Biofertilizers and Organic Fertilizers in Fertilizer (Control) Order, 1985

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Department of Agriculture and Cooperation, Ministry of Agriculture, Govt of India,
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Regional Centre of Organic Farming, Bengaluru
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Regional Centre of Organic Farming, Panchkula
Regional Centre of Organic Farming, Imphal
Regional Centre of Organic Farming, Jabalpur
Regional Centre of Organic Farming, Nagpur

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Biofertilizers and Organic Fertilizers Covered in Fertilizer (Control) Order, 1985
(Amendments - March 2006, November, 2009 and June 2012)

General Rules

Ministry of Agriculture, Department of Agriculture and Cooperation, Government of India, New Delhi, vide their order Dated 24th March, 2006 included biofertilizers and organic fertilizers under section 3 of the Essential Commodities Act, 1955 (10 of 1955), in Fertilizer (Control) Order, 1985. These rules were further amended in respect of applicability, specifications and testing protocols vide Gazette notification 3 November, 2009.

Abstract details of these rules, including the amendment are as follows:

Definition

(Clause 2 a, aa)
Biofertilizers  - Biofertilizers means the product containing carrier based (solid or liquid) living microorganisms which are agriculturally useful in terms of nitrogen fixation, phosphorus solubilization or nutrient mobilization, to increase the productivity of the soil and/or crop.

(Clause 2 oo)
Organic Fertilizer - Organic fertilizer means substances made up of one or more unprocessed material(s) of a biological nature (plant/animal) and may include unprocessed mineral materials that have been altered through microbiological decomposition process”.

(Clause 2 nna)
“Non-edible de-oiled cake fertilizer” means substance obtained as residue after oil extraction (by expeller and/or through solvent extraction) from crushed seeds of non-edible oilseeds (such as castor, neem) for use in soil as fertilizer.

Biofertilizers and organic fertilizers are also covered under the broad term of fertilizers
As per the clause 2(h) “Fertiliser means any substance used or intended to be used as a fertilizer of the soil and/or crop specified in Part A of Schedule – I and includes a mixture of fertilizer, special mixture of fertilizer, biofertilizers specified in Schedule III, organic fertilizers specified in Schedule IV and non-edible de-oiled cake fertilizers specified in schedule V”.
Prescribed standards

(Clause 2(q) [iv])
Prescribed standards in relation to a Biofertilizer means the prescribed limit of components included in column 1 of part A of Schedule III, and the standard set out in the corresponding entry in column 2, subject to the limits of permissible variation as specified in part B of that schedule”.

Clause 2(q){v}
Prescribed standards in relation to a Organic fertilizer means the prescribed limit of components included in column 1 of part A of Schedule IV, and the standard set out in the corresponding entry in column 2, subject to the limits of permissible variation as specified in part B of that schedule.

Clause 2(q){vi}
In relation to a Non-edible, De-oiled cake meal fertilizer specified in column (2) of part A of schedule V, the standard set out in the corresponding entry in column (2) of the said part, subject to the limits of permissible variation as specified in part B of that schedule”.

Clause 3
Fixation of prices
Not applicable in case of Biofertilizers and organic fertilizers.

Clause 4
Display of stock position and price list of fertilizers
Every dealer, who makes or offers to make a retail sale of any fertilizer (including biofertilizer and organic fertilizer), shall prominently display in his place of business:-

(a)  The quantities of opening stock of different fertilisers held by him on each day;
(b)  A list of prices or rates of such fertilisers (including biofertilizer and organic fertilizer)

Clause 5
Issue of cash/credit memorandum
Every dealer shall issue a cash or credit memorandum to a purchaser of a biofertilizer and organic fertilizer in Form M.

Clause 6
Allocation of fertilisers to various States
Not applicable in case of Biofertilizers and organic fertilizers.
Clause 7
Registration of Industrial dealers and authorization of other dealers
No person shall sell, offer for sale or carry on the business of selling of biofertilizer or organic fertilizer at any place as wholesale dealer or retail dealer except under and in accordance with clause 8: Provided that a State Government may, if it considers it necessary or expedient, by notification in the Official Gazette, exempt from the provisions of this clause any person selling fertilizer to farmers in such areas and subject to such conditions as may be specified in that notification.

Clause 8
Application for intimation or registration
1. Every person intending to sell or offer for sale or carrying on the business of selling of fertilizer as Industrial Dealer shall obtain a certificate of registration from the controller by making an application in Form A together with the fee prescribed under clause 36 and a Certificate of source in Form O.

2. Every person including a manufacturer, an importer, a pool handling agency, wholesaler and a retail dealer intending to sell or offer for sale or carrying on the business of selling of fertilizer shall make a Memorandum of Intimation to the Notified Authority, in Form A1 duly filled in, in duplicate, together with the fee prescribed under clause 36 and certificate of source in Form O.

3. On receipt of a Memorandum of Intimation, complete in all respects, the Notified Authority shall issue an acknowledgement of receipt in Form A2 and it shall be deemed to be an authorization letter granted and the concerned person as authorised dealer for the purposes of this Order.

Provided that a certificate of registration granted before the commencement of the Fertiliser (Control) Amendment Order, 2003, shall be deemed to be an authorization letter granted under the provisions of this Order.

Provided further that where the applicant is a State Government, a manufacturer or an importer or a pool-handling agency, it shall not be necessary for it or him to submit Form O.

Provided also that a separate Memorandum of Intimation shall be submitted by an applicant for whole sale business or retail dealership, as the case may be.

Provided also that where fertilizers (BFs and OFs) are obtained for sale from different sources, a certificate of source from each such source shall be furnished in Form O."
Provided also that where the manufacturer of organic fertilizer is a State Government or municipality, it shall not be necessary for it to obtain the authorisation letter:

Provided also that where the manufacturer of vermin-compost, other than a State Government or municipality, has annual production capacity less than 50 metric tonnes, it shall not be necessary for him to obtain the authorisation letter.

Clause 9
Grant or refusal of certificate of registration
The Controller, shall grant a certificate of registration in Form 'B' within thirty days of the receipt of application to any person who applies for it under clause 8; Provided that no certificate of registration shall be granted to a person:

a. if his previous certificate of registration is under suspension; or
b. if his previous certificate of registration has been cancelled within a period of one year immediately preceding the date of application; or
c. if he has been convicted of an offence under the Act, or any Order made there under within three years immediately preceding the date of making the application; or
d. if he fails to enclose with the application a certificate of source; or
e. if the application is incomplete in any respect; or
f. if he makes an application for obtaining the certificate of registration for industrial dealer and, excepting if he is a manufacturer, importer or pool handling agency, holds an authorization letter for wholesale dealer or retail dealer or both, and as the case may be, the vice-versa.

Clause 10
Period of validity of certificate of registration and letter of authorization
Every certificate of registration granted under clause 9 and every authorization letter issued under clause 8 shall, unless renewed, suspended or cancelled, be valid for a period of three years from the date of its issue.

Clause 11
Renewal of certificates of registration and authorization letters
(1) Every holder of a certificate of registration granted under clause 9 or authorization letter granted or deemed to have been granted under clause 8, desiring to renew such certificate or authorization letter shall, before the date of expiry of such certificate of registration or authorization letter, as the case may be, make an application for
renewal to the Controller, in Form C, or to the Notified Authority in Form A1, respectively, in duplicate, together with the fee prescribed under clause 36 for such renewal and a certificate of source as required under clause 8.

(2) On receipt of an application under sub-clause (1), together with such fee and certificate of source, the controller may renew the certificate of registration or the Notified Authority, as the case may be shall issue acknowledgement receipt of renewal in form A 2. Provided that a certificate of registration shall not be renewed if the holder of the same did not sell any fertilizer during the period of one year immediately preceding the date of expiry of the period of validity.

(3) If any application for renewal is not made before the expiry of the period of validity of the certificate of registration or, as the case may be, the authorization letter but is made within one month from the date of such expiry, the certificate of registration or, as the case may be, the authorization letter shall be dealt as provided in sub-clause (2) on payment of such additional fee as may be prescribed under clause 36 in addition to the fee for renewal.

(4) Where the application for renewal of certificate of registration is made within the time specified in sub-clause (1) or sub-clause (3), the applicant shall be deemed to have held a valid certificate of registration until such date as the controller passes orders on the application for renewal.

(5) If an application for renewal of a certificate of registration or authorization letter is not made within one month from the date of expiry of their period of validity, the same shall be deemed to have lapsed on the date on which its validity expired and any business carried on after that date shall be deemed to have been carried on in contravention of clause 7.

Clause 12
Restriction on preparation
No person shall carry on the business of preparing any Biofertilizers or Organic fertilisers except under and in accordance with the terms and conditions of a certificate of manufacture granted to him under clauses 15 or 16.

Clause 13
Conformity in manufacture (Clause 13 b and c)
  b. No person shall manufacture any Bio-fertilizer unless such Bio-fertilizer conforms to the standards set out in the part A of schedule-III.
c. No person shall manufacture any Organic fertilizer unless such organic fertilizer conforms to the standards set out in the part A of schedule-IV.

Clause 14
Requirement for certificate of manufacture
(3) Every person desiring to obtain a Certificate of Manufacture for preparation or organic fertilizer or biofertiliser shall make an application in Form D, in duplicate, together with a fee prescribed therefore under clause 36, to Registering authority.

“Provided that where the manufacturer of organic fertilizer is a State Government or a municipality, it shall not be necessary for it to obtain the Certificate of Manufacture:

Provided further that where the manufacturer of vermi-compost, other than a State Government or municipality, has annual production capacity less than fifty metric tonnes, it shall not be necessary for him to obtain the Certificate of Manufacture for preparation of vermi-compost.”

Clause 15
Grant or refusal of certificate of manufacture
Sub-Clause (1) - On receipt of an application under Clause 14 the registering authority shall, by order in writing, either grant or refuse to grant the certificate of manufacture in respect of biofertiliser or organic fertilizers and shall, within 45 days from the date of receipt of application, furnish to the applicant a copy of the order so passed.

Sub-Clause (2) - Where an application for a certificate of manufacture for biofertilisers or organic fertilizers is not refused under sub-clause (1), the registering authority, shall within 45 days from the date of receipt of the application, grant a certificate of manufacture in Form F.

Clause 16
Not applicable to Biofertilizers and organic fertilizers.

Clause 17
Period of validity of a certificate of manufacture for preparation of biofertilizers or organic fertilizers
Every certificate of manufacture under Clause 15 for preparation of Biofertilizers or organic fertilizers, shall, unless suspended or cancelled, be valid for a period of three years from the date of issue.
Clause 18
Renewal of certificate of manufacture
1. Every holder of certificate of a manufacture for preparation of Biofertilizer or organic fertilizers desiring to renew the certificate, shall before the date of expiry of the said certificate of manufacture make an application to the registering authority in Form D in duplicate, together with the fee prescribed for this purpose under clause 36.

2. On receipt of application for renewal as provided in sub-clause (1) and keeping in view the performance of applicant and other relevant circumstances, the registering authority may, if he so decides, renew the certificate of manufacture by endorsement on Form F and in case the certificate of manufacture is not renewed, the registering authority shall record in writing his reasons for not renewing the certificate of manufacture.

3. Application made after the expiry but within one month can also be renewed on payment of additional fee.

4. Where application is made within the stipulated period, the applicant deemed to have valid certificate of manufacture till registering authority passes order on the application.

5. If application for renewal is not made within the stipulated period the certificate of manufacture shall be deemed to have expired immediately on the expiry of its validity period and any business carried after that date shall be deemed to have been carried on in contravention of clause 12.

Clause 19
Restriction on manufacture/import, sale and distribution of fertilizers (BFs and OFs)
No person shall himself or by any other person on his behalf:-

a. manufacture/import for sale, sell, offer for sale, stock or exhibit for sale or distribute any fertiliser which is not of prescribed standard;

b. manufacture/Import for sale, sell, offer for sale, stock or exhibit for sale, or distribute any mixture of fertilisers, which is not of prescribed standard** (subject to such limits of permissible variation as may be specified from time to time by the Central Government) or special mixture of fertilisers which does not conform to the particulars specified In the certificate of manufacture granted to him under this Order in respect of such special mixture.

c. sell, offer for sale, stock or exhibit for sale or distribute:-
i. any fertiliser the container whereof is not packed and marked in the manner laid down in this Order;

ii. any fertiliser which is an imitation of or a substitute for another fertiliser under the name of which it is sold;

iii. Any fertilizer which is adulterated;

iv. any fertiliser the label or container whereof bears the name of any individual firm or company purporting to be manufacturer/Importer of the fertiliser, which individual, firm or company is fictitious or does not exist.

v. any fertiliser, the label or container whereof or anything accompanying therewith bears any statement which makes a false claim for the fertiliser or which is false or misleading in any material particular.

vi. any substance as a fertiliser which substance is not, in fact, a fertiliser; or

vii. any fertilizer without exhibiting the minimum guaranteed percentage by weight of plant nutrient.

Provided that specifications of city compost in Schedule IV shall, in case of municipalities, be applicable only when it is traded in packaged form for use in agriculture:

Provided further that the specifications of vermi-compost in Schedule IV shall be applicable only in such cases where it is sold in packaged form and for agricultural purposes.

Provided also that the specifications of non-edible de-oiled cake fertilizer in schedule V shall be applicable only in such cases where it is sold in packaged form for agricultural purposes.

Clause 20
Not applicable on Biofertilizers and organic fertilizers.

Clause 21
aa. Requirement of packing and marking
Every container in which any Biofertiliser, Organic fertilizer and non-edible de-oiled cake fertilizer is packed shall conspicuously be superscribed with the word “BIO-FERTILISER/ ORGANIC FERTILISER / NON-EDIBLE DE-OILED CAKE FERTILIZER” and shall bear only such particulars and unless otherwise required under any law nothing else, as may from time to time, be specified by the Controller in this behalf.
Clause 21 A
Manufacturers to comply with certain requirements for laboratory facilities
Every manufacturer shall, in order to ensure quality of their product, possess minimum laboratory facility, as may be specified from time to time by the controller.

Clause 24
Manufacturers/Pool handling agencies to appoint officers responsible with compliance of the Order
Every manufacturing organization, **importer and pool handling agency shall appoint in that organization and in consultation with the Central Government, an officer, who shall be responsible for compliance with the provisions of this Order.

Clause 25
Not applicable on Biofertilizers and organic fertilizers

Clause 26
Appointment of registering authority
The State Government may, by notification in the Official Gazette, appoint such number of persons, as it thinks necessary, to be registering authorities for the purpose of this Order for industrial dealers, and may, in any such notification define the limits of local area within which each such registering authority shall exercise his jurisdiction.

Appointment of Notified authorities
The State Government may, by notification in the Official Gazette, appoint such number of persons, as it thinks necessary, to be Notified Authorities for the purpose of this Order and define the local limits within which each such Notified Authority shall exercise his jurisdiction.

Clause 27
Appointment of inspectors
The State Government or Central Government may by notification in the official Gazette appoint such number of persons, as it thinks necessary to be inspectors of biofertilisers for the purpose of this order and define the limits of local area within which each inspector shall exercise his jurisdiction.

Clause 27 B
Qualifications for appointment of inspectors for biofertilizer, organic fertilizer and non-edible de-oiled cake fertilizer
No person shall be eligible for appointment as inspector of biofertilisers/organic fertilizers under this order unless he may possess the following qualifications namely:
1. Graduate in Agriculture or science with Chemistry/ Microbiology as one of the subjects and
2. Training or experience in the field of quality control of biofertilisers/ organic fertilizers/ non-edible de-oiled cake fertilizer.

Clause 28
Powers of Inspectors
(1) An inspector may, with a view to securing compliance with this Order:
   (a) require any manufacturer, importer, pool handling agency, wholesale dealer or retail dealer to give any information in his possession with respect to the manufacture, storage and disposal of any fertilizer manufactured or, in any manner handled by him.

   (b) Provided that the importer shall prepare the sampling details in duplicate in form J1 and handover one copy of the same to the dealer or his representative from whom the sample has been drawn.
   (ba) draw samples of any biofertilisers in accordance with the procedure of drawals of samples laid down in schedule III.
   (bb) draw samples of any organic fertilisers in accordance with the procedure of drawal of samples laid down in schedule IV.
   (bc) draw samples of any Non-edible De-oiled cake meal fertilizer in accordance with procedure of drawl of samples laid down in schedule V.”

   (c) enter upon and search any premises where any fertiliser is manufactured/ Imported or stored or exhibited for sale, if he has reason to believe that any fertiliser has been or is being manufactured/imported, sold, offered for sale, stored, exhibited for sale or distributed contrary to the provisions of this Order;

   (d) seize or detain any fertiliser in respect of which he has reason to believe that a contravention of this Order has been or is being or is [attempted] to be committed;

   (e) seize any books of accounts or documents relating to manufacture, storage or sale of fertilisers, etc. in respect of which he has reason to believe that any contravention of this Order has been or is being or is about to be committed;

   (f) Provided that the Inspector shall give a receipt for such fertilisers or books of accounts or documents so seized to the person from whom the same have been seized; Provided further that the books of accounts or documents so seized shall be returned to the person from whom they were seized after copies thereof or extracts therefrom as certified by such person, have been taken.
(2) Subject to the proviso to paragraphs (d) and (e) of sub-clause (1), the provisions of the Code of Criminal Procedure, 1973 (2 of 1974) relating to search and seizure shall, so far as may be, apply to searches and seizures under this clause. Provided also that the inspector shall give the stop sale notice in writing to the person whose stocks have been detained and initiate appropriate action as per the provisions of this order within a period of twenty one days. If no action has been initiated by the inspector within the said period of twenty one days from the date of issue of the said notice, the notice of stop sale shall be deemed to have been revoked.

(3) Where any fertiliser is seized by an inspector under this clause, he shall forthwith report the fact of such seizure to the collector whereupon the provisions of sections 6A, 6B, 6C, 6D and 6E of the Act, shall apply to the custody, disposal and confiscation of such fertilisers.

(4) Every person, if so required by an inspector, shall be bound to afford all necessary facilities to him for the purpose of enabling him to exercise his powers under sub-clause (1).

Clause 29 1A and 1B
Analysis of samples
1A. A biofertiliser sample drawn by an inspector shall be analysed in accordance with the instructions laid down in schedule III in the National Centre of Organic Farming, Ghaziabad or Regional Centres of Organic Farming at Bangalore, Bhubaneshwar, Hissar, Imphal, Jabalpur and Nagpur or any other laboratory notified by Central or State Government.

1B. A organic fertilizer and non-edible de-oiled cake fertilizer sample drawn by an inspector shall be analysed in accordance with the instructions laid down in schedule IV in the National Centre of Organic Farming, Ghaziabad or Regional Centres of Organic Farming at Bangalore, Bhubaneshwar, Hissar, Imphal, Jabalpur and Nagpur or any other laboratory notified by Central or State Government.

(2) Every laboratory referred to in sub-clause (1) shall, in order to ensure accurate analysis, of samples, possess minimum equipment and other laboratory facilities, as may be specified from time to time by the Controller in this behalf.

Clause 29 C
Laboratories for referee Analysis of Biofertiliser
1. National Centre of Organic Farming, Ghaziabad or Regional centre of Organic Farming at Bangalore, Bhubaneshwar, Hissar
Panchkula), Imphal, Jabalpur and Nagpur and every laboratory referred
to in clause 29(1A) shall be designated as referee laboratory for the
purpose of analysis of any sample of Biofertiliser provided that no such
laboratory which carried out the first analysis of fertilizer sample shall be
so designated in respect of that sample.

Provided further that in respect of any sample the analysis of which has
been challenged may be sent for referee analysis to any one of the
other laboratories except those which are located in the state or where
the first analysis has been done, provided that National Centre of
Organic Farming, Ghaziabad and Regional Centre of Organic Farming
at Bangalore, Bhubaneshwar, Hissar (shifted to Panchkula), Imphal,
Jabalpur and Nagpur shall be considered as one group of laboratories
and a sample first analysed by any one of them, shall not be sent for
referee analysis to any other in that group, but only to any other
laboratory notifyed by a State Government or Central Government.

2. Notwithstanding anything contained in this order, the Appellate
Authority as specified in sub-clause 32A in case of sample collected by
the state Government laboratory, or the Controller, in case of sample
collected by National Centre of Organic Farming, Ghaziabad or
Regional Centre of Organic Farming at Bangalore, Bhubaneshwar,
Hissar (shifted to Panchkula), Imphal, Jabalpur and Nagpur, as the
case may be, shall decide and send, one of the two remaining samples,
for reference analysis as provided under sub-clause (1).

Clause 29D
Laboratories for referee analysis of Organic fertilizer

1. National Centre of Organic Farming, Ghaziabad or Regional Centre of
Organic Farming at Bangalore, Bhubaneshwar, Hissar (shifted to
Panchkula), Imphal, Jabalpur and Nagpur and every laboratory referred
to in clause 29 (1A) shall be designated as referee laboratory for the
purpose of analysis of any sample of Organic fertilizer, provided that no
such laboratory which carried out the first analysis of fertilizer sample
shall be so designated in respect of that sample.

