table, Bidders are requested to take note of the following additional requirements, conditions, and related services pertaining to the fulfillment of the requirements:

Delivery Term	DAP Kyiv					
[INCOTERMS 2010]	The products shall be supplied to the Central Warehouse of MoH					
(Pls. link this to price schedule)	or designated by them entity appointed by UNDP. Exact location					
	of the warehouse will be notified at the time of contracting.					
		_				
		vnership right from seller to buyer occurs				
	•	the transfer of risk of goods loss or damage				
	_	s are delivered to the named warehouse.				
	Partial delivery is ac	ceptable.				
Mode of Transport Preferred	⊠AIR	⊠LAND				
	⊠SEA	□OTHER [pls. specify]				
Shipping documents	Commercial inv	oice – 2 originals.				
	 Packing list – 1 	• •				
		Certificate of Analysis for each batch – copies				
		ne stamp of the Supplier.				
		rigin, if goods are being imported				
	 Air Way Bill (air shipments)/Bill of Lading (sea shipments), if goods are being imported 					
	 Registration Certificate issued by State Administration of Pharmaceutical Products and Drug Control Service of Ukraine (if applicable). 					
	Declaration of 0 regulations (if a	Conformity with the requirements of technical pplicable).				
Customs, if needed, clearing shall be done by:	UNDP					
Ex-factory / Pre-shipment		pection may be carried out by UNDP or its				
inspection	representative for verification of quality, quantity, packing, labelling, marking and sampling. In cases when pre-shipment inspection is required, the corresponding Purchase Order will					
Inspection upon delivery	specify this conditio MoH/UNDP will	conduction inspection upon delivery.				
inspection upon delivery	IVIOTI/ OINDE WIII	conduction inspection upon delivery.				
	Quality Control may	be required upon discretion of UNDP/MoH.				

Payment Terms	Within 30 calendar days after delivery subject to written acceptance of goods delivery, duly signed and stamped by UNDP/MoH and provision of original invoice.
	In case testing is required, satisfactory testing results is a prerequisite for payment release.
	Progress payments could be provided in case of partial delivery.

5. SHELF LIFE

Products must have a minimum of 75% of the total product shelf life or should have 15 months' shelf life remaining at the time of delivery and must bear the dates of manufacture and expiry. Shelf life shall be indicated for all products quoted in the offer submitted.

6. PACKAGING, LABELLING, DELIVERY

- 1) Upon receipt of an incoming batch, UNDP follow a thorough quality control procedure, which includes review of Certificates of Analysis (CoA) for each batch of finished product to be supplied, Registration Certificate issued by the Ministry of Health of Ukraine or Declaration of Conformity, inspection against UNDP specifications, labelling and packaging.
- 2) Medical products shall be transported and stored in accordance with the temperature mode specified in the product instruction. All temperature restricted commodities must be shipped with clear marking the corresponding temperature conditions. It is the responsibility of the Bidder to provide complete packing as required for transportation. Bidders shall explain their capabilities and experience to handle temperature control items where applicable.

All temperature restricted commodities must be supplied with USB data loggers for monitoring temperature and humidity as well as with dry ice / cold packs/ ice packs / wet ice, as applicable, which must last at least 72 hours. The box with temperature restricted items should be clearly marked as "keep cool (2-8°C)" or "keep frozen (-20°C)" or "keep 2-30°C".

The numbers of dataloggers should be 1 if shipment has 5 or less boxes, 2 if shipment has more than 5 boxes. If products are shipped in containers, each container should have 2 dataloggers.

Dataloggers should be activated, set up with adequate alarm levels and placed inside a box with the

Dataloggers should be activated, set up with adequate alarm levels and placed inside a box with the products. The boxes with dataloggers should be clearly identified with bright color stickers (ideally orange).

The minimum technical requirements for dataloggers are as follows:

- Measures temperature (from -30° to 70°c, with accuracy +/- 0.3°c).
- Readings to include time and date
- Single or multiple use
- Direct USB interface, without need for additional cable
- Automatically creates PDF report when connected to computer.
- Rapid data download to graph
- Alarm levels set up before shipping according to manufacturer's storage requirements

- LCD featuring up to 1 decimal point readings
- Alarm indication on LCD screen
- Sampling rate: at least 1 measure per hour
- Push button to activate and stop logging.
- Easy to understand user's guide & instructions

All cases should be marked with/prominently indicate the following:

- A. Shipping marks;
- B. The generic name of the product;
- C. The dosage form (tablet, ampoule, syrup);
- D. Strength/ concentration of the product;
- E. Number of registration certificate
- F. Date of manufacture and expiry (in clear language not code);
- G. Batch number;
- H. Quantity per case;
- I. Special instructions for storage;
- J. Name of manufacturer;
- K. Carton numbering e.g. carton 1/40;
- L. Any additional cautionary statements.
- 3) The individual packages shall be packed in carton boxes. Each carton shall contain only one product and one batch. Packing must be sufficiently strong to withstand rough handling and exposure to extreme temperatures and air moisture.
- 4) Labelling of package at the moment of supply must correspond to the one in the product's state registration record or Declaration of Conformity.
- 5) Packaging must preserve quality, safety and stability of the product it contains. The entire package must be duly sealed and protected from spoiling. Each product shall contain instructions for the use of the medicinal product in Ukrainian (preferably) or the original language. The instruction for use must indicate clearly the reference of the version.

