



Detection of highly abundant small molecules in the stratum corneum of healthy young women using desorption electrospray ionization-mass spectrometry imaging

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1	Manuscript type: Research paper
2	Detection of highly abundant small molecules in the stratum corneum of healthy young
3	women using desorption electrospray ionization-mass spectrometry imaging
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27 Abstract

Aging reflects skin appearance drastically, which reduces skin juvenescence. However, the 28 29 small molecules underlying skin juvenescence have not been well studied. We aimed to 30 explore the molecules potentially responsible for young-looking skin. Eleven healthy women 31 aged 24–59 years were recruited and divided into young and middle-aged groups. Multiple 32 layers of the stratum corneum from the cheek area were taken by tape-stripping, followed by 33 desorption electrospray ionization mass spectrometry imaging (DESI-MSI). Overall, five 34 molecules (*m*/*z* 284.33, 340.39, 488.39, 628.37, and 863.65) were highly abundant in young 35 subjects. Among them, *m*/*z* 284.33 and 340.39 were dominantly detected in each layer of all 36 young subjects. Interestingly, *m*/*z* 488.39 and 628.37 were prominent in young subject 4. All 37 of these molecules were gradually decreased in the successive layers of the stratum corneum 38 in subjects 3 and 4 of the young group. These molecules could be endogenous, co-related with 39 youthful skin, or retained from topical cosmetics. Extensive research is indispensable to 40 characterize them and find the relationship between these molecule's retention capacity in 41 the stratum corneum with different skin parameters. Our findings provide a novel perspective 42 on young skin that could be advantageous in future cosmetic formulations to improve skin 43 juvenescence.

44

45 **Keywords:** Stratum corneum, DESI-MSI, molecules, skin juvenescence.

47 Introduction

Skin is the most visible external attribute of the human being¹⁾. It is one of the critical 48 indicators of aging^{2,3)}. The aging process significantly impacts the skin's young appearance^{4,5)}. 49 50 The appearance of the skin is highly correlated to the stratum corneum, which is the outer 51 layer of the skin. Stratum corneum, in turn, consists of multiple layers in it⁶⁾. Improved skin 52 texture and blood flow, prevention of pigmented spot formation, and increased moisture in 53 the stratum corneum are the external parameters for skin to have beautiful, young-looking 54 skin⁷⁾. Keratin is a major component of stratum corneum and constitute more than 85% of the total proteins in it⁸⁾. Keratin and associated proteins are deformed over the aging process^{9,10)}. 55 56 However, until now very few studies on the other small molecules in the stratum corneum 57 underlying skin juvenescence have been done. In this connection, we aimed to discover the potential small molecules in the stratum corneum that might be responsible for skin 58 59 juvenescence. While it is feasible to investigate human skin that has been surgically resected 60 during some diseases, however in healthy individuals, skin sampling is desired to be noninvasive¹¹⁾. Several studies have recently been employed in normal conditions to analyze 61 skin^{7,12,13)}. Recently, several techniques have been employed to analyze skin conditions by 62 imaging the skin based on its external appearance^{7,14-16)}. Apart from these, mass 63 64 spectrometry-based studies are a more advanced technique due to providing molecular information as well^{17,18}. Among the mass spectrometric methods, the mass spectrometry 65

66	imaging (MSI) analysis adds a new dimension to the biomolecular distribution analysis in the
67	skin ¹⁹⁻²³⁾ . Currently, multiple MSI modalities are in use, like matrix-assisted laser
68	desorption/ionization (MALDI)-MSI, secondary-ion mass spectrometry (SIMS) imaging, and
69	DESI-MSI. Compared to other MSI modalities, DESI-MSI is a rapidly in-situ molecular imaging
70	technique because it does not require matrix coating on the sample, unlike MALDI-MSI.
71	Moreover, DESI-MSI is used to detect compounds and visualize their spatial distribution in the
72	sample in ambient conditions $^{11,24-27)}$. For straightforward, rapidness, and direct surface
73	analysis capacity in ambient condition, in this study, we employed DESI-MSI to detect small
74	molecules of stratum corneum, which are highly abundant in young-aged skin ²⁸⁾ .
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75 76 77 78 79	Materials and Methods Reagent and chemicals Acetonitrile, methanol, formic acid, ultrapure water, and isopropanol were purchased from FUJIFILM Wako pure chemical industries (Osaka, Japan). Leucine enkephalin was purchased from Waters Corporation (Milford, MA, USA). Sodium formate was obtained from

