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Laboratory of Environmental Biology
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Division of Environmental Health Sciences
Albany, New York

NYS DOH LEB-603

Preparation of Samples for Medical Marijuana Testing
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1.0. Scope and Application

1.1. This method, NYS DOH LEB-603, Preparation of Samples for Medical Marijuana Testing (ELAP Method ID 9926) describes the preparation of samples of medical marijuana products for the detection of microbial contaminants, including bile tolerant gram negative bacteria, *Escherichia coli*, *Clostridium botulinum*, *Pseudomonas* species, *Enterococcus* species, *Salmonella* species, *Penicillium* species, *Aspergillus* species, *Mucor* species, and thermophilic actinomycetes as required in Title 10 (Health), Subpart 55-2.15 and Chapter XIII, Section 1004.14 of the official Compilation of Codes, Rules, and Regulations, of the State of New York. Numbers of fungi and aerobic bacteria are also determined.

1.2. Protocols for the identification of these organisms in samples of medical marijuana products can be found in the NYS DOH LEB-600 series. See Appendix A for Medical Marijuana Microbial Testing Plan flowcharts.

2.0. Summary of the Method

2.1. Twenty doses of individual lots of medical marijuana product are pooled. Either 10 or 15 doses are diluted at least 1:10 in PBST, and the remaining doses are archived. Samples containing materials that are not soluble in water are diluted with isopropyl myristate prior to adding PBST. Matrix spikes of microbial contaminants under test and positive and negative controls are prepared and analyzed in parallel with sample aliquots.

3.0. Definitions

3.1. PBST stands for Phosphate Buffered Saline, pH 7.2, containing 0.1% Tween® 80.

3.2. Polysorbate 80 (Tween® 80) is a nonionic surfactant and emulsifier.

3.3. Batched product refers to the combining of individual sample aliquots from a single lot of product into one sample to be tested.

3.4. BioBalls® are small water-soluble balls containing a known number of microorganisms that are used for quality control.

3.5. RO stands for an organization that is registered to manufacture and dispense medical marijuana in New York State.


4.0. Health and Safety Warnings

4.1. Microbiological analyses involve the culturing of potentially pathogenic organisms.

4.1.1. All microbiologically contaminated materials, including media, shall be autoclaved after use.

4.1.2. Contaminated glassware and plastic ware shall be decontaminated prior to washing.

4.1.3. Laboratory equipment and benches shall be disinfected using either Envirocid®®, 10% bleach, or a minimum concentration of 70% ethanol before and after use.

4.1.4. Mouth pipetting is prohibited.
4.1.5. All accidents, particularly those which may result in infection, shall be reported according to laboratory-specific policies and procedures.

5.0. Shipping Conditions, Receiving, Preservation and Storage

5.1. Sample Shipping Conditions

5.1.1. The medical marijuana products from the Registered Organizations (ROs) are shipped as per manufacturer’s specifications and must adhere to all regulatory requirements.

5.2. Sample Receipt

5.2.1. Medical marijuana products from the RO are received, verified and documented ensuring that method, regulatory and Accreditation Body requirements are met.

5.3. Method Holding Times

5.3.1. Once samples are diluted, they are immediately analyzed according to NYS DOH LEB-604 and NYS DOH LEB-605.

5.4. Preservation

5.4.1. Samples diluted in PBST that are not required for analyses are stored refrigerated until it has been determined that they are not needed for additional microbiological evaluation.

5.4.2. Unused, reconstituted BioBalls® can be stored for one week at 1.0-8.0°C.

5.5. Storage

5.5.1. If storage is required prior to analysis, samples are refrigerated within a box having double locks in a locked refrigerator.

5.5.2. Once analysis is complete, remaining doses are maintained at room temperature within a box having double locks, in a locked room, until destroyed

6.0. Interferences

6.1. Some components of medical marijuana products, e.g., ethanol, may inhibit the growth of microorganisms.

7.0. Apparatus and Materials

7.1. Equipment and Supplies

7.1.1. Water bath, set at 35.0-40.0°C

7.1.2. Erlenmeyer flasks, foam plugged, various sizes, sterile

7.1.3. Graduated cylinders, various sizes, sterile

7.1.4. 50mL conical tubes, sterile – Krackeler Scientific cat. no. 3-352098, or equivalent

7.1.5. 1.7mL microcentrifuge tubes, sterile – Krackeler Scientific cat. no. 383-MCT175C, or equivalent

7.1.6. 250mL centrifuge bottles, sterile – Fisher Scientific cat. no 0553853, or equivalent