In case medical products are delivered in original packaging with instructions for the use in the original language, Ukrainian translation of instruction for the use shall be provided in the electronic format at the time of supply.

6) UNDP reserves the right to have at any time the items inspected, tested for quality assurance and rejected if found not in compliance with the requested specifications.

7) Pre-shipment inspection

When all the goods from a specific purchase order are ready for shipment with their final packing and marking, a pre-shipment inspection may be carried out by UNDP or its representative for verification of quality, quantity, packing, labelling, marking and sampling.

In cases when pre-shipment inspection is required, the corresponding Purchase Order will indicate this.

For this purpose, the Supplier will have to submit the applicable documentation to UNDP or its representative and allow UNDP or its representative access to all the goods. At least the packing list showing also the batch numbers per product and the full address of inspection should be made available to UNDP or its representative 7 working days before the pre-shipment inspection is requested to be carried out. Inspection/testing by UNDP or its representative in no way relieves the Supplier from the performance of full contractual obligations to UNDP. The cost of the pre-shipment inspection will be borne by UNDP. However, it is the responsibility of the supplier to assure that all facilities, to carry out a proper inspection are made available at their expense, and the goods for one shipment are presented at one location and on the date requested by UNDP or its representative. Furthermore, UNDP or its representative will charge the Supplier for the repeat, supplementary or abortive inspection visits necessitated by the fault of the supplier. UNDP or its representatives may inspect the production premises and the process of the manufacture.

In case of the detection of a defective product either in the quality of a product or other defects such as packaging, the Supplier will be requested to replace the complete batch at its own cost within one (1) month. In the event of a dispute by the Supplier, a counter analysis will be carried out by an independent neutral laboratory agreed by both UNDP and the Supplier. If the counter analysis confirms the defect, the cost of such analysis will be borne by the Supplier as well as the replacement and disposal of the defective batch. In the event of the independent analysis confirming the quality of the product, UNDP will meet all costs for such analysis.

- 8) Stipulations concerning Supplier responsibility for Quality, Packaging and Warranty
- a) UNDP shall have the right to make claims under the above warranty for three months after the Goods have been delivered to the final destination indicated in the Purchase Order. Upon receipt of a written notice from UNDP, the Supplier shall, with all reasonable speed, replace the defective Goods without cost to Purchaser at the final location. The Supplier will be required to remove, at his own risk and cost, the defective Goods once the replacement Goods have been delivered. If the defective Goods are not removed within 30 days, UNDP will dispose on the Supplier's costs.
- b) The Supplier's responsibility for labelling and quantities of goods for every Purchase Order extends to the point at which the goods are inspected by UNDP or its representative and, if required, a Clean Report of Findings (CRF) is issued by UNDP or its representative, upon delivery, for the specific PO. Where discrepancies are found by UNDP or its representative in labelling and/or quantities, these shall be rectified promptly by the Supplier at its own cost.
- c) The Supplier is responsible for the intrinsic quality of the finished product and for the intrinsic quality of the primary packaging of the product, prior to and after the CRF is issued. The Supplier's responsibility will be according to the Incoterms 2010 standards specified in the PO.
- 9) Stipulations concerning Recalls: In the event any of the Goods are recalled either by the National Regulatory Authority (NRA) of the country of production, the NRA of the recipient country or the

Manufacturer, after the CRF related to the PO(s) covering the same Goods is issued, the Supplier shall notify UNDP within fourteen (14) days, providing full details of the reason for the recall and replace affected goods within one (1) month, at its own cost, the items covered by the recall with Goods that fully meet the requirements of the Technical Specifications and original PO(s) against which they were supplied, and arrange for collection or destruction of any defective Goods. If the Supplier fails to fulfill its recall obligation promptly, UNDP will, at the Supplier's expense, carry out the recall.

10) Quality Assurance

Prior to shipment or upon arrival at the destination, some batches of the product may be tested (randomly) to ensure that the products meet Quality Assurance according to agreed contractual standards and requirements. Such tests might include, using an independent laboratory as service provider and or in-house quality checks and any consignment or batch(es) of goods not meeting the above mentioned standards would be rejected.

SECTION 4

Criteria for award and checklist of documents required

Following documents should be attached to the filled-in sections #-8

Please ensure that all documents necessary to enable objective evaluation are attached to your response to this ITB:

Award Criteria	Corresponding document	Yes	No	Reference
Compliance of Bidder with Qualifi	cations Requirements			
Minimum 3 years of experience in similar nature and minimum 2	1. Certificate of Registration of the business, including Articles of Incorporation, or equivalent			
similar contracts fulfilled over	document if Bidder is not a corporation			
the past 3 years	2. Statement of Satisfactory Performance (Reference letters) from the Top 3 Clients in terms of Contract Value the past 3 years. Please provide reference letters to prove experience in similar nature of contracts			
Minimum annual turnover over	3. Latest Audited Financial Statement (Income			
the past 2 years shall equal to no	Statement and Balance Sheet) including			
less than 75% of the total	Auditor's Report for the past 2 years			
amount to be contracted				
Compliance of product/quoted wi	th product standards and requirements (please co	mplete	checkl	ist for each
product quoted)				
1.1. Medical products must be	Medical Device license (Canada), OR			
produced and controlled in	EC Full Quality Assurance Certificate or EC			
accordance with product	Production Quality Assurance Certificate cate or			
standards and quality system	EC Type-Examination Certificate (CE/ Conformité			
standards recommended by the	Européenne mark) or / Conformité Européenne			
World Health Organization	92/42 or CE/ Conformité Européenne 98/79, OR			
(WHO) AND/OR the International	TGA Production Quality Assurance Certificate or			
Medical Devise Regulators Forum	TGA Type-Examination Certificate or TGA Full			
(IMDRF) (former Global	Quality Assurance Certificate issued by			
Harmonization Task Force	Therauptic Goods Administration), OR			
(GHTF). In order to be compliant	PMDA (Pharmaceuticals and Medical Devices			
with this criterion bidders will be	Agency) approval or JMHLW (Japan Ministry of			
requested to provide one of the	Health, Labour and Welfare) Minister's approval,			
following pre-market approval(s)	OR			
/ market clearance(s) (please				
refer for details to Section 3 of				
ITB).				
	PMA (pre-market approval) letter or BLA license			
	(Biologics License Application) or 510k device			
	letter issued by US Food and Drug			
	Administration			
2.1. The proposed product(s)	Registration Certificate issued by the State			
must have registered/certified	Administration on Pharmaceutical Products and			
for the use in Ukraine (please	Drugs Service of Ukraine, OR			

refer for details to Section 3 of ITB)	Declaration of Conformity with the requirements of technical regulations issued for the use in Ukraine (Resolution of the Cabinet of Ministers of Ukraine #753, #754, #755) AND Evidence of at least one successfully completed supply of this product in the similar volume in/to Ukraine within the past three years confirmed by medical institution/diagnostic center/laboratory center (since March 2014)		
1.2.a./2.2.a. Suppliers and manufacturers must provide an evidence of conformity* to at least one of the following conformity with the following Quality Management System standards as recognized by GHTF standard	ISO 13485, OR ISO 9001, OR United States QS (21 CFR part 820), OR Japan QS Standard for medical devices		
1.2.b./2.2.b. Suppliers and manufacturers must provide an evidence of GMP certification of manufacturing site by PIC/S authorities	A copy of valid GMP Certificate issued by PIC/S authorities for the manufacturing site(s) of the proposed product(s)		
Availability of valid registration/certification in Ukraine at the time of supply as defined in Section 3, (if, at the moment of the bid submission, the quoted medical products are not registered in Ukraine but comply with the quality requirements of this ITB (OPTION	Option A: A copy of a valid registration certificate for every medicinal product quoted issued by the Ministry of Health of Ukraine AND/OR Declaration of Conformity with the requirements of technical regulations (Resolution of the Cabinet of Ministers of Ukraine #753, #754, #755 dd. 02.10.2013)		
1), a Commitment letter shall be provided)	Option B: If, at the moment of the bid submission, the quoted medicinal products are not registered/certified in Ukraine but comply with the quality requirements of this ITB, a Commitment letter (Annex 2) from the bidder acknowledging acceptance of the terms and conditions for undertaking a registration/certification procedure (see Section 3, para #3 Registration/Authorization for use in Ukraine for details). By submitting the Bid, the Bidder automatically		
	agrees to maintain and renew registration/certification for these products until their shelf life expiration.		

Compliance with shelf life, packing and labelling requirements (please refer for details to Section 3 of ITB).	Please provide Information on shelf life in the Form 7 Technical Bid Form		
Acceptability of the	Please provide Information on delivery schedule		
Transportation/Delivery	in the Form 7 Technical Bid Form		
Schedule (please refer for details			
to Section 3 of ITB)			

List of other documents required for evaluation of Bidder	Yes	No	Reference
List of Shareholders and Other Entities Financially Interested in the Firm owning 5% or more of the stocks and other interests, or its equivalent if Bidder is not a corporation			
Valid Certificate of Authorization to act on behalf of the Manufacturer in case the bidder is not a Manufacturer.			
All information regarding any past and current litigation during the last five (5) years, in which the bidder is involved, indicating the parties concerned, the subject of the litigation, the amounts involved, and the final resolution if already concluded.			
Quality Certificate (e.g., ISO, etc.) and/or other similar certificates, accreditations, awards and citations received by the Bidder, if any			
Environmental Compliance Certificates, Accreditations, Markings/Labels, and other evidences of the Bidder's practices which contributes to the ecological sustainability and reduction of adverse environmental impact (e.g., use of non-toxic substances, recycled raw materials, energy-efficient equipment, reduced carbon emission, etc.), either in its business practices or in the goods it manufactures, if any available			

List of other documents required for evaluation of product quoted (please complete			
checklist for each product quoted)	Yes	No	Reference
Instruction for the use in accordance with the legislation of Ukraine. In case quoted medicinal products are not registered/certified in Ukraine, instructions for the use in the original language shall be provided.			
Technicalspecification of product quoted			
Safety data sheet (SDS) of the product			
Patent Registration Certificate/s, (if applicable) or relevant license/s (if available)			

BRIEF SUMMARY

1. On the simplified procedure of state registration of medicinal products procured with involvement of the international organizations

The procedure of state registration (re-registration) of medicinal products was approved by the Cabinet of Ministers of Ukraine Resolution No. 376 of 26.05.2005.