Medicine (Code: 19-123). Study subjects were provided with sufficient information and signed informed consent forms before collecting tape-stripped stratum corneum samples. Each subject's stratum corneum sample was anonymized by assigning a unique numerical subject number.

87 Study subjects and tape stripped stratum corneum samples collection

88	In this study, 11 healthy Japanese female volunteers participated. Only individuals who
89	participated in this testing schedule could consent to the stratum corneum sampling and other
90	test items, and not suffering from skin diseases like eczema or other pathological conditions
91	were selected. Subjects who recently received special facial care or medication that may affect
92	the sample collection site (face) were excluded from this study. In addition, only women were
93	chosen as subjects to eliminate sexual differences between men and women. Six subjects
94	were categorized as young, aged between 20-30s. (subject number 1 to 6 in the young group).
95	The other five subjects were between the ages of 40-60s and categorized as middle-aged
96	(subject number 7 to 11 in the middle-aged group) (Table 1). As a sampling area, we selected
97	the cheek area of the face. Redness, pores, and spots were scored by photo evaluation, while
98	skin transparency was scored by visual evaluation. The redness, pores

Table 1. Demographic data of the donors of tape stripped stratum corneum sample

Subject group	Subject ID	Age (years)	Redness (0 to 2 scale)	(0 to 2	(0 to 2	Skin transparency (0 to 4 scale)	
	1	27	1.5	1.5	0.5	3.5	Redness, many bumps due to acne scars, normal skin tone
	2	30	1.5	1.5	0.5	3.5	Many bumps due to worsening acne, normal skin tone
Young-	3	31	0.5	0.5	0.5	3.5	Many bumps due to acne scars, normal skin tone
aged	4	25	0.5	0.5	0	0	Brighter skin tone, almost no pigmentation, less uneven skin surface
	5	28	0	0	0.25	1	Lighter skin tone with unevenness due to fine pigmentation, less uneven skin surface
	6	24	0	0	0	1	Lighter skin tone, few bumps on the skin surface, but bumps near the nose
	7	59	0	0	2	3.5	Uneven skin tone due to hyperpigmentation, skin surface irregularity is not worse than that of the same age group, skin tone is darker
Middle-	8	55	0	2	2	3.5	Many uneven colors due to pigmentation, uneven skin surface due to flowing texture, dark skin tone Less unevenness in skin tone due to pigmentation than
aged	9	45	0.5	0.5	0.5	2	the same age group, skin surface irregularity about the same as the same age group, lighter skin tone
	10	41	0.5	0.5	0.5	2	Uneven skin tone due to pigmentation, less uneven skin surface than same age group, lighter skin tone
	11	43	0	0	0.25	2	Lighter skin tone with very little unevenness due to pigmentation