7.1.7. Automatic pipetters and sterile aerosol-resistant micropipette tips

7.1.8. Pipette-aid
7.1.9. Pipettes, sterile disposable, 5mL, 10mL, 25mL
7.1.10. Balance
7.1.11. NIST traceable weights
7.1.12. Forceps, either metal or disposable individually wrapped and sterile (Krackeler Scientific, Albany, NY, cat. no. 8-F37944-0000-PK, or equivalent)
7.1.13. Spatulas, either metal or disposable individually wrapped and sterile (Krackeler Scientific, Albany, NY, cat. no. 83-3012, or equivalent).
7.1.14. Pasteur pipettes, sterile
7.1.15. Pliers
7.1.16. Bunsen burner, or equivalent
7.1.17. Biosafety cabinet with HEPA filter

7.2. Reagents and Chemicals
7.2.1. Phosphate Buffered Saline pH 7.2, 100 mL and 1L aliquots in tubes/bottles, containing 0.1% Tween 80 (PBST).
7.2.2. Isopropyl myristate, Sigma Chemical Company, cat. no. M0757 or equivalent, pre-warmed to 35-40°C.
7.2.3. *Aspergillus brasiliensis* (also known as *A. niger*) ATCC 16404, BioBall® MultiShot 550, bioMérieux cat. no. 56001 or 56011.
7.2.4. *Clostridium sporogenes* ATCC 11437, BioBall® MultiShot 550, bioMérieux cat. no. 56004 or 56014.
7.2.5. *Enterococcus faecalis* ATCC 29212, either lab prepared spiking solution or BioBall® MultiShot 550, bioMérieux cat. no. 56005 or 56015.
7.2.6. *Escherichia coli* ATCC 8739, either lab prepared spiking solution or BioBall® MultiShot 550, bioMérieux cat. no. 56006 or 56016.
7.2.7. *Klebsiella pneumoniae* ATCC 13883, lab prepared spiking solution.
7.2.8. *Pseudomonas aeruginosa* ATCC 9027, either lab prepared spiking solution or BioBall® MultiShot 550, bioMérieux cat. no. 56007 or 56017.
7.2.9. *Salmonella typhimurium* ATCC 14028, either lab prepared spiking solution or BioBall® SingleShot, bioMérieux cat. no. 413775 or 56044.
7.2.10. Ethanol for flame-sterilization
7.2.11. Disinfectants such as Envirocide® (Fisher Scientific cat. no. 19898220), 70% ethanol, and/or Clorox.

7.3. Forms
7.3.1. Medical Marijuana Sample Preparation Log (e.g., LEB-RS-603A)
7.3.2. Medical Marijuana Controlled Substance Inventory Log (e.g., LEB-RS-603B)
7.3.3. Summary of Sample Preparation and Analysis Set-Up (e.g., LEB-AP-603A)
7.3.4. Medical Marijuana Microbial Testing Plan (e.g., LEB-AP-603B)
7.3.5. Medical Marijuana Sample Types and Preparation Instructions (e.g., LEB-AP-603C)
7.3.6. Balance Calibration Record
8.0. Quality Control/Assurance
8.1. Method Detection Limits
8.1.1. Method Detection Limits are product-specific and are determined in accordance with relevant standards, regulations and Accreditation Body requirements.

8.2. Calibration and Standardization
8.2.1. Temperatures of the cold room and refrigerator are observed and recorded twice daily separated by at least 4 hours on either the Cold Room or Refrigerator Temperature Records.
8.2.1.1. If the cold room or refrigerator does not stay within 1.0-8.0°C, follow laboratory-specific corrective actions.
8.2.1.2. The optimum temperature range for a cold room or refrigerator is 1.0-4.0°C
8.2.1.3. If the cold room or refrigerator was in a defrost cycle at the time that the temperature was recorded, and the temperature does not reach 8.0°C, re-testing of media is not required.
8.2.1.4. Media may be re-tested for quality, depending on the number of degrees and the amount of time that the cold room temperature was out of compliance, at the discretion of the laboratory.
8.2.2. Temperature of the freezer is observed and recorded twice daily separated by at least 4 hours on the Freezer Temperature Record.
8.2.2.1. If the temperature on the freezer exceeds -15.0°C, follow laboratory-specific corrective actions.
8.2.3. Water bath temperatures shall be observed and recorded twice daily, separated by at least 4 hours.
8.2.3.1. Temperature of the 35.0-40.0°C water bath is observed and recorded on the Water Bath Temperature Record.
8.2.3.1.1. If water bath temperature does not stay within 35.0-40.0°C, follow laboratory-specific corrective actions. Analytical results may be invalidated if the temperature exceeds 40.0°C, at the discretion of the laboratory.
8.2.4. Max/min temperatures are recorded when twice-daily temperature measurements are not possible, such as on holidays and weekends.
8.2.4.1. Max/min temperatures of freezers may show transient excursions of temperature above 0.0°C due to defrost cycles.
8.2.4.2. This is acceptable, provided that the quality of BioBalls® are unaffected.
8.2.5. Thermometers must be calibrated against a NIST-certified thermometer as prescribed by the Accreditation Body and in accordance with relevant regulations and standards.
8.2.6. Sterility of disposable loops, spreaders, and spatulas is determined as prescribed by the Accreditation Body and in accordance with relevant regulations and standards.