State registration of the medicinal products procured by international organizations is provided by the Ministry of Health of Ukraine pursuant to an application and subject to an opinion of the MoH State Expert Centre (hereinafter referred to as the Centre) drawn up on the basis of results of the expert examination of the registration materials for their authenticity, conducted according to the procedure specified by the MoH of Ukraine.

Information on the procedures for state registration may be found under the below links:

- 1. Law of Ukraine "On Medicines" http://zakon2.rada.gov.ua/laws/show/123/96-%D0%B2%D1%80
- 2. Decree of the Cabinet of Ministers of Ukraine dated 26.05.2005 № 376 http://zakon5.rada.gov.ua/laws/show/376-2005-%D0%BF
- 3. Decree of MOH of Ukraine dated 03.11.2015 № 721 http://zakon2.rada.gov.ua/laws/show/z1453-15
- 2. On additional relevant information on VAT for bidders. This information is provided for references only and UNDP should not be hold accountable for any details. Bidders are encouraged to check the details with relevant authorities directly.

Operations on supply (transfer) of pharmaceuticals and medical products shall be temporarily, until March 31, 2019, exempt from VAT, if importation and/or supply is done under contracts with specialized procurement organizations listed in the Law of Ukraine 'On Public Procurement', concluded with the objective of implementing agreements between the central executive body of Ukraine in charge of developing and implementing the national health policy and a relevant specialized procurement organization within the framework of budget programmes for implementation of public health action plans and/or comprehensive programme activities in the health sector.

Provided VAT exemption condition may not be applied under the Ukrainian legislation. VAT amount should be clearly indicated in a separate line (if applicable).

For more information on VAT exemption, please refer to the following legislation documents:

- 1. Tax Code of Ukraine, Chapter #XX Transitional Provisions, Section #2; Paragraph #38 on conditions of temporary VAT exemption of medicines and medical products that are procured by specialized organizations for the National Public Health Programme to the Ministry of Health (MoH) in Ukraine: http://zakon2.rada.gov.ua/laws/show/2755-17/page45
- 2. Decree of the Cabinet of Ministers of Ukraine #1153 dated 02.12.2015 on the procedures of

importation, supply and targeted use of medicines, medical devices that are VAT exempted: http://zakon3.rada.gov.ua/laws/show/1153-2015-%D0%BF

Prices specified shall remain firm and not be increased. In case Offeror increase price after awarding contract, UNDP will consider this as a ground for contract termination, liquidating Bid or Performance Security amount and either awarding the next qualified Bidder or initiating a new bidding process.

Commitment letter

(This should be written in the Letterhead of the Bidder. Except for indicated fields, no changes may be made in this template.)

Insert: Location

То:	[insert: Name and Address of UNDP focal point]
	/Madam: We, the undersigned, hereby offer to supply the goods required for [insert: title of goods and services d as per ITB] in accordance with your Invitation to Bid dated We hereby commit to register/certify the below listed products as per current Ukrainian legislation is.
	Products: 1 2 3
	We fully understand and recognize that UNDP is not bound to accept this Bid, that we shall bear all costs red with its preparation and submission, registration fees and that UNDP will in no case be responsible or or those costs, regardless of the conduct or outcome of the evaluation.
	We remain, Yours sincerely, Authorized Signature [In full and initials]: Name and Title of Signatory: Name of Firm:
Contact	Details:
	[please mark this letter with your corporate seal, if available]

Certificate of Authorization to act on behalf of the Manufacturer in case the Bidder is not a Manufacturer

(This should be written in the Letterhead of the Manufacturer. Certificate shall cover all items for which the company is bidding)

		Insert: Location
To:	[insert: Name and Address of UNDP focal point]	msert. Date
Dear Si	ir/Madam:	
[name	e undersigned, who is established manufacturer or producer of [insert name of producer of Bidder] to submit a Bid, and subsequently sign and implement the stitle of goods and services required as per ITB] for the supply of following product Products: 1.	contract, against the
	 	
For and	d on behalf of Manufacturer or Producer:	
	Yours sincerely, Authorized Signature [In full and initials]: Name and Title of Signatory: Name of Firm:	
Contac	t Details:	

Section 5: Bid Submission Form²

(This should be written in the Letterhead of the Bidder. Except for indicated fields, no changes may be made in this template.)

Insert: Location Insert: Date

To: [insert: Name and Address of UNDP focal point]

Dear Sir/Madam:

We, the undersigned, hereby offer to supply the goods and related services required for [insert: title of goods and services required as per ITB] in accordance with your Invitation to Bid dated .We are hereby submitting our Bid, which includes the Technical Bid and Price Schedule.

We hereby declare that:

- a) All the information and statements made in this Bid are true and we accept that any misrepresentation contained in it may lead to our disqualification;
- b) We are currently not on the removed or suspended vendor list of the UN or other such lists of other UN agencies, nor are we associated with, any company or individual appearing on the 1267/1989 list of the UN Security Council;
- c) We have no outstanding bankruptcy or pending litigation or any legal action that could impair our operation as a going concern; and
- d) We do not employ, nor anticipate employing, any person who is or was recently employed by the UN or UNDP.