100 and spots scores were scored on a 5-point scale from 0 to 2 with 0.5 increments. The lower 101 score of redness, pores, and spots indicates better skin condition and is considered to have a 102 youthful appearance of the skin. The skin transparency score was scored on a 9-point scale 103 from 0 to 4 with 0.5 increments. Lower scores indicated higher skin transparency. The skin 104 with more increased skin transparency is typically considered to have a youthful appearance 105 of the skin. These redness, pores, spots and skin transparency score evaluation was constructed based on the "Skin Aging Atlas" exclusive to Asian type²⁹. Before stratum 106 107 corneum sample collection, each subject's face was thoroughly washed with the same 108 (commercial) mild cleanser (Table S1) using tap water. After face washing, the subjects were 109 allowed to acclimate in a constant temperature and humidity-maintained room (room 110 temperature 20 \pm 2 °C, humidity 50 \pm 5%) for 15 minutes at least. The stratum corneum 111 samples (horny layers of the epidermis) were then collected by 1 cm² tape-stripping by 112 cellophane tape (CELLOTAPE[™], CT-18, NICHIBAN Co., Ltd., Tokyo, Japan) from the cheek four 113 times from the same spot. Although the thickness of stripped stratum corneum samples was 114 not measured, we tried collecting stratum corneum samples from each subject with a similar 115 method to minimize the removed stratum corneum thickness variation between subjects. The 116 first stripping refers to the stratum corneum's outermost layer, whereas subsequent strippings 117 refer to the stratum corneum's inner layers. In all cases, the tape-stripped samples were 118 preserved at -80 °C until mass spectrometric analysis.

119 DESI-MSI analysis

For DESI-MSI of the stratum corneum sample, tape-stripped samples were mounted on regular glass slides (Matsunami Glass Ind., Ltd., Kishiwada, Japan). A double-sided tape (Conductive tape assy, 241-08728-92, Shimadzu Corporation, Kyoto, Japan) was attached to the glass slide before attaching the non-adhesive side of the stripped tape. The analysis was

124 conducted with a quadrupole time-of-flight (Q-TOF) mass spectrometer (Xevo G2-XS Q-TOF, 125 Waters Corporation, Milford, MA, USA) in positive ionization mode. The selected areas on the 126 glass slide were scanned with a scan rate and pixel size of 200 μ m/sec and 200 μ m × 200 μ m, 127 respectively. A solvent pump (ACQUITY UPLC Binary Solvent Manager, Waters Corporation, 128 Milford, MA, USA) was used to supply the solvent (98:2 methanol/water, v/v) at a flow rate of 129 2 μL/min. Mass resolving power and mass window were set at 20000 and 0.02 Da, respectively. 130 The DESI source conditions were optimized as follows: (i) capillary voltage of 3.0 kV, (ii) 131 nitrogen gas pressure of 0.4 MPa, and (iii) inlet temperature of 120 °C. Analyzer mode was set 132 as "sensitivity." Mass spectra were collected in a mass range of m/z 100 to 1000. The sodium 133 formate solution (500 μ M) in isopropanol: water (90:10, v/v) was used to calibrate the DESI 134 mass spectra externally, and the detector setup was performed using leucine enkephalin 135 solution (500 µM). The lock mass correction option was used for mass accuracy corrections 136 using m/z 309.2036 (Na⁺ adduct of diisopropyl sebacate, a typical background peak in mass 137 spectrometry).