8.2.7. The volumetric accuracy of automatic pipettors and serological pipettes is as prescribed by the Accreditation Body and in accordance with relevant regulations and standards.

8.2.8. The intensity and efficacy of the UV light in the biosafety cabinet is measured quarterly.

8.2.9. Biosafety cabinets are certified annually as prescribed by the Accreditation Body and in accordance with relevant regulations and standards.

8.3. Quality Control

8.3.1. The top-loading balance is calibrated prior to pooling and results are recorded.

8.3.2. Lab prepared spikes are counted at a minimum of once a week in as prescribed by the Accreditation Body and in accordance with relevant regulations and standards.

8.3.3. Comparative recovery and sterility between lots of PBST are determined.

8.3.4. Liquid media is stored in tightly-capped bottles in the dark at 4°C for up to 3 months from the date of preparation.

8.3.5. The use test for deionized water is performed annually, when cartridges are changed, or repairs are made to the deionized water systems.

8.4. Corrective/Preventive Actions

8.4.1. The laboratory will initiate non-conformances and/or corrective/preventive actions in accordance with laboratory-specific procedures and as prescribed by the Accreditation Body and in accordance with relevant regulations and standards.

9.0. Procedure

9.1. General

9.1.1. Aseptic technique is used for all procedures.

9.1.1.1. Aseptic technique can be found in a general microbiology textbook or on-line.

9.1.2. All work surfaces are disinfected prior to preparation of spiking solutions, sample pooling/batching, and sample preparation.

9.1.3. Preparation of spiking solutions are performed in a physically different location than sample preparation and initial sample analyses to prevent cross-contamination of incoming products.

9.2. Matrix Spikes

9.2.1. Every RO has different brands of products which can come in different forms. For example, an RO may have a “Extra Strength” brand that comes in capsules, tinctures, and vaporizers. Each one of the forms under
the “Extra Strength” brand is considered a separate product type and is analyzed as such.

9.2.2. Every new product type analyzed for microbial contaminants will have corresponding matrix spikes for the first three times that specific product type is analyzed or until results are consistent. After at least three consistent matrix spike results, that specific product type will only need corresponding matrix spikes every 20\textsuperscript{th} time it is received for analysis.

9.3. Preparation of Spiking Solutions

9.3.1. Prior to any sample preparation or analysis, the spiking solutions must be prepared either in the laboratory or by using BioBalls®.

9.3.2. Laboratory Prepared Spiking Solutions

9.3.2.1. Follow laboratory specific procedures.

9.3.2.2. Lab prepared spikes are counted at a minimum of once a week.

9.3.3. BioBall® Rehydration

9.3.3.1. For each BioBall® label a 1.7mL sterile microcentrifuge tube with the organism name and date of rehydration.

9.3.3.2. Add 1.1mL of PBST to each tube.

9.3.3.3. Remove the BioBall® vial from the freezer, slowly uncap, and tip the BioBall into the tube containing PBST.

9.3.3.4. Once the BioBall® dissolves, invert the tube several times to mix gently.

9.3.3.5. Incubate the tubes for 4 hours at room temperature.

9.3.3.6. Vortex to mix prior to use.

9.3.3.7. The reconstituted BioBall® can now be used in 100µL aliquots as a 50 CFU spike.

9.3.3.8. BioBall® preparations of \textit{K. pneumoniae} and \textit{T. vulgaris} are not available, so laboratory prepared spikes must be used.

9.4. Pooling/Batching Accepted Samples

9.4.1. Pooling of sample aliquots is performed in a biological safety cabinet. Prior to pooling, place all supplies needed for pooling in the biological safety cabinet, close the sash, and UV disinfect the cabinet for 15 minutes.

9.4.2. When the samples arrive, the containers are disinfected using either Envirocide®, 10% bleach, or a minimum concentration of 70% ethanol prior to placing them in the biosafety cabinet.

9.4.3. Aseptically add 20 doses from the same production date/lot number to a sterile, labeled, vessel that has been tared, and record the weight on the sterile vessel.

9.4.3.1. Pool doses from each of the containers submitted for the lot of product being analyzed.

9.4.3.2. If 20 doses are not available, record the number of doses (e.g., LEB-RS-603A).
9.4.3.3. Occasionally a sample matrix is too viscous to accurately batch 20 doses. In this instance, more than 20 doses can be pooled to ensure there are enough doses for analysis.

