China - Peoples Republic of

Post: Beijing

National Food Safety Standard - Aerobic Plate Count

Report Categories:
- FAIRS Subject Report

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Report Highlights:
On November 20, 2009, China notified the WTO of "National Food Safety Standard of the People’s Republic of China for the Microbiological Examination of Foods - Aerobic Plate Count" as SPS/N/CHN/189. The date for submission of final comments to the WTO is January 1, 2010. The proposed date of entry into force has not been specified.

Executive Summary:
On November 20, 2009, China notified the WTO of "National Food Safety Standard of the People’s Republic of China for the Microbiological Examination of Foods - Aerobic Plate Count" as SPS/N/CHN/189. The date for submission of final comments to the WTO is January 1, 2010. The proposed date of entry into force has not been specified.
Thanks go to the consortium of industry and 3rd country Embassies in Beijing for their assistance in translating and reviewing this standard.

This report contains an UNOFFICIAL translation of National Food Safety Standard of the People’s Republic of China for the Microbiological Examination of Foods - Aerobic Plate Count.

**General Information:**

BEGIN TRANSLATION

National Food Safety Standard of the People’s Republic of China

GB 4789.2----XXXX
Substitute GB/T 4789.2-2008

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**Microbiological Examination in Foods - Aerobic Plate Count**

(Draft for Comment)

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Issued by XXXX-XX-XX
Implemented by XXXX-XX-XX

Issued by the Ministry of Health
of the People’s Republic of China

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Preface


The differences between the first method of this Standard and the method of FDA/BAM lie in that:
- The applicable scope of plate count is changed from 25CFU ~250CFU to 30CFU~300CFU;
- The cultivation temperature is changed from 35±1°C to 36±1°C;
- Dilution by 10 times, changed from adding 10ml sample solution into 90ml diluents, to adding 1ml sample solution into 9ml diluents;
- Spiral plate count is not applied.
The difference between Method II and AOAC 990.12 lies in that: the cultivation temperature is changed from 35±1°C to 36±1°C.
This Standard substitutes the GB/T 4789.2-2003 “Microbiological examination for food hygiene - Aerobic plate count”.
The Standard is proposed and put under centralized management by the Ministry of Health of the People’s Republic of China.
The replaced former editions are:

Microbiological Examination of Foods - Aerobic Plate Count

1. Scope
This Standard defines the determination method of aerobic plate count in foods.
This Standard is applicable to the determination method of aerobic plate count in all kinds of foods.

2. Terms and Definitions
The following terms and definitions apply to this Standard.
Aerobic plate count: The aerobic plate count obtained from 1ml (or 1g) of sample under certain cultivation conditions (such as the ingredients of culture medium, cultivation temperature and time, pH, and aerobic, etc) after proper treatment.
The results obtained under the cultivation conditions prescribed in this Standard only include the aerobic plate count of a group of mesophilic aerobic bacterium or facultative anaerobe grown on the agar plate.

3. Equipment and Materials
In addition to conventional sterilization and cultivation equipment in microbiological laboratory, other equipment and materials are as follows:
3.1 Thermostatic cultivator: 36±1°C, 30±1°C
3.2 Refrigerator: 2°C~5°C.
3.3 Thermostatic water bath: 46±1°C
3.4 Balance: accuracy of 0.1g.
3.5 Homogenizer.
3.6 Oscillator.
3.7 Sterile pipette: 1ml (with a scale of 0.01ml), 10ml (with a scale of 0.1ml) or micropipettor and tips.
3.8 Sterile conical beaker: 250ml, 500ml.
3.9 Sterile culture plate: with a diameter of 90mm.
3.10 pH meter or pH colorimetric tube or precise pH indicator paper.
3.11 Magnifying glass (and) bacterial colony counter or Petrifilm™ automatic interpretoscope.
- Petrifilm™ is the trade name of the products provided by 3M Company. This information is for the convenience of the user of the Standard, not necessarily indicate that the product is recognized. If any other equivalent product has the same effect, these equivalents shall be applied.

4. Culture Media and Reagents
4.1 Agar Culture Medium for plate count: please refer to Section A.1.
4.2 Phosphate buffer solution: please refer to Section A.2.

4.3 Sterile normal saline solution: measure 8.5g NaCl to dissolve in 1000ml distilled water, and autoclave at 121°C for 15 minutes.
4.4 1mol/L NaOH: measure 40g NaOH to dissolve in 1000ml distilled water.
4.5 1mol/L HCl: measure 90mL of concentrated hydrochloric acid, and dilute to 1000mL with distilled water.
4.6 Petrifilm™ testing plate and cover plate for aerobic plate count.

Method I Plate Count Method
5. Examination Procedures
For the examination procedures of aerobic plate count, please refer to Fig.1.