138

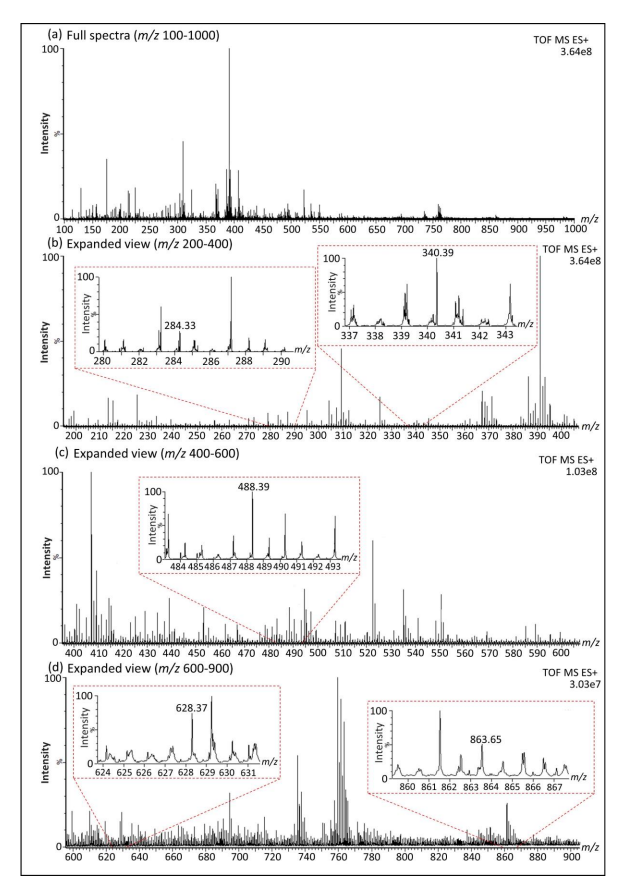
139 Data analysis

140 The data acquisition and processing were made using the MassLynx (Version 4.1; Waters 141 Corporation, Milford, MA, USA) program for DESI-MSI experiments. Raw data derived from the DESI-MSI experiment was imported into HDImaging software (Version 1.4; Waters 142 143 Corporation, Milford, MA, USA). The candidate selection was performed from the list of 300 144 most abundant *m*/zs generated in HDImaging software and manually compared the DESI-MSI 145 ion image between groups (Fig. S3). Thus, we have considered the average intensity between 146 groups in picking candidates. Background peaks were excluded considering the molecule's 147 distribution in the negative control tape sample. This negative control tape sample was the only tape sample attached to the glass slide, hence free from the stratum corneum sample.
Region of interest (ROIs) was manually drawn on each sample area to calculate average
intensity using HDImaging software. Average signal intensities of individual pixels of the ROIs
(Fig. S1b) were then compared for relative abundance.

152 Results

153 DESI-MSI detected highly abundant molecules in young-aged subjects

154 In positive ion mode, we analyzed the DESI ion distribution of tape-stripped stratum 155 corneum samples. We explored the most abundant 300 DESI-MSI peaks between m/z 100 to 156 m/z 1000 and, on average, found five molecules (m/z 284.33, m/z 340.39, m/z 488.39, m/z 157 628.37, and m/z 863.65) are highly abundant in the stratum corneum of the young-aged subject compared to the middle-aged subjects (Fig. 1b to d and Fig. 2b to k). Among them, 158 *m/z* 284.33 was 122%, *m/z* 340.39 was 196%, *m/z* 488.39 was 345%, *m/z* 628.37 was 265%, 159 160 and m/z 863.65 was 137%, highly abundant in average intensity in young-aged subjects 161 compared to middle-aged subjects. *m*/z 414.43 is a molecule shown here as a control, which 162 was not much different in distribution among the young and middle-aged groups (only 7% higher in the young group) (Fig. 2I, m). Among all detected candidate molecules, *m*/z 284.33 163 164 and m/z 340.39 were prominently detected in each layer of all young subjects (Fig. 2b, d and 165 Fig. S2a to h).