9.4.4. On the Medical Marijuana Sample Preparation Log (e.g., LEB-RS-603A) record the sample accession number, sample type, dose amount, combined pool weight, analyst initials, and date.

9.4.4.1. If the viscosity of the pooled doses will require the addition of isopropyl myristate prior to dilution, check the corresponding box.

9.4.5. Determine the number of number of doses that have been pooled and record it on the Medical Marijuana Sample Preparation Log (e.g., LEB-RS-603A).

9.4.5.1. If the number of doses is less than or more than 20, the number of doses batched is calculated by dividing the total weight or volume by the weight or volume of one dose.

9.4.5.1.1. For example, if one dose of a product is 4.5mg, and the weight of the pooled product is 99mg, 99/4.5 = 22 doses.

9.4.6. Determine the volume of the diluted aliquot and record it on the Medical Marijuana Sample Preparation Log (e.g., LEB-RS-603A).

9.4.6.1. The diluted aliquot is typically a 1:10 dilution of a single dose in PBST and is calculated by multiplying the weight/volume of one dose by 10.

9.4.6.1.1. For example, if one dose of a product is 4.5mg, 4.5mg x 10 = 45µLs of diluted aliquot.

9.4.6.2. Occasionally a product will require more than a 1:10 dilution to allow for accurate dispensing for sample analyses. See the Medical Marijuana Sample Types and Preparation Instructions (e.g., LEB-AP-603C).

9.4.6.3. One diluted aliquot is used in each of the subsequent analyses.

9.4.7. Once 20 doses have been pooled, either 10 or 15 doses are removed for the subsequent analyses according to sections 9.5.-9.6. and the remaining doses are archived according to section 9.8.

9.4.7.1. The number of doses needed for analysis depends on if there are corresponding matrix spikes. See LEB-AP-603A for a summary of sample preparation and analyses set up.

9.5. Preparation of Products Soluble in Water

9.5.1. Aseptically add either 10 or 15 pooled doses from the same production date/lot number to a labeled, sterile vessel.

9.5.1.1. Viscous samples with small doses (less than 50µL or 50mg) may prevent accurate pipetting because a small portion of the sample may stick to the insides of pipette tips. In this case it is recommended that all 20 doses are diluted prior to sample
analysis to ensure there is enough volume for all sample analyses. See the Medical Marijuana Sample Types and Preparation Instructions (e.g., LEB-AP-603C).

9.5.2. Calculate the volume of PBST required to produce the necessary dilution (weight/volume for pills or solids or volume/volume for liquid samples) by multiplying the combined weight/volume of either 10 or 15 doses by the dilution factor (typically 10) and subtracting the combined weight/volume of the doses.

9.5.2.1. For example, if the product requires a 1:10 dilution and the combined volume of 15 doses is 67.5µL (4.5µL/dose x 15 doses), (67.5µL x 10) – 67.5µL = 607.5µL. So 67.5µL of the combined 15 doses are added to 607.5µL of PBST.

9.5.2.2. Occasionally a product will require more than a 1:10 dilution to allow for accurate dispensing for sample analyses Medical Marijuana Sample Types and Preparation Instructions (e.g., LEB-AP-603C).

9.5.3. Add the calculated volume of PBST to the batched sample and vortex to mix.

9.5.3.1. Pills or capsules may require heating to enhance dissolution.

9.5.3.1.1. If needed, use PBST preheated to 35.0-40.0°C and heat the diluted aliquots to 35.0-40.0°C in a water bath with intermittent vortexing until the material is dissolved.

9.6. Preparation Products Insoluble in Water

9.6.1. Aseptically add either 10 or 15 pooled doses from the same production date/lot number to a labeled, sterile vessel.

9.6.1.1. Viscous samples with small doses (less than 50µL or 50mg) may prevent accurate pipetting because a small portion of the sample may stick to the insides of pipette tips. In this case it is recommended that all 20 doses are diluted prior to sample analysis to ensure there is enough volume for all sample analyses. Medical Marijuana Sample Types and Preparation Instructions (e.g., LEB-AP-603C).

9.6.2. Calculate the volume of PBST required to produce the necessary dilution (weight/volume for pills or solids or volume/volume for liquid samples) by multiplying the combined weight/volume of either 10 or 15 doses by the dilution factor (typically 10) and subtracting the combined weight/volume of the doses.

9.6.2.1. For example, if the product requires a 1:10 dilution and the combined volume of 15 doses is 67.5µL (4.5µL/dose x 15 doses), (67.5µL x 10) – 67.5µL = 607.5µL. So 67.5µL of the combined 15 doses are added to 607.5µL of PBST.

9.6.2.2. The use of PBST preheated to 35.0-40.0°C can assist with creating an emulsion.
9.6.2.3. Occasionally a product will require more than a 1:10 dilution to allow for accurate dispensing for sample analyses Medical Marijuana Sample Types and Preparation Instructions (e.g., LEB-AP-603C).