We confirm that we have read, understood and hereby fully accept the Schedule of Requirements and Technical Specifications describing the duties and responsibilities required of us in this ITB, and the General Terms and Conditions of UNDP's Standard Contract for this ITB.

We agree to abide by this Bid for 120 days.

We undertake, if our Bid is accepted, to initiate the supply of goods and provision of related services not later than the date indicated in the Data Sheet.

We fully understand and recognize that UNDP is not bound to accept this Bid, that we shall bear all costs associated with its preparation and submission, and that UNDP will in no case be responsible or liable for those costs, regardless of the conduct or outcome of the evaluation.

We remain,

Yours sincerely,

Authorized Signature [In full and initials]	:
Name and Title of Signatory:	
Name of Firm:	
Contact Details:	

 $^{^2}$ No deletion or modification may be made in this form. Any such deletion or modification may lead to the rejection of the Bid.

Section 6:

Documents Establishing the Eligibility and Qualifications of the Bidder Bidder Information Form³

Date: [insert date (as day, month and year]of Bid Submission]

		ITB No.: [insert num	iber of bidding	process
		Page	of	pages
1. Bidder's Legal Name [insert Bidd	er's legal name]			
2. In case of Joint Venture (JV), legal	name of each party: [insert legal na	ime of each party in J	IV]	
3. Actual or intended Country/ies of	Registration/Operation: [insert acto	ual or intended Coun	try of Registrat	ion]
4. Year of Registration in its Location	n: [insert Bidder's year of registration	1]		
5. Countries of Operation	6. No. of staff in each Country	7.Years of Operation	on in each Cour	ntry
8. Legal Address/es in Country/ies or registration]	f Registration/Operation:[insert Bidd	ı der's legal address in	country of	
9. Value and Description of Top three	e (3) Biggest Contract for the past fiv	ve (5) years		
10. Latest Credit Rating (Score and S	ource, if any)			
 Brief description of litigation his outcomes, if already resolved. 	tory (disputes, arbitration, claims, e	etc.), indicating curre	nt status and	
12. Bidder's Authorized Representat	ive Information			
Name: [insert Authorized Represei	-			
Address: [insert Authorized Repres	sentative's Adaress] uthorized Representative's telephor	ne/fav numhersl		
Email Address: [insert Authorized I		ic/jux numbersj		
13. Are you in the UNPD List 1267.1	· · · · · · · · · · · · · · · · · · ·	or □ NO		

³The Bidder shall fill in this Form in accordance with the instructions. Apart from providing additional information, no alterations to its format shall be permitted and no substitutions shall be accepted.

14. Attached are copies of original documents of:
□All eligibility document requirements listed in the Data Sheet
☐ If Joint Venture/Consortium – copy of the Memorandum of Understanding/Agreement or Letter of Intent to form a JV/Consortium, or Registration of JV/Consortium, if registered
\Box If case of Government corporation or Government-owned/controlled entity, documents establishing legal and financial autonomy and compliance with commercial law.

Joint Venture Partner Information Form (if Registered)⁴

Date: [insert date (as day, month and year) of Bid Submission]
ITB No.: [insert number of bidding process]

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1. Bidder's Legal Name: [insert E	Bidder's legal name]		
2. JV's Party legal name: [insert JV's Party legal name]			
3. JV's Party Country of Registration: [insert JV's Party country of registration]			
4. Year of Registration: [insert Part	ty's year of registration]		
5. Countries of Operation	6. No. of staff in each Country	7.Years of Operation in each Country	
8. Legal Address/es in Country/ies registration]	of Registration/Operation: [insert Pa	rty's legal address in country of	
9. Value and Description of Top the	ree (3) Biggest Contract for the past fi	ve (5) years	
10. Latest Credit Rating (if any) :Cl	ick here to enter text.		
,	history (disputes, arbitration, claims, I. Click here to enter text.	etc.), indicating current status and	

⁴The Bidder shall fill in this Form in accordance with the instructions. Apart from providing additional information, **n**o alterations to its format shall be permitted and no substitutions shall be accepted.

13. JV's Party Authorized Representative Information
Name: [insert name of JV's Party authorized representative]
Address: [insert address of JV's Party authorized representative]
Telephone/Fax numbers: [insert telephone/fax numbers of JV's Party authorized representative]
Email Address: [insert email address of JV's Party authorized representative]
14. Attached are copies of original documents of: [check the box(es) of the attached original documents]
□All eligibility document requirements listed in the Data Sheet
□Articles of Incorporation or Registration of firm named in 2.
□In case of government owned entity, documents establishing legal and financial autonomy and compliance with commercial law.