166 Fig. 1. Average DESI-MSI mass spectrum from tape stripped stratum corneum.

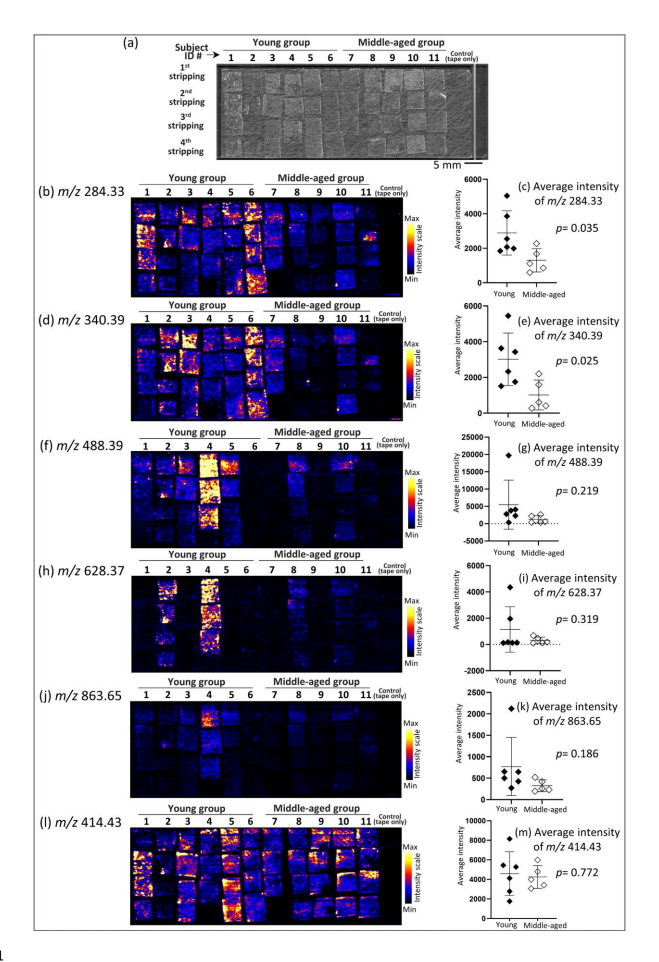
167 Young-aged subject 4 showed the most prominent distribution of m/z 488.39 and m/z 628.37

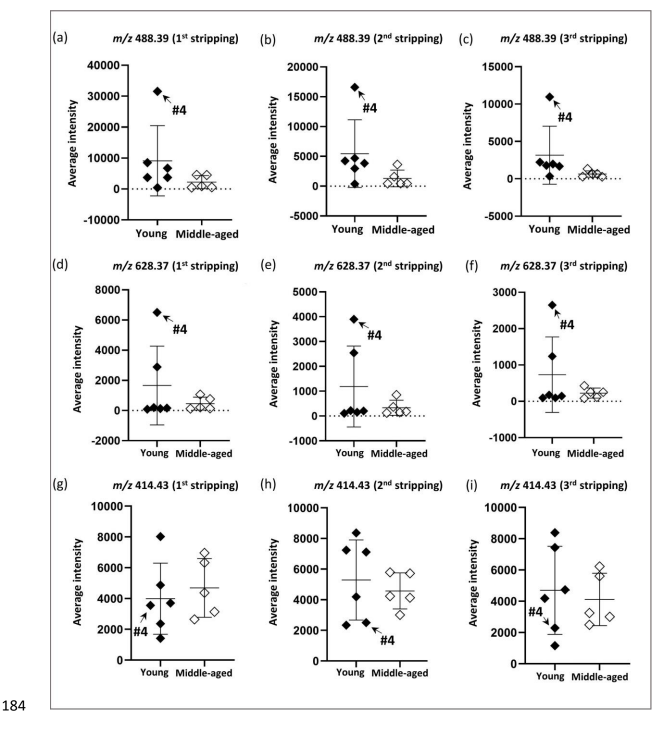
- 168 Although the candidate molecules were highly abundant on average in the young subjects,
- 169 we discovered the most prominent distribution of m/z 488.39 (Fig. 3a to c) and m/z 628.37
- 170 (Fig. 3d to f) in subject number 4 from the young-aged panel. By analyzing the average
- 171 intensity of each stratum corneum layer, we also discovered that the intensity was gradually
- decreasing in the deeper layers of the stratum corneum (Fig. 2f, h).
- 173