Provided further that in respect of any sample the analysis of which has
been challenged, may be sent for referee analysis to any one of the
other laboratories except those which are located in the state or where
the first analysis has been done.

Provided that National Centre of Organic Farming, Ghaziabad and
Regional Centre of Organic Farming at Bangalore, Bhubaneshwar,
Hissar (shifted to Panchkula), Imphal, Jabalpur and Nagpur shall be
considered as one group of laboratories and a sample first analysed by
any one of them, shall not be sent for referee analysis to any other in
that group, but only to any other laboratory notified by a State Government or Central Government.

2. Notwithstanding anything contained in this order, the Appellate Authority as specified in sub-clause 1 of clause 32A in case of sample collected by the state Government laboratory, or the Controller, in case of sample collected by National Centre of Organic Farming, Ghaziabad or Regional Centre of Organic Farming at Bangalore, Bhubaneshwar, Hissar (shifted to Panchkula), Imphal, Jabalpur and Nagpur, as the case may be, shall decide and send, one of the two remaining samples, for reference analysis as provided under sub-clause (1).

Clause 30
Time limit for analysis and communication of results
1. Where sample of a organic fertilizer, biofertilizer and non-edible de-oiled cake fertilizer has been drawn, the same shall be dispatched along with a memorandum in form K-1 to the laboratory for analysis within a period of seven days from the date of its drawl.

2. The laboratory shall analyse the sample and forward the analysis report in case of organic fertilizers, biofertilizers and non-edible de-oiled cake fertilizer in Form L-1, L2 and L3 respectively within 30 days from the date of receipt of the sample in the laboratory to the authority specified in the same memorandum.

3. The authority to whom the analysis report is sent, shall communicate the result of the analysis to the dealer/manufacturer etc from whom the sample was drawn within 15 days from the date of receipt of the analysis report of the laboratory.

Clause 31
Suspension cancellation or debarment
(1) A notified authority, registration authority, or as the case may be, the controller may, after giving the authorized dealer or the holder of certificate of registration or certificate of manufacture or any other certificate granted under this order, an opportunity of being heard, suspend such authorization letter or certificate or debar the dealer from carrying on the business of fertilizer/biofertilizer or organic fertilizer on one or more of the following grounds, namely:-

(a) That the authorization letter or certificate of registration or certificate of manufacture, as the case may be has been obtained by willful suppression of material facts or by misrepresentation of relevant particulars:
(b) That any of the provisions of this order or any terms and conditions of the memorandum of intimation or certificate of registration or
certificate of manufacture, as the case may be, has been contravened or not fulfilled:

Provided that while debarring from carrying on the business of fertilizer/biofertilizer/organic fertilizer or canceling the certificate, the dealer or the certificate holder thereof may be allowed for a period of 30 days to dispose of the balance stock, if any, held by him.

Provided further that the stock lying with the dealer after the expiry of the said period of 30 days shall be confiscated.

(2) Where the contravention alleged to have been committed by a person is such as would, on being proved, justify his debarment from carrying on the business of fertilizer/biofertilizer/organic fertilizer or, cancellation of authorization letter or certificate of manufacture or any other certificate granted under this order to such person the notified authority or registering authority or, as the case may be, the controller may, without any notice, suspend such certificate, authorization letter, as an interim measure.

Provided that the registering authority, notified authority, or as the case may be, the controller shall immediately furnish to the affected person details and the nature of contravention alleged to have been committed by such person and, after giving him an opportunity of being heard, pass final orders either revoking the order of suspension or debarment within 15 days from the date of issue of the order of suspension.

Provided further that where in final order is passed within the period as specified above, the order of interim suspension shall be deemed to have been revoked without prejudice, however, to any further action, which the registering authority, notified authority or as the case may be, the controller may take against the affected person under sub-clause (1).

(3) Wherever an authorization or certificate is suspended, cancelled or the person is debarred from carrying on the business under these orders, the notified authority or registering authority or, as the case may be, the controller shall record a brief statement of the reasons for such suspension, or as the case may be, cancellation or debarment and furnish a copy thereof to the person whose certificate or authorization letter has been suspended or cancelled or business has been debarred.

(4) When the person alleged to have committed the contravention is an industrial dealer, the notified authority may take action against the holder of such certificate of registration under sub-clause (1) and sub-clause (2):
Provided that where such certificate is suspended or cancelled, the notified authority shall, within a period of 15 days from the date of issue of such order of suspension or cancellation, furnish to the controller also, besides sending the same to the person whose certificate has been suspended or cancelled, a detailed report about the nature of contravention committed and a brief statement of the reasons for such suspension or, as the case may be, cancellation.

Provided further that the controller, shall, in case of the order for suspension passed by the Notified authority, on receipt of the detailed report and after giving the person an opportunity of being heard, pass final order either revoking the order of suspension or canceling the certificate of registration, within 15 days from the date of receipt of the detailed report from the Notified Authority, failing which the order of interim suspension passed by the notified authority shall be deemed to have been revoked, without prejudice however, to further action which the controller may take against the holder of certificate under sub-clause (1):

Provided also that the order of cancellation passed by the notified authority shall remain effective as if it had been passed by the controller till such time the controller, on receipt of the detailed report from the notified authority, and if deemed necessary, after giving the person a fresh opportunity of being heard, pass the final order either revoking or confirming the order of cancellation.

Clause 32
Appeals at Central Government Level
(1) In any State, where the fertilizer allocation is made by the Central Government under this order and if the suspension or cancellation of authorization letter of the manufacture and or pool handling agency or debarment of business, in any way has an effect of dislocating the said allocation and if the Central Government is of the opinion that it is necessary or expedient so to do for maintaining the supplies, may direct the concerned State Government to furnish detailed report about the nature of contravention and a brief statement of the reasons for such suspension or cancellation and pass such order as it may think fit, confirming, modifying or annulling the order of State Government:

Provided that if the report called by the Central Government is not received from the State Government within a period of 15 days from the date of issue of the communication, the Central Government may decide the case without the report on merit.
(Clause 32(1) is not applicable on Biofertilizers and Organic fertilizers as there is no provision of their allocation by the Central Government).

(2) Any person aggrieved by the analysis report of the Notified Testing Laboratory or its Regional Laboratories may appeal to the controller from referee analysis of such sample within a period of 30 days from the receipt of analysis report.

Clause 32(A)
Appeal at the State Government Level

(1) The State Government shall, by notification in the Official Gazette, specify such authority as the Appellate Authority before whom the appeals may be filed by any person, except by an Industrial dealer, aggrieved by any of the following orders or action of registering authority or a notified authority, namely:-

(i) refusing to grant a certificate of manufacture or
(ii) suspending or canceling a certificate of manufacture or
(iii) suspending or canceling authorization letter or debarring from carrying on the business of selling or
(iv) non-issuance of Authorization letter or certificate of manufacture within the stipulated period or
(v) Non-issuance of amendment in authorization letter within the stipulated period.

(2) Any person aggrieved by analysis report of Testing Laboratory notified by State Governments, may appeal to the appellate Authority appointed under sub-clause (1) for reference analysis of such sample within thirty days from the date of receipt of analysis report.

Clause 33
Grant of duplicate copies of authorization letter or certificate of manufacture, certificate of registrations, etc.

Where an authorization letter or a certificate of registration or a certificate of manufacture or any other certificate granted or as the case may be, renewed under this Order is lost or defaced, the Notified Authority, the registering authority, as the case may be, the controller may, on an application made in this behalf, together with the fee prescribed for this purpose under Clause 36, grant a duplicate copy of such certificate.

Clause 34
Amendment of authorization letter, certificate of manufacture and certificate of registration
The Notified Authority, registering authority or controller as the case may be, on application being made by the holder of an authorization letter, a certificate of registration or certificate of manufacture, together with the fee prescribed for this purpose under clause 36, amend an entry in such authorization letter or a certificate of registration or a certificate of manufacture or as the case may be.

Clause 35
Maintenance or records and submission of returns, etc
(1) The controller may, by an order made in writing direct the dealers, manufactures, importers, and pool handling agencies:-

(a) To maintain such book of accounts, records etc relating to their business in Form N and
(b) To submit to such authority, returns and statements in such form and containing such information relating to their business and with in such time as may be specified in that order.

(2) Where a person holds certificate of registration for retail sale and wholesale sale, he shall maintain separate books of accounts for these two types of sales made by him.

(3) Where a State Government, a manufacturer, an importer and a pool handling agency holds valid certificates of registration for sale of fertilizers in, wholesale or retail or both and also for sale for industrial use, he shall maintain separate books of accounts for these two or three types of sales made by him.

(4) Every importer shall inform the Director of Agriculture of the State in which he intends to discharge the imported fertilizer, under intimation to the Central Government, before the import is made or within a period of 15 days after an indent for import is placed, the following details, namely

(i) name of fertilizer
(ii) name of country of import
(iii) name of manufacturer
(iv) quantity to be imported
(v) date of arrival of the consignment
(vi) name of the discharge port
(vii) other information

Clause 36
Fees
(1) The fees payable for grant, amendment or renewal of an authorization letter or certificate of registration or certificate of manufacture, a duplicate of such certificates or, renewal thereof under this order shall be such as the State Governments may, from time to time fix, subject to the maximum fees fixed for the purpose by the Central Government and different fees fixed for different purposes or for different classes of dealers for different types of fertilizers.

(2) The authority to whom and the manner in which the fee fixed under sub-clause (1) shall be paid, shall be such as may be specified by the State Government by notification in the Official Gazette.

(3) Any fee paid under sub-clause (1) shall not be refundable unless the grant or renewal of any certificate registration or certificate of manufacture or duplicate copy of such certificate or renewal under this order has been refused.

(4) The fee payable for grant, amendment, renewal or duplicate copy of certificate of registration for industrial dealer and the authority to whom and the manner in which such fee shall be paid, shall be such as may be specified by the Controller from time to time by notification in the Official Gazette.

Clause 37
Service of Orders and Directions
Any order or direction made or issued by the Controller or by any other authority under this Order shall be served in the same manner as provided in sub-section (5) of section (3) of the Act.

Clause 38
Advisory Committee
(1) The Central Government may by notification in the Official Gazette and on such terms and conditions as may be specified in such notification, constitute a committee called the “Central Fertilizer Committee” consisting of a Chairman and not more than 10 other persons having experience or knowledge in the field, who shall be member of this committee, to advise the Central Government regarding:
   (i) inclusion of a new fertilizer under this Order;
   (ii) Specifications of various fertilizers;
   (iii) Grades/formulations of physical/granulated mixture of fertilizers that can be allowed to be prepared in a State;
   (iv) Requirement of laboratory facilities in a manufacturing unit, including a unit manufacturing physical/ granulated mixtures of fertilizers;
(v) Methods of drawl and analysis of samples;
(vi) Any other matter referred by the Central Government to the committee.

(2) The Committee may, subject to the previous approval of the Central Government, make bye-laws fixing the quorum and regulating its own procedures and the conduct of all business to be transacted by it.

(3) The Committee may co-opt such number of experts and for such purposes or periods as it may deem fit, but any expert so co-opted shall not have the right to vote.

(4) The committee may appoint one or more sub-committees, consisting wholly of members of the committee or partly of the members of the committee and partly co-opted members as it thinks fit, for the purpose of discharging such of its functions as may be delegated to such sub-committee or sub-committees by the Central Fertilizer Committee.

(5) The State Government may by notification in the Official Gazette and on such terms and conditions as may be specified in such notification constitute a committee called the State Fertilizer Committee consisting of a Chairman and not more than 4 members, having experience of knowledge in the field, including a representative from State Agricultural University, the Fertilizer Industry and Indian Micro Fertilizers Association to advise the State Government regarding the grades/formulations of, mixtures of fertilizers.

Clause 39
Repeal and saving
(1) The Fertilizer (Control) Order, 1957 is hereby repealed except as respects things done or omitted to be done under the said order before the commencement of this order.

(2) Notwithstanding such repeal, the order made by any authority, which is in force, immediately before the commencement of this order and which is consistent with this order, shall continue in force and all appointments made, prices fixed, certificates granted and directions issued under repealed order and in force immediately before such commencement shall likewise continue in force and be deemed to be made, fixed, granted or issued in pursuance of this order till revoked.
**Schedule III**

[See clause 2(h) and (q)]

**PART – A**

Specification of Biofertilizers

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### 1. Rhizobium

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<table>
<thead>
<tr>
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<tbody>
<tr>
<td>(i)</td>
<td>Base</td>
<td>Carrier based* in form of moist/dry powder or granules, or liquid based</td>
</tr>
<tr>
<td>(ii)</td>
<td>Viable cell count</td>
<td>CFU minimum $5 \times 10^7$ cell/g of powder, granules or carrier material or $1 \times 10^8$ cell/ml of liquid.</td>
</tr>
<tr>
<td>(iii)</td>
<td>Contamination level</td>
<td>No contamination at $10^2$ dilution</td>
</tr>
<tr>
<td>(iv)</td>
<td>pH</td>
<td>6.5-7.5</td>
</tr>
<tr>
<td>(v)</td>
<td>Particles size in case of carrier based material</td>
<td>All material shall pass through 0.15-0.212mm IS sieve</td>
</tr>
<tr>
<td>(vi)</td>
<td>Moisture percent by weight, maximum in case of carrier based</td>
<td>30-40%</td>
</tr>
<tr>
<td>(vii)</td>
<td>Efficiency character</td>
<td>Should show effective nodulation on all the species listed on the packet.</td>
</tr>
</tbody>
</table>

*Type of carrier:* The carrier materials such as peat, lignite, peat soil, humus, wood charcoal or similar material favouring growth of organism.

---

### 2. Azotobacter

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<table>
<thead>
<tr>
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</thead>
<tbody>
<tr>
<td>(i)</td>
<td>Base</td>
<td>Carrier based* in form of moist/dry powder or granules, or liquid based</td>
</tr>
<tr>
<td>(ii)</td>
<td>Viable cell count</td>
<td>CFU minimum $5 \times 10^7$ cell/g of powder, granules or carrier material or $1 \times 10^8$ cell/ml of liquid.</td>
</tr>
<tr>
<td>(iii)</td>
<td>Contamination level</td>
<td>No contamination at $10^2$ dilution</td>
</tr>
<tr>
<td>(iv)</td>
<td>pH</td>
<td>6.5-7.5</td>
</tr>
<tr>
<td>(v)</td>
<td>Particles size in case of carrier based material</td>
<td>All material shall pass through 0.15-0.212mm IS sieve</td>
</tr>
<tr>
<td>(vi)</td>
<td>Moisture percent by weight, maximum in case of carrier based</td>
<td>30-40%</td>
</tr>
<tr>
<td>(vii)</td>
<td>Efficiency character</td>
<td>The strain should be capable of fixing at least 10 mg of nitrogen per g of sucrose consumed.</td>
</tr>
</tbody>
</table>

*Type of carrier:* The carrier material such as peat, lignite, peat soil, humus, wood charcoal or similar material favouring growth of the organism.

---

### 3. Azospirillum

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<tbody>
<tr>
<td>(i)</td>
<td>Base</td>
<td>Carrier based* in form of moist/dry powder or granules, or liquid based</td>
</tr>
<tr>
<td>(ii)</td>
<td>Viable cell count</td>
<td>CFU minimum $5 \times 10^7$ cell/g of powder, granules or carrier material or $1 \times 10^8$ cell/ml of liquid.</td>
</tr>
<tr>
<td>(iii)</td>
<td>Contamination level</td>
<td>No contamination at $10^2$ dilution</td>
</tr>
<tr>
<td>(iv)</td>
<td>pH</td>
<td>6.5-7.5</td>
</tr>
<tr>
<td>(v)</td>
<td>Particles size in case of carrier based material</td>
<td>All material shall pass through 0.15-0.212mm IS sieve</td>
</tr>
<tr>
<td>(vi)</td>
<td>Moisture percent by weight, maximum in case of carrier based</td>
<td>30-40%</td>
</tr>
<tr>
<td>(vii)</td>
<td>Efficiency character</td>
<td>Formation of white pellicle in semisolid N-free bromothymol blue media.</td>
</tr>
</tbody>
</table>

---
**Type of carrier:-** The carrier material such as peat, lignite, peat soil, humus, wood Charcoal or similar material favouring growth of the organism.

### 4. Phosphate solubilising Bacteria

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<table>
<thead>
<tr>
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</thead>
<tbody>
<tr>
<td>(i)</td>
<td>Base</td>
<td>Carrier based* in form of moist/dry powder or granules, or liquid based</td>
</tr>
<tr>
<td>(ii)</td>
<td>Viable cell count</td>
<td>CFU minimum $5 \times 10^7$ cell/g of powder, granules or carrier material or $1 \times 10^8$ cell/ml of liquid.</td>
</tr>
<tr>
<td>(iii)</td>
<td>Contamination level</td>
<td>No contamination at $10^5$ dilution</td>
</tr>
<tr>
<td>(iv)</td>
<td>pH</td>
<td>6.5-7.5 for moist/dry powder, granulated carrier based and 5.0 – 7.5 for liquid based</td>
</tr>
<tr>
<td>(v)</td>
<td>Particles size in case of carrier based material.</td>
<td>All material shall pass through 0.15-0.212mm IS sieve</td>
</tr>
<tr>
<td>(vi)</td>
<td>Moisture percent by weight, maximum in case of carrier based.</td>
<td>30-40%</td>
</tr>
<tr>
<td>(vii)</td>
<td>Efficiency character</td>
<td>The strain should have phosphate solubilizing capacity in the range of minimum 30%, when tested spectrophotometrically. In terms of zone formation, minimum 5mm solubilization zone in prescribed media having at least 3mm thickness.</td>
</tr>
</tbody>
</table>

*Types of Carrier:-* The carrier material such as peat, lignite, peat soil, humus, wood Charcoal or similar material favouring growth of the organism.

### 5. Mycorrhizal Biofertilizers

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<table>
<thead>
<tr>
<th></th>
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</thead>
<tbody>
<tr>
<td>i.</td>
<td>Form/base</td>
<td>Fine Powder/ tablets/ granules/ root biomass mixed with growing substrate</td>
</tr>
<tr>
<td>ii.</td>
<td>Particle size for carrier based powder formulations</td>
<td>90% should pass through 250 micron IS sieve (60 BSS)</td>
</tr>
<tr>
<td>iii.</td>
<td>Moisture content percent maximum</td>
<td>8 -12</td>
</tr>
<tr>
<td>iv.</td>
<td>pH</td>
<td>6.0 to 7.5</td>
</tr>
<tr>
<td>v.</td>
<td>Total viable propagules/ gm of product, minimum</td>
<td>100 /gm of finished product</td>
</tr>
<tr>
<td>V.</td>
<td>Infectivity potential</td>
<td>80 infection points in test roots/gm of mycorrhizal inoculum used</td>
</tr>
</tbody>
</table>

### 6. Potassium Mobilizing Biofertilizers (KMB)

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<table>
<thead>
<tr>
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</thead>
<tbody>
<tr>
<td>1.</td>
<td>Base</td>
<td>Carrier based* in form of moist/dry powder or granules, or liquid based</td>
</tr>
<tr>
<td>2.</td>
<td>Viable cell count</td>
<td>CFU minimum $5 \times 10^7$ cells/g of powder, granules, or carrier material on dry weight basis or $1 \times 10^8$ cell/ml of liquid</td>
</tr>
<tr>
<td>3.</td>
<td>Contamination</td>
<td>No contamination at $10^5$ dilution</td>
</tr>
<tr>
<td>4.</td>
<td>pH</td>
<td>6.5-7.5 for carrier based in form of powder or granules and 5.0-7.5 for liquid based</td>
</tr>
<tr>
<td>5.</td>
<td>Particle size in case of carrier based moist powder</td>
<td>Powder material shall pass through 0.15 to 0.212 mm IS sieve</td>
</tr>
<tr>
<td>6.</td>
<td>Moisture per cent, by weight, maximum in case of powder based</td>
<td>30-40</td>
</tr>
<tr>
<td>7.</td>
<td>Efficiency character</td>
<td>Maximum 10 mm solubilization zone in prescribed media having at least 3mm thickness.</td>
</tr>
</tbody>
</table>
*Type of carrier – The carrier material such as peat, lignite, peat soil, humus, talc or similar material favouring growth of microorganisms.

