

# Augmentation of Sebaceous Lipogenesis by an Extract of Grifola frondosa (Maitake Mushroom) In Vivo and In Vitro



augments



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lipogenic actions for sebocytes.

# Introduction

Sebaceous lipids (sebum) play an important role in maintaining physiological functions by forming a biological barrier in the skin (Fig. 1). The decrease of sebum levels in the skin is thought to depress the barrier functions, and thereafter may be associated with the development of dry skin (xerosis) with a variety of complaints including a rough or scaly skin surface, and pruritus (Fig. 1). Therefore, a novel aspect to control sebaceous lipogenesis might be beneficial for the prevention of dry skin and sequential itching. In the present study, we investigated the effects of an ethanol extract of Grifola frondosa (Maitake) fruit body (Gripin® (Fig. 2) on sebum production in hamsters and humans and compared these effects with those of an ethanol extract of Agaricus blazei murrill (Agaricus) in vivo and in vitro.

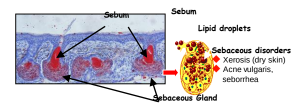


Fig. 1 Sebaceous glands in the skin

Left panel, oil red O and hematoxylin staining in the skin of hamster.





These results provide novel evidence that Gripin®

sebaceous lipogenesis in hamsters and humans in vivo and in

vitro. Thus, Gripin<sup>®</sup> is likely to be a unique anti-dry skin agent with

Fig. 3 Gripin<sup>®</sup> augments the formation of lipid droplets in hamster sebocytes A, control; B, Gripin<sup>®</sup> (400 µg/ml); and C, the ethanol extract prepared from Agaricus blazei murrill (400  $\mu$ g/ml). The cells were stained with oil red O.

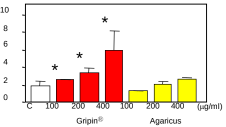
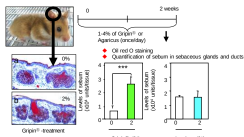
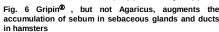


Fig. 4 Gripin<sup>®</sup> increases the intracellular levels of TG in hamster sebocytes

, significantly different from untreated cells (C) (p<0.05).</p>





Treatments of  $Gripin^{\mathbb{R}}$ and Agaricus at 1-4% were performed and similar results were obtained under these experimental conditions. Thus, typical data of 2% Gripin® and Agaricus treatment are shown. \*\*\*, significantly different from vehicle-treated cells (p<0.001).

Diacytglycerol acyttransferase (DGAT) activity. DGAT activity in hanster sebocytes treated with Gripin® was measured using 1,2-diolecyl glycerol, and [<sup>14</sup>C]palmitoyl-CoA as previously described (3, 4). In-vivo experiments in hansters. Auricles of 3 week-old male golden hamsters were topically treated with 4% Gripin® or Agaricus extract in 55% ethanol ad5% glycerol, or with the same volume of which for 14 days. After the treatments, the frozen tissue sectors were stained with 0.5% oil red 0 and counterstained with Mayer's hematoxylin estimation as described above. The relative intervisity (unterlisticus) of oil red 0 staining in selection and because due to the days and the days and the days of the posterior anabove. The experimental protocol was approved by the Committee of Animal Care and Use of Tayo University of Patamaray and Life Sciences. In-vivo trial of Gripin® cream in volunteers. Gripin® creams (0, 1 and 1%) and vehicle new ere topically treated three mass a day for 1 days on the posterior anabovariatis of seven healthy volunteers (six male and one female: age: 22-41 years). The skin was dues that because of the active surface was extracted twice with actore using stainless cups at 1 h after the wijng. The sebum extracts were subjected to latoscan and the mean using in components were measured as described above.

described above. **Statistical analysis.** Data are presented as mean  $\pm$  standard deviation (SD), and were analyzed by a one-way analysis of variance (ANOVA) and by the Fisher test for multiple comparisons. A value of p<0.05 was considered to indicate a statistically significant

