File No. 31026/36/2016-MD
Government of India
Ministry of Chemicals and Fertilizers
Department of Pharmaceuticals

Janpath Bahwan, New Delhi
Dated, the 15 March, 2018

Subject: Guidelines for implementation of “Public procurement (Preference to Make in India) Order – 2017”- reg.

Department of Industrial Policy and Promotion (DIPP), Ministry of Commerce and Industry, Government of India vide Order No. P-45021/2/2017-B.E.-II dated 15.06.2017 has issued Public Procurement (Preference to Make in India), Order 2017. Department of Pharmaceuticals has been identified as Nodal Department for implementation of this Order in respect of Medical Devices Sector.

2. In this connection, this Department has prepared draft guidelines for implementation of this Order which are enclosed. All stakeholders are requested to furnish their comments/ views on these guidelines within 21 days from the date of uploading of the same on this Department website. The comments could be provided at Email: parveen.19@gov.in

Encl: As above

(Parveen Kumar)
Under Secretary to the Government of India
Telephone No. 23352298

To:

Director (NIC)/DoP with request to upload the guidelines on the Department website immediately
GUIDELINES

Subject: Public Procurement (Preference to Make in India) Order (PPO), 2017 – Guidelines for Public procurement of Medical Devices–reg.

No. 31026/36/2016-MD: Whereas Department of Industrial Policy and Promotion (DIPP), pursuant to Rule 153(iii) of the General Financial Rules 2017, has issued Public Procurement(Preference to Make in India) Order (PPO), 2017 vide no. P-4502/2/2017-B.E.-II dated 15.06.2017.

Whereas DIPP, in order to facilitate the implementation of the PPO, 2017, vide D.O. No. P-45021/2/2017-BE-II dated 14.08.2017 has identified Department of Pharmaceuticals (DoP) as the Nodal Department for implementing the provisions of the PPO 2017 relating to goods &services related to pharmaceuticals sector.

Whereas Para 3 of PPO, 2017 makes it mandatory for procuring entities to give purchase preference to local suppliers,Para 5 of PPO, 2017 empowers Nodal Ministry to prescribe percentage and the manner of calculation of minimum local content in respect of any particular item relating to Pharmaceutical sector and Para 9 of PPO, 2017 deals with verification of local content.

Now, therefore, DoP issues the following guidelines for implementation of the provisions of the PPO, 2017 with respect to public procurement of medical devices:

1) Percentage of Minimum Local Content: Medical Device Industry (MDI) is a multi-product industry responsible for provisioning of wide variety of crucial medical products ranging from simple tongue depressors & glucometer strips to large radiology & electronic medical equipments. The medical devices industry can be broadly classified as consisting of (a) medical disposables and consumables; (b) medical electronics, hospital equipment, surgical instruments; (c) Implants; and (d) In-Vitro Devices/Diagnostic Reagents. Individually there are around 5000 different kinds of medical devices and it is not practical to prescribe the local content and percentage of preference in domestic procurement for each medical device.

Moreover, DoP is in the process of collecting accurate and reliable data regarding total capacity and production of various categories of medical devices in India, regarding the level of competition in the market in different segment of medical devices and regarding the processes involved in the manufacture of medical devices for prescribing the percentage of minimum local content for each category of medical devices, for purchase of supplies only from local suppliers where the estimated value of procurement is Rs. 50 Lakhs or less and for determining the manner of calculation of local content in the medical devices to be procured by the public agencies. The percentage of local content, the manner
of calculation of the local content and the provision of supplies to be procured from local suppliers only where the estimated value of procurement is Rs. 50 Lakhs or less may be revised after one year or as soon as the relevant data in this regard becomes available whichever is earlier.

However for the time being, based on the present level of the understanding of the medical device market in India and discussion with various industry representatives, the following percentages of minimum local content in domestic medical devices for public procurement are prescribed for the various segments of medical devices:

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<thead>
<tr>
<th>Category of Medical Devices</th>
<th>% of Local Content</th>
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<tbody>
<tr>
<td>Medical disposables and consumables</td>
<td>50%</td>
</tr>
<tr>
<td>Medical electronics, hospital equipment, surgical instruments</td>
<td>25%</td>
</tr>
<tr>
<td>Implants</td>
<td>40%</td>
</tr>
<tr>
<td>Diagnostic Reagents/IVDs</td>
<td>25%</td>
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2) **Manner of calculation of Local Content:**
   i. Local content of Medical Device shall be computed on the basis of the cost of domestic components in the device compared to the total cost of the device. The total cost of product shall be the cost incurred for the production of the medical device including direct component i.e. material cost, manpower cost and overhead costs including profit but excluding taxes and duties.
   ii. The determination of local content cost shall be based on the following:
      a) In the case of direct component (material), based on the country of origin
      b) In the case of manpower, based on domestic manpower
   iii. The calculation of local content of the combination of several kinds of goods shall be based on the ratio of the sum of multiplication of local content of each goods with the acquisition price of each goods to the acquisition price of combination of goods.
   iv. Format of calculation of local content shall be as contained in Enclosure-I.