Section 7: Technical Bid Form⁵

INSERT TITLE OF THE ITB			
Ī			
(PERTISE OF FIRM/ OR	GANISATION		
This section should fully explain the Bidder's resources in terms of personnel and facilities necessary for the performance of this requirement.			
1.1 Brief Description of Bidder as an Entity: Provide a brief description of the organization / firm submitting the Bid, its legal mandates/authorized business activities, the year and country of incorporation, and approximate annual budget, etc. Include reference to reputation, or any history of litigation and arbitration in which the organization / firm has been involved that could adversely affect or impact the delivery of goods and/or performance of related services, indicating the status/result of such litigation/arbitration.			
1.2. Financial Capacity: Based on the latest Audited Financial Statement (Income Statement and Balance Sheet) describe the financial capacity (liquidity, stand-by credit lines, etc.) of the bidder to engage into the contract. Include any indication of credit rating, industry rating, etc.			
rating, etc.		•	
rating, etc. e following information or relevant to those re		gage into the contract.	
e following information		gage into the contract.	
e following information or relevant to those red of Types of ity activities	equired for this Co	rate experience within ntract. References Contact Details (Name, Phone,	
rreet	RPERTISE OF FIRM/ OR sources in terms of personances activities, the yearce to reputation, or a ed that could adversely ting the status/result or ited Financial Statemer	RPERTISE OF FIRM/ ORGANISATION sources in terms of personnel and facilities revide a brief description of the organizations activities, the year and country tence to reputation, or any history of litigated that could adversely affect or impact ting the status/result of such litigation/ar	

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 $^{^{5}\}mbox{Technical Bids not submitted in this format may be rejected.}$

SUBSECTION 3.2 - SCOPE OF SUPPLY, TECHNICAL SPECIFICATIONS, AND RELATED SERVICES

This section should demonstrate the Bidder's responsiveness to the specification by identifying the specific components proposed, addressing the requirements, as specified, point by point; providing a detailed description of the essential performance characteristics proposed; and demonstrating how the proposed bid meets or exceeds the specifications.

- <u>2.1. Scope of Supply</u>: Please provide a detailed description of the goods to be supplied, indicating clearly how they comply with the technical specifications required by the ITB (please see Annex 4 Annex 4 should be submitted in both PDF and EXCEL formats); describe how the organization/firm will supply the goods and any related services, keeping in mind the appropriateness to local conditions and project environment.
- 2.1.1 Please describe the Freight Forwarder details and Arrangements. Ability to provide/coordinate necessary shipping services, including air, sea and cold chain delivery (if required)
- 2.1.2 Please provide the detailed Implementation Schedule.

<u>Delivery lead time is a factor of a crucial importance in this project. Please make all possible efforts to propose supply of all requested quantities within shortest timeframe possible. In case partial delivery is proposed, please provide suggested time schedule.</u>

A supporting document with full details may be annexed to this section.

- <u>2.2 Statement of Full Disclosure</u>: This is intended to disclose any potential conflict in accordance with the definition of "conflict" under Section 5 of this document, if any.
- 2.3 Other: Any other comments or information regarding the bid and its implementation.

Section 8: Price Schedule Form

The Bidder is required to prepare the Price Schedule as indicated in the Instruction to Bidders.

Please refer to Annex 5 (excel sheet) with the Price Schedule Form.

Section 9: FORM FOR BID SECURITY

(This must be finalized using the official letterhead of the Issuing Bank. Except for indicated fields, no changes may be made in this template.)

To: UNDP

WHEREAS (hereinafter called "the Bidder") has submitted a Bid to UNDP dated , to deliver goods and execute related services for(hereinafter called "the Bid"):

AND WHEREAS it has been stipulated by you that the Bidder shall furnish you with a Bank Guarantee by a recognized bank for the sum specified therein as security in the event that the Bidder:

- a) Fails to sign the Contract after UNDP has awarded it;
- b) Withdraws its Bid after the date of the opening of the Bid;
- c) Fails to comply with UNDP's variation of requirement, as per ITB Section 3; or
- d) Fails to furnish Performance Security, insurances, or other documents that UNDP may require as a condition to rendering the contract effective.

AND WHEREAS we have agreed to give the Bidder such this Bank Guarantee:

NOW THEREFORE we hereby affirm that we are the Guarantor and responsible to you, on behalf of the Bidder, up to a total of [amount of guarantee] [in words and numbers], such sum being payable in the types and proportions of currencies in which the Price Bid is payable, and we undertake to pay you, upon your first written demand 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 for the sum specified therein.

This guarantee shall be valid until 30 days after the date of validity of the bids.

SIGNATURE AND SEAL OF THE GUARANTOR BANK

Date

Name of Bank

Address

Section 10: FORM FOR PERFORMANCE SECURITY

(This must be finalized using the official letterhead of the Issuing Bank. Except for indicated fields, no changes may be made in this template.)

To:	UNDP [Insert contact information as provided in Data Sheet]
	WHEREAS [name and address of Contractor] (hereinafter called "the Contractor") has rtaken, in pursuance of Contract No dated, to deliver the goods and execute ed services (hereinafter called "the Contract"):
	AND WHEREAS it has been stipulated by you in the said Contract that the Contractor shall h you with a Bank Guarantee by a recognized bank for the sum specified therein as security impliance with his obligations in accordance with the Contract:
	AND WHEREAS we have agreed to give the Contractor such a Bank Guarantee:
sum I payak argun	NOW THEREFORE we hereby affirm that we are the Guarantor and responsible to you, on f of the Contractor, up to a total of [amount of guarantee] [in words and numbers], such being payable in the types and proportions of currencies in which the Contract Price is ole, and we undertake to pay you, upon your first written demand and without cavil or nent, any sum or sums within the limits of [amount of guarantee as aforesaid] without your ng to prove or to show grounds or reasons for your demand for the sum specified therein.
certif	This guarantee shall be valid until a date 30 days from the date of issue by UNDP of a icate of satisfactory performance and full completion of services by the Contractor.
	SIGNATURE AND SEAL OF THE GUARANTOR BANK
Date	
Name	e of Bank
Addre	ess