9.6.3. If, due to matrix viscosity, an emulsion cannot be created when the batched sample is diluted, add an equal volume of filter-sterilized isopropyl myristate (pre-warmed to 35.0-40.0°C) to the pooled sample prior to dilution with PBST.

9.6.3.1. For example, if the combined volume of 15 doses is 67.5µL (4.5µL/dose x 15 doses), add 67.5µL of pre-warmed isopropyl myristate.

9.6.3.2. If an equal volume of isopropyl myristate was added, the combined volume of 15 doses is 135µL (4.5µL/dose plus 4.5µL of isopropyl myristate x 15 doses), (135µL x 10) - 135µL = 1215µL. So 135µL of the combined 15 doses with isopropyl myristate is added to 1215µL of PBST preheated to 35.0-40.0°C.

9.6.4. Add the calculated volume of PBST to the batched sample and vortex to mix.

9.7. Analysis of product

9.7.1. Once the samples have been pooled and diluted, and the BioBalls® have rehydrated for 4 hours, immediately proceed with NYS DOH LEB-604 and NYS DOH LEB-605.

9.7.1.1. Sample analyses required are summarized in the Summary of Sample Preparation and Analysis Set-Up (e.g., LEB-AP-603A).

9.7.2. Un-used aliquots of rehydrated BioBalls® can be reused when stored at 1.0-8.0°C for up to one week.

9.8. Product archival and re-analysis

9.8.1. After pooling the 20 doses (see section 9.4), the pool is stored refrigerated in a double-lock box until analyzed.

9.8.2. When storing the pool, the accession ID, weight of the pool, date of storage, and the initials of the analyst and witness are recorded on the Medical Marijuana Controlled Substance Inventory Log (e.g., LEB-RS-603B).

9.8.3. After removing either 10 or 15 doses for sample analyses, the remaining doses are stored refrigerated in a double-locked safe until analyses are completed.

9.8.4. When archiving the remaining doses the weight of the archived doses, date of archival, and the initials of the analyst and witness are recorded on the Medical Marijuana Controlled Substance Inventory Log (e.g., LEB-RS-603B).

9.8.4.1. If all the doses are used for sample analyses (ie, viscous samples insoluble in water that require the addition of isopropyl
myristate) then “n/a” is recorded in the archived weight section on the Medical Marijuana Controlled Substance Inventory Log (e.g., LEB-RS-603B) as they will not be stored. The box “All used for initial testing” on LEB-RS-603B can be checked to indicate this.

9.8.5. The recording of the initial archived weights and archiving the remaining doses are witnessed by a second analyst and both analyst initials are recorded in the “Analyst/Witness Initials” section on the Medical Marijuana Controlled Substance Inventory Log (e.g., LEB-RS-603B).

9.8.6. If archived samples are analyzed after initial testing is completed, analysts first record the weight of the archived samples on the Medical Marijuana Controlled Substance Inventory Log (e.g., LEB-RS-603B) in the “Second Archived Weight/Date” section.

9.8.6.1. This is to indicate if there was any sample evaporation while the samples were archived.

9.8.7. Analysts then remove the amount of product needed for analysis and record the weight of the remaining doses on the Medical Marijuana Controlled Substance Inventory Log (e.g., LEB-RS-603B) in the “Second Use Archived Weight/Date” section.

9.8.8. The recording of the archived and second use weights are witnessed by a second analyst and both analyst initials and the date are recorded on the Medical Marijuana Controlled Substance Inventory Log (e.g., LEB-RS-603B).

9.8.9. In the comments section of the Medical Marijuana Controlled Substance Inventory Log (e.g., LEB-RS-603B) indicate that the archived doses are being used for re-analysis.

9.9. Product Destruction

9.9.1. When analysis of a lot of product is completed and released, and re-analysis is not necessary, the archived doses are destroyed in accordance with relevant regulations and standards.

10.0. Data Acquisition, Reduction, Analysis, Calculations, Acceptance Criteria and Documentation

10.1. Record the sample accession number, sample type, total amount of sample received in grams or mLs, final sample volume in PBST or isopropyl myristate/PBST, expiration dates of PBST and isopropyl myristate, analyst and witness initials and date on the Medical Marijuana Sample Preparation Log (e.g., LEB-RS-603A).

10.2. Record the date, target weight, measured weight, calculated error, pass/fail, and initials on the Balance Verification Form.

10.3. Record the accession number, initial weight, initial archive weight, second archive weight, second use archived weight, dates and analyst and witness initials on the Medical Marijuana Controlled Substance Inventory Log (e.g., LEB-RS-603B).
11.0. Method Performance

11.1. Demonstration of Capability

11.1.1. Prior to acceptance and use of this method for data reporting, a satisfactory initial demonstration of capability (DOC) is required. Thereafter, an ongoing DOC is to be performed annually.