### 7. Zinc Solubilizing Biofertilizers (ZSB)

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<tbody>
<tr>
<td>1.</td>
<td>Base</td>
</tr>
<tr>
<td></td>
<td>Carrier based in form of moist/dry powder or granules, or liquid based</td>
</tr>
<tr>
<td>2.</td>
<td>Viable cell count</td>
</tr>
<tr>
<td></td>
<td>CFU minimum 5x10^7 cells/g of powder, granules, or carrier material on dry weight basis or 1x10^8 cell/ml of liquid</td>
</tr>
<tr>
<td>3.</td>
<td>Contamination</td>
</tr>
<tr>
<td></td>
<td>No contamination at 10^5 dilution</td>
</tr>
<tr>
<td>4.</td>
<td>pH</td>
</tr>
<tr>
<td></td>
<td>6.5-7.5 for carrier based in form of powder or granules and 5.0-7.5 for liquid based</td>
</tr>
<tr>
<td>5.</td>
<td>Particle size in case of carrier based moist powder</td>
</tr>
<tr>
<td></td>
<td>Powder material shall pass through 0.15 to 0.212 mm IS sieve</td>
</tr>
<tr>
<td>6.</td>
<td>Moisture per cent, by weight, maximum in case of powder based</td>
</tr>
<tr>
<td></td>
<td>30-40</td>
</tr>
<tr>
<td>7.</td>
<td>Efficiency character</td>
</tr>
<tr>
<td></td>
<td>Maximum 10 mm solubilization zone in prescribed media having at least 3mm thickness.</td>
</tr>
</tbody>
</table>

### 8. Acetobacter

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<tbody>
<tr>
<td>1.</td>
<td>Base</td>
</tr>
<tr>
<td></td>
<td>Carrier based in form of moist/dry powder or granules, or liquid based</td>
</tr>
<tr>
<td>2.</td>
<td>Viable cell count</td>
</tr>
<tr>
<td></td>
<td>CFU minimum 5x10^7 cells/g of powder, granules, or carrier material or 1x10^8 cells/ml of liquid</td>
</tr>
<tr>
<td>3.</td>
<td>Contamination level</td>
</tr>
<tr>
<td></td>
<td>No contamination at 10^5 dilution</td>
</tr>
<tr>
<td>4.</td>
<td>pH</td>
</tr>
<tr>
<td></td>
<td>5.5-6.0 moist/dry powder, granulated or carrier based and 3.5-6.0 for liquid</td>
</tr>
<tr>
<td>5.</td>
<td>Particle size in case of carrier based material</td>
</tr>
<tr>
<td></td>
<td>All material shall pass through 0.15 to 0.212 mm IS sieve</td>
</tr>
<tr>
<td>6.</td>
<td>Moisture per cent, by weight, maximum in case of carrier based</td>
</tr>
<tr>
<td></td>
<td>30-40%</td>
</tr>
<tr>
<td>7.</td>
<td>Efficiency character</td>
</tr>
<tr>
<td></td>
<td>Formulation of yellowish pellicle in semisolid medium N free medium</td>
</tr>
</tbody>
</table>

*Type of carrier – The carrier material such as peat, lignite, peat soil, humus, wood charcoal or similar materials favouring growth of organism.

### 9. Carrier Based Consortia

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Base</td>
</tr>
<tr>
<td></td>
<td>Carrier based in form of moist powder or granules</td>
</tr>
<tr>
<td>2.</td>
<td>Viable cell count</td>
</tr>
<tr>
<td></td>
<td>CFU minimum in a mixture of any 2 or maximum three of following microorganisms: CFU minimum Rhizobium or Azotobacter or Azospirillum 1x10^7 per g CFU minimum PSB 1x10^7 per g CFU minimum KSB 1x10^7 per g</td>
</tr>
<tr>
<td>3.</td>
<td>Particle size in case of carrier based moist powder</td>
</tr>
<tr>
<td></td>
<td>All material shall pass through 0.15 to 0.212 mm IS sieve</td>
</tr>
<tr>
<td>4.</td>
<td>Total viable count of all the biofertiliser organisms in the product</td>
</tr>
<tr>
<td></td>
<td>CFU minimum – 5x10^7 cells per gm of carrier/powder</td>
</tr>
<tr>
<td>5.</td>
<td>Moisture per cent, by weight, maximum in case of carrier based</td>
</tr>
<tr>
<td></td>
<td>30-40%</td>
</tr>
</tbody>
</table>
6. **Contamination**
   - No contamination at $10^{-4}$ dilution for carrier based / granule based inoculants

7. **Efficiency character**
   - **Azotobacter**
     - The strain should be capable of fixing at least 10 mg of Nitrogen fixation/g of C-source
   - **Azospirillum**
     - The strain should be capable of fixing at least 10 mg of N-fixation/g of malate applied
   - **PSB**
     - Minimum 5mm zone of solubilization zone on PSB media having at least 3mm thickness
   - **KMB**
     - Minimum 5mm zone of solubilization on KSB media having at least 3mm thickness
   - **Rhizobium**
     - Nodulation test positive

10. **Liquid Consortia**

   1. **Individual Viable count in Liquid based**
      - CFU minimum in a mixture of any 2 or more of following microorganisms:
        - CFU minimum Rhizobium or Azotobacter or Azospirillum $1 \times 10^8$ per ml
        - CFU minimum PSB $1 \times 10^8$ per ml
        - CFU minimum KSB $1 \times 10^8$ per ml
   
   2. **Total viable count of all the biofertiliser organisms in the product**
      - CFU minimum $5 \times 10^5$ cells per ml of liquid based

   3. **Contamination**
      - No contamination at any dilution

4. **pH**
   - 5.0–7.0

5. **Efficiency character**
   - **Azotobacter**
     - The strain should be capable of fixing at least 10 mg N-fixation/g of C-source
   - **Azospirillum**
     - The strain should be capable of fixing at least 10 mg of N-fixation/g of malate applied
   - **PSB**
     - Minimum 5mm zone of solubilization zone on PSB media having at least 3mm thickness
   - **KMB**
     - Minimum 5mm zone of solubilization on KSB media having at least 3mm thickness
   - **Rhizobium**
     - Nodulation test positive

**Part-B**

**Tolerance limit of Biofertilizers**

1. In case of Rhizobium, Azotobacter, Azospirillum and Phosphate solubilizing bacteria, the total viable counts shall not be less than $1 \times 10^7$ CFU/gm of carrier material in the form of powder or granules or $5 \times 10^7$ CFU/ml in case of liquid formulations during the entire period of shelf life.
2. In case of Mycorrhizal Biofertilizers, the viable propagules shall not be less than 80.

PART C
Procedure for drawl of samples of Biofertilizers
Procedure for sampling of biofertilizers

1. General Requirements of Sampling

1.0 In drawing, preparing and handling the samples, the following precautions and directions shall be observed.

1.1 Sampling shall be carried out by a trained and experienced person as it is essential that the sample should be representative of the lot to be examined.

1.2 Samples in their original unopened packets should be drawn and sent to the laboratory to prevent possible contamination of sample during handling and to help in revealing the true condition of the material.

1.3 Intact packets shall be drawn from a protected place not exposed to dampness, air, light, dust or soot.”

2. Scale of Sampling

2.1 Lot
All units (containers in a single consignment of type of material belonging to the same batch of manufacture) shall constitute a lot. If a consignment consists of different batches of the manufacture the containers of the same batch shall be separated and shall constitute a separate lot.

2.2 Batch
All inoculant prepared from a batch fermentor or a group of flasks (containers) constitute a batch.

2.3 For ascertaining conformity of the material to the requirements of the specification, samples shall be tested from each lot separately.

2.4 The number of packets to be selected from a lot shall depend on the size of the lot and these packets shall be selected at random and in order to ensure the randomness of selection procedure given in IS 4905 may be followed.”

3. Drawal of Samples

3.1 The Inspector shall take three packets as sample from the same batch. Each sample constitutes a test sample.

3.2 These samples should be sealed in cloth bags and be sealed with the Inspector’s seal after putting inside Form P. Identifiable details
such as sample number, code number or any other details which enable its identification shall be marked on the cloth bags.

3.3 Out of the three samples collected, one sample so sealed shall be sent to in-charge of the laboratory notified by the State Government under clause 29 or to National Centre for Organic Farming or to any of its Regional Centres. Another sample shall be given to the manufacturer or importer or dealer as the case may be. The third sample shall be sent by the inspector to his next higher authority for keeping in safe custody. Any of the latter two samples shall be sent for referee analysis under subclause (2) of clause 29B.

3.4 The number of samples to be drawn from the lot

<table>
<thead>
<tr>
<th>Lot/Batch Number of Samples</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Upto 5,000 packets</td>
<td>03</td>
</tr>
<tr>
<td>5,001-10,000 packets</td>
<td>04</td>
</tr>
<tr>
<td>More than 10,000 packets</td>
<td>05</td>
</tr>
</tbody>
</table>
1. A. METHODS OF ANALYSIS OF RHIZOBIUM BIOFERTILISERS

1. Apparatus

1.1 Pipettes Graduated 1 ml and 10 ml.
1.2 Dilution Bottles or flasks
1.3 Petri Dishes Clear, Uniform, flat-bottomed.
1.4 Hot Air Oven - Capable of giving uniform and adequate temperature, equipped with a thermometer calibrated to read up to 250° C and with vents suitably located to assure prompt and uniform heating.
1.5 Autoclave
1.6 Incubator
1.7 Hand Tally or Mechanical counting Device
1.8 pH meter

2. Reagents

2.1 Congo Red-one percent aqueous solution
2.2 Medium

Use a plating medium of the following composition:

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>Agar</td>
<td>20 g</td>
</tr>
<tr>
<td>Yeast Extract</td>
<td>1 g</td>
</tr>
<tr>
<td>Mannitol</td>
<td>10 g</td>
</tr>
<tr>
<td>Potassium hydrogen phosphate (K₂HPO₄)</td>
<td>0.5g</td>
</tr>
<tr>
<td>Magnesium sulphate (MgSO₄. 7H₂O)</td>
<td>0.2g</td>
</tr>
<tr>
<td>Sodium Chloride (NaCl)</td>
<td>0.1g</td>
</tr>
<tr>
<td>Congo red</td>
<td>2.5 ml</td>
</tr>
<tr>
<td>Distilled water</td>
<td>1000 ml</td>
</tr>
<tr>
<td>pH</td>
<td>7.0</td>
</tr>
</tbody>
</table>

2.3 Sterilizing and preparation procedure for plates:

2.3.1. Sterilize the sampling and plating equipment with dry heat in a hot air oven at not less than 160° C for not less than 2 hours.

2.3.2 Sterilize the media by autoclaving at 120° C for 20 min. To permit passage of steam into and from closed container when autoclaved, keep stoppers slightly loosened or plugged with cotton. Air from with in the chamber of the sterilizer should be ejected allowing steam pressure to rise.
Preparation of Plating Medium and Pouring

2.3.3 Prepare growth medium in accordance with the composition of the specific Biofertilizers.

2.3.4 Melt the required amount of medium in boiling water or by exposure to flowing steam in partially closed container but avoid prolonged exposure to unnecessarily high temperature during and after melting. Melt enough medium which will be used within 3h. Re-sterilization of the medium may cause partial precipitation of ingredients.

2.3.5 When holding time is less than 30 min promptly cool the molten medium to about 45°C, and store until used, in a water bath or incubator at 43°C to 45°C. Introduce 12 to 15 ml of liquefied medium or appropriate quantity depending on size of the Petri dish at 42 to 44°C into each plate. Gently lift the cover of the dish just enough to pour in the medium. Sterilize the lips of the medium container by exposure to flame.

a. Immediately before pouring.
b. Periodically during pouring, and
c. When pouring is completed for each batch of plates, if portion of molten medium remain in containers and are to be used without subsequent sterilization for pouring additional plates. As each plate is poured thoroughly mix the medium with test portions in the Petri dish.

2.3.6 By rotating and tilting the dish and without splashing the medium over edge, spread the medium evenly over the bottom of the plate. Provide conditions so that the medium solidifies with reasonable promptness (5-10min) before removing the plates from surface.

3. Preparation of serial dilution for plate count

3.1 Dispense 30 g of Inoculants to 270 ml of sterile distilled/demineralized water and shake for 10 min on a reciprocal shaker or homogenizer. Make serial dilutions up to $10^{-9}$ by suspending 10 ml aliquot of previous dilution to 90ml of water. Take 0.1 ml or suitable alliquotes of $10^{-5}$ to $10^{-9}$ dilutions using sterile pipettes and deliver to Petri dishes containing set medium as given in 2.1 and spread it uniformly with a spreader. Invert the plates and promptly place them in the incubator.
4. Incubation of plates
4.1. Label the plates and incubate at 28±2°C for 3 to 5 days for fast growing Rhizobia and 5 to 10 days for slow-growing ones.

4.2. Colony counting aids
Count the colonies with the aid of magnifying lens under uniform and properly controlled, artificial illumination. Use a colony counter, equipped with a guide plate and rules in Centimeter Square. Record the total number of colonies with the hand tally. Avoid mistaking particles of undissolved medium or precipitated matter, in plates for pinpoint colonies. To distinguish colonies from dirt, specks and other foreign matter, examine doubtful objects carefully.

4.3. Count all plates but consider for the purpose of calculation plates showing more than 30 and less than 300 colonies per plate. Disregard colonies, which absorb Congo red and stand out as reddish colonies. Rhizobium stands out as white, translucent, glistening and elevated colonies. Count such colony numbers and calculate figures in terms of per gram, of carrier. Also check for freedom from contamination at 10⁻⁵ dilution.

5. Test for nodulation
5.1 Pot culture test
Plant nutrient solution

<table>
<thead>
<tr>
<th>Sl. No</th>
<th>Composition</th>
<th>Conc.</th>
<th>g/l</th>
</tr>
</thead>
<tbody>
<tr>
<td>a</td>
<td>Potassium chloride</td>
<td>0.001M</td>
<td>0.0745</td>
</tr>
<tr>
<td>b</td>
<td>Di Potassium hydrogen Phosphate (K₂HPO₄)</td>
<td>0.001M</td>
<td>0.175</td>
</tr>
<tr>
<td>c</td>
<td>Calcium sulphate (CaSO₄ 2H₂O)</td>
<td>0.002M</td>
<td>0.344</td>
</tr>
<tr>
<td>d</td>
<td>Magnesium sulphate (MgSO₄ 7H₂O)</td>
<td>0.001M</td>
<td>0.246</td>
</tr>
<tr>
<td>e</td>
<td>Trace elements solution:</td>
<td>-</td>
<td>0.5ml</td>
</tr>
<tr>
<td>1. Copper sulphate (CuSO₄ 5H₂O)</td>
<td>0.01mg/kg</td>
<td>0.78</td>
<td></td>
</tr>
<tr>
<td>2. Zinc Sulphate (ZnSO₄ 7H₂O)</td>
<td>0.25 mg/kg</td>
<td>2.22</td>
<td></td>
</tr>
<tr>
<td>3. Ammonium molybdate ((NH₄)₆Mo₇O₂₄ 4H₂O)</td>
<td>0.0025 mg/kg</td>
<td>0.01</td>
<td></td>
</tr>
<tr>
<td>4. Magnesium sulphate (MgSO₄ 7H₂O)</td>
<td>0.25 mg/kg</td>
<td>2.03</td>
<td></td>
</tr>
<tr>
<td>5. Boric acid (H₃BO₄)</td>
<td>0.125 mg/kg</td>
<td>1.43</td>
<td></td>
</tr>
<tr>
<td>6. Water</td>
<td></td>
<td>1 l</td>
<td></td>
</tr>
</tbody>
</table>

Prepare the solution no (e) consisting of trace elements in one liter of stock solution and add to final nutrient solution at the rate of 0.5 ml per liter.

<table>
<thead>
<tr>
<th>f</th>
<th>Iron solution:</th>
<th>0.5 ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ferrous sulphate</td>
<td>5</td>
<td></td>
</tr>
<tr>
<td>Citric acid</td>
<td>5</td>
<td></td>
</tr>
<tr>
<td>Water</td>
<td>100 ml</td>
<td></td>
</tr>
</tbody>
</table>

Prepare the solution no. (f) As 100 ml of stock solution and add final nutrient solution at the rate of 0.5 ml per liter.
5.2 Preparation
Prepare the nutrient solution by weighing out substances (a), (b) and (d) and dissolving them in a liter of water. To this solution add 0.5 ml of trace elements solution and 0.5 ml of iron solution. Grind in a mortar 0.344 g of calcium sulphate (c) to a fine consistency and add to the final nutrient solution. Autoclave the nutrient solution thus prepared, at 120°C for 20 min.

Notes
1. The nutrient solution may be prepared in the tap water provided the water is soft.
2. The nutrient solution should be shaken well to disperse calcium sulphate before dispensing.
3. If the solution is made up with distilled water, the pH is about 7.2 before autoclaving and falls to 5.5 on autoclaving and rises slowly on standing to about 5.8. However, there is no need to adjust pH. For most tropical legumes; pH of about 6.0 is adequate.

5.3 Procedure
5.3.1 Immerse the seeds in 95 percent alcohol and follow by surface sterilization in freshly prepared chlorine water (for 15 to 20 min) or 0.1 percent mercuric chloride solution 3 min in a suitable container such as a screw-capped bottle or a test tube with a rubber hung. In case of seeds with tough seed coat, concentrated sulphuric acid may be used as a surface sterilants for 20 to 30 min. It is recommended that the seeds should be placed overnight in a desiccators containing calcium chloride before surface sterilization with sulphuric acid. Pour out the Sterilants and wash the seeds in several changes of sterile water and wash the seeds in several changes of sterile water (at least ten times) to get rid of the Sterilants. Fill earthenware or glazed pot with soil (2 parts soil and 1 part washed coarse sand) (pH 6 to 7) and autoclave for 2 h at 120°C. After two days incubation at room temperature, repeat autoclaving to ensure complete sterility of soil. Inoculate surface sterilized seeds with water slurry of the inoculant taken from a culture packet (15 to 100 g seeds per gram of inoculants depending on the size of the seed) and sow the seeds. Keep a set of pots with Uninoculated seeds as control and also a set of pots with ammonium nitrate at the rate of 100 kg N/ha as control aid incubate them in a pot-culture house during appropriate seasons for appropriate plants, taking care to separate the inoculated pots from the control pots. If growth rooms or cabinets having facilities to adjust temperature and light are
available, the pots may be incubated in such controlled environmental conditions. Sterilize the nutrient solution at 120°C for 20 min and irrigate each pot once to the moisture holding capacity of soil. Subsequently, water the seedling periodically with sterilized water preferably through a plastic tube, taking care to prevent splashing of water from inoculated pots to uninoculated ones. Maintain required number of replicated pots (4 to 6) for each botanical species for statistical analysis.

5.3.2 After two to three weeks of growth, thin down the number of plants in each pot to four uniform plants. At the end of 6 to 8 weeks, take one set of pots from both the control and inoculated series and, separate the plants carefully from the soil under slow running water. Obtain data on the number, colour (effective nodules are pink or red) and mass of nodules. At the end of 6 to 8 weeks, harvest the shoot system, dry at 60°C for 48 h and determine dry mass. For the above purpose, maintain adequate replications of pots (4 to 16).

5.3.3 Record the nodulation data regarding formation of pink colour of nodules as revealed visually when nodules are cut open by razor blade. After computing the data, based on the dry mass of plants and nodulation data decide the effectiveness of culture. If good effective pink nodulation is obtained in inoculated plants together with local absence or sometimes presence of stray nodules in controls and if there is a 50 percent increase in the dry mass of plants over the Uninoculated control without nitrate, it may be concluded that the culture is of the require quality.

1.B. METHOD OF ANALYSIS OF AZOTOBACTER BIO-FERTILISER

1. Apparatus - same as Rhizobium
2. Reagents:
   2.1. Medium
   Use a plating medium of the following composition
   
<table>
<thead>
<tr>
<th>Ingredient</th>
<th>Quantity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Agar</td>
<td>20g</td>
</tr>
<tr>
<td>Sucrose (C_{12} H_{26}O_{11})</td>
<td>20.0 g</td>
</tr>
<tr>
<td>Ferric sulphate Fe_{2} (SO_{4})_{3}</td>
<td>0.1 g</td>
</tr>
<tr>
<td>Dibasic potassium phosphate (K_{2}HPO_{4})</td>
<td>1.0 g</td>
</tr>
<tr>
<td>Magnesium sulphate (MgSO_{4}, 7H_{2}O)</td>
<td>0.5 g</td>
</tr>
<tr>
<td>Sodium Chloride (NaCl)</td>
<td>0.5 g</td>
</tr>
<tr>
<td>Calcium carbonate (CaCO_{3})</td>
<td>2.0g</td>
</tr>
<tr>
<td>Sodium Molybdate (Na_{2}MoO_{4})</td>
<td>0.005gms</td>
</tr>
<tr>
<td>Distilled water</td>
<td>1000ml</td>
</tr>
<tr>
<td>pH</td>
<td>6.8 to 7.2</td>
</tr>
</tbody>
</table>
2.2. Sterilization and preparation procedure for plates: Same as Rhizobium

3. Preparation of plating medium and pouring
   Preparation of serial dilution for plate counts
   3.1. Dispense 30 g of Inoculants to 270 ml of sterile distilled/demineralized water and shake for 10 min on a reciprocal shaker or homogenizer. Make serial dilutions up to $10^{-9}$ by suspending 10 ml aliquot of previous dilution to 90 ml of water. Take 0.1 ml or suitable aliquotes of $10^{-5}$ to $10^{-9}$ dilutions using sterile pipettes and deliver to Petri dishes containing set medium as given in 2.1 and spread it uniformly with a spreader. Invert the plates and promptly place them in the incubator.

4. Incubation of plates: Same as Rhizobium.
   4.1. Label the plates and incubate at 28 +/- 3° C for 4 to 6 days.
   4.2. Colony counting aids: Same as Rhizobium.