174 Candidate molecules were gradually decreased in deeper layers of the stratum corneum of

175 young subjects 3 and 4

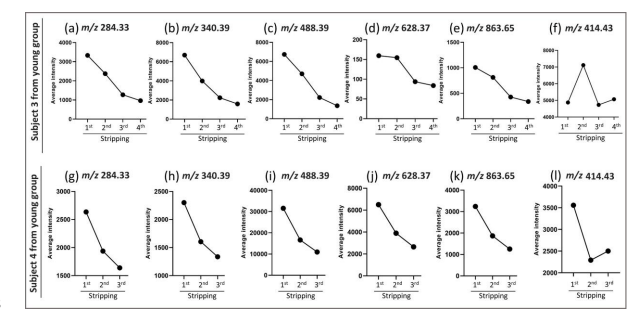
176 Although, on average, all candidate molecules are rich in young-aged subjects, 177 interestingly, they are most abundant in the first stripped stratum corneum of the young 178 subject 3 (Fig. 4a to e) and 4 (Fig. 4f to j). These molecule's abundance is gradually decreased 179 in the subsequent strippings. That refers to the gradual decrease of the candidate molecules 180 in the deeper layers of the stratum corneum of the young-aged subjects 3 and 4.





185 Fig. 3. Average intensity of *m*/*z* 488.39, *m*/*z* 628.37, and *m*/*z* 414.43 in each time of tape





188

Fig. 4. Stratum corneum's layer-wise average intensity of the candidate molecules in subject
3 and subject 4 from the young group.

192 Discussion

193 In this investigation, we could detect abundant small molecules from the stratum corneum of young, healthy individuals using the robust measurement ability of DESI-MSI^{30,31)}. Unlike 194 195 earlier studies that employed DESI-MSI to explore stratum corneum molecular abundance, our study focused on the cheek area of the face as a representative part of the body to study 196 age-dependent molecular abundance²²⁾. Among the candidate molecules, m/z 284.33 and 197 *m*/*z* 340.39 prominently exist in all stratum corneum of all young subjects (Fig. 2b, d and Fig. 198 S2a to h). Our finding surmises that these molecules might be highly related to skin 199 200 juvenescence. However, our study is limited to positive-ion mode-based detection of the 201 potential candidate molecules highly relevant for young-aged skin. The addition of negative 202 ion mode, molecular identification by comprehensive analysis using liquid chromatography-203 mass spectrometry, liquid chromatography with tandem mass spectrometry, and further 204 functional studies are required to determine these molecule's relationship with young-aged 205 skin³²⁾. Interestingly subject 4 of the young-aged subjects showed a most prominent 206 distribution of *m*/*z* 488.39 and *m*/*z* 628.37 in her stratum corneum (Fig. 3a to f). This most 207 prominent abundance reflects the unique properties of this young subject, who has a very 208 smooth skin surface with no pigmentation (Table 1). In the future, it would be beneficial to 209 deepen the research on this subject to find out the exact relationship between its skin 210 parameters and the retention capacity of those molecules. Also, we observed a gradual 211 decrease of the candidate molecules only (Fig. S4 a to f) in the subsequent tape strippings of 212 subjects 3 and 4. We surmised that some specialty, like reduced cell numbers in the deeper layers of the stratum corneum of these two subjects, might reflect such distributions²²⁾. In the 213 214 future, it would be informative to study the exact relationship between these subject's 215 stratum corneum and the candidate molecule's gradual decrease.

This preliminary study shows the differential abundance of some specific molecules in the stratum corneum of young-aged subjects. This is also supported by the previous studies where it was evident that the property of the stratum corneum, like age and skin condition, plays a role in the compositional difference in the skin^{9,10)}. This study is the groundwork for the relationship between skin condition and underlying molecular distribution in the stratum

221	corneum. We successfully detected higher distribution of some small molecules in the young
222	subjects. In essence, it is indispensable to identify and characterize the candidate molecules
223	to consider them for inclusion in product development for promoting skin juvenescence. In
224	the future, our findings may support establishing novel candidate molecules for the skin care
225	product industry to promote or retain skin juvenescence.
226	
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231	Conflicts of Interest
232	The authors declare no conflict of interest. The funders remain neutral in study design,
233	data collection and analysis, manuscript writing, or publication decisions.

