

COPYRIGHT NOTICE & PERMISSION TO USE

3M Food Safety has purchased the right to make AOAC® *Official Method of Analysis* **990.12** <http://www.eoma.aoac.org/> available to you in electronic form on Condition that you accept the following terms of use. Please read and respect these terms of use.

Copyright

AOAC INTERNATIONAL, Rockville, MD, USA owns the copyright of this Official Method publication. All rights are reserved.

Permission to Use

You may download an electronic file of this method publication for the purposes of viewing and/or printing one copy of the method. Neither the electronic file nor a hard copy print may be reproduced. You may not remix, transform, build upon or modify this material in any way for redistribution.

17.2.07

AOAC Official Method 990.12
Aerobic Plate Count in Foods
Dry Rehydratable Film
(Petrifilm Aerobic Count Plate) Method
First Action 1990
Final Action 1994

Results of the interlaboratory study supporting acceptance of the method:

Flour: $s_r = 0.225$; $s_R = 0.246$; $RSD_r = 5.3\%$; $RSD_R = 5.8\%$
Nuts: $s_r = 0.272$; $s_R = 0.674$; $RSD_r = 7.4\%$; $RSD_R = 18.4\%$
Shrimp: $s_r = 0.540$; $s_R = 0.615$; $RSD_r = 9.8\%$; $RSD_R = 11.1\%$
Spice: $s_r = 0.274$; $s_R = 0.303$; $RSD_r = 6.0\%$; $RSD_R = 6.6\%$
Turkey: $s_r = 0.278$; $s_R = 0.348$; $RSD_r = 5.3\%$; $RSD_R = 6.6\%$
Vegetables: $s_r = 0.310$; $s_R = 0.454$; $RSD_r = 6.3\%$; $RSD_R = 9.2\%$

A. Principle

See [989.10A](#) (see 17.3.03).

B. Apparatus

See [989.10B\(a\)](#) and (c)–(e) (see 17.3.03).

C. Reagent

Dilution water.—To prepare stock solution, dissolve 34 g KH_2PO_4 in 500 mL H_2O , adjust to pH 7.2 with 1M NaOH (ca 175 mL), and dilute to 1 L with water. To prepare buffered water for dilutions, dilute 1.25 mL stock solution to 1 L with boiled and cooled water. Autoclave 15 min at 121°C.

D. Preparation of Test Suspension

See [966.23B](#) (see 17.2.01).

E. Determination

Place dry-film aerobic count plate on flat surface. Lift top film and inoculate 1 mL test suspension onto center of film base. Carefully place top film down on inoculum. Distribute suspension over prescribed growth area with downward pressure in center of plastic spreader device (recessed side down). Leave plate undisturbed 1 min to permit gel to solidify. Incubate plates 48 ± 3 h at $35^\circ \pm 1^\circ\text{C}$.

In incubator, place plates in horizontal position, clear side up, in stacks not exceeding 20 units. Count plates promptly after incubation period. After incubation is complete, plates may be stored frozen (-15°C) up to 7 days. Avoid this as a routine practice.

Use standard colony counter for counting purposes. Magnifier-illuminator may also be used to facilitate counting. Colonies stain in various shades of red. Count all colonies in countable range (30–300 colonies).

To compute bacterial count, multiply total number of colonies per plate (or average number of colonies per plate if counting duplicate plates of same dilution) by reciprocal of dilution used. When counting colonies on duplicate plates of consecutive dilutions, compute mean number of colonies for each dilution before determining average bacterial count. Estimated counts can be made on plates with >300 colonies and should be reported as estimated counts. In making such counts, circular growth area can be considered to contain ca twenty 1 cm squares. To isolate colonies for further identification, lift top film and pick colony from gel.

Reference: *JAOAC* **73**, 242(1990).

Revised: March 2002