Comparison of Different Temperature Conditions for Microbial Enumeration of *Aspergillus brasiliensis* in the Preservative Efficacy Test

JUN USUKURA *

Fundamental Research Institute Division, MANDOM Corp. 5-12 Juniken-cho, Chuo-ku, Osaka 540-8530, Japan

Received 30 April, 2020/Accepted 3 August, 2020

The preservative efficacy test is an important method for assessing the antimicrobial effect of cosmetic products. In this study, the optimum conditions for the efficient microbial enumeration of *Aspergillus brasiliensis* were investigated. Cosmetic products, inoculated with *A. brasiliensis* spore suspensions, were cultivated at 22.5°C, 32.5°C, or 40°C and the detection rate and the number of colonies were determined using the pour culture method. There was no difference in the viable counts of visible colonies among different temperature conditions. However, the viable counts after 3 days of culture were significantly greater for the cultures maintained at 32.5°C or 40°C compared with those maintained at 22.5°C. This effect was attenuated in products containing fatty acids, which could inhibit fungal growth. Overall, these results demonstrate that cultivating *A. brasiliensis* at 32.5°C reduces the time required for enumeration in the preservative efficacy test. Thus, the results of this study are expected to help improve and expedite microbiological quality control in the cosmetic industry.

**Key words**: *Aspergillus brasiliensis* / Cosmetics / Preservative efficacy test.
The spores were then collected with a cotton bar and placed in a saline solution containing 2% Tween 80 and $10^6$ cells/mL, and filtered using sterilized gauze. Each cosmetic product, about 20 g, was weighed in a glass container and then inoculated with the prepared spore solution at a concentration of approximately $10^4$ cfu/g. The microbial enumeration of each sample was determined on days 7, 14, and 21 of storage at 22.5 °C. Microbial enumeration was conducted by pouring 1 g or 1 mL of the specimen into glucose peptone agar medium, containing lecithin and polysorbate. Microbial enumeration was calculated by counting the number of colonies on day 1, 2, 3, 5, or 7 after pouring the culture at 22.5°C, as the conventional test condition, or under 32.5°C and 40°C as the potentially improved condition.

Representative examples of typical plates for the lotion cultured under each temperature condition on day 21 (or day 7 for the facial wash) after A. brasiliensis spore suspension inoculation are shown in Fig. 1. At 22.5°C, the conventional condition for culturing A. brasiliensis, small colonies were visible on the plate by day 2 for the lotion and day 3 for the facial wash, which became more clearly visible by the next day. At 32.5°C or 40°C, the colonies tended to form about 1 day earlier than at 22.5°C. However, the colonies that were formed at 40°C tended to be smaller than those at 22.5°C or 32.5°C. Based on a previous report showing that A. brasiliensis colonies cannot grow bigger when cultured at temperatures above 30°C (Koide and Yosokawa, 1991), we predicted that culturing at a higher temperature than the typical conditions might reduce the colony size owing to growth inhibition.

However, there were no significant differences in the viable counts calculated from day 7 after pour culture was performed under each temperature condition. These results suggested that temperature influenced the

| Genus | Species | Lower limit | Optimum | Upper limit |
|-------|---------|-------------|---------|------------|
| Aspergillus | clavatus | 8 | 30 | 38 |
| | flavus | 8 | 35 | 42 |
| | fumigatus | 10 | 37 | 52 |
| | niger | 5 | 33 | 42 |
| | restrictus | 6 | 30 | 37 |

**TABLE 1.** Culture temperatures of Aspergillus species.

| Product category | n= | Addition |
|------------------|----|----------|
| Facial wash      | 3  | Fatty acid |
| Body wash        | 1  | Other    |
| Lotion           | 3  |          |
| Milk/Cream/Hair Wax | 3 |          |

**TABLE 2.** Test specimens.

**FIG. 1.** Appearance of A. brasiliensis colonies on days 1–7 after pour culture. Left: lotion on day 21 after inoculation; right: facial wash on day 7 after inoculation.
growth rate of colonies, but not the viable counts, based on the final number of the colonies.