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320	

321 Figure legends

322 Fig. 1. Average DESI-MSI mass spectrum from tape stripped stratum corneum.

(a) full positive-ion mass spectrum ranged *m/z* 100-1000. (b) Expanded view of masses ranged
200-400 showing detection of *m/z* 284.33, and *m/z* 340.39. (c) Expanded view of masses
ranged *m/z* 400-600 showing detection of *m/z* 488.39, and (d) Expanded view of masses
ranged *m/z* 600-800 showing detection of *m/z* 628.37, and *m/z* 863.65.

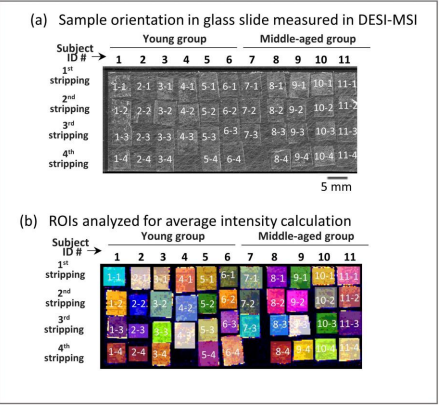
Fig. 2. Prominent distributions of highly abundant molecules in the tape-stripped stratum corneum samples from young and middle-aged subjects.

(a) Stratum corneum sample orientation in glass slide. (b) DESI-MSI ion images of m/z 284.33, 329 330 (c) average intensity comparison of m/z 284.33 between young and middle-aged subjects. (d) 331 DESI-MSI ion images of m/z 340.39, (e) average intensity comparison of m/z 340.39 between 332 young and middle-aged subjects. (f) DESI-MSI ion images of *m/z* 488.39, (g) average intensity 333 comparison of m/z 488.39 between young and middle-aged subjects. (h) DESI-MSI ion images 334 of m/z 628.37, (i) average intensity comparison of m/z 628.37 between young and middle-335 aged subjects. (j) DESI-MSI ion images of m/z 863.65, (k) average intensity comparison of m/z336 863.65 between young and middle-aged subjects. (I) DESI-MSI ion images of m/z 414.43, a 337 compound showing similar distribution in young and middle-aged subjects, (m) average 338 intensity comparison of m/z 414.43 between young and middle-aged subjects. All values are 339 presented as mean ± SD and p values were calculated between groups by Student's unpaired 340 two-tailed t-test.

Fig. 3. Average intensity of *m/z* 488.39, *m/z* 628.37, and *m/z* 414.43 in each time of tape stripped stratum corneum samples.

(a) Average intensity of *m/z* 488.39 in first, (b) second, and (c) third stripped stratum corneum.
Average intensity of *m/z* 628.37 in first (d), second (e), and third (f) stripped stratum corneum.

- Average intensity of *m*/*z* 414.43 in first (g), second (h), and third (i) stripped stratum corneum
- as control. All values are presented as mean ± SD.
- Fig. 4. Stratum corneum's layer-wise average intensity of the candidate molecules in subject
- 348 **3 and subject 4 from the young group.**
- 349 (a) Average intensity of *m/z* 284.33, (b) *m/z* 340.39, (c) *m/z* 488.39, (d) *m/z* 628.37, (e) *m/z*
- 350 863.65, and (f) *m/z* 414.43 (as control) in young subject 3. Average intensity of (g) *m/z* 284.33,
- 351 (h) *m/z* 340.39, (i) *m/z* 488.39, (j) *m/z* 628.37, (k) *m/z* 863.65, and (l) *m/z* 414.43 (as control)
- in young subject 4.



355 Fig. S1. ROIs drawn to measure average intensity. (a) sample orientation in glass slide

356 measured in DESI-MSI. (b) ROIs drawn.