11.1.2. An initial DOC shall be made prior to using any method, and at any time there is a change in instrument type, personnel or method or any time that a method has not been performed by the laboratory or analyst in a twelve (12) month period.

11.1.3. All DOCs shall be documented, and all data applicable to the demonstration shall be retained and readily available at the laboratory.

11.1.4. Consult relevant standards, regulations and Accreditation Body requirements for additional information on performing DOCs for microbial contaminants.

11.2. Laboratory Detection Limits

See 8.1.

12.0. Waste Management/Pollution Prevention

12.1. It is the responsibility of the laboratory to comply with all federal, state and local regulations governing waste management, particularly the hazardous waste identification rules and land disposal restrictions.

12.2. Bacterial/fungal cultures and contaminated or potentially contaminated disposable materials are disposed of in biohazardous waste cans and autoclaved prior to discarding.

12.3. Dispose of non-hazardous water waste in the laboratory sink followed by flushing with tap water.

12.4. Dispose of glassware in appropriately labeled boxes. Be sure that, whenever possible, the glass has been thoroughly rinsed and is contaminant-free before disposal.

12.5. Consult federal, state and local regulations for additional information or for information on the disposal of products not described in this method.

12.6. Unused products are disposed of according to section 9.9.

13.0. References


13.2. Standard Methods for the Examination of Water and Wastewater, 22nd edition, section 9050C.

13.3. NYS DOH LEB-604, Microbial Presence/Absence Test for Medical Marijuana Samples

13.4. NYS DOH LEB-605, Aerobic Bacteria and Mold Plate Counts for Medical Marijuana Testing

13.5. Title 10 (Health), Subpart 55-2.15 and Chapter XIII, Section 1004.14 of the official Compilation of Codes, Rules, and Regulations, of the State of New York
14.0. Appendices
## Summary of Sample Preparation and Analysis Set-Up (LEB-AP-603A)

<table>
<thead>
<tr>
<th>Analysis Number</th>
<th>Product Needed</th>
<th>Target</th>
<th>Matrix Spike Organism</th>
<th>Medium</th>
<th>Temperature</th>
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<tr>
<td>1</td>
<td>Yes</td>
<td>Aerobic bacteria</td>
<td>n/a</td>
<td>TSA</td>
<td>30-35°C</td>
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<tr>
<td>2</td>
<td>Yes</td>
<td>Aerobic bacteria, MS</td>
<td><em>E coli</em></td>
<td>TSA</td>
<td>30-35°C</td>
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<td>3</td>
<td>NO</td>
<td>Aerobic bacteria, positive control</td>
<td><em>E coli</em></td>
<td>TSA</td>
<td>30-35°C</td>
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<td>Aerobic bacteria, negative control</td>
<td>n/a</td>
<td>TSA</td>
<td>30-35°C</td>
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<td>5</td>
<td>Yes</td>
<td>Fungi</td>
<td>n/a</td>
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<td>20-25°C</td>
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<td>6</td>
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<td>20-25°C</td>
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<td>NO</td>
<td>Fungi, positive control</td>
<td><em>A. brasiliensis</em></td>
<td>SDA</td>
<td>20-25°C</td>
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<td>8</td>
<td>NO</td>
<td>Fungi, negative control</td>
<td>n/a</td>
<td>SDA</td>
<td>20-25°C</td>
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<td>9</td>
<td>Yes</td>
<td>Bile tolerant gram-negative organisms</td>
<td>n/a</td>
<td>EEBM</td>
<td>30-35°C</td>
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<td>10</td>
<td>Yes</td>
<td>Bile tolerant gram-negative organisms, MS</td>
<td><em>K. pneumoniae</em></td>
<td>EEBM</td>
<td>30-35°C</td>
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<td>11</td>
<td>Yes</td>
<td><em>E coli, Salmonella spp., Pseudomonas spp.</em></td>
<td>n/a</td>
<td>TSB</td>
<td>30-35°C</td>
</tr>
<tr>
<td>12</td>
<td>Yes</td>
<td><em>E coli, Salmonella spp., Pseudomonas spp., MS</em></td>
<td><em>E coli, S. typhimurium, Ps. aeruginosa</em></td>
<td>TSB</td>
<td>30-35°C</td>
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<tr>
<td>13</td>
<td>Yes</td>
<td>Clostridium botulinum</td>
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<td>RCM</td>
<td>30-35°C, anaerobic</td>
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<td>14</td>
<td>Yes</td>
<td>Clostridium botulinum*</td>
<td>n/a</td>
<td>RCM</td>
<td>30-35°C, anaerobic</td>
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<td>15</td>
<td>Yes</td>
<td>Clostridium botulinum, MS</td>
<td><em>Clostridium sporogenes</em></td>
<td>RCM</td>
<td>30-35°C, anaerobic</td>
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<td>16</td>
<td>Yes</td>
<td>Enterococcus spp.</td>
<td>n/a</td>
<td>TSB</td>
<td>30-35°C, 5% CO₂</td>
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<tr>
<td>17</td>
<td>Yes</td>
<td>Enterococcus spp., MS</td>
<td><em>E. faecalis</em></td>
<td>TSB</td>
<td>30-35°C, 5% CO₂</td>
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<td>18</td>
<td>Yes</td>
<td>Thermophilic actinomycetes</td>
<td>n/a</td>
<td>TSB</td>
<td>50-55°C</td>
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<tr>
<td>19</td>
<td>Yes</td>
<td>Thermophilic actinomycetes, MS</td>
<td><em>T. vulgaris</em></td>
<td>TSB</td>
<td>50-55°C</td>
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<td>Yes</td>
<td>Archived sample</td>
<td>n/a</td>
<td>n/a</td>
<td>Room temp</td>
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<td>Yes</td>
<td>Archived sample</td>
<td>n/a</td>
<td>n/a</td>
<td>Room temp</td>
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<td>22</td>
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<td>Archived sample</td>
<td>n/a</td>
<td>n/a</td>
<td>Room temp</td>
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<td>Yes</td>
<td>Archived sample</td>
<td>n/a</td>
<td>n/a</td>
<td>Room temp</td>
</tr>
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<td>24</td>
<td>Yes</td>
<td>Archived sample</td>
<td>n/a</td>
<td>n/a</td>
<td>Room temp</td>
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</tbody>
</table>