Figure 1. The Examination Procedures of plate count
6. Operation Procedures
6.1 Dilution of samples
6.1.1 Solid and semi solid samples: measure 25g sample into a sterile homogenizing cup containing 225ml phosphate buffer solution or normal saline solution, homogenize with 8000-10000r/min for 1-2min, or place into sterile homogenizing bag containing 225ml diluent, beating with slapping type homogenizer for 1min to 2min, and then formulate into 1:10 sample solution.
6.1.2 Liquid Sample: Measure 25ml sample with sterile pipette into a sterile conical beaker containing 225ml phosphate buffer solution or normal saline solution (proper amount of sterile beads are placed in the beaker in advance), and then homogenize and formulate into 1:10 sample solution.
6.1.3 Absorb 1ml of 1:10 sample solution with 1ml sterile pipette or micropipettor, drip the solution into the sterile tube containing 9ml diluent along the wall of the tube (it is noted that the tip of the pipette shall not touch the diluent solution surface), shake up the test tube or place a piece of sterile pipette, blow repeatedly to homogenize, and then formulate into 1:100 sample solution.
6.1.4 Follow the operation procedures in 6.1.3, formulate the sample solution with series of dilution of
10 times. For each dilution, one piece of 1ml sterile pipette or tip is replaced.
6.1.5 As per the estimation of contamination status of samples, select 2 to 3 sample solutions with proper dilution (for liquid sample, original liquid shall be applied), when carrying out the escalating 10 times series of dilution, for each dilution, 1ml of sample solution is placed into two sterile plates. At the same time, measure 1ml of diluents into two sterile plate respectively to serve as blank controls.
6.1.6. Timely cool down the agar culture medium plates with 15-20ml content in each plate to 46℃ (which are placed into 46±1℃ water bath), decant the plates, and then rotate the plates to homogenize.
6.2 Cultivation
6.2.1 After the solidification of agar, turn the plates up-side-down, cultivate at 36±1℃ for 48h±2h. For aquatic products, cultivate at 30±1℃ for 72h±3h.
6.2.2 If the samples possibly contain bacteria that could spread growing on the surface of agar culture medium, a thin layer of agar culture medium is covered on the agar surface after solidification (about 4ml), and then turn the plate up-side-down after solidification, and cultivate it as per 6.2.1.
6.3 Plate Count
It could be observed with naked eyes, apply magnifying glass or bacteria colony counter when necessary, and record the dilution times and corresponding plate count. Plate count number is represented by colony-forming units (CFU).
6.3.1 Select the plates for total plate count with colony number between 30-300CFU, and without spreading growth on the plate. For plate with plate count under 30CFU, the number of colony is recorded, while for plate count over 300, it shall be recorded as uncountable. For each dilution degree, the average number of two plates shall be applied.
6.3.2 For those plates with large piece of colony growing, they shall not be applied. However, the plates without large piece of colony growth shall be applied for plate count; If the piece of colony covers less than one half of the plate area, and the colonies on the remaining half of the plate area scatter evenly, it shall be counted of this half of the plate and then multiply by 2, to represent the entire plate count.
6.3.3 When chain like growth occurs on the plate without evident border line between colonies on the plate, each chain shall be calculated as one colony.

7. Representation of Results
7.1 Calculation method for aerobic plate count
7.1.1 If there is only one dilution degree whose plate count fall in the proper counting scope, the average plate count of both plate shall be calculated, and then multiply the average value by corresponding dilution times, to serve as the total plate count in one gram (or ml) of sample.
7.1.2 If there are two continuous dilution degrees whose plate count falls in the proper counting scope, they shall be calculated as in Formula (1):

\[ N = \frac{\sum C}{(n_1 + 0.1n_2)d} \]  \hspace{1cm} (1)

Where,
N - Plate Count in sample;
\( \sum C \) - The total number of colonies on the plates (including the plates within the range of proper plate count);
N_1 - The number of colonies on the plates of the first proper dilution degree;
N_2 - The number of colonies on the plates of the second proper dilution degree;
d - Dilution Factor (the first dilution degree).
Example:

<table>
<thead>
<tr>
<th>Dilution degree</th>
<th>1:100 (the first dilution degree)</th>
<th>1:1000 (the second dilution degree)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of colonies</td>
<td>232,244</td>
<td>33,35</td>
</tr>
</tbody>
</table>

\[ N = \sum \frac{C}{(n_1 + 0, 1n_2) d} \]

\[ = \frac{232 + 244 + 33 + 35}{\left[2 + (0.1 \times 2)\right] \times 10^{-2}} = \frac{544}{0.022} = 24727 \]

The values mentioned above are round-up, and then represented as 25000 or 2.5×10^4.