The colony detection rates under each condition were calculated using the following formula: viable counts determined on each cultivating date/viable counts determined on day 7. The products containing fatty acid soap were distinguishable from other products with an evident delay in colony growth due to the inhibitory effect of fatty acids on microbial growth (Fig. 2, 3). Compared with cultures maintained at 22.5°C, the detection rate on each cultivating date was greater for cultures maintained at 32.5°C in all product categories. On day 2 of culture, the colony detection rates under high-temperature conditions (32.5°C or 40°C) were greater than 90%, whereas those at 22.5°C were less than 80%. In particular, the detection rate on day 3 of culture in the “other” products category (i.e., not fatty acids) significantly differed between 22.5°C and 32.5°C or 40°C culture conditions (p < 0.05). These results suggested that the cultivating temperature could

**FIG. 2.** Detection rates of *A. brasiliensis* in fatty acid-containing products on day 7 after culture in each temperature condition (◆: 22.5°C, ■: 32.5°C, ▲: 40°C). Vertical axis shows detection rate which is calculated from the following formula (viable count determined on each cultivating date) / (viable count determined on day 7). **: p < 0.05, *: p < 0.10

**FIG. 3.** Detection rate of *A. brasiliensis* in products without fatty acids (Others) on day 7 after culture in each temperature condition (◆: 22.5°C, ■: 32.5°C, ▲: 40°C). Vertical axis shows detection rate which is calculated from the following formula (viable count determined on each cultivating date) / (viable count determined on day 7). **: p < 0.05
increase the colony detection rate of *A. brasiliensis*. In addition, the growth rate at 40°C was higher than that at 22.5°C, suggesting that high temperature inhibited the colony size while promoting germination or growth (Fig. 2 and Fig. 3).

Overall, these results confirmed that the number of days to achieve visible detection of colony formation in all cosmetic product categories could be shortened by culturing at higher temperatures, such as 32.5°C or 40°C. Additionally, the final viable counts determined under high-temperature conditions on day 7 after inoculation were equivalent to those detected at the conventional culture temperature of 25°C.

These results suggest that microbial enumeration at 32.5°C may be the most suitable condition for determining the viable counts more rapidly when conducting the preservative efficacy test of *A. brasiliensis*. The results of this study can efficiently shorten microbiological quality control, thereby contributing to expediting the development of cosmetic products.

REFERENCES

Dawson, N. L., and Reinhardt, D. J. (1981) Microbial flora of in-use, display eye shadow testers and bacterial challenges of unused eye shadows. *Appl. Environ. Microbiol.*, **42**(2), 297-302.

Devilieghere, F., De Loy-Hendrickx, A., Rademaker, P. M., Pipeiros, Crozier, A., De Beets, B., Joly, L. and Keromen, S. (2015) A new protocol for evaluating the efficacy of some dispensing systems of a packaging in the microbial protection of water-based preservatives-free cosmetic products. *Int. J. Cosmet. Sci.*, **37**, 627-635.

International Organization for Standardization. (2012) ISO 11930. Cosmetics – Microbiology – Evaluation of the antimicrobial protection of a cosmetic product.

Ja’nos, V., Sa’ndor, K., Be a’ta, T., Jens, C.F., Giancarlo, P., Antonia, S., Martin, M. and Robert, A. S. (2007) *Aspergillus brasiliensis* sp. nov., a biseriate black *Aspergillus* species with world-wide distribution. *IJSSEM*, **57**, 1925-1932.

Japanese Pharmacopoeia, Seventeenth Edition. (2016) JP17. 2499-2501.

Koide, S., and Yasokawa, D. (2008) Growth prediction of mycelial mat and fruiting zone diameters of *Aspergillus niger* subjected to temperature changes (in Japanese). *Nippon. Shokuhin Kagaku Kogaku Kaishi*, **55**(7), 338-344.

Takatori, K. (1991) *Hitome de wakaru zusetsu kabi kensa·sousa manual* (in Japanese). pp. 20-21. TECHNO SYSTEM, Tokyo.