Azotobacter chroococcum colonies are gummy, raised with or without striations, viscous and often sticky. The pigmentation varies from very light brown to black. Count the colony number and observe the cyst formation as given below and calculate number per gram of the carrier material.

Grow the vegetative cells at 30° C on Burks agar medium comprising sucrose 20 g, dipotassium hydrogen phosphate 0.64 g, dihydrogen potassium phosphate 0.20 g, sodium chloride 0.20 g, calcium sulphate 0.05 g, sodium molybdate 0.001 g, ferric sulphate 0.003 g, agar 20 g and distilled water 1000 ml. Look for vegetative cells after 18 to 24 h either by simple staining method or through a phase contrast microscope.

Grow the cyst cells on Burks agar medium as given above with 0.3 percent n-butanol in place of the carbon source. Look for cyst formation after 4 to 5 days incubation.

5. Test for Nitrogen fixation in pure cultures
   5.1. Pure culture medium
   5.1.1. Prepare medium as given for Azotobacter (2.1 under 1B), excluding agar.
   5.2. Procedure
      Select from each Azotobacter colony, of the type that has been counted as Azotobacter chroococcum. Pick up one colony and plate on the medium given in. Use this pure culture for inoculating the broth for nitrogen fixation. For this purpose, take 50 ml aliquots of broth in 250 ml conical flasks for inoculation. After 12 days growth at 28° C, test the contents of the flasks for purity by streaking on fresh medium and concentrating over water bath (50 to 60° C) to dryness. Wash the dried culture and
take it as a sample. The contents of the flasks in inoculated control series should be similar manner.

5.3. Determination by Kjeldahl Method

(i) Reagents

(ii) Sulphuric acid-93-98 percent, N-free

(iii) Digestion mixture- Mix copper sulphate and potassium sulphate in the ratio 1: 10 and grind them to a fine powder.

(iv) Sodium hydroxide pellets or solution, N-free- For solution, dissolve about 450 g of sodium hydroxide in water, cool, and dilute 1 liter (sp gr of the solution should be at least 1.36)

(v) Zinc granules-reagent grade.

(vi) Indicators: -

(a) Methyl red indicator - Dissolve 1g of methyl red in 200ml of Ethanol.

(b) Mixed indicator- Prepare mixed indicator by Dissolving 0.8 of methyl red and 0.2 g of methyl blue in 500 ml of ethanol.

(vi) Hydrochloric or sulphuric acid – Standard solution 0.5 or 0.1 N when amount of nitrogen is small.

(vii) Sodium hydroxide standard solution – 0.1 N (or other specified concentration)

Note : Ratio of salt to acid (m/v) should be about 1:1 at the end of the digestion for proper temperature control. Digestion may be incomplete at a lower ratio, and nitrogen may be lost at higher ratio. Each gram of fat consumes 10 ml of sulphuric acid and each gram of carbohydrate 4.0 ml of sulphuric acid during digestion.

5.4. Apparatus

(i) For digestion - Use kjeldahl’s flasks of hard, moderately thick, well annealed glass with total capacity approximately 500 to 800 ml. Conduct digestion over heating device adjust to bring 250 ml of water at 25° C to rolling boil in about 5 minutes. To test the heaters, preheat for 10 minutes in the case of gas burners and for 30 minutes in the case of electric heaters. Add 3 to 4 boiling chips to prevent superheating.
(ii) For distillation - Use 500 to 800ml kjeldahl's flask fitted with rubber stopper through which passes the lower end of an efficient scrubber bulb or trap to prevent mechanical carry-over of sodium hydroxide during distillation. Connect the upper end of the bulb tube to a condenser by rubber tubing. Trap the outlet of the condenser in such a way as to ensure absorption of ammonia distilled over with the receiver.

5.5. Procedure: -

(a) Place 0.25 g of the sample in the digestion flask. Add 0.7 gm mercuric oxide, 15 gm potassium sulphate followed by 25 ml of sulphuric acid. Shake, let stand for about 30 minutes and heat carefully until frothing ceases. Boil briskly until the solution clears and continue boiling further for 90 minutes. Cool, add about 200 ml of water cool to room temperature and add a few zinc granules.

(b) Tilt the flask and carefully add 50 ml of sodium hydroxide solution without agitation. Immediately connect the flask to the distillation bulb on the condenser whose tip is immersed in 50 ml of standard 0.1 N acid in the receiving flasks. Rotate the digestion flask carefully to mix the content. Heat until 150 ml of the distillate collects and titrate excess acid with 0.1 N base using methyl-red or mixed indicator. Carry out blank determination on reagents. Note: Check the ammonia recording periodically, using inorganic nitrogen control, for example, ammonium sulphate.

(c) Calculation: -

(i) Nitrogen content, percent by mass =

\[
\frac{\text{Milliliters of 0.1 N acid for sample} - \text{milliliters of 0.1 N acids for blank}}{\text{Mass of sample taken}} \times 0.14
\]

(ii) Total nitrogen in culture = Total dry mass of sample \times \% nitrogen.

(d) Take a 1.0 g of accurately weighed sample each from the inoculated series and from the controls. Put them separately in 250 ml volumetric flask, add 150 ml water, mix the content and make up the volume to 250 ml water. Shake for 5 minutes and centrifuge for 15 minutes at 10000 rpm. Estimate glucose in the supernatant in triplicate. The difference between the two provides the data of actual amount of glucose consumed. Calculate the amount of nitrogen fixed per gram of sucrose consumed.
Determination of Glucose: - From the supernatant, draw suitable aliquots and estimate reducing sugars (glucose) as follows:

Reagents.

(ii) Soxhelt modification of Fehling solution: - Prepare by mixing equal volumes of solution A and solution B immediately before using.

(iii) Copper sulphate solution (Solution A)- Dissolve 34.639 g of copper sulphate crystals (CuSO4 5H2O) in water, dilute to 500ml and filter through glass wool or filter paper.

**Standardization of copper sulphate solution:** - Using separate pipettes, pipette accurately 5 ml of solution A and 5 ml of solution B into a conical flask of 250 ml capacity. Heat this mixture to boiling on asbestos gauze and add standard invert sugar solution from a burette, about 1 ml less than the expected volume, which will reduce the Fehling solution completely (about 48 ml). Add 1 ml of methylene blue indicator while keeping the solution boiling. Complete the titration within 3 min, the end point being indicated by change of colour from blue to red. From the volume of invert sugar solution used, calculate the strength(s) of the copper sulphate solution by multiplying the titre value by 0.001 (mg/ml of the standard invert sugar solution). This would give the quantity of invert sugar required to reduce the copper in 5 ml of copper sulphate solution.

(iv) Potassium sodium tartrate (Rochelle salt) solution (solution B): - Dissolve 173 g of potassium sodium tartrate and 50 g of sodium hydroxide in water, and dilute to 500 ml. Let the solution stand for a day, and filter.

(v) Hydrochloric acid – sp gr 1.18 at 20° C (approximately 12 N)

(vi) Standard invert sugar solution –Weigh accurately 0.95 g of sucrose and dissolve it in 500 ml of water. Add 32 ml of concentrated hydrochloric acid, boil gently for 30 min and keep aside for 24 hours. Neutralize with sodium carbonate and make the final volume to 1000ml; 50 ml of this solution contains 0.05 g of invert sugar.

(vii) Methylene blue indicator- 0.2 percent in water.
(viii) Procedure: - Place about 1 g(M), accurately weighed, of the prepared sample of AI into a 250ml volumetric flask and dilute with about 150 ml of water. Mix thoroughly the contents of the flask and make the volume of 250 ml with water. Using separate pipettes, take accurately 5 ml each of solution A and solution B in a porcelain dish. Add about 12 ml of AI solution from a burette and heat to boiling over an asbestos gauze. Add 1 ml of methylene blue indicator and while keeping the solution boiling complete the titration within 3 minutes, the end point being indicated by change of colour from blue to red. Note the volume (H) in ml of AI solution required for the titration.

(ix) Calculation
Total reducing sugars, percent by mass = \( \frac{250 \times 100 \times S}{H \times M} \)

Where
S = strength of copper sulphate solution,
H = volume in ml of AI solution required for titration, and
M = mass in g of AI taken for the test.

5.6 Determination of sucrose
(i) Procedure: - To 100 ml of the stock AI solution, add 1 ml of concentrated hydrochloric acid and heat the solution to near boiling. Keep aside overnight. Neutralize this solution with sodium carbonate and determine the total reducing sugars as described in.
(ii) Calculation
(a) Sucrose, percent by mass = (reducing sugars after inversion, percent by mass) –(reducing sugars before inversion, percent by mass) x 0.95
(b) Nitrogen, mg per gram of sucrose consumed = 2(a-b)-C
Where
a= initial quantity of sucrose taken for the test
b=mass of sucrose as calculated in (a),and
c= amount of nitrogen fixed per gram of glucose.

1.C. Method of Analysis of Azospirillum Biofertilisers
1. Apparatus: same as Rhizobium

2. Reagents
2.1 Medium

Use N-free semisolid medium (Nfb) of the following composition for preparation of MPN tubes
DL-Malic acid                  5.0  
K₂HPO₄                      0.5  
MgSO₄ 7H₂O            0.2  
NaCl                       0.1  
CaCl₂                      0.02  
Trace element Soln.        2.0 ml  
Fe EDTA (1.64% Soln.)      4.0 ml  
Vitamin soln.              1.0 ml  
KOH                        4.0 ml  
Bromothymol blue (0.5% aq.) 2.0 ml  
Adjust pH to 6.8-7.0 with KOH 
For semi solid add agar    1.75 g  
For solid medium add agar   15.0 g

2.1.1 **Trace element solution (g/litre)**
Na₂MoO₄ 2H₂O                  0.2  
MnSO₄ H₂O                     0.235  
H₃BO₃                       0.28  
CuSO₄ 5H₂O                   0.008  
ZnSO₄ 7H₂O                   0.024  
Distilled water             1000 ml  
Use 2 ml of this solution in one litre of Nfb media

**Vitamin solution (g/litre)**
Biotin                      0.01  
Pyridoxin                   0.02  
Distilled water            1000 ml  
Use one ml of this sol. in one litre of Nfb media

2.2 **Sterilization and preparation of MPN tubes**
2.2.1 Prepare Nitrogen free Bromothymol Blue malate medium as mentioned at paragraph 2.1. Boil to dissolve agar. Quickly dispense 10 ml molten medium in 15 x 150 ml test tubes or screw capped culture tubes and close either with cotton plugs or screw caps. Minimum of 25 such tubes shall be needed for each sample.

2.2.2 Sterilize the tubes by autoclaving at 121°C for 20 minutes, as in Rhizobium at paragraph 2.3.2.

3. **Preparation of serial dilution for MPN count**
Dispense 30 g of Azospirillum biofertilizers in 270 ml of sterile water and shake for 10 minutes on a reciprocal shaker. Make serial dilutions up to 10⁻⁸ dilution. Pipette out 1 ml aliquots of 10⁻⁴ to 10⁻⁸ dilution and deliver it to screw cap tubes or test tubes containing N-free semi solid Nfb media.
4. **Incubation of tubes**

Label the tubes and incubate at 36 ± 1°C for 3-4 days in vertical position in a test tubes stand. Do not disturb the medium during the entire period of incubation.

<table>
<thead>
<tr>
<th>P_1</th>
<th>P_2</th>
<th>Most probable number for indicated values of P_3</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
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</tr>
<tr>
<td>0</td>
<td>0</td>
<td>-</td>
</tr>
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<td>0</td>
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<tr>
<td>0</td>
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</tr>
<tr>
<td>0</td>
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<td>0.094</td>
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<tr>
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<tr>
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<tr>
<td>1</td>
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<tr>
<td>1</td>
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<tr>
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</tr>
<tr>
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<td>4</td>
<td>1.3</td>
</tr>
<tr>
<td>5</td>
<td>5</td>
<td>2.4</td>
</tr>
</tbody>
</table>

*Most Probable Numbers for use with 10 fold dilution and 5 tubes per dilution (Cochran, 1950)*

5. **Counting**

5.1 Count the tubes which have turned blue and have developed typical white sub-surface pellicle.
5.2 Count the tubes as +ve or –ve for the presence of sub-surface pellicle and consider for the purpose of calculation.

5.3 **Method for Estimating MPN Count**

5.3.1 To calculate the most probable number of organisms in the original sample, select as \( P_1 \) the number of positive tubes in the least concentrated dilution in which all tubes are positive or in which the greatest number of tubes is +ve, and let \( P_2 \) and \( P_3 \) represent the numbers of positive tubes in the next two higher dilutions.

5.3.2 Then find the row of numbers in Table 1 in which \( P_1 \) and \( P_2 \) correspond to the values observed experimentally. Follow that row of numbers across the table to the column headed by the observed value of \( P \).

5.3.3 The figure at the point of intersection is the most probable number of organisms in the quantity of original sample represented in the inoculum added in the second dilution. Multiply this figure by the appropriate dilution factor to obtain the MPN value.

5.3.4 **Azospirillum count/g of carrier =** \( \frac{\text{Value from MPN table} \times \text{Dilution level}}{\text{Dry mass of product}} \)

**1.D. METHOD OF ANALYSIS OF PHOSPHATE SOLUBULISING BACTERIAL BIOFERTILIZER**

1. **Apparatus** - Same as Rhizobium

2. **Reagents**
   2.1 Medium
   Use a plating medium of the following composition:
   
<table>
<thead>
<tr>
<th>Ingredient</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glucose</td>
<td>10.0g</td>
</tr>
<tr>
<td>Tri-calcium phosphate</td>
<td>5.0 g</td>
</tr>
<tr>
<td>Ammonium sulphate</td>
<td>0.5 g</td>
</tr>
<tr>
<td>Magnesium sulphate</td>
<td>0.1 g</td>
</tr>
<tr>
<td>Sodium Chloride</td>
<td>0.2 g</td>
</tr>
<tr>
<td>Yeast extracts</td>
<td>0.5 g</td>
</tr>
<tr>
<td>Manganese sulphate</td>
<td>Trace</td>
</tr>
<tr>
<td>Ferrous sulphate</td>
<td>Trace</td>
</tr>
<tr>
<td>Distilled water</td>
<td>1000ml</td>
</tr>
<tr>
<td>Agar</td>
<td>15.0 g</td>
</tr>
<tr>
<td>pH adjusted to</td>
<td>7 ± 0.2</td>
</tr>
</tbody>
</table>

2.2 Sterilizing & preparation procedure for plates:
   Same as Rhizobium
2.3 Preparation of Plating Medium and Pouring
Same as Rhizobium

3. Preparation of Serial Dilution for Plate Counts:
Same as Rhizobium.

4. Incubation of Plates:
   4.1. Label the plates and incubate at 28± 1° C for 4 to 6 days.
   4.2. Colony counting aids: Same as Rhizobium

   **Counting**
   Count the total number of colonies on the plates including colonies with solubilisation zone with the help of a colony counter.

   **Methods of counting solubilisation zones**
   a. Take 10 g of PSBI (BF) in 90 ml in water
   b. Make a ten fold dilution series up to 10^7.
   c. Take 0.2 ml aliquote of 10^5 to 10^7 dilutions using sterile pipettes and delivered to petri dishes containing Pikowskya media.
   d. Spread it uniformly. Invert the plates and incubate them up to 2 weeks at 28 ±2°C.
   e. Count the colonies showing hallow cones and measure their diameter.

5. Determination of soluble phosphorus using Ascorbic acid method:
   5.1 Apparatus
   Spectrometer capable of transmission measurement at 840 to 880 mm.
   Extractant: It is Olsen extract.

   5.2 Reagents
   Ammonium Molybdate [(NH₄)₆Mo₇O₂₄·4H₂O]
   L- Ascorbic Acid
   p-Nitro phenol
   4NH₂SO₄

   5.3 Preparation of reagents
   5.3.1 Sulphomolybdic Acid:-
   - Take 20 g of ammonium molybdate and dissolve in 300 ml of distilled water.
   - Add slowly 450 ml of 10N - H₂SO₄.
   - Cool the above mixture and add 100 ml of 0.5 percent solution of antimony Potassium tartrate.
   - Cool and make the volume to one liter. Store in glass bottle away from direct sunlight.
5.3.2 Preparation of Mixed Reagent
Add 1.5 g of L-ascorbic acid in 100ml of the above stock solution and mix. Add 5ml of this solution to develop colour. Mixed reagent is to be prepared fresh as it does not keep for more than 24 h.

5.3.3 Procedure
(i) Preparation of Sample
Pure culture medium same as at 2.1 above excluding agar. Prepare broth medium in 100 ml aliquots in 6 no., 250 ml conical flasks and sterilize in autoclave at 121°C for 20 min.

(ii) Inoculation of Medium
Select one PSB colony of the type that has been counted as PSB (showing sufficient zone of solubilization) and streak on set medium as described at 2.1 in a Petri dish. Use this pure culture for inoculating the broth. Inoculate 3 flasks and keep 3 flasks as uninoculated control. Incubate the flasks over rotary shaker for 12 days at 28+1°C.

After 12 days, filter the contents of each flask separately through Whatman No. 42 filter paper or centrifuge at 10,000 rpm for 15 min.

(iii) Add 10 ml of filtrate/centrifugate to 50 ml of olsen extractant and shake for 30 min over rotary shaker.

(iv) Filter the suspension through Whatman filter paper No. 40. If the filtrate is coloured then add a tea spoon of Dacro-60 (activated phosphorous free carbon), reshake and filter.

(v) Take a known aliquot (5 to 25 ml) of the extract in a 50 ml volumetric flask.

(vi) Add 5 drops of p-nitrophenol indicator (1.5 per cent solution in water) and adjust the pH of the extract between 2 and 3 with the help of 4NH₂SO₄. The yellow colour will disappear when the pH of the solution becomes 3. Swirl gently to avoid loss of the solution along with the evolution of CO₂.

(vii) When the CO₂ evolution has subsided, wash down the neck of the flask and dilute the solution to about 40 ml.

(viii) Add 5 ml of the sulphotomolybdic acid mixed reagent containing ascorbic acid, swirl the content and make up the volume.
(ix) Measure the transmission after 30 min at 880 nm using red filter. The blue colour developed remains stable upto 60 minutes.

(x) Record the concentration of phosphorous (P) in the extract form the standard curve and calculate the concentration of soluble phosphorous as follows:”

5.3.4. Calculations

(a) Weight of the substance taken = x g
(b) Volume of the extract added = 50 ml
(c) Volume of the extract taken for P determination = y ml
(d) Volume made after colour developed = 50 ml
(e) Reading from the standard curve against percent transmission recorded = z ppm
(f) Soluble Phosphorous percent p = \( \frac{Z \times 50 \times 10^{-6} \times 50 \times 100}{Y \times X} \)

5.3.4 Preparation of standard Curve
Prepare standard curve using 0.1 to 0.6 ppm P in 50 ml volumetric flask. Plot the standard curve by taking concentration of soluble P on x- axis and percent T on y- axis using a semi log graph paper. It is a straight line relationship between the soluble P and percent T when plotted on a semi-log graph paper.

1. E. METHODS OF ANALYSIS FOR MYCORRHIZAL BIOFERTILIZERS

1. Estimation of pH
   As per methodology specified in Schedule IV Part D at Serial Number 1

2. Estimation of moisture contents
   As per methodology specified in schedule IV Part D at serial number of FCO 1985

3. Estimation of total viable propagules
   3.1. Harvesting of spores from finished product

   (a) By sieving

   Equipment and Reagent
   Stalking sieves with nylon or stainless steel mesh and a large range of pore sizes for isolating spores from the carrier or soil sample

   - 40-50 micron (0.04 mm) sieve for small sized spores
   - 100 micron (0.10 mm) sieve for medium sized spores
b. **Procedure** - Mix the Soil in a substantial volume of water and decant through a series of sieves arranged in descending order of mesh size. Roots and coarse debris are collected on a coarse (60-ISS) sieve, while spores are captured on one or more finer sieves. Vigorous washing with water is necessary to free spores from aggregates of clay or organic materials. Collect the sieving in jars. Transfer the sieving onto the grided petri dishes/plate and observe under stereomicroscope. Count the number of spores in plate/dish and express it as spores/g of the sample.

c. **By sucrose gradient**

i. Collect the sieving by the method described above. Transfer the sieving into centrifuge tubes and centrifuge for 5 minutes at 1750 rpm in a horizontal rotor.

ii. Decant the supernatant liquid carefully and resuspend pellet in 60% sucrose solution. Again centrifuge for 2–5 minutes

iii. Pour the supernatant (with spores) onto a 300BSS sieve size and rinse with water to remove the sugar. Transfer the sieving onto the grided petri dishes/plate and observe under stereomicroscope. Count the number of spores in plate/dish and express it as spores/g of the sample.