Section 11: General Terms and Conditions for Goods

1. ACCEPTANCE OF THE PURCHASE ORDER

This Purchase Order may only be accepted by the Supplier's signing and returning an acknowledgement copy of it or by timely delivery of the goods in accordance with the terms of this Purchase Order, as herein specified. Acceptance of this Purchase Order shall effect a contract between the Parties under which the rights and obligations of the Parties shall be governed solely by the terms and conditions of this Purchase Order, including these General Conditions. No additional or inconsistent provisions proposed by the Supplier shall bind UNDP unless agreed to in writing by a duly authorized official of UNDP.

2. PAYMENT

- 2.1.1 UNDP shall, on fulfillment of the Delivery Terms, unless otherwise provided in this Purchase Order, make payment within 30 days of receipt of the Supplier's invoice for the goods and copies of the shipping documents specified in this Purchase Order.
- 2.1.2 Payment against the invoice referred to above will reflect any discount shown under the payment terms of this Purchase Order, provided payment is made within the period required by such payment terms.
- 2.1.3 Unless authorized by UNDP, the Supplier shall submit one invoice in respect of this Purchase Order, and such invoice must indicate the Purchase Order's identification number.
- 2.1.4 The prices shown in this Purchase Order may not be increased except by express written agreement of UNDP.

3. TAX EXEMPTION

- 3.1 Section 7 of the Convention on the Privileges and Immunities of the United Nations provides, inter alia, that the United Nations, including its subsidiary organs, is exempt from all direct taxes, except charges for utilities services, and is exempt from customs duties and charges of a similar nature in respect of articles imported or exported for its official use. In the event any governmental authority refuses to recognize UNDP's exemption from such taxes, duties or charges, the Supplier shall immediately consult with UNDP to determine a mutually acceptable procedure.
- 3.2 Accordingly, the Supplier authorizes UNDP to deduct from the Supplier's invoice any amount representing such taxes, duties or charges, unless the Supplier has consulted with UNDP before the payment thereof and UNDP has, in each instance, specifically authorized the Supplier to pay such taxes, duties or charges under protest. In that event, the Supplier shall provide UNDP with written evidence that payment of such taxes, duties or charges has been made and appropriately authorized.

4. RISK OF LOSS

Risk of loss, damage to or destruction of the goods shall be governed in accordance with DDU Incoterms 2000, unless otherwise agreed upon by the Parties on the front side of this Purchase Order.

5. EXPORT LICENCES

Notwithstanding any INCOTERM 2000 used in this Purchase Order, the Supplier shall obtain any export licenses required for the goods.

6. FITNESS OF GOODS/PACKAGING

The Supplier warrants that the goods, including packaging, conform to the specifications for the goods ordered under this Purchase Order and are fit for the purposes for which such goods are ordinarily used and for purposes expressly made known to the Supplier by UNDP, and are free from defects in workmanship and materials. The Supplier also warrants that the goods are contained or packaged adequately to protect the goods.

7. INSPECTION

- 1. UNDP shall have a reasonable time after delivery of the goods to inspect them and to reject and refuse acceptance of goods not conforming to this Purchase Order; payment for goods pursuant to this Purchase Order shall not be deemed an acceptance of the goods.
 - 2. Inspection prior to shipment does not relieve the Supplier from any of its contractual obligations.

8. INTELLECTUAL PROPERTY INFRINGEMENT

The Supplier warrants that the use or supply by UNDP of the goods sold under this Purchase Order does not infringe any patent, design, trade-name or trade-mark. In addition, the Supplier shall, pursuant to this warranty, indemnify, defend and hold UNDP and the United Nations harmless from any actions or claims brought against UNDP or the United Nations pertaining to the alleged infringement of a patent, design, trade-name or trade-mark arising in connection with the goods sold under this Purchase Order.

9. RIGHTS OF UNDP

In case of failure by the Supplier to fulfil its obligations under the terms and conditions of this Purchase Order, including but not limited to failure to obtain necessary export licenses, or to make delivery of all or part of the goods by the agreed delivery date or dates, UNDP may, after giving the Supplier reasonable notice to perform and without prejudice to any other rights or remedies, exercise one or more of the following rights:

- a) Procure all or part of the goods from other sources, in which event UNDP may hold the Supplier responsible for any excess cost occasioned thereby.
- b) Refuse to accept delivery of all or part of the goods.
- c) Cancel this Purchase Order without any liability for termination charges or any other liability of any kind of UNDP.

10. LATE DELIVERY

Without limiting any other rights or obligations of the parties hereunder, if the Supplier will be unable to deliver the goods by the delivery date(s) stipulated in this Purchase Order, the Supplier shall (i) immediately consult with UNDP to determine the most expeditious means for delivering the goods and (ii) use an expedited means of delivery, at the Supplier's cost (unless the delay is due to Force Majeure), if reasonably so requested by UNDP.