MS = matrix spike, TSB = tryptic soy broth, TSA = tryptic soy agar, SDA = Sabouraud Dextrose agar, EEBM = Enterobacterial Enrichment Broth Mossel, RCM = Reinforced Clostridial Medium.

Matrix spikes are prepared according to LEB-603 section 9.3, and should contain less than 100 organisms.

*Heat sample at 80°C for 10 minutes*
Medical Marijuana Microbial Testing Plan (LEB-AP-603B)

Medical Marijuana Microbial Testing Plan

Target Organisms

Pellet Sample

- Arthrobacter
  - Bile Tolerant Gram Negative
  - Citrobacter freundii
  - Enterobacteriaceae spp.
  - Enterococcus spp.
  - E. coli
  - Escherichia coli
  - Pseudomonas spp.
  - Salmonella spp.

- Acinetobacter
  - Aerobic Plate Counts

- Listeria monocytogenes
  - Aerobic Plate Counts

- Aspergillus spp.
  - Mucor spp.
  - Penicillin spp.

- Thermophilic Actinomycetes
  - Mold Plate Counts

- Gram Positive Staphylococcus aureus (ATCC 6538)

- Thermophilic Actinomycetes

- Slime producing colonies

- Pseudomonas sp.

- Slime producing colonies
Medical Marijuana Microbial Testing Plan

General Set-Up

- Aerobes
- Bile Tolerant Gram Negative
- C. jejuni
- Enterococcus spp.
- Thermophilic Actinomycetes
- Aerobic Plate Counts
- Mold Plate Counts
- Extra Doses

- Unspiked Sample
- EC, PA, ST Spiked Sample
- KP Spiked Sample
- EF Spiked Sample
- TV Spiked Sample
- EC Spiked Sample
- Positive Control
- Negative Control
- AB Spiked Control

- AB Spiked Sample
- Unspiked Sample
- Unspiked Sample
- Unspiked Sample
- Unspiked Sample
- Unspiked Sample
- AB Spiked Sample

- 20 doses pooled
- 10-15 doses tested
- 5-10 doses archived

AB – Aspergillus brasiliensis ATCC 16404
KP – Klebsiella pneumoniae ATCC 13883
CS – Clostridium sporogenes ATCC 11437
EC – Escherichia coli ATCC 8739
PA – Pseudomonas aeruginosa ATCC 9027
ST – Salmonella typhimurium ATCC 14028
EF – Enterococcus faecalis ATCC 29212
TV – Thermactinomyces vulgaris ATCC 43649
Medical Marijuana Microbial Testing Plan

Organism Identification Assays

- Aerobes
  - API 20E
  - API 20NE
  - API 20NE

- Bile Tolerant Gram Negative
  - API 20E

- C. Influenzae
  - API 20NE

- Enterococcus spp.
  - API 20NE
  - Gram Stain

- Therapeutic Actinomycetes
  - API 20NE

- Aerobic Plate Counts
  - RT-PCR

- Mold Plate Counts
  - RT-PCR
  - RT-PCR
Medical Marijuana Microbial Testing Plan