### 3.2. Spore staining

**a. Equipment and Reagent**

1. Equipments and reagents for spore extraction as described previously.
2. 2,5-diphenyl-2N-tetrazolium bromide (MTT).
3. Distilled water
4. Eppendorf
5. Stereomicroscope
6. Petri-dishes
b. Procedure

- Prepare 0.25% solution of MTT (2,5-diphenyl-2N-tetrazolium bromide.
- Avoid exposure of MTT solution to light, as the stain is light sensitive.
- Add freshly collected AMF spores (approximately 100 in number) collected by any of the two methods described above, to the staining solution and incubate at 27°C in sterile eppendorf in dark.
- Observe the spores for different colour reactions using stereomicroscope under dark field after 24 hours, 48 hours and 72 hours of incubation.
- Spores, which stained red or pink, are treated as viable.

\[
\% \text{Spore viability} = \frac{\text{No. of spores which stained red or pink}}{\text{Total number of spores}} \times 100
\]

4. Assessment of Infectivity Potential

The bioassay is used to determine the number of infective propagules present in the product. Once the infective propagules (spores, mycelia and vesicles in the root fragments) come in contact with the host roots they give out a turgid mycelial structure - the appressoria, which is the initial step in the penetration event. This appressoria enters the root through an ‘entry point’. This entry point can be visualized by staining and enumerated as a measure of the infectivity of the inoculum. Host plants are grown from pre germinated seeds and a known weight of the inoculum is applied to experimental host plant in pots. These pots are maintained for 14 days after which they are harvested, the root length measured and then stained. The resulting entry points are counted to ascertain the infectivity potential.

(A) Equipments and Reagents

a. Pots (5 x 7 cm in size)
b. Sorghum seeds (Sorghum vulgare)
c. Scissors and needles
d. Petri dish (grided)
e. Water bath
f. Glass slides and cover slips
g. Compound microscope
h. Coarse sieve to prevent root loss during washing/changing solutions
i. Plastic vials with tight-sealing lids for storage of stained samples in 50% glycerol
j. Potassium hydroxide solution (5-10%)
k. Alkaline $\text{H}_2\text{O}_2$ (25% Ammonia solution: 3 ml + 10% $\text{H}_2\text{O}_2$: 30 ml + Distilled water: 67 ml)

l. 1% HCl

m. 50% glycerol-water (v/v) solution for de-staining and storage of stained roots.

n. Lactoglycerol (Lactic acid: 876 ml + Glycerine: 64 ml + Distilled water: 60 ml)

(B) Staining solutions

a. 0.01 % acid fuschin: 0.01 g acid fuschin in 100ml acetoglycerol.

b. 0.05% trypan blue: 0.05 g trypan blue in 100ml acetoglycerol.

c. 0.03% Chlorozol black E (CBE) in lactoglycerol (1:1:1 lactic acid, glycerol and water).

d. Dissolve CBE in water before adding equal volumes of lactic acid and glycerol.

(C) Procedure

a. Place 100 g test sample in a pot

b. Dilute the inoculum with sterilized sand if the inoculum is very rich

c. Plant 10-12 pre germinated seeds of Sorghum and grow for 14 days

d. Harvest the pots and recover roots (fine roots can be rescued using sieve) completely.

e. Chop the roots equally 1 cm in length.

f. Measure and record the root length (using grid line intersect method described below) from each sample/dilution.

g. Clear the roots in KOH solution and stain the root pieces (described below)

h. Count the number of infection points/entry points formed on randomly picked 100 segments

i. Calculate the average number of entry points formed in 1 cm segment

j. Calculate the total number of infection points/inf ective propagules (IP) by multiplying the average number of entry points formed in 1 cm segment by the total root length.

k. Extrapolate the IP present as numbers per gram of substrate/inoculum

(D) Estimation of root length (Tennant D 1975)

(1) Equipment

a. Scissors

b. Petri dish (9 cm in size consisting 1.33 cm x 1.33 cm grids)

c. Wash bottle

d. Stereo zoom microscope
(2) The lines intersect (Tennant 1975) method is used to estimate the length of hyphae and roots. Root length is measured by dispersing roots against a grid of squares on the bottom of a tray. The roots are spread apart from one another over a grid in 2 mm to 10-mm depth of water. The eyes of the observer are cast along all the horizontal and vertical lines of the grid and root is counted using a hand held click counter.

The root length is calculated as follows:

Root length = No. of intersects x 11/14 x grid size

Where, 11/14 is a constant, and the size of the grid is the length of one side of one square of the grid.

Counting root intersections
a. Randomly disperse root in dish with grid lines.
b. Count the intersects on roots across the horizontal and vertical lines.
c. An example of 10 root segments is presented to show how the root length is calculated

Total number of intersects = HI+VI = 8 + 7 (example) = 15

Thus, the root length = 11/14 X 15 X 1 (as the grid size is 1cm) = 11 cm (example)
(3) Clearing and staining root specimens
Clearing and staining procedures requires root samples that should be washed free of soil. It is important that KOH and staining solution volumes are sufficient for the amount of roots being processed and that, roots are not tightly clumped together for uniform contact with solutions. To ensure uniform staining, the roots should be chopped in to smaller (1-2 cm) segments.

(a) Wash root specimens under running tap water thoroughly. Place them in beaker containing 5-10% KOH solution for about 15-30 minutes. The concentration of KOH and time of incubation of roots depend upon the age and tenderness of the roots.

(b) Pour off the KOH solution and rinse the roots well in a beaker using at least three complete changes of tapwater or until no brown colour appears in the rinse water.

(c) Cover the roots with alkaline H$_2$O$_2$ at room temperature for 10 minutes or until roots are bleached.

(d) Rinse the roots thoroughly using at least three complete changes of tap water to remove the H$_2$O$_2$.

(e) Cover the roots with 1% HCl and soak for 3-4 min. And then pour off the solution. DO NOT rinse after this step because the specimens must be acidified for proper staining.

(f) Incubate the roots with staining solution (0.01% acid fuchsine in lactoglycerol or 0.05% trypan blue in lacto phenol) and keep them overnight for staining.

(g) Place the root specimens in glass petriplate /multiwell plate for destaining. The destaining solution (50% glycerol) is the standard used in step 4, but of course, without the stain.

(4) Sample storage and slide preparation
If clearing and staining is not possible immediately then fresh roots can be kept moist and stored at 5 °C (for several days), or may be preserved in 50% ethanol for months together in tightly sealed vials.

Staining quality is subsequently improved by destaining roots in 50% glycerol for several months prior to observation to allow excess stain to leach from roots. Semi-permanent slides of stained roots can be made with PVLG mountant. For temporary slide the stained roots can be observed in plain lactoglycerol.
1F. METHOD OF ANALYSIS FOR POTASH SOLUBILISING BIOFERTILIZERS (KSB)

1. Estimation of total viable count and contamination

1. Apparatus-

1.1 Pipettes graduated 1 ml and 10 ml
1.2 Dilution bottles or flasks
1.3 Petri dishes clear, uniform, flat-bottomed
1.4 Hot-air oven: Capable of giving uniform and adequate temperature, equipped with a thermometer, calibrated to read up to 250°C and with venus suitably located to assure prompt and uniform heating.
1.5 Autoclave
1.6 Incubator
1.7 Handy tally or mechanical counting device
1.8 Ph meter

2. Reagents

2.1 Medium
Use plating medium of the following composition for total viable count and contamination

Medium for analysis of total viable count and contamination (ingredients g/lit)

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<tr>
<th>Ingredient</th>
<th>Quantity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manitol</td>
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</tr>
<tr>
<td>Yeast extract</td>
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</tr>
<tr>
<td>Peptone</td>
<td>2.0</td>
</tr>
<tr>
<td>Agar</td>
<td>18.5</td>
</tr>
<tr>
<td>Trace element solution</td>
<td>1 ml</td>
</tr>
<tr>
<td>Distilled Water</td>
<td>1000 ml</td>
</tr>
</tbody>
</table>

Trace element solution (ingredients g/lit)

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>Quantity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sodium molybdate</td>
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</tr>
<tr>
<td>Boric acid</td>
<td>0.28</td>
</tr>
<tr>
<td>Manganese sulphate</td>
<td>0.23</td>
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<tr>
<td>Copper sulphate</td>
<td>0.01</td>
</tr>
<tr>
<td>Zinc sulphate</td>
<td>0.03</td>
</tr>
<tr>
<td>Distilled Water</td>
<td>1000 ml</td>
</tr>
</tbody>
</table>
Medium for studying zone of solubilization in KSB
(Ingredients g/lit)

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>Amount</th>
</tr>
</thead>
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<tr>
<td>Glucose</td>
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<td>Ferric chloride</td>
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<td>Calcium carbonate</td>
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<tr>
<td>Potassium mineral (mica powder)</td>
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</tr>
<tr>
<td>Calcium phosphate</td>
<td>2.0</td>
</tr>
<tr>
<td>Distilled Water</td>
<td>1000</td>
</tr>
</tbody>
</table>

2.2 Sterilizing and preparation procedure for plates

2.2.1 Sterilize the sampling and plating equipment with dry heat in a hot air oven at less than 160°C for not less than 2 hours;

2.2.2 Sterilize the media by autoclaving at 120°C for 20 min. To permit passage of steam into and from closed containers when autoclaved, keep stoppers slightly loosened or plugged with cotton. Air from within the chamber of the sterilizer should be ejected allowing steam pressure to rise.

2.3 Preparation of plating medium and pouring

2.3.1 Prepare growth medium in accordance with the composition of the specific biofertiliser.

2.3.2 Melt the required amount of medium in boiling water or by exposure to flowing steam in partially closed container but avoid prolonged exposure to unnecessarily high temperature during and after melting. Melt enough medium which will be used within 3 h. Re-sterilization of the medium may cause partial precipitation of ingredients.

2.3.3 When holding time is less than 30 min. promptly cool the molten medium to about 45°C, and store until used, in a water bath or incubator at 43 to 45°C. Introduce 12 to 15 ml of liquefied medium or appropriate quantity depending on size of the Petri dish at 42 to 44°C into each plate. Gently lift the cover of the dish just enough to pour in the medium. Sterilizes the lips of the medium containers by exposure to flame.

(a) Immediately before pouring.
(b) Periodically during pouring, and
(c) When pouring is complete for each batch of plates, if portions of molten medium remain in containers and are to be used without subsequent sterilization for pouring additional plate. As each plate is poured thoroughly mix the medium with test portions in the petri dish.

2.3.4 By rotating and tilting the dish and without splashing the medium over edge spread the medium evenly over the bottom of the plate. Provide conditions so that the medium solidifies with reasonable promptness (5-10 min) before removing the plates from level surface.

3. Preparation of Serial Dilution for Plate Counts

3.1 Dispense 10 g of inoculants to 90 ml of sterile distilled dematerialized water and shake for 10 min on a reciprocal shaker or homogenizer. Make serial dilutions upto 10^{10}. Take 1:0 ml or suitable aliquots of 10^6 to 10^9 dilutions using sterile pipettes and deliver to petri dishes containing set medium as given in 2.1 and spread it uniformly with a spreader or use droplet method. Invert the plates and promptly place them in the incubator.

4. Incubation of Plates

4.1 Label the plates and incubate at 28± 2^0C for 4 to 6 days.

4.2 Colony counting aids
   Count the colonies with the aid of magnifying lens under uniform and properly controlled, artificial illumination. Use a colony counter, equipped with a guide plate and rules in centimeter square. Record the total number of colonies with the hand tally. To distinguish colonies from dirt, specks and other foreign matter, examine doubtful objects carefully.

4.3 Count all plates but consider for the purpose of calculation plates showing more than 30 and less than 300 colonies per plate. Disregard colonies which absorb congo red and stand out as reddish colonies. *Fraturia aurentia* (KMB) stand out as white-opaque glistening and domed colonies. Count such colony numbers and calculate figures in terms of per litre, of carrier. Also check for freedom from contamination at 10^5 dilution.
4.4 Counting
Count the total number of colonies on the plates including colonies with solubilization zone with the help of a colony counter.

5 Method of estimation of K solubilization zones

5.1 Take 10 g of KSB in 90 ml sterile distilled water

5.2 Make a tenfold dilution series up to $10^7$

5.3 Take 1.0 ml aliquot of $10^5$ to $10^7$ dilutions using sterile pipettes and deliver to petri dishes containing K-solubilization zone media.

5.4 Spread it uniformly, invert the plates and incubate for up to 2 weeks at $28\pm2^0C$.

5.5 Count the colonies showing solubilization zones and measure the diameter of solubilization zone. Calculate average zone of solubilization in mm.

1G METHOD OF ANALYSIS OF FOR ZINC SOLUBILIZING BIOFERTILIZERS

1. Estimation of total viable count and contamination

2. Apparatus

2.1 Pipettes graduated 1ml and 10 ml
2.2 Dilution bottles or flasks
2.3 Petri dishes clear, uniform, flat-bottomed
2.4 Hot-air oven: Capable of giving uniform and adequate temperature, equipped with a thermometer, calibrated to read upto $250^0C$ and with venus suitably located to assure prompt and uniform heating.
2.5 Autoclave
2.6 Incubator
2.7 Hand tally or mechanical counting device
2.8 pH meter

3. Reagents

3.1 Medium
Use plating medium of the following compostion for total viable count and contamination
Medium for analysis of Total Viable Count, Contamination and zone of solubilization for Zn solubilisation for Zn solubilizing biofertilizer

<table>
<thead>
<tr>
<th>Ingredients</th>
<th>g/lit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glucose</td>
<td>10.0</td>
</tr>
<tr>
<td>Zinc oxide</td>
<td>1.0</td>
</tr>
<tr>
<td>Amm sulphate</td>
<td>0.5</td>
</tr>
<tr>
<td>Potassium chloride</td>
<td>0.2</td>
</tr>
<tr>
<td>Yeast extract</td>
<td>0.5</td>
</tr>
<tr>
<td>Ferrous sulphate</td>
<td>0.01</td>
</tr>
<tr>
<td>Manganese sulphate</td>
<td>0.01</td>
</tr>
<tr>
<td>Di Pot Hyd. Phosphate</td>
<td>0.5</td>
</tr>
<tr>
<td>Distilled Water</td>
<td>1000 ml</td>
</tr>
</tbody>
</table>

3.2 Sterilizing and preparation of plates

3.2.1 Sterilize the sampling and plating equipment with dry heat in a hot air oven at less than 160°C for not less than 2 hours;

3.2.2 Sterilize the media by autoclaving at 120°C for 20 min. To permit passage of steam into and from closed containers when autoclaves, keep stoppers slightly loosened or plugged with cotton. Air from within the chamber of the sterilizer should be ejected allowing steam pressure to rise.

3.3 Preparation of plating medium and pouring

3.3.1 Prepare growth medium in accordance with the composition of the specific biofertilizer.

3.3.2 Melt the required amount of medium in boiling water or by exposure to flowing steam in partially closed container but avoid prolonged exposure to unnecessarily high temperature during and after melting. Melt enough medium which will be used within 3 hours. Re-sterilization of the medium may cause partial precipitation of ingredients.

3.3.3 When holding time is less than 30 min. promptly cool the molten medium to about 45°C, and store until used, in a water bath or incubator at 43- to 45°C. Introduce 12 to 15 ml of liquefied medium or appropriate quantity depending on size of the petri dish at 42 to 44°C into each plate. Gently lift the cover of the dish just enough to pour in the medium. Sterilize the lips of the medium containers by exposure to flame.

(a) Immediately before pouring.
(b) Periodically during pouring, and
(c) When pouring is complete for each batch of plates, if portions of molten medium remain in containers and are to be used without subsequent sterilization for pouring additional plate. As each plate is poured thoroughly mix the medium with test portions in the petridish.

3.3.4 By rotating and tilting the dish and without splashing the medium over edge spread the medium evenly over the bottom of the plate. Provide conditions so that the medium solidifies with reasonable promptness (5-10 min) before removing the plates from level surface.

4. Preparation of Serial Dilution for Plate Counts

4.1 Dispense 10 g of inoculants to 90 ml of sterile distilled dematerialized water and shake for 10 min on a reciprocal shaker or homogenizer. Make serial dilutions upto $10^{10}$. Take 1:0 ml or suitable aliquots of $10^6$ to $10^9$ dilutions using sterile pipettes and deliver to petri dishes containing set medium as given in 2.1 and spread it uniformly with a spreader or use droplet method. Invert the plates and promptly place them in the incubator.

5. Incubation of Plates

5.1 Label the plates and incubate at $28\pm 2^0C$ for 4 to 6 days.

5.2 Colony counting aids : Count the colonies with the aid of magnifying lens under uniform and properly controlled, artificial illumination. Use a colony counter, equipped with a guide plate and rules in centimeter square. Record the total number of colonies with the hand tally.

5.3 Count all plates but consider for the purpose of calculation plates showing more than 30 and less than 300 colonies per plate. Disregard colonies which absorb congo red and stand out as reddish colonies. Zinc solubilizing biofertilisers stand out as white, translucent, glistening and elevated colonies. Count such colony numbers and calculate figures in terms of per litre, of carrier. Also check for freedom from contamination at $10^5$ dilution.

5. Counting
Count the total number of colonies on the plates including colonies with solubilization zone with the help of a colony counter.
6. Method of estimation of Zinc solubilisation zones

6.1 Take 10 g of ZSB in 90 ml sterile distilled water
6.2 Make a tenfold dilution series upto $10^7$
6.3 Take 1.0 ml aliquot of $10^5$ to $10^7$ dilutions using sterile pipettes and deliver to petri dishes containing Zinc solubilization zone media.
6.4 Spread it uniformly, invert the plates and incubate for up to 2 weeks at $28\pm2^0C$.
6.5 Count the colonies showing solubilization zones and measure the diameter of solubilization zone. Calculate average zone of solubilization in mm.

1.H. METHOD OF ANALYSIS OF ACETOBACTER (spp.)

1. Apparatus - Same as Rhizobium
2. Reagents -
   2.1 Medium
       Use plating medium of the following composition for viable count and contamination :-

       Medium for analysis of total viable count and contamination (ingredients gram/litre)
       Sucrose                               100 g
       $K_2HPO_4$ (Di-Potassium Hydrogen Phosphate) 0.4 g
       $KH_2PO_4$ (Potassium di-hydrogen phosphate) 0.6 g
       MgSO$_4$ (Magnesium Sulphate) 0.2 g
       Calcium Chloride 0.02 g
       Sodium Molybdate 0.02 g
       Ferric Chloride 0.01 g
       Bromothymol blue solution (0.5% in 0.2 m KOH) 5.0 ml
       Distilled water 1000ml
       pH 5.5 g
       agar agar 18.5 g
3. Sterilizing & preparation procedure for plates:
   Same as Rhizobium
   3.1 Preparation of Plating Medium and Pouring
       Same as Rhizobium
4. Preparation of Serial Dilution for Plate Counts:
   Same as Rhizobium.
5. Incubation of Plates:
   Same as Rhizobium.
5.1. Colony counting Aids: Count the colonies with the aid of magnifying lens under uniform and properly controlled, artificial illumination. Use a colony counter, equipped with guide plate and rules in centimeter square. Record the total number of colonies with hand tally. Avoid mistaking particles of undissolved medium or precipitated matter in plates for pin point colonies. To distinguish colonies from dirt, specks and foreign matter, examine doubtful objects carefully.

5.2 Count all plates but consider for the purpose of calculation only those plates showing more than 30 and less than 300 colonies per plate. Acetobacter a nitrogen fixing bacteria stand out as irregular 2-3 mm diameter, smooth flat with bright yellow or yellow with orange centre colour. Count such colony numbers and calculate figures in terms of per litre, of carrier. Also check freedom from contamination at $10^5$.

6. Test for confirmation:

6.1 Apparatus
Same as Azospirillum.

6.2 Reagents

6.2.1 Medium: Semi solid for pellicle formation (ingredients gm per liter)

- Sucrose: 100 g
- $K_2HPO_4$ (Di-Potassium Hydrogen Phosphate): 0.4 g
- $KH_2PO_4$ (Potassium di-hydrogen phosphate): 0.6 g
- $MgSO_4$ (Magnesium Sulphate): 0.2 g
- Calcium Chloride: 0.02 g
- Sodium Molybdate: 0.02 g
- Ferric Chloride: 0.01 g
- Bromothymol blue solution (0.5% in 0.2 m KOH): 5.0 ml
- Distilled water: 1000 ml
- pH: 5.5 g
- agar agar: 1.75 g

6.3 Sterilization and preparation of MPN tubes:
Same as Azospirillum

6.4 Preparation of serial dilution for MPN count:
Same as Azospirillum

6.5 Incubation of tubes:
Same as Azospirillum

6.6 Counting: Yellowish pellicle formation below 1 mm of upper surface of nitrogen free semi solid media. Counting the tubes or plates which have turned yellowish in colour after inoculation and ascertained the
presence of pellicle in undistributed medium. To determine usual contamination on the same examine doubtful objects carefully.

6.7 Method for estimating MPN count:
Count all tubes which have turned yellowish and consider them for purpose of calculation. Count such type of tubes and tally this count with MPN table (as specified in the method of analysis of Azospirillum) to get the number of cells per gram of carrier or number of cells per ml of liquid.