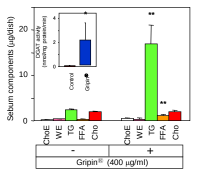


Fig. 5 Gripin® preferentially augments TG synthesis by increasing DGAT activity in hamster sebocytes Inserted panel, DGAT activity. \* and \*\*, significantly different

from untreated cells (Control) (p<0.05 and 0.01, respectively). ChoE, cholesterol ester; WE, wax ester; TG, triacylglycerols; FFA, free-fatty acids; and Cho, cholesterol,

#### Table 1 The augmentation of sebum components in Gripin® cream-treated skin of healthy volunteers

Sebum components	The number of volunteers whose lipid levels in the skin were increased	
	0.1% Gripin®	1% Gripin <sup>®</sup>
Squalene	6/7	6/7
Wax ester	4/7	4/7
Triacylglycerols	3/7	3/7
Free-fatty acids	4/7	6/7

When 0.1 and 1% Gripin® cream, and vehicle cream were topically treated three times a day for 14 days on the antebrachial regions of seven healthy volunteers (six males and one female, age: 22-41 years) , a thin-layer chromatographic analysis was performed to quantify the levels of sebum components on the skin surface.

### References

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(Maitake) fruiting body (Gripin®) , Grifola frondosa (Maitake); B, dry powder of Maitake; and C, ethanol extract of Maitake, designated Gripin®

Fig. 2 Ethanol extract of Grifola frondosa

## Results

- When hamster sebocytes were treated with 1. Gripin® , the intracellular lipid droplet formation was augmented in dose dependent manner (100-400 µg/ml) (Fig. 3).
- 2 Gripin® preferentially augmented the synthesis of triacylglycerols (TG), a major sebum component, in hamster sebocytes (Figs. 4 and 5). The augmentation of TG production resulted from the increase of cellular DGAT activity in hamster sebocytes
- (Fig. 5, inserted panel). The topical treatment of hamster auricles with З 1-4% Gripin® augmented sebum accumulation in sebaceous glands (Fig. 6).
- Another ethanol extract prepared from Agaricus blazei murrill (Agaricus) showed less or no effect on sebaceous lipogenesis in hamsters in vivo and in vitro (Figs. 3, 4, and 6)
- In most of the volunteers treated with 0.1 and 1% Gripin  ${}^{\mathbb{R}}$  cream, the levels of sebum components including TG, squalene, freefatty acids, and cholesterol were augmented when compared with the vehicle cream treated skin.

# Materials & Methods

Treatment. Hamster sebocytes were treated with Gripin<sup>®</sup> (100-400 µg/ml) or the ethanol extract of Agaricus blazei murrill (Agaricus) (100-400 µg/ml) in DMEMF12 supplemented with the serumes for up to 7 days. Oil red O stalming. After the treatment of sebocytes with Gripin<sup>®</sup> or Agaricus, cells

Of red Stating. After the treatment of sebocytes with Gripin<sup>®</sup> or Agaricus, cells were stained with 0.3% oil red 0 in isopropanoidstilled H<sub>2</sub>O (32, vd·vel), and then viewed with a light microscope furnished with a digital camera. The cells were also counterstained with Mayer's hematoxylin solution. Quantitative analysis of sebaceous lipids. The sonicated-cell lysates were subjected to an automatic thin-layer chromatography, latorscan, as previously described (1). Otherwise, the sonicated-cell lysates were used to measure the layed oil microcellular TG using Liquitech TG-II according to the manufacture's instructions. The intracellular DNA content was measured using automatic measure the seried oil microcellular G distribution to enter the series the layed oil microcellular G distribution to the series the layed oil microcellular G distribution to the series the layed oil microcellular G distribution to a series the layed of microcellular G distribution to a series the layed of the manufacture's instructions. The intracellular DNA content was measured using automation taken to perm DNA (62:100 µg/m) and 3.5-diaminobenzoic acid dihydrochloride as previously described (2).

Gripin<sup>®</sup> (%)