Incubation Conditions

[Diagram showing various incubation conditions for samples, including temperatures and time frames.]
Medical Marijuana Microbial Testing Plan

Media Needed

Pooled Sample

- Trpionase Soy Broth
- Eosin Methenamine
- Reinforced Clostridial Medium
- Trpionase Soy Broth
- Trpionase Soy Broth
- Aerobic Plate Count
- Motil Plate Count

MacConkey Broth
MacConkey Agar

Cetamide Agar

RV Enrichment Broth
XLD Agar

Violet Red Bile Glucose Agar

Columbia Agar

Staphylococcal Selective Agar

Trypticase Soy Agar

Trypticase Soy Agar

Salmonella
Denitrific Agar

Salmonella
Denitrific Agar

Salmonella
Denitrific Agar
# Medical Marijuana Sample Types and Preparation (LEB-AP-603C)

<table>
<thead>
<tr>
<th>Product Type</th>
<th>Isopropyl Myristate Added</th>
<th>PBST Dilution</th>
<th>Heat and Vortex</th>
<th>Comments</th>
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<tbody>
<tr>
<td>Balm</td>
<td>X</td>
<td>1:10</td>
<td>X</td>
<td>Dilute all 20 doses</td>
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<tr>
<td>Capsules – Cellulose Fill</td>
<td></td>
<td>1:20</td>
<td>X</td>
<td></td>
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<tr>
<td>Capsules – Oil Fill</td>
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<td>1:10</td>
<td>X</td>
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<tr>
<td>Syrup</td>
<td></td>
<td>1:10</td>
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<tr>
<td>Tincture/Oral Solution/Oral Spray – Ethanol Base</td>
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<td>1:10</td>
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<td></td>
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<tr>
<td>Tincture/Oral Solution/Oral Spray – Oil Base</td>
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<td>1:10</td>
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<tr>
<td>Vaporizer – Liquid viscosity</td>
<td>X</td>
<td>1:10</td>
<td>X</td>
<td>Dilute all 20 doses</td>
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<tr>
<td>Vaporizer – Waxy viscosity</td>
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<td>1:10</td>
<td>X</td>
<td>Dilute all 20 doses</td>
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<td>Lotion</td>
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<td>1:10</td>
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<td>Dilute all 20 doses</td>
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<tr>
<td>Lozenge</td>
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<td>1:10</td>
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<td>Powder</td>
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<td>1:10</td>
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<td>Plant Material</td>
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<td>1:100</td>
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<td>Dilute all 20 doses</td>
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<td>Suppository</td>
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<td>1:10</td>
<td>X</td>
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<tr>
<td>Tablets – Cellulose Base</td>
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<td>1:20</td>
<td>X</td>
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# Medical Marijuana Sample Preparation Log (LEB-RS-603A)

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<tr>
<th>Accession #</th>
<th>Sample Type</th>
<th>Dose Amount</th>
<th>No. of Batched Doses</th>
<th>Combined Pool Weight</th>
<th>Diluted Aliquot Volume</th>
<th>Analyst Initials/Date</th>
<th>Comments</th>
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Medical Marijuana Controlled Substance Inventory Log (LEB-RS-603B)

<table>
<thead>
<tr>
<th>Accession #</th>
<th>Pooled Weight Initials/Date</th>
<th>Initial Archived Weight Initials/Date</th>
<th>Second Archived Weight</th>
<th>Second Use Archived Weight</th>
<th>Analyst/Witness Initials</th>
<th>Comments</th>
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<td></td>
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<td></td>
<td>□ All used for initial testing □ Transfer for destruction □ Other (see back)</td>
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<tr>
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<td></td>
<td></td>
<td></td>
<td>□ All used for initial testing □ Transfer for destruction □ Other (see back)</td>
</tr>
<tr>
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<td></td>
<td></td>
<td></td>
<td>□ All used for initial testing □ Transfer for destruction □ Other (see back)</td>
</tr>
<tr>
<td></td>
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<td></td>
<td></td>
<td></td>
<td>□ All used for initial testing □ Transfer for destruction □ Other (see back)</td>
</tr>
<tr>
<td></td>
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<td></td>
<td></td>
<td></td>
<td>□ All used for initial testing □ Transfer for destruction □ Other (see back)</td>
</tr>
<tr>
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<td></td>
<td></td>
<td></td>
<td>□ All used for initial testing □ Transfer for destruction □ Other (see back)</td>
</tr>
<tr>
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<td></td>
<td></td>
<td></td>
<td>□ All used for initial testing □ Transfer for destruction □ Other (see back)</td>
</tr>
</tbody>
</table>

*Archived samples are re-weighed upon removal from the lock-box to document any losses due to evaporation during storage.