1.I. METHOD OF ANALYSIS OF CARRIER BASED CONSORTIA OF BIOFERTILISER AND LIQUID CONSORTIA OF BIOFERTILISER

1. Methods of Analysis of Rhizobium Biofertiliser – Same as specified for Rhizobium at 1.A.
2. Methods of Analysis of Azotobacter – same as specified for Azotobacter at 1.B.
3. Methods of Analysis Azospirillum – same as specified for Azospirillum as 1.C.
4. Methods of Analysis of Phosphate Solubilising Bacteria PSB – same as specified for Phosphate Solubilising Bacteria at 1.D.
5. Methods of Analysis of Potash Mobilising Bacteria (KMB) – same as specified for Phosphate Solubilising Bacteria at 1.F.

MAINTENANCE AND PREPARATION OF CULTURE AND QUALITY CONTROL AT BROTH STAGE

RHIZOBIUM:
1. Maintenance of pure cultures
   Maintain pure culture of rhizobia on yeast extract mannitol agar (YEMA) slants to the following composition.

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mannitol</td>
<td>10.0g</td>
</tr>
<tr>
<td>Potassium hydrogen phosphate (K_2HPO_4)</td>
<td>0.5 g</td>
</tr>
<tr>
<td>Magnesium sulphate (MgSO_4 7 H_2O)</td>
<td>0.2g</td>
</tr>
<tr>
<td>Sodium chloride (NaCl)</td>
<td>0.1 g</td>
</tr>
<tr>
<td>Calcium carbonate (CaCO_3)</td>
<td>1.0 g</td>
</tr>
<tr>
<td>Yeast extracts</td>
<td>1.0g</td>
</tr>
<tr>
<td>Agar</td>
<td>18g</td>
</tr>
<tr>
<td>Distilled water</td>
<td>1 liter</td>
</tr>
<tr>
<td>pH</td>
<td>6.8-7.0</td>
</tr>
</tbody>
</table>

Transfer a loopful of the pure culture to each of the agar slant aseptically in an inoculation room and incubate at 28+/- 2° C for 3 to
10 days depending upon the species of Rhizobium. Always keep culture at 4° C.

2. Preparation of Inoculums Cultures
   2.1 Prepare yeast extract mannitol broth of the composition as given in 1.1 minus the agar.
   2.2 Transfer a loop full of the culture.

3. Quality Control Test Recommended at Broth Stage:
   3.1 Qualitative Tests
   3.1.1 Check for freedom from the visible contaminants
   3.1.2 The pH of the bacterial broth shall normally be between 6.5 and 7.5.
   3.1.3 Smear and Gram stain
   3.1.3.1 Reagents
   a. Ammonium oxalate crystal violet stain- weigh 0.2 g of crystal violet and dissolve in 20 ml of 95 percent ethyl alcohol. Dissolve separately 0.8 g of ammonium oxalate in 30 ml of distilled water. Mix the two solution and filter through a filter paper.
   b. Iodine solution
      Iodine 1.00 g
      Potassium iodide 2.00 g
      Distilled water 300 ml
      Weigh the ingredient and dissolve in water. Filter through a filter paper.
   c. Erythrosine
      Erythrosine 1.00 g
      Phenol 5.00 g
      Distilled water 100 ml
      Weigh the ingredient, dissolve in distilled water and filter through a filter paper.
   3.1.3.2 Procedure
      Prepare a smear on a clean microscope slide, fix over a flame by gently and intermittent heating, air cool and flood with ammonium oxalate crystals violet stain for 1 min. After removing the excess of ammonium oxalate crystals violet, wash the slide under a gentle stream of running tap water. Flood the slide with iodine solution for half a minute remove excess stain wash 95 percent ethyl alcohol and finally wash under a gentle stream of running tap water. Flood the slide with erythrosine stain for about 3 min, wash under a gentle stream of running tap water and dry between the folds of a filter paper. Examine the slide under a compound microscope using an oil immersion objective.
Note:- A smear prepared from undiluted broths should be free from Gram positive cells. The presence of a few gram positive cells in occasional fields which may be due to dead cells in the medium may be disregarded.

3.1.4 Absence of Growth on Glucose- Peptone Agar
The composition of the glucose –peptone agar is as follows:
- Glucose 10.0 g
- Peptone 20.0 g
- Sodium chloride (NaCl) 5.0 g
- Agar (IS6850) 15.0 g
- Distilled water 100 ml
- Bromocresol purple 10 ml
- Alcoholic solution (1.6%) 
- pH 7.2

Note:- when a loopful of the broth is streaked into this Medium and incubated at 28±2°C for 24 h, the purple-violet colour of the medium (due to the indicator bromo-cresol purple) shall not change. If the colour changes to yellow (acidic reaction) or blue (alkaline reaction) the broth is grossly contaminated. Hence the broth should be rejected.

3.1.5. Streak on yeast extract mannitol agar with congo-red
When a loopful of broth culture is streaked to a plate of this medium and incubated at 28± 2°C for 3 - 10 days, it shows colonies of bacteria with growth characteristics same as that of the pure culture used in the preparation of the broth, otherwise the broth should be rejected.

3.2 Quantitative Test
3.2.1 Viable or plate counts - Serially dilute one milliliter of the broth to obtain dilutions of the order of $10^6$ to $10^9$. Plate 0.2 ml aliquots of the dilution on YEMA plates and incubate at 28±2°C for 2 to 6 days, depending on the species of Rhizobium. The counts of viable Rhizobium in the final broth from shake culture or fermenters shall be not less than $10^8$ to $10^9$ cells/ ml. Otherwise, the broth should be rejected.

**AZOSPIRILLUM**

1. **Maintenance of pure cultures.**
   Maintain pure culture of Azospirillum on nitrogen free bromothymol blue medium and maintain as solid medium.
Transfer a loopful of pure culture to each of the agar culture tube aseptically in an inoculation room and incubate 37± 2°C for three days and keep in undisturbed. Always keep pure culture below 5°C.

2. **Preparation of Inoculums culture and Mass culture:**
   Inoculums culture and mass culture of this standard shall be prepared as described for Rhizobium of this standard.

3. **Quality Control Test Recommended at Broth Stage**
   3.1. **Quality Test**
      3.1.1 Check for free from contaminants by preparing slide and observing under microscope.
      3.1.2 The pH of bacterial broth shall normally be between 7.0 to 8.0.
      3.1.3 Gram staining test shall be carried out as described for Rhizobium of this standard.
      3.1.4. See the colour change in the media after 24 hours from inoculation. The colour will change from green to blue.
      3.1.5 Watch the pellicle just below the surface of the media. It is checked on the third day after keeping inoculated broth undisturbed.

3.2 **Quantitative Test**
   3.2.1 Most probable Number (MPN) as given in Annexure E. The counts of Azospirillum in the final broth from shake culture or fermenter shall be not less than 10⁸ to 10⁹ cells/ ml. Otherwise the broth should be rejected.

**AZOTOBACTOR**

1. Maintenance of pure cultures.
   Maintain pure culture of Azotobacter on slants of the following composition
   
<table>
<thead>
<tr>
<th>Ingredient</th>
<th>Quantity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Agar</td>
<td>20 g</td>
</tr>
<tr>
<td>Sucrose</td>
<td>20 g</td>
</tr>
<tr>
<td>Ferrous Sulphate</td>
<td>0.1 g</td>
</tr>
<tr>
<td>Dibasic Potassium Phosphate</td>
<td>1.0 g</td>
</tr>
<tr>
<td>Magnesium Sulphate</td>
<td>0.5 gm</td>
</tr>
<tr>
<td>Calcium carbonates</td>
<td>2.0 gm</td>
</tr>
<tr>
<td>Sodium Molybdate</td>
<td>0.005gm</td>
</tr>
</tbody>
</table>

   Transfer a loopful pure culture to each of agar slants aseptically in an inoculation room and incubate at 28±2°C for 3 to 10 days depending up on the species of Azotobactor. Always keep pure culture at 5°C.
2. Preparation of inoculums culture
   2.1 Prepare Jensen’s media broth of the compositor as given in
       1.1 minus the agar.
   2.2 Transfer a loop full of the culture into a 100 ml/250 ml conical
       flask containing the broth. Incubate the flasks at 28±2°C on a
       rotary shaker for 2 to 6 days.

3 Quality control Tests recommended at Broth stage.
   3.1 Quality test
       3.1.1. Check for free from contaminants by preparing slide and
               observing under microscope.
       3.1.2 The pH by bacterial broth shall normally be between 6.5 to
               7.0
       3.1.3 Gram staining test shall be carried out as described for
               Rhizobium of this standard.

   3.2 Quantitative test
       3.2.1 Viable cell count same as Rhizobium

4. Packing, Marking, Storage and Use

   4.1 Packing - Biofertilizers shall be packed in suitable plastic bags/packets,
       thickness of which shall not be less than 75-100 micron or in suitable
       plastic bottles.

   4.2 Marking - Each polyethylene pack shall be marked legibly and indelibly
       with the following information:
       (a) Name of the product,
       (b) Name and address of the manufacturer,
       (c) Crops for which intended;
       (d) Type of the carrier used;
       (e) Batch number;
       (f) Date of manufacture;
       (g) Expiry date which shall not be less than 6 months from the date
           of manufacture in case of carrier based powdered/granulated
           formulation of Rhizobium, Azotobacter, Azospirillum and PSB
           biofertilisers and liquid based Rhizobium biofertiliser, while it
           shall not be less than 12 months from the date of manufacture
           in case of liquid based Azotobacter, Azospirillum and PSB
           biofertilisers;
       (h) Net mass in kg/gram and area meant for;
       (i) Storage instruction worded as under;
           “STORE IN COOL PLACE AWAY FROM DIRECT SUN LIGHT
           AND HEAT”
(j) Any other information required under the standards of weights and Measure (Packaged Commodities) Rule.1977.

4.3 Items (c),(f) and (g) shall be printed on a coloured ink background.

4.4 Direction for use of biofertiliser shall be printed briefly on the packets as given below.

“The contents of the packet are sufficient enough for seed treatment on to the given area to be broadcasted or given seedling for root dipping depending on the specified crops as denoted on the packet. Mix the inoculants with seeds gently with the minimum amount of water taking care to avoid damage to seed coat. Dry the inoculated seed under shade over clean surface gunny bag and sow them immediately.

Use only for the crops mentioned. Use before the expiry date and do not expose to direct sunlight or heat.

Biofertiliser is not a chemical fertilizer hence do not mix inoculated seeds or inoculants with agro-chemicals.”

4.5 Storage

Inoculants shall be stored by the manufacturer in a cool and dry place away from direct heat preferably at temperature of 20°C. It shall also be the duty of the manufacturer to instruct the retailers and in turn, the users about the precaution to be taken during storage.
Schedule –IV  
[See clause 2(h) and (q)]  
Part – A  
Specifications of Organic Fertilizers  

1. City compost:  

<table>
<thead>
<tr>
<th>(i)</th>
<th>Moisture, per cent by weight</th>
<th>15.0-25.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>(ii)</td>
<td>Colour</td>
<td>Dark brown to black</td>
</tr>
<tr>
<td>(iii)</td>
<td>Odour</td>
<td>Absence of foul odour</td>
</tr>
<tr>
<td>(iv)</td>
<td>Particle size</td>
<td>Minimum 90% material should pass through 4.0 mm IS sieve</td>
</tr>
<tr>
<td>(v)</td>
<td>Bulk density (g/cm$^3$)</td>
<td>&lt;1.0</td>
</tr>
<tr>
<td>(vi)</td>
<td>Total organic carbon, per cent by weight, minimum</td>
<td>12.0</td>
</tr>
<tr>
<td>(vii)</td>
<td>Total Nitrogen (as N), per cent by weight, minimum</td>
<td>0.8</td>
</tr>
<tr>
<td>(viii)</td>
<td>Total Phosphates (as P$_2$O$_5$), per cent by weight, minimum</td>
<td>0.4</td>
</tr>
<tr>
<td>(ix)</td>
<td>Total Potash (as K$_2$O), per cent by weight, minimum</td>
<td>0.4</td>
</tr>
<tr>
<td>(x)</td>
<td>C:N ratio</td>
<td>&lt;20</td>
</tr>
<tr>
<td>(xi)</td>
<td>pH</td>
<td>6.5 - 7.5</td>
</tr>
<tr>
<td>(xii)</td>
<td>Conductivity (as dsm$^{-1}$), not more than</td>
<td>4.0</td>
</tr>
<tr>
<td>(xiii)</td>
<td>Pathogens</td>
<td>Nil</td>
</tr>
<tr>
<td>(xiv)</td>
<td>Heavy metal content, (as mg/Kg), maximum</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Arsenic as (As$_2$O$_3$)</td>
<td>10.00</td>
</tr>
<tr>
<td></td>
<td>Cadmium (as Cd)</td>
<td>5.00</td>
</tr>
<tr>
<td></td>
<td>Chromium (as Cr)</td>
<td>50.00</td>
</tr>
<tr>
<td></td>
<td>Copper (as Cu)</td>
<td>300.00</td>
</tr>
<tr>
<td></td>
<td>Mercury (as Hg)</td>
<td>0.15</td>
</tr>
<tr>
<td></td>
<td>Nickel (as Ni)</td>
<td>50.00</td>
</tr>
<tr>
<td></td>
<td>Lead (as Pb)</td>
<td>100.00</td>
</tr>
<tr>
<td></td>
<td>Zinc (as Zn)</td>
<td>1000.00</td>
</tr>
</tbody>
</table>
### 2. Vermicompost

<table>
<thead>
<tr>
<th>(i)</th>
<th>Moisture, per cent by weight</th>
<th>15.0-25.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>(ii)</td>
<td>Colour</td>
<td>Dark brown to black</td>
</tr>
<tr>
<td>(iii)</td>
<td>Odour</td>
<td>Absence of foul odour</td>
</tr>
<tr>
<td>(iv)</td>
<td>Particle size</td>
<td>Minimum 90% material should pass through 4.0 mm IS sieve</td>
</tr>
<tr>
<td>(v)</td>
<td>Bulk density (g/cm$^3$)</td>
<td>0.7 - 0.9</td>
</tr>
<tr>
<td>(vi)</td>
<td>Total organic carbon, per cent by weight, minimum</td>
<td>18.0</td>
</tr>
<tr>
<td>(vii)</td>
<td>Total Nitrogen (as N), per cent by weight, minimum</td>
<td>1.0</td>
</tr>
<tr>
<td>(viii)</td>
<td>Total Phosphate (as P$_2$O$_5$), per cent by weight, minimum</td>
<td>0.8</td>
</tr>
<tr>
<td>(ix)</td>
<td>Total Potassium (as K$_2$O), per cent by weight, minimum</td>
<td>0.8</td>
</tr>
<tr>
<td>(x)</td>
<td>Heavy metal content, (as mg/Kg), maximum</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cadmium (as Cd)</td>
<td>5.0</td>
</tr>
<tr>
<td></td>
<td>Chromium (as Cr)</td>
<td>50.00</td>
</tr>
<tr>
<td></td>
<td>Nickel (as Ni)</td>
<td>50.00</td>
</tr>
<tr>
<td></td>
<td>Lead (as Pb)</td>
<td>100.00</td>
</tr>
</tbody>
</table>

### 3. Phosphate Rich Organic Manure (PROM)

<table>
<thead>
<tr>
<th>(i)</th>
<th>Moisture per cent by weight, maximum</th>
<th>15.0-25.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>(ii)</td>
<td>Particle size-minimum 90% material should pass through 4.0 mm IS sieve</td>
<td></td>
</tr>
<tr>
<td>(iii)</td>
<td>Bulk density (g/cm$^3$)</td>
<td>1.646</td>
</tr>
<tr>
<td>(iv)</td>
<td>Total organic carbon per cent by weight, minimum</td>
<td>7.87</td>
</tr>
<tr>
<td>(v)</td>
<td>Total nitrogen (as N) per cent by weight, minimum</td>
<td>0.42</td>
</tr>
<tr>
<td>(vi)</td>
<td>Total phosphates (as P$_2$O$_5$) per cent by weight, minimum</td>
<td>10.42</td>
</tr>
<tr>
<td>(vii)</td>
<td>Total potash (as K$_2$O) per cent by weight, minimum</td>
<td>-</td>
</tr>
<tr>
<td>(viii)</td>
<td>C:N ratio</td>
<td>18:73:1</td>
</tr>
<tr>
<td>(ix)</td>
<td>pH( 1:5 solution) maximum</td>
<td>6.72</td>
</tr>
<tr>
<td>(x)</td>
<td>Conductivity (as dSm$^{-1}$) not more than</td>
<td>8.27</td>
</tr>
<tr>
<td>(xi)</td>
<td>Heavy metal content (as mg/kg), maximum</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Arsenic (as As$_2$O$_3$)</td>
<td>10.0</td>
</tr>
<tr>
<td></td>
<td>Cadmium (as Cd)</td>
<td>5.0</td>
</tr>
<tr>
<td></td>
<td>Chromium (as Cr)</td>
<td>50.0</td>
</tr>
<tr>
<td></td>
<td>Copper (as Cu)</td>
<td>300.0</td>
</tr>
<tr>
<td></td>
<td>Mercury (as Hg)</td>
<td>0.15</td>
</tr>
<tr>
<td></td>
<td>Nickel (as Ni)</td>
<td>50.0</td>
</tr>
<tr>
<td></td>
<td>Lead (as Pb)</td>
<td>100.0</td>
</tr>
<tr>
<td></td>
<td>Zinc (as Zn)</td>
<td>1000.0</td>
</tr>
</tbody>
</table>
4. Organic Manure

<table>
<thead>
<tr>
<th>(i)</th>
<th>Moisture per cent by weight, maximum</th>
<th>25.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>(ii)</td>
<td>Particle size</td>
<td></td>
</tr>
<tr>
<td>(iii)</td>
<td>Bulk density (g/cm²)</td>
<td>&lt;1.0</td>
</tr>
<tr>
<td>(iv)</td>
<td>Total organic carbon per cent by weight, minimum</td>
<td>14.0</td>
</tr>
<tr>
<td>(v)</td>
<td>Total nitrogen (as N) per cent by weight, minimum</td>
<td>0.5</td>
</tr>
<tr>
<td>(vi)</td>
<td>Total phosphates (as P₂O₅) per cent by weight, minimum</td>
<td>0.5</td>
</tr>
<tr>
<td>(vii)</td>
<td>Total potash (as K₂O) per cent by weight, minimum</td>
<td>0.5</td>
</tr>
<tr>
<td>(viii)</td>
<td>NPK nutrients – Total N, P₂O₅ and K₂O nutrient should not be less than 3%</td>
<td></td>
</tr>
<tr>
<td>(ix)</td>
<td>C:N ratio</td>
<td>&lt;20</td>
</tr>
<tr>
<td>(x)</td>
<td>pH</td>
<td>6.5-7.5</td>
</tr>
<tr>
<td>(xi)</td>
<td>Conductivity (as dSm⁻¹) not more than</td>
<td>4.0</td>
</tr>
<tr>
<td>(xii)</td>
<td>Pathogen</td>
<td>Nil</td>
</tr>
<tr>
<td>(xiii)</td>
<td>Heavy metal content, (as mg./kg), maximum</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Arsenic (as As₂O₃)</td>
<td>10.0</td>
</tr>
<tr>
<td></td>
<td>Cadmium (as Cd)</td>
<td>5.0</td>
</tr>
<tr>
<td></td>
<td>Chromium (as Cr)</td>
<td>50.0</td>
</tr>
<tr>
<td></td>
<td>Copper (as Cu)</td>
<td>300.0</td>
</tr>
<tr>
<td></td>
<td>Mercury (as Hg)</td>
<td>0.15</td>
</tr>
<tr>
<td></td>
<td>Nickel (as Ni)</td>
<td>50.0</td>
</tr>
<tr>
<td></td>
<td>Zinc (as Zn)</td>
<td>1000.0</td>
</tr>
</tbody>
</table>

**Note**: The source of organic manure is any of the plant biomass/animal biomass/animal Excreta)

**Part –B**

**TOLERANCE LIMIT OF ORGANIC FERTILISER**

A sum total of nitrogen, phosphorus and potassium nutrients shall not be less than 1.5% in City Compost and shall be not less than 2.5% in case of vermicompost.

**Part- C**

**PROCEDURE FOR DRAWL OF SAMPLE OF ORGANIC FERTILISER**

(As per methodology as mentioned under schedule-ii, part-A of FCO,1985)
The Inspector shall draw any sample of Organic fertilizer in accordance with the procedure of drawl mentioned under schedule-ii, Part-A.
1. **General requirements of sampling**

In drawing samples, the following measures and precautions should be observed:

(a) Samples shall not be taken at a place exposed to rain/sun.
(b) The sampling instruments shall be clean and dry when used.
(c) The material being sampled, the sampling instruments and the bags of samples should be free from any adventitious contaminations.
(d) To draw a representative sample, the contents of each bag selected for sampling should be mixed as thoroughly as possible by suitable means.
(e) The sample should be kept in suitable, clean dry and airtight glass or screwed hard polythene bottle of about 400gm capacity or in a thick-gauged polythene bag. This should be put in a cloth bag, which may be sealed with the Inspector’s seal after putting inside the detailed description as specified in Form ‘P’. Identifiable details may also be put on the cloth bag like sample No./Code No. or any other details which enables its identification.
(f) Each sample bag should be sealed air tight after filling and marked with details of sample and type of fertilizer and the name of Inspector who has collected sample.