11. ASSIGNMENT AND INSOLVENCY

- 11.1. The Supplier shall not, except after obtaining the written consent of UNDP, assign, transfer, pledge or make other disposition of this Purchase Order, or any part thereof, or any of the Supplier's rights or obligations under this Purchase Order.
- 11.2. Should the Supplier become insolvent or should control of the Supplier change by virtue of insolvency, UNDP may, without prejudice to any other rights or remedies, immediately terminate this Purchase Order by giving the Supplier written notice of termination.

12. USE OF UNDP OR UNITED NATIONS NAME OR EMBLEM

The Supplier shall not use the name, emblem or official seal of UNDP or the United Nations for any purpose.

13. PROHIBITION ON ADVERTISING

The Supplier shall not advertise or otherwise make public that it is furnishing goods or services to UNDP without specific permission of UNDP in each instance.

14. CHILD LABOUR

The Supplier represents and warrants that neither it nor any of its affiliates is engaged in any practice inconsistent with the rights set forth in the Convention on the Rights of the Child, including Article 32 thereof, which, inter alia, requires that a child shall be protected from performing any work that is likely to be hazardous or to interfere with the child's education, or to be harmful to the child's health or physical, mental, spiritual, moral or social development.

Any breach of this representation and warranty shall entitle UNDP to terminate this Purchase Order immediately upon notice to the Supplier, without any liability for termination charges or any other liability of any kind of UNDP.

15. MINES

The Supplier represents and warrants that neither it nor any of its affiliates is actively and directly engaged in patent activities, development, assembly, production, trade or manufacture of mines or in such activities in respect of components primarily utilized in the manufacture of Mines. The term "Mines" means those devices defined in Article 2, Paragraphs 1, 4 and 5 of Protocol II annexed to the Convention on Prohibitions and Restrictions on the Use of Certain Conventional Weapons Which May Be Deemed to Be Excessively Injurious or to Have Indiscriminate Effects of 1980.

Any breach of this representation and warranty shall entitle UNDP to terminate this Purchase Order immediately upon notice to the Supplier, without any liability for termination charges or any other liability of any kind of UNDP.

16. SETTLEMENT OF DISPUTES

16.1 Amicable Settlement

The Parties shall use their best efforts to settle amicably any dispute, controversy or claim arising out of, or relating to this Purchase Order or the breach, termination or invalidity thereof. Where the Parties wish to seek such an amicable settlement through conciliation, the conciliation shall take place in accordance with the UNCITRAL Conciliation Rules then obtaining, or according to such other procedure as may be agreed between the Parties.

16.2 Arbitration

Unless, any such dispute, controversy or claim between the Parties arising out of or relating to this Purchase Order or the breach, termination or invalidity thereof is settled amicably under the preceding paragraph of this Section within sixty (60) days after receipt by one Party of the other Party's request for such amicable settlement, such dispute, controversy or claim shall be referred by either Party to arbitration in accordance with the UNCITRAL Arbitration Rules then obtaining, including its provisions on applicable law. The arbitral tribunal shall have no authority to award punitive damages. The Parties shall be bound by any arbitration award rendered as a result of such arbitration as the final adjudication of any such controversy, claim or dispute.

17. PRIVILEGES AND IMMUNITIES

Nothing in or related to these General Terms and Conditions or this Purchase Order shall be deemed a waiver of any of the privileges and immunities of the United Nations, including its subsidiary organs.

18. SEXUAL EXPLOITATION:

- 18.1 The Contractor shall take all appropriate measures to prevent sexual exploitation or abuse of anyone by it or by any of its employees or any other persons who may be engaged by the Contractor to perform any services under the Contract. For these purposes, sexual activity with any person less than eighteen years of age, regardless of any laws relating to consent, shall constitute the sexual exploitation and abuse of such person. In addition, the Contractor shall refrain from, and shall take all appropriate measures to prohibit its employees or other persons engaged by it from, exchanging any money, goods, services, offers of employment or other things of value, for sexual favors or activities, or from engaging in any sexual activities that are exploitive or degrading to any person. The Contractor acknowledges and agrees that the provisions hereof constitute an essential term of the Contract and that any breach of this representation and warranty shall entitle UNDP to terminate the Contract immediately upon notice to the Contractor, without any liability for termination charges or any other liability of any kind.
- 18.2 UNDP shall not apply the foregoing standard relating to age in any case in which the Contractor's personnel or any other person who may be engaged by the Contractor to perform any services under the Contract is married to the person less than the age of eighteen years with whom sexual activity has occurred and in which such marriage is recognized as valid under the laws of the country of citizenship of

such Contractor's personnel or such other person who may be engaged by the Contractor to perform any services under the Contract.

19.0 OFFICIALS NOT TO BENEFIT:

The Contractor warrants that no official of UNDP or the United Nations has received or will be offered by the Contractor any direct or indirect benefit arising from this Contract or the award thereof. The Contractor agrees that breach of this provision is a breach of an essential term of this Contract.

20. AUTHORITY TO MODIFY:

Pursuant to the Financial Regulations and Rules of UNDP, only the UNDP Authorized Official possess the authority to agree on behalf of UNDP to any modification of or change in this Agreement, to a waiver of any of its provisions or to any additional contractual relationship of any kind with the Contractor. Accordingly, no modification or change in this Contract shall be valid and enforceable against UNDP unless provided by an amendment to this Agreement signed by the Contractor and jointly by the UNDP Authorized Official.