The effect of saturated phospholipids on human skin assessed with shotgun lipidomic analysis

Dorothea Gutekunst 1*, Christian Klose 2, Christoph Heidecke 3, Peter van Hoogevest 1

1 Phospholipid Research Center, Germany; *dgutekunst@phospholipid-institute.com; 2 Lipotype GmbH, Germany; 3 Lipoid GmbH, Germany

Introduction

Phospholipids (PLs), as natural and well-tolerated excipients, are widely used in dermal drug delivery systems. Unsaturated PLs have a transition temperature (Tm) below 0°C, whereas the Tm of saturated PLs is much higher. When applying PLs topically, the temperature of skin dictates the physical state and influences the penetration as well as delivery properties of phospholipid-based formulations. Saturated PLs (e.g. PHoSPHoLIPon® 90 H) are in a gel-like state on the skin, caused by their Tm around 50°C [1].

Regarding their rigid structure it can be assumed, that saturated PLs accumulate in upper epidermal layers. Until now, this behaviour is not well investigated for semi-solid phospholipid-based formulations, like oil-in-water (o/w) emulsions, and especially not with respect to structural properties of PLs. One reason might be the lack of suitable analytical methods to identify and quantify the exogenous PLs besides endogenous skin PLs and to follow the metabolism of the PLs in the skin.

Materials & Methods

TEST FORMULATION

o/w cream with 6 % PHoSPHoLIPon® 90 H

PHoSPHoLIPon® 90 H composition:
Phosphatidylcholine (hydrogenated) ≥ 90 %
Σ (C16:0, C18:0) ≥ 98 %
Σ (C18:1, C18:2, C18:3) ≤ 2 %

IN-VIVO PILOT STUDY

This study was performed in accordance with the Ethical Consideration for cosmetic products. The test formulation was manufactured in compliance with cosmetic Good Manufacturing Practice and can therefore be considered as safe for human testing.

Performance:

Healthy female volunteers (18 – 45 years of age)
Application 2x/d
Tape stripping

Tapes five and ten were analysed with Lipotype shotgun lipidomics technology, according to the following workflow [2]:

Sample preparation
Sample infusion
MS analysis
Lipid identification
Data analysis

Results

RECOVERY OF APPLIED SATURATED PHOSPHOLIPIDS

The left graph presents an overview of PC-subspecies characterised by different fatty acid composition. After treatment with PHoSPHoLIPon® 90 H-based formulation, the volunteers (1–3) presented a similar profile of saturated fatty acid species (B). The test formulation induced an increased concentration of characteristic subspecies such as 16:0;0 – 18:0:0 and 18:0:0 – 18:0:0. An unequivocal discrimination between exogenous and endogenous PC was possible because of the small amount of stearic and palmitic acid in skin-derived PC species (A).

Consideration of the natural variability of the skin lipid composition, the results show a good reproducibility. On average, 60 mol % of exogenous PC was detected in the fifth skin strip. The detected concentration of exogenous PC decreased with increasing depth. 48 mol % were finally measured in tape 10.

Discussion & Conclusion

This study successfully confirms that hydrogenated PLs penetrate into the SC. A high amount in upper skin layers and a decrease in deeper layers demonstrate an accumulation of saturated PC in upper epidermal layers. This skin distribution was already observed by Blume [2000] [3] and can be used to support the skin barrier function or to influence the distribution of a co-formulated drug substance.

For the first time the detection of exogenous PLs besides endogenous skin lipids was demonstrated using the Lipotype shotgun lipidomics technology. It offers the opportunity for further investigations and will be used to assess the penetration properties as well as metabolism of (un)saturated PLs.

References