2. **Sampling from bagged material**

(i) **Scale of sampling**

(a) **Lot (for manufacturers/importers)** - All bags in a single consignment of the material of the same grade and type drawn from a single batch of the manufacturer/importer shall constitute a lot. If a consignment is declared to consist of different batches of manufacturer/importer, all the bags of each batch shall constitute a separate lot. In the case of a consignment drawn from a continuous process, 2000 bags (or 100 tonnes) of the material shall constitute a lot.

(b) **Lot (for dealer)** - The lot is an identifiable quantity of same grade and type of fertilizer stored at an identifiable place subject to a maximum limit of 100 tones. The inspector based on visible appearance of bags, their packing and storage conditions shall identify the lot. The stock of less than 100 tones with a dealer may also constitute one or more lots, if the material (fertiliser) of different sources and brand is available in such quantities.

(c) **Selection of bags for sampling** - The number of bags to be chosen from a lot shall depend upon the size of the lot as given below:

<table>
<thead>
<tr>
<th>Lot size (No. of bags) (N)</th>
<th>No. of bags to be selected for sampling (n)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Upto 10</td>
<td>1</td>
</tr>
<tr>
<td>11-100</td>
<td>2</td>
</tr>
</tbody>
</table>
All the bags of a lot should be arranged in a systematic manner. Start counting from any bag randomly; go on counting as 1, 2, 3, .......... up to R and so on. R being equal to the integral of N/n. Thus every 4th bag counted shall be withdrawn and all bags shall constitute the sample bags from where the sample is to be drawn for preparing a composite sample.

(ii) **Sampling from big godowns/high stackings** - If the procedure given in Para 2 (i)(c) is not possible to be adopted, the sample should be drawn from the randomly selected fertilizer bags from different layers, from top and from all open sides in a zig zag fashion.

(iii) **Sampling from small godowns** - All the fertilizer bags of the same grade and type of each manufacturer though received on different dates shall be segregated and properly stacked. All bags of same grade and type of fertilizer manufactured by a particular manufacturing unit may be considered as one lot based on their physical conditions and the sample shall be drawn as per procedure laid down in para 2 (i) (c) and 4.

(iv) **Sampling from damaged stocks**
   (a) In case of torn or lumpy bags, damaged fertilizer bags or sweepings, the stock should be arranged according to identifiable lots. From each lot the number of bags shall be selected as per procedure 2 (i) (c). If the bags allow the use of sampling probe conveniently, the samples should be drawn from sampling probe.
   (b) In case it is not possible to use the sampling probe, the bags may be opened and fertilizer material mixed together uniformly by hammering the big lumps or putting pressure, if required, and then samples drawn by using suitable sample device.

3. **Sampling probe**
   (i) An appropriate sampling instrument to be used by the Inspector for collection of a representative sample is called sampling probe. The probe may comprise of a slotted single tube with solid cone tip made of stainless steel or brass. The length of the probe may be approximately 60 to 65 cms and the diameter of the tube may be approximately 1.5 cm and the slot width may be 1.2 to 1.3 cms. The

| 101-200 | 3 |
| 201-400 | 4 |
| 401-600 | 5 |
| 601-800 | 6 |
| 801-1000| 7 |
| 1001-1300| 8 |
| 1301-1600| 9 |
| 1601-2000| 10 |
probe may be used if the physical condition of the fertilizers and the packing material permits its use.

(ii) In case of high-density polythene packing and also when the fertilizer material is not in free flowing condition, the use of sampling probe may not be possible. In such case, selected bags for drawing samples may be opened and the fertilizers may be taken out of the bags and spread on a clean surface and samples drawn with the help of a suitable sampling device, which may be made of stainless steel or brass cup.

4. Drawal of samples from bags
(i) Drawal of sample and preparation of composite samples.
Draw, with an appropriate sampling instrument, (sampling probe) small portions of the material from the selected bags as per procedure in para 2(i) (b), 2(ii), 2(iii) and 2(iv) (a). The sampling probe shall be inserted in the bag from one corner to another diagonally and when filled with fertilizer, the probe is withdrawn and fertilizer is emptied in a container/ or on polythene sheet/ or on a clean hard surface and made into one composite sample.

(ii) If the bags do not permit the use of sampling probe, empty the contents of the bags on a level, clean and hard surface and draw a composite sample by the process of quartering as described under para 3(ii) or 5.

4A. Weight of one Sample
One sample of organic fertilizer shall have approximate 400gms weight.

5. Preparation of composite sample
If the composite sample collected from the different selected bags is larger than required weight, its size shall be reduced by method of quartering as detailed below:-
Spread the composite sample on a level, clean, hard surface, flatten it out and divide it into four equal parts. Remove any diagonally opposite parts. Mix the two remaining parts together to form a cone, flatten out the cone and repeat the operation of quartering till a composite sample of required weight is obtained.

6. Preparation of test sample and reference sample
(i) The composite sample obtained above shall be spread out on a clean, hard surface and divided into three approximately equal portions each of the weight as specified in Para 4A. Each of these samples shall constitute the test sample.

(ii) Each test sample shall be immediately transferred to a suitable container as defined under para 1(e). The slip with detailed
description may be put inside the sample bag. Each bag shall also be properly labeled as mentioned in para 1(f).

(iii) Each test sample container shall then be sealed with seals of the inspector, if possible, seal of the manufacturer/ importer/ dealer or purchaser as the case may be also be affixed.

(iv) Out of the three samples collected, one sample so sealed shall be sent to the Incharge of the laboratory notified by the state Government under clause 29 or National/Regional Centre of Organic Farming, Ghaziabad, Bangalore, Bhubaneshwar, Hisar, Imphal, Jabalpur or Nagpur for analysis. Another sample shall be given to the manufacturer or importer or dealer or the purchaser, as the case may be. The third sample shall be sent by the Inspector to his next higher authority for keeping in safe custody. Any of the latter two samples may be sent for referee analysis as provided for under sub-clause (2) of clause 29B.

Part- D

METHODS OF ANALYSIS OF ORGANIC FERTILISERS

1. Estimation of pH

- Make 25 g of compost into a suspension in 50 ml of distilled water and shake on a rotary shaker for 2 hours.
- Filter through Whatman No. 1 or equivalent filter paper under vacuum using a Buchner funnel.
- Determine pH of the filtrate by pH meter.

2. Estimation of moisture

Weigh to the nearest mg about 5 gm of the prepared sample in a weighed clean, dry Petri Dish. Heat in an oven for about 5 hours at 65°C ±1°C to constant weight. Cool in a dessicator and weigh. Report percentage loss in weight as moisture content.

Calculation:

\[
\text{Moisture percent by weight} = \frac{100(B-C)}{B-A}
\]

A = Weight of the Petri Dish
B = Weight of the Petri dish plus material before drying
C = Weight of the Petri Dish plus material after drying

3. Estimation of Bulk density

Requirements
Method
- Weigh a dry 100ml cylinder (W1gm)
- Cylinder is filled with the sample up to the 100 ml mark. Note the volume (V1 ml)
- Weigh the cylinder along with the sample (W2 gm)
- Tap the cylinder for two minutes.
- Measure the compact volume (V2 ml)

Calculation
Bulk density = \( \frac{\text{Weight of the sample taken (W2-W1)}}{\text{Volume (V1-V2)}} \)

4. Estimation of conductivity
Requirements
- 250 ml flask, Funnel [OD-75mm]
- 100ml Beaker, Analytic Balance
- Potassium Chloride [AR grade] , Filter paper
- Conductivity meter [With temperature compensation system]

Method
- Pass fresh sample of organic fertilizer through a 2-4mm sieve.
- Take 20 gm of the sample and add 100 ml of distilled water to it to give a ratio of 1:5
- Stir for about an hour at regular intervals.
- Calibrate the conductivity meter by using 0.01 M potassium chloride solution.
- Measure the conductivity of the unfiltered organic fertilizer suspension.

Calculation
Express the results as millimho’s or dsm\(^{-1}\) at 25°C specifying the dilution of the organic fertilizer suspension viz, 1:5 organic fertilizer suspensions.

5. Estimation of Organic Carbon
Apparatus
(i) Silica/Platinum crucible 25 g cap.
(ii) Muffle Furnace

Procedure
Accurately weigh 10 gm of sample dried in oven at 105°C for 6 hrs, in a pre weighed crucible and ignite the material in a Muffle furnace at 650 – 700°C
for 6-8 hrs. Cool to room temperature and keep in Desiccator for 12 hrs. Weigh the contents with crucible

**Calculation**
Calculate the total organic carbon by the following formulae:-

\[
\text{Total Organic matter \%} = \frac{\text{Initial wt} - \text{final wt}}{\text{wt. of sample taken}} \times 100
\]

\[
\text{Total C\%} = \frac{\text{Total organic matter \%}}{1.724}
\]

6. **Estimation of total Nitrogen**
As mentioned under Schedule – II, Part-B, 3 (v) of FCO,1985.

**Apparatus**
1. Suitable Kjeldahl assembly consisting of 500-800 ml round bottom, digestion flask and Kjeldahl distillation assembly consisting of 500-800 ml distillation flask, splash head tube and condenser, all with appropriate glass joints. The length of the condenser's delivery tube should be long enough to keep immersed in a flask for ammonia absorption.
2. Kjeldahl digestion unit with heating control, suitable for 500-800 ml flasks.

**Reagents**
- Sulphuric acid – 93-98% H\(_2\)SO\(_4\), N-free
- Salicylic acid, reagent grade, N-free
- Sodium thiosulphate (Na\(_2\)S\(_2\)O\(_3\) 5H\(_2\)O), reagent grade
- Zinc dust- impalpable powder
- Copper sulphate
- Potassium or sodium sulphate
- 45% NaOH solution. Dissolve 450 gm of Sodium hydroxide pellets in distilled water and make up the volume to 1000ml
- Methyl red indicator – Dissolve 1gm methyl red in 200 ml alcohol
- Hydrochloric or sulphuric acid standard solution – 0.1N or as per requirement
- Sodium hydroxide standard solution 0.1N or as per requirement.

**Procedure**
1. Place weighed sample (0.7-2.2gm) in digestion flask.

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2. Add 40 ml H$_2$SO$_4$ containing 2 grams salicylic acid. Shake until thoroughly mixed and let stand, with occasional shaking, 30 minutes or more.

3. Then add (i) 5 grams Na$_2$S$_2$O$_3$.5H$_2$O or (ii) 2 grams zinc dust (as impalpable powder not granulated zinc or filing).

4. Shake the flask and let it stand for five minutes then heat over low flame until frothing ceases.

5. Turn off heat, add 0.7 grams copper sulphate, 15 gm powdered K$_2$SO$_4$ (or anhydrous Na$_2$SO$_4$), and boil briskly until solution clears, continue boiling for another at least 2 hours.

6. Remove from burner and cool, add 200 ml of water and swirl the flask to dissolve all the contents.

7. Transfer to 500 ml volumetric flask, giving several washings with water to the digestion flask. Make up the volume to 500 ml.

8. Take 25 ml aliquot in the distillation flask, add 300 ml water and a pinch of zinc dust.

9. Take 20 ml of standard acid solution in the receiving conical flask, add 4 drops of methyl red indicator and keep the flask at the lower end of the condenser in such a way that the lower tip of the condenser is fully immersed in acid solution.

10. Add 30 ml of 45% NaOH to the distillation flask, gently so that the contents do not mix.

11. Immediately connect the flask to distillation assembly and swirl to mix the contents. Heat until all the ammonia is distilled (at least 150 ml distillate).

12. Remove from receiving flask. Rinse outlet tube into receiving flask with a small amount of distilled water.

13. Titrate the contents in the receiver conical flask with standard NaOH solution.

14. Determine blank on reagents using same quantity of standard acid in receiving conical flask.

**Calculation**

\[
\text{Nitrogen } \% \text{ by weight} = \frac{1.401(V_1N_1-V_2N_2)-(V_3N_1-V_4N_2) \times df}{W}
\]

where

- \(V_1\) = Volume in ml of standard acid taken in receiver flask for sample
- \(V_2\) = Volume in ml of standard NaOH used in titrating standard acid in receiver flask after distillation of test sample
- \(V_3\) = Volume in ml of standard acid taken in receiver flask for blank
- \(V_4\) = Volume in ml of standard NaOH used in titrating standard acid in receiver flask after distillation in blank
- \(N_1\) = Normality of standard acid

\(W\) = Weight of sample taken for analysis.
N₂ = Normality of standard NaOH
W = Weight in gm of sample taken
df = Dilution factor of sample

7. Estimation of C: N Ratio
Method
Calculate the C:N ratio by dividing the organic carbon value with the total nitrogen value.

8. Estimation of phosphate
Preparation of sample - Accurately weigh 10 gm oven dried sample in 50 g cap. silica crucible and ignite it to 650° – 700°C for 6-8 hrs to obtain ash. Cool and keep in a Dessicator. Transfer the contents to a 100 ml beaker. Add 30 ml 25% HCl. Wash the crucible with 10 ml 25% HCl twice and transfer the contents to Beaker. Heat over hot plate for 10-15 min. Keep for 4 hrs. Filter through Whatman No.1 filter paper. Wash with distilled water 4-5 times (till acid free). Make up the volume of filtrate to 250 ml in a volumetric flask. Estimate total P by gravimetric quinoline molybdate method as described under Schedule – II, Part B, 4(ii) of FCO 1985.

Gravimetric quinoline molybdate method for determination of total phosphorus
Reagents
1. Citric molybdic acid reagent – Dissolve 54 gm, 100% molybdic anhydride (Mo₃) and 12 gm NaOH with stirring in 400 ml hot water and cool. Dissolve 60 gm citric acid in mixture of 140 ml HCl and 300 ml water and cool. Gradually add molybdic solution to citric acid solution with stirring. Cool, filter and dilute to 1 lit. (solution may be green or blue colour depending on exposure to light) If necessary add 0.5% KBrO₃ solution drop by drop until green colour becomes pale. Store in dark in polyethylene bottle.
2. Quinoline solution – Dissolve 50 ml synthetic quinoline with stirring in mixture of 60 ml HCl and 300 ml water. Cool dilute to 1 lit and filter. Store in polyethylene bottle.
3. Quimociac reagent – Dissolve 70 gm of sodium molybdate dehydrate in 150 ml water. Dissolve 60 gm citric acid in mixture of 85 ml HNO₃ and 150 ml water and cool. Gradually add molybdate solution to citric acid-nitric acid mixture with stirring. Dissolve 5 ml synthetic quinoline in mixture of 35 ml HNO₃ and 100 ml water. Gradually add this solution to molybdate –citric-nitric acid solution mix and let it stand for 24 hr. Filter, add 280 ml acetone, dilute to 1 lit with water and mix well. Store in polyethylene bottle.

Procedure
1. Digest 1 gm sample as described above and dilute to 200 ml.
2. In 500 ml Erlenmeyer flask pipette aliquot containing not more than 25mg \(P_2O_5\) dilute to approximately 100 ml with water. Proceed with one of the following method.

a. Add 30 ml citric-molybdic acid reagent and boil gently for 3 min (solution must be precipitate free at this stage). Remove from heat and swirl carefully. Immediately add from burette 10 ml quinoline solution with continuous swirling (add first 3-4 ml drop wise and remainder in steady stream) or

b. Add 50 ml quimaciac reagent, cover with watch glass place on hot plate in well ventilated hood and boil for 1 min.

After treatment with a or b cool to room temperature, swirl carefully 3-4 time during cooling, filter through sintered glass Gooch crucible Grade 4 (30 ml capacity), previously dried at 250\(^\circ\)C and weighed, and wash 5 times with 25 ml portion of water. Dry crucible and contents for 30 min. at 250\(^\circ\)C. Cool in dessicator to constant weight as \((C_9H_7N)_3H_3PO_4\cdot12MoO_3\). Subtract blank weight. Multiply by 0.03207 to obtain weight of \(P_2O_5\). Report as percent \(P_2O_5\).

9. Estimation of Potassium

**Flame photometry method:** Total Potassium are usually determined by dry ashing at 650-700 Degree Centigrade and dissolving in concentrated hydrochloric acid.

**Reagent and Standard curve**

(1) Potassium chloride standard solution: Make a stock solution of 1000 ppm K by dissolving 1.909 g. of AR grade potassium chloride (dried at 60\(^\circ\)C for 1 h) in distilled water 1; and diluting up to 1 litre. Prepare 100 ppm standard by diluting 100 ml of 1000 ppm stock solution to 1 litre with extracting solution.

(2) Standard curve: Pipette 0.5, 10, 15 and 20 ml of 100 ppm solution into 100 ml volumetric flasks and make up the volume upto the mark. The solution contain 0, 5, 15 & 20 ppm K respectively.

**Procedure:**

- Take 5g sample in a porcelain crucible and ignite the material to ash at 650-700 C in a muffle furnace.
- Cool it and dissolve in 5 ml concentrated hydrochloric acid, transfer in a 250 ml beaker with several washing of distilled water and heat it.
- Again transfer it to a 100 ml volumetric flask and make up the volume.
- Filter the solution and dilute the filtrate with distilled water so that the concentration of K in the working solution remains in the range of 0 to 20 ppm, if required.
- Determine K by flame photometer using the K- filter after necessary setting and calibration of the instrument.
• Read similarly the different concentration of K of the standard solution in flame photometer and prepare the standard curve by plotting the reading against the different concentration of the K.
• Calculation: Potash (K) % by weight = R X 20 X diluting factor, where R = ppm of K in the sample solution (obtained by extra plotting from standard curve).

10. Estimation of Cadmium, Copper, Chromium, Lead, Nickel and Zinc

Material Required
1. Triacid mixture: Mix 10 parts of HNO₃ (Nitric acid), 1 part of H₂SO₄ (Sulphuric Acid) and 4 parts of HClO₃ (Perchloric Acid)
2. Conical flask, 250ml
3. Hot plate
4. Whatman filter paper No.42
5. Atomic Absorption Spectrophotometer (AAS)

Processing of sample
Take 5.0 g or suitable quantity of oven dried (105°C) sample thoroughly ground and sieved through 0.2 mm sieve in a conical flask. Add 30 ml triacid mixture, cover it with a small glass funnel for refluxing. Digest the sample at 200°C on a hot plate till the volume is significantly reduced with a whitish residue. After cooling, filter the sample with Whatman No. 42 filter paper and make up to 100 ml in a volumetric flask.

Preparation of working standards
Cadmium - As mentioned under Schedule – II, Part B, 8(x) of FCO (1985)
Copper - As mentioned under Schedule – II, Part B, 8(iv) of FCO (1985)
Chromium - Dilute 1, 2, 3 and 4 ml of standard 100 ppm Chromium standard solution with doubled distilled water in volumetric flasks and make up the volume to 100 ml to obtain standards having concentrations of 1, 2, 3, 4 ppm
Lead - As mentioned under Schedule – II, Part B, 8(v) of FCO (1985)
Nickel - Dilute 1, 2, 3 and 4 ml of standard 100 ppm Nickel standard solution with doubled distilled water in volumetric flasks and make up the volume to 100ml to obtain standards having concentrations of 1, 2, 3, 4 ppm
Zinc - As mentioned under Schedule – II, Part B, 8(ii ) of FCO (1985)
(Alternatively dilute 1, 2, 3 and 4 ml of 100ppm standard stock solution of respective element with doubled distilled water in volumetric flasks and make up the volume to 100ml to obtain standards having concentrations of 1, 2, 3, 4 ppm)
Measurement of Result
Estimate the metal concentrations of Cd, Cu, Cr, Pb, Ni, Zn by flaming the standard solution and samples using atomic absorption spectrophotometer (AAS) as per the method given for instrument at recommended wavelength for each element. Run a blank following the same procedure.

Expression of Result
Express the metal concentration as mg/kg on oven dry weight basis in 3 decimal units.
(Reference: Manual for Analysis of Municipal Solid Waste (compost): Central Pollution Control Board).

11. Estimation of Mercury
Reagents:
(a) Concentrated Nitric acid (HNO₃)
(b) Concentrated Sulphuric acid (H₂SO₄)
(c) Potassium persulphate (5% solution): Dissolve 50g of K₂S₂O₈ in 1 litre of distilled water.
(d) Potassium permagnate (5% solution): Dissolve 50g of KMnO₄ in 1 litre of distilled water.
(e) Hydroxylamine sodium chloride solution: Dissolve 120 g of Hydroxylamine salt and 120 g of sodium chloride (NaCl) in 1 litre distilled water.
(f) Stannous chloride (20%): Dissolve 20 g of SnCl₂ in 100 ml distilled water.