7.1.3 If the colony numbers on the plates of all dilution degrees are all over 300, count the plates with the maximum dilution degree. For other plates, they shall be recorded as uncountable, and the results shall be obtained by multiplying the average colony number by the maximum dilution times.

7.1.4 If the colony numbers on the plates of all dilution degree are all less than 30, it shall be calculated by multiplication of average colony number on the minimum dilution degree plates by the dilution times.

7.1.5 If, for plates of all dilution degrees (including the original liquid samples), there is no colony growth, then it shall be calculated as multiplying the minimum dilution degree by a factor smaller than 1.

7.1.6 If, for plates of all dilution degrees, the colony number falls outside the range between 30 and 300, part of which are less than 30 or more than 300, then it shall be calculated for the plates whose colony number is closest to 30 or 300, as the average colony number multiply by dilution times.

7.2 Report of plate count

7.2.1 When the plate count falls within 100, it shall be rounded up and reported as two significant digits.

7.2.2 When the plate count is larger than or equal to 100, the third digit shall be rounded up, and take the first two digits, while the following digits are replaced by 0; it could also be indicated as exponential of 10, round-up and then take the two significant digits.

7.2.3 When all the plates are covered by spreading colonies, making it unable to calculate, it shall be reported as colony spreading.

7.2.4 When there are colonies growing on the blank control, the examination result is invalid.

7.2.5 For sampling by weight, CFU/g is applied as the report unit, while for sampling by volume, CFU/ml is applied as the report unit.

Method II Petrifilm™ Aerobic Count Plate Testing Method

8. Examination Procedures
Except that the agar culture medium plate is replaced by Petrifilm™ count plate, the others are the same as in Section 5.

9. Operation Procedures

9.1 Dilution of sample
Follow 6.1.1 and 6.1.2 to prepare 1:10 sample solution, then adjust the pH value to 6.6~7.2 with 1mol/L NaOH or 1mol/L HCl.

9.2 Inoculation
As per the estimation of sample contamination status, select 2 to 3 sample solutions with proper dilution degree for examination. Place the count plate on the even laboratory table, and remove the upper film, and absorb 1ml sample solution with pipette or micropipettor, drip perpendicularly in the
center of the count plate, then cover the upper film, it is allowed to let the upper film drop directly. However, it is forbidden to roll the upper film. Then, place the pressure pad (with the concave side facing downward) in the center of the upper film, press gently, to cover the sample solutions on the round culture film, and it is forbidden to twist the pressure pad. Remove the pressure pad, keep still for at least 1 min to solidify the culture medium. Two count plates are inoculated for each dilution degree.

9.3 Cultivation
Place the count plates with transparent side facing upward, horizontally in the incubator, which could be piled up to 20 pieces, with the same cultivation temperature and time as that in 6.2.

9.4 Counting
9.4.1 Counting is immediately carried out by the end of cultivation, it could be counted either with naked eyes or with bacteria colony counter, magnifying glass, Petrifilm™ automatic interpretoscope. Select the count plates with colony number between 30 and 300 for counting.

9.4.2 Count all the red colonies. When the concentration of bacteria is very high, the whole count plate shall become red or pink, then it shall be recorded as “uncountable”.

9.4.3 Sometimes, when the concentration of bacteria is very high, there is no detectable colony in the center of the count plate, however along the brim of the round cultivation film, there are many small colonies, whose results shall be recorded as “uncountable”; after further dilution of the sample, accurate reading could be obtained.

9.4.4 Some bacteria may liquefy the gel, causing local spreading or smear of the colony. If the liquefying phenomena interfere with counting, it could be counted of colonies in the area that did not liquefy to estimate the colony number.

10. Representation of Results
The same as that of Section 7.

Appendix A
(Normative Appendix)

Culture Media and Reagents
A.1 Plate count agar (PCA) culture medium
A.1.1 Ingredients
Tryptone 5.0g
Yeast Extract 2.5g
Glucose 1.0g
Agar 15.0g
Distilled Water 1000ml
pH 7.0±0.2
A.1.2 Formulation method
Add the above mentioned ingredients into distilled water, boil for dissolving, and then adjust the pH value. Distribute into tubes or conical beakers, autoclave at 121°C for 15 min.

A.2 Phosphate Buffer Solution
A.2.1 Ingredients:
KH₂PO₄ 34.0g
Distilled Water 500ml
pH 7.2
A.2.2 Formulation Method

Stock Solution: Measure 34.0 g KH$_2$PO$_4$ to dissolve in 500ml distilled water, adjust the pH value to 7.2 with about 175ml of 1mol/L NaOH solution, then dilute with distilled water to a volume of 1000ml, and then store in the refrigerator.

Diluent Solution: Measure 1.25ml of the Stock Solution, dilute with distilled water to 1000ml, distribute into proper containers, and then autoclave at 121℃ for 15min.