3) **Requirement of Purchase Preference:** Purchase preference shall be given to local suppliers by all procuring entities as per provisions laid down in para 3 of PPO, 2017 subject to the condition that para 3(a) of the PPO 2017 shall be applicable only when there are two or more than two local suppliers for any tender of value upto Rs. 50 Lakhs and they certify that they can supply the desired medical devices in the required quantities.

4) **Verification of Local Content:**
   a) The local supplier at the time of tender, bidding or solicitation shall be required to furnish self-certification of local content in the format as contained in Enclosure-II.
   b) In each tender, procuring entity shall clearly mention the details of its competent authority which is empowered to look into procurement related complaints and the fees for such complaints, relating to implementation of PPO, 2017.
   c) In case a complaint is received by the procuring entity against the claim of a bidder regarding domestic value addition in medical device, the procuring entity shall have full rights to inspect and examine all the related documents and take a decision. In case any clarification is needed, matter may be referred to DoP.
   d) Any complaint referred to the procuring entity shall be submitted along with all necessary documentation in support of the complaint regarding domestic value addition.
claimed in medical device and shall be disposed of within 4 weeks of the reference by the procuring entity.

e) In case, the complaint is referred to DoP by a bidder or procuring entity, the grievance redressal committee to be set up under DoP for the purpose shall dispose of the complaint.

f) In case, the matter is referred to DoP, the grievance redressal committee shall dispose of the complaint within 4 weeks of its reference and receipt of all documents from the bidder after taking in consideration, the view of the procuring entity. The bidder shall be required to furnish the necessary documentation in support of the local content claimed in medical devices to the grievance redressal committee under DoP within 2 weeks of the reference of the matter. If no information is furnished by the bidder, the grievance redressal committee may take further necessary action, in consultation with procuring entity to establish the bonafides of the claim.

g) In case of reference of any complaint to DoP by the concerned bidder, there would be a fee of Rs. 2 Lakh or 1% of the value of the medical devices being procured (subject to a maximum of Rs. 5 Lakh), whichever is higher, to be paid by way of a Demand Draft to be deposited with the procuring entity, along with the complaints by the complainant. In case, the complaint is found to be incorrect, the complaint fee shall be forfeited. In case, the complaint is upheld and found to be substantially correct, deposited fee of the complainant would be refunded without any interest.

5) All other provisions of PPO, 2017 shall be applicable as such and shall be adhered to by all procuring agencies for procurement of any medical device.

(Parveen Kumar)
Under Secretary to the Govt. of India
Ph. 23352298
Calculation of Local Content

| Name of manufacturer | Calculation by Manufacturer  
|-----------------------|-----------------------------|

| Cost Component | Calculation by Manufacturer  
|----------------|-----------------------------|

| Cost Component | Calculation by Manufacturer  
<table>
<thead>
<tr>
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<tbody>
<tr>
<td>Cost (Domestic Component) a</td>
<td>Cost (Imported Component) b</td>
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</table>

Note:

I. **Cost (Domestic Component):** Sum of the costs of all inputs which go into the product (including duties and taxes levied on procurement of inputs except those for which credit/set-off can be taken) which have not been imported directly or through a domestic trader or an intermediary.

II. **Cost (Imported Component):** Sum of the costs of all inputs which go into the product (including duties and taxes levied on procurement of inputs except those for which credit/set-off can be taken).
Format for Affidavit of Self Certification regarding Local Content in a Medical Device to be provided on Rs. 100/- Stamp Paper

Date: ______________

I ___________________________________________________________________________, S/o, D/o, W/o ____________________________________________________________________________, do hereby solemnly affirm and declare as under:

That I will agree to abide by the terms and conditions of the policy of Government of India issued vide Notification No:

That the information furnished hereinafter is correct to best of my knowledge and belief and I undertake to produce relevant records before the procuring entity or any authority so nominated by the Department of Pharmaceuticals, Government of India for the purpose of assessing the local content.

That the local content for all inputs which constitute the said medical device has been verified by me and I am responsible for the correctness of the claims made therein.

That in the event of the domestic value addition of the product mentioned herein is found to be incorrect and not meeting the prescribed value-addition norms, based on the assessment of an authority so nominated by the Department of Pharmaceuticals, Government of India for the purpose of assessing the local content, action will be taken against me as per Order No. P-45021/2/2017-B.E.-II dated 15.06.2017 and Notification No. 31026/36/2016-MD dated …………

I agree to maintain the following information in the Company’s record for a period of 8 years and shall make this available for verification to any statutory authority:

i) Name and details of the Domestic Manufacturer (Registered Office, Manufacturing unit location, nature of legal entity)

ii) Date on which this certificate is issued

iii) Medical devices for which the certificate is produced

iv) Procuring entity to whom the certificate is furnished

v) Percentage of local content claimed

vi) Name and contact details of the unit of the manufacturer

vii) Sale Price of the product

viii) Ex-Factory Price of the product

ix) Freight, insurance and handling

x) Total Bill of Material

xi) List and total cost value of inputs used for manufacture of the medical device

xii) List and total cost of inputs which are domestically sourced. Value addition certificates from suppliers, if the input is not in- house to be attached.

xiii) List and cost of inputs which are imported, directly or indirectly

For and on behalf of __________________________ (Name of firm/entity)

Authorized signatory (To be duly authorized by the Board of Director)