Materials required
(a) Water bath
(b) Flameless atomic absorption spectrophotometer or cold vapour mercury analyzer.
(c) BOD bottle, 300 ml

Processing of sample:
(a) Take 5 g (finely ground but not dried) sample in an oven at a temperature of 105°C for 8 hours for moisture estimation.
(b) Take another 5 g sample (finely ground but not dried) in a BOD bottle, add to it 2.5 ml of conc. HNO₃, 5ml of cone. H₂SO₄ and 15 ml of 5% KMnO₄.
(c) After 15 minutes add 8 ml of 5% K₂S₂O₈.
(d) Close the bottle with the lid and digest it on a water bath at 95°C for 2 hours.
(e) After cooling to room temperature add 5 ml hydroxylamine sodium chloride soln.
**Measurement:**
Reduction of the digested sample is brought out with 5 ml of 20% SnCl₂ immediately before taking the reading, using a cold vapour mercury analyzer.

**Expression of results:**
Express the mercury concentration as mg/Kg on oven dry weight basis in 3 decimal units.
(Reference: Manual for Analysis of Municipal Solid Waste (compost); Central Pollution Control Board).

**12. Estimation of Arsenic**

**Processing of sample** – Suspend 10 gm finely ground sample in 30 ml aqua regia (HNO₃ + HCl in a ratio of 1:3) in a beaker. Keep on hot plate till moist black residue is obtained (do not dry). Add 5 ml aqua regia and allow to dry on a hot plate till residue is moist. Dissolve the residue in 30 ml conc. HCl and filter through Whatman No.1 filter paper in 100 ml volumetric flask. Wash filter paper 3-4 times with double distilled water. Make up the volume to 100 ml. Take 1 ml of this solution in 100 ml volumetric flask, add 5 ml conc. HCl and 2 gm KI and make up the volume to 100 ml. Prepare standards having concentration of 0.05, 0.1 and 0.2 ppm by diluting 0.05, 0.1 and 0.2 ml, respectively of standard Arsenic solution with double distilled water in volumetric flask and make up the volume to 100 ml.

**Measurement** – Estimate Arsenic using vapour generation assembly attached to Atomic Absorption Spectrophotometer as per the procedure given for the instrument.

**13. Pathogenicity Test**

**Apparatus**
1. Samples of Compost
2. Lactose Broth of Single and Double Strength
3. Culture Tubes
4. Durham Tubes
5. Bunsen Burner
6. Sterile Pipettes
7. Incubator, Autoclaves,
8. Petri-Plates
9. Inoculation Loops

**Preparation of Culture Media**

**A. For Presumptive Test**

1. **Lactose Broth**
   - Beef Extract : 6.0 g
   - Peptone : 10.0 g
   - Lactose : 10.0 g
   - D.W. : 1000 ml
B. For Confirmative Test

1. Eosine Methylene Blue Agar Media (EMB Media)

Peptone : 10.0 g  
Lactose : 5.0 g  
Sucrose : 5.0 g  
K₂HPO₄ : 2.0 g  
Eosine Y : 0.4 g  
Methylene Blue : 0.06 g  
Agar : 15.0 g  
D.W. : 1000 ml

C. For Completed Test

1. Nutrient Agar

Beef Extract : 3.0 g  
Peptone : 5.0 g

Procedures

A. Presumptive Test

1. Prepare 12 tubes of lactose broth for each sample and close the tube with cotton plugs/caps and autoclave at 121°C for 20 min.
2. Fill Durham tubes with sterilized distilled water and keep in beaker and autoclave at 121°C for 20 min.
3. Suspend 30 g of compost sample in 270 ml of sterile distilled water and serially dilute up to 10⁻⁴ dilution as per Schedule III, Part D, serial number 3 of FCO (1985)
4. Suspend 1 ml suspension from 10⁻¹ to 10⁻⁴ in 3 tubes for each dilution
5. Insert distilled water filled Durham tube in inverted position in each tube and close the tube again
6. Inoculate tubes at 36°C for 24h in incubator

Result

- Production of gas within 24h - Confirms the presence of coliforms in the sample
- Production of gas within 48h - Doubtful Test.
- No Gas Production - Negative Test

B. Confirmative Test

Confirmative test is for differentiating the coliforms from non-coliforms as well as Gram negative and Gram positive bacteria. In this test, the EMB agar plates are inoculated with sample from positive tubes producing gas. Emergence of small colonies with dark centres confirms the presence of Gram negative, lactose fermenting coliform bacteria. Sometimes some of the non-coliforms also produce gas, therefore, this test is necessary.
1. Prepare EMB agar plates with the composition as per the method given above
2. Inoculate plates with the help of inoculation loop with streaking of samples showing positive/doubtful tests in the presumptive test
3. Incubate plates at 30±1°C for 12 h in incubator
4. Dark centred or nucleated colonies appear which may differentiate between *E. coli* and *E. aerogenes* based on size of colonies and metallic sheen

**Result**

*E. coli* colonies on this medium are small with metallic sheen, where as *E. aerogenes* colonies are usually large and lack the sheen.

**C. Completed Test**

This test is required for further confirmation.

**Procedure**

1. Pick up a single colony from EMB agar plate
2. Inoculate it into lactose broth and streak on a nutrient agar slant
3. Incubate the slants
4. Perform Gram reaction after attaining the growth

**Result**

Gram-negative nature of bacteria is indicative of a positive completed test.”
**Schedule V**
[see clause 2(h) and (q)]

**Part – A**

Specifications of Non-edible De-oiled cake fertilizer

1. Castor de-oiled cake

<table>
<thead>
<tr>
<th>Sl. No.</th>
<th>Parameter</th>
<th>Requirement</th>
</tr>
</thead>
<tbody>
<tr>
<td>i</td>
<td>Moisture percent by weight</td>
<td>maximum 12</td>
</tr>
<tr>
<td>ii</td>
<td>Colour</td>
<td>Brown to black</td>
</tr>
<tr>
<td>iii</td>
<td>Odour</td>
<td>Typical oily odour specific to the oil of that seed and no foul odour,</td>
</tr>
<tr>
<td>iv</td>
<td>Ash content % by weight</td>
<td>(maximum) 15.0</td>
</tr>
<tr>
<td>v</td>
<td>Total Organic Carbon, % by weight, minimum</td>
<td>25</td>
</tr>
<tr>
<td>vi</td>
<td>Total Nitrogen (as N) % by weight, Minimum</td>
<td>4.5</td>
</tr>
<tr>
<td>vii</td>
<td>Total Phosphates (as P₂O₅) % by weight Minimum</td>
<td>1.0</td>
</tr>
<tr>
<td>viii</td>
<td>Total Potash (as K₂O) % by weight, Minimum</td>
<td>1.0</td>
</tr>
<tr>
<td>ix</td>
<td>C: N ratio</td>
<td>&lt; 10</td>
</tr>
<tr>
<td>x</td>
<td>pH</td>
<td>6.0-8.0</td>
</tr>
<tr>
<td>xi</td>
<td>Conductivity (as dsm⁻¹) not more than</td>
<td>4.0</td>
</tr>
<tr>
<td>xii</td>
<td>Particle size</td>
<td>Not less than 75% of the material shall pass through 4 mm IS sieve</td>
</tr>
</tbody>
</table>

**Part – B**

Tolerance limit
0.5 Unit for Nitrogen, Phosphorous and Potassium Nutrients combined.

**Part- C**

PROCEDURE FOR DRAWL OF SAMPLE OF ORGANIC FERTILISER

The Inspector shall draw any sample of Non-edible De-oiled cake meal fertilizer in accordance with the procedure of drawl mentioned under schedule-II, part-A of FCO, 1985.

**Part- D**

METHODS OF ANALYSIS FOR NON-EDIBLE DE-OILED CAKE MEAL FERTILIZERS

1. Estimation of pH
   As mentioned in Schedule IV Part D (1) of FCO 1985
2. **Estimation of moisture**  
   As mentioned in Schedule IV Part D (2) of FCO 1985

3. **Estimation of ash Content**  
   **Apparatus**  
   i) Silica/Platinum crucible 25 g cap.  
   ii) Muffle Furnace  
   iii) Desiccator  
   
   Weigh to the nearest mg about 5 gm of oven dried powdered sample in a weighed clean, dry Petri Dish. Incinerate in a muffle furnace for about 6-8 hours at 650-700°C to constant weight. Cool in a desiccator and weigh. Report percentage of ash content obtained.

   **Calculation:**  
   Ash content in percent by weight = \[ \frac{100 \times (C - A)}{B - A} \]  
   
   A = Weight of the empty crucible  
   B = Weight of the empty crucible plus material before ashing  
   C = Weight of the empty crucible plus material after ashing

4. **Estimation of conductivity**  
   As mentioned in Schedule IV Part D (4) of FCO 1985

5. **Estimation of organic carbon**  
   As mentioned in Schedule IV Part D (5) of FCO 1985

6. **Estimation of total nitrogen**  
   **Apparatus**  
   1. Suitable Kjeldahl assembly consisting of 500-800 ml round bottom, digestion flask and Kjeldahl distillation assembly consisting of 500-800 ml distillation flask, splash head tube and condenser, all with appropriate glass joints. The length of the condenser's delivery tube should be long enough to keep immersed in a flask for ammonia absorption.  
   2. Kjeldahl digestion unit with heating control, suitable for 500-800 ml flasks.

   **Reagents**  
   a. Sulphuric acid – 93-98% H₂SO₄, N-free  
   b. Salicylic acid, reagent grade, N-free  
   c. Sodium thiosulphate (Na₂S₂O₃ 5H₂O), reagent grade  
   d. Zinc dust- impalpable powder  
   e. Copper sulphate
f. Potassium or sodium sulphate

g. Selanium powder

h. Red Mercury oxide (HgO)

i. 45% NaOH solution. Dissolve 450 gm of Sodium hydroxide pellets in distilled water and make up the volume to 1000ml

j. Methyl red indicator – Dissolve 1gm methyl red in 200 ml alcohol

k. Hydrochloric or sulphuric acid standard solution – 0.1N or as per requirement

l. Sodium hydroxide standard solution 0.1N or as per requirement.

**Procedure**

1. Place weighed finely powdered sample (0.5-1.0 gm) in digestion flask.

15. Add 1 gm digestion mixture (20 g CuSO₄ + 3 gm selenium dust + 1 gm HgO)

16. Add 50 ml conc H₂SO₄

17. Shake the flask and let it stand for five minutes then heat over low flame until frothing ceases.

18. Turn off heat, add 15 - 20 gm powdered K₂SO₄ (or anhydrous Na₂SO₄), and boil briskly until solution clears.

19. Add 5 gm Sodium thiosulphate continue boiling for another at least 2-3 hours.

20. Remove from burner and cool, add 200 ml of water and swirl the flask to dissolve all the contents.

21. Transfer to 500 ml volumetric flask, giving several washings with water to the digestion flask. Make up the volume to 500 ml.

22. Take 25 ml aliquot in the distillation flask, add 300 ml water and a pinch of zinc dust

23. Take 20 ml of standard acid solution (N/10 H₂SO₄) in the receiving conical flask, add 4 drops of methyl red indicator and keep the flask at the lower end of the condenser in such a way that the lower tip of the condenser is fully immersed in acid solution.

24. Add 40 ml of 45% NaOH to the distillation flask, gently so that the contents do not mix.

25. Immediately connect the flask to distillation assembly and swirl to mix the contents. Heat until all the ammonia is distilled (at least 150 ml distillate).

26. Remove from receiving flask. Rinse outlet tube into receiving flask with a small amount of distilled water.

27. Titrate the contents in the receiver conical flask with standard NaOH solution.

28. Determine blank on reagents using same quantity of standard acid in receiving conical flask.
Calculation

Nitrogen % by weight = \frac{1.401(V_1N_1-V_2N_2)-(V_3N_1-V_4N_2) \times df}{W}

where

\begin{align*}
V_1 &= \text{Volume in ml of standard acid taken in receiver flask for sample} \\
V_2 &= \text{Volume in ml of standard NaOH used in titrating standard acid in receiver flask after distillation of test sample} \\
V_3 &= \text{Volume in ml of standard acid taken in receiver flask for blank} \\
V_4 &= \text{Volume in ml of standard NaOH used in titrating standard acid in receiver flask after distillation in blank} \\
N_1 &= \text{Normality of standard acid} \\
N_2 &= \text{Normality of standard NaOH} \\
W &= \text{Weight in gm of sample taken} \\
df &= \text{Dilution factor of sample}
\end{align*}

7. Estimation of C:N ratio

Method

Calculate the C:N ratio by dividing the organic carbon value with the total nitrogen value

8. Estimation of Phosphate

As mentioned in Schedule IV Part D (8) of FCO 1985

9. Estimation of Potassium

As mentioned in Schedule IV Part D (9) of FCO 1985
MEMORANDUM TO ACCOMPANY ORGANIC FERTILIZER/BIOFERTILIZER/NON-EDIBLE DE-OILED CAKE FERTILIZER SAMPLE FOR ANALYSIS

No
From

To
Incharge
Organic Fertilizer/ Biofertilizer/ Non-edible de-oiled cake fertilizer Quality Control Laboratory

The Organic Fertilizer/ Biofertilizer/ Non-edible de-oiled cake fertilizer sample as per the details given below are sent for analysis:-

1. Name of Organic Fertilizer/ Biofertilizer/ Non-edible de-oiled cake fertilizer
2. Date of sampling
3. Physical condition
4. Code number of sample

The analysis report may be forwarded to

Place

Date
Signature and metallic seal

Inspector

Impression of Fertilizer Inspector
ANALYSIS REPORT OF ORGANIC FERTILIZER SAMPLE

No._________________

Government of ______________________________

_______________________________________________

(Name of the Laboratory)

Date ______________________________

To

The Fertiliser Inspector

_______________________________________________

The analysis report of the organic fertilizer sample forwarded vide your reference No.______________ dated ________ is as per details given below:

(1) Name of organic fertilizer ______________________________

(2) Date of sampling ______________________________

(3) Code No. of sample as indicated by _______________________

the Inspector Date of receipt of the sample in ________________

laboratory

(4) Laboratory sample No. ______________________________

(5) Date of analysis of sample ______________________________

(6) Analysis of organic fertilizer (on fresh weight basis)

<table>
<thead>
<tr>
<th>Sl. No.</th>
<th>Specification as per FCO</th>
<th>Composition as analysis</th>
<th>Variation</th>
<th>Permissible tolerance limit</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td></td>
<td></td>
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<td></td>
</tr>
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<td>2</td>
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<tr>
<td>3</td>
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<tr>
<td>4</td>
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</tr>
<tr>
<td>5</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(A) Physical Characteristics –
   i. Moisture content ____________________________
   ii. Bulk density ______________________________
   iii. Particle size _____________________________

(B) Chemical Characteristics –
i. Total Organic Carbon ________________________________
ii. Total Nitrogen ________________________________
iii. C:N ______________________________________
iv. Phosphorus _________________________________
v. Potassium _________________________________
vi. pH _______________________________________
vii. Conductivity ________________________________

(C) Heavy Metal
i. Cadmium _________________________________
ii. Chromium ________________________________
iii. Copper _________________________________
iv. Mercury _________________________________
v. Nickel _________________________________
vi. Lead _________________________________
vii. Zinc _________________________________

Remarks: The sample is/is not according to specification and fails in _____
___________________________________________________________
___________________________________________________________

Signature of the Incharge
(Testing Laboratory)

Copy to:
Director of Agriculture
________________________
________________________
ANALYSIS REPORT OF BIOFERTILISER SAMPLE

No._________________      Date _________

Government of ________________________

____________________________________

____________________________________

(Name of the Laboratory)

To
The Fertiliser Inspector

____________________________________

The analysis report of the biofertiliser sample forwarded vide your reference No.___________ dated __________ is as per details given below :

(1) Name of biofertilizer

(2) Date of sampling

(3) Code No. of sample as indicated by the Inspector

(4) Date of receipt of the sample in laboratory

(5) Laboratory sample No.

(6) Date of analysis of sample

(7) Analysis of Biofertiliser (on fresh weight basis)

<table>
<thead>
<tr>
<th>Sl. No.</th>
<th>Specification as per FCO (Rhizobium, Azotobacter, Azospirillum, PSM)</th>
<th>Composition as per analysis (Rhizobium, Azotobacter, Azospirillum, PSM)</th>
<th>Variation</th>
<th>Permissible tolerance limit</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>5</td>
</tr>
</tbody>
</table>

(A) Physical Characteristics –
   i.  Moisture content

85
ii. Particle size

(B) Chemical Characteristics –
i. pH

(C) Microbial Characteristics
i. Viable Cell Count
ii. Contamination Level

(D) Efficiency Characteristics
*(i) Nodulation Test
**(ii) Nitrogen fixed (mg)/g of sucrose consumed
***(iii) Formation of white pellicle in semi solid Nitrogen free bromothymol blue media
+(iv) (a) Solubilization zone (mm)
(b) P-phosphorus (%) Spectrophotometer

Remarks: The sample is/is not according to specification and fails in 
____________________________________________________________________
____________________________________________________________________

Signature of the Incharge
(Testing Laboratory)

Copy to:
Director of Agriculture

____________________________________________________________________
ANALYSIS REPORT OF NON-EDIBLE DE-OILED CAKE FERTILISER SAMPLE

No._________________      Date __________

Government of ________________________
____________________________________
____________________________________
(Name of the Laboratory)

To
The Fertiliser Inspector
___________________________________
___________________________________

The analysis report of the organic fertilizer sample forwarded vide your reference No.__________________ dated ________________ is as per details given below:

(1) Name of Non-edible de-oiled Cake fertiliser
(2) Date of sampling _____________________________
(3) Code No. of sample as indicated by ____________________________
    the Inspector
(4) Date of receipt of the sample in ________________ laboratory
(5) Laboratory sample No. _____________________________
(6) Date of analysis of sample _____________________________
(7) Analysis of Non-edible de-oiled cake fertilizer (on fresh weight basis)

<table>
<thead>
<tr>
<th>Sl. No.</th>
<th>Specification as per FCO</th>
<th>Composition as per analysis</th>
<th>Variation</th>
<th>Permissible tolerance limit</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td></td>
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<td>2</td>
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</tr>
<tr>
<td>3</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(A) Physical Characteristics –
   i. Moisture content
   ii. Particle size

87
(B) Chemical Characteristics –

<table>
<thead>
<tr>
<th>i.</th>
<th>ii.</th>
<th>iii.</th>
<th>iv.</th>
<th>v.</th>
<th>vi.</th>
<th>vii.</th>
<th>viii.</th>
<th>ix.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total Organic Carbon</td>
<td>Total Nitrogen</td>
<td>C:N</td>
<td>Phosphorus</td>
<td>Potassium</td>
<td>pH</td>
<td>Conductivity</td>
<td>Total Ash</td>
<td>Others</td>
</tr>
</tbody>
</table>

Remarks: The sample is/is not according to specification and fails in _____

________________________________________________________________________

________________________________________________________________________

Signature of the Incharge
(Testing Laboratory)

Copy to:
Director of Agriculture

____________________
____________________
Minimum Laboratory Requirements for Quality Testing Laboratory for Biofertilizer and Organic Fertilizers (S.O. 2724E) Gazette Notification Dated 8th November 2010

In pursuance of sub-clause (2) of clause 29 of the Fertiliser (Control) Order, 1985, the Controller with a view to ensure accurate analysis of fertilizer samples, hereby specifies that every laboratory notified for testing the samples of biofertilizer and organic fertilizer under sub-clause (1) shall possess within one year from the date of publication of this notification, the following minimum laboratory equipment and other laboratory facilities, namely:-

(A) Equipment and other facilities for testing samples of biofertilizers;

1. **Hot air oven** (up to 250°C)  
   Chamber size minimum 24” x 24” x 24”
2. **Vertical Autoclave**  
   16” Dia x 24” height
3. **BOD incubator**  
   capacity min-9cft., 50°C- 50°C
4. **pH Meter**
5. **Laminar Air Flow**  
   minimum size 3 x 2 x 2ft.
6. **Reciprocal/Orbital incubator shaker**
7. Colony counter or Mechanical counting device
8. **Serological water bath**
9. **Electronic balance**
10. **Water distillation unit**
11. **Kjeldahl digestion unit**
12. **Kjeldahl distillation assembly**
13. **Standard IS sieve**
14. **Spectrophotometer**  
   360-960nm
15. **Binocular research microscope**  
   40 and 100X objective,  
   preferably with phase contrast attachment having 10X,  
   40X and 100 X phase objectives
16. **Flame photometer**
17. **Rotary shaker**
18. **Vacuum filtration device**
19. **Vacuum pump**
20. **Earthen Pots**
21. **Cooling storage cabinet for samples**  
   10-25°C
22. **Rotary vacuum dryer**

(B) Equipments and other facilities for testing samples of organic fertilizer laboratories:

1. **Hot air oven**  
   chamber size minimum 24”x 24” x 24”
2. **Rotary shaker**
3. **Vacuum filtration device**
4. Vacuum pump
5. pH meter
6. Electronic Weighing balance
7. Conductivity meter
8. Flame Photometer

9. Spectrophotometer 360-960nm
10. Muffel furnace
11. Hot plate cum stirrer
12. Atomic Absorption Spectrophotometer (AAS)
13. Cold vapour Mercury analyzer or vapour generator Assembly for AAS
14. Kjedahl digestion unit
15. Kjeldahl distillation assembly
16. Fume hood with exhaust facility
17. Soxhlet apparatus for refluxing