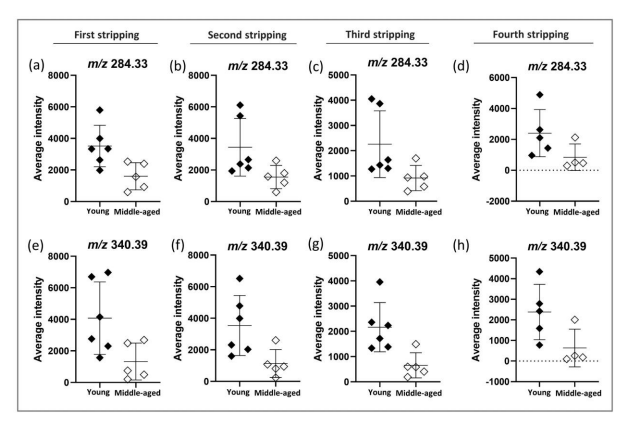


Fig. S2. *m/z* 284.33 and *m/z* 340.39 were prominently detected in each layer of all young
subjects. All values are presented as mean ± SD.

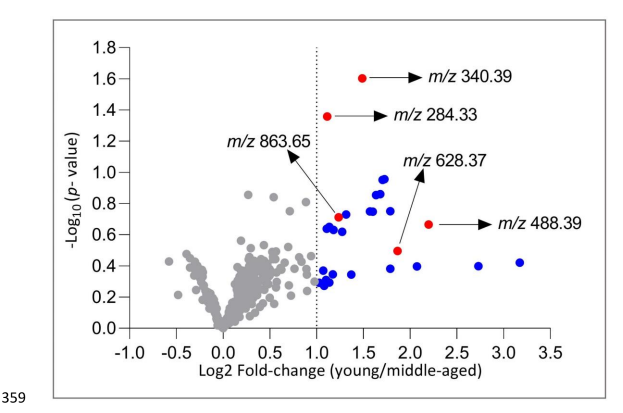


Fig. S3. Volcano plot compares the molecules from young and middle-aged subjects. Log2 fold changes and their corresponding –Log₁₀ *p*-values of 300 most abundant molecules generated by HDImaging software were taken for construction of the volcano plot. Molecules increased more than 2-fold are depicted in blue dots, among them which molecules are detected in all samples and average intensity was higher in young groups are depicted in red dots. All other molecules are not found to be relevant to this study and depicted as gray dots. Student's unpaired two-tailed t-test was used for the volcano plot construction.

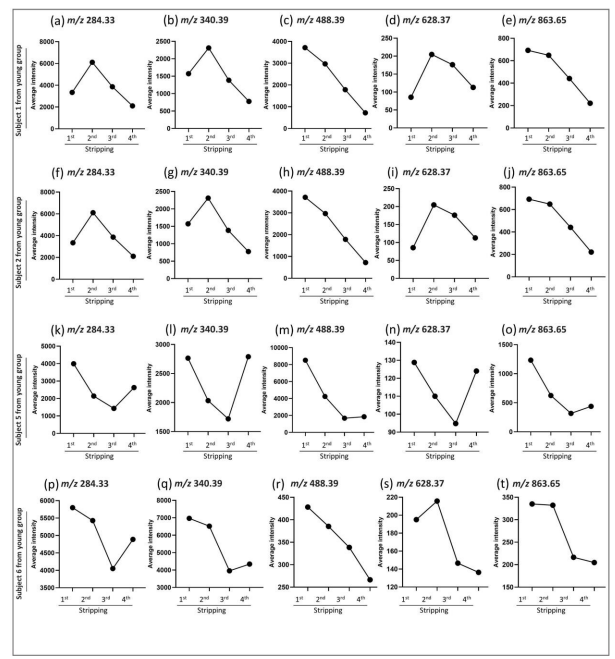


Fig. S4. Stratum corneum's layer-wise average intensity of the candidate molecules in
subject 1, 2, 5 and 6 from young group. Average intensity of *m/z* 284.33 (a), *m/z* 340.39 (b), *m/z* 488.39 (c), *m/z* 628.37 (d), and *m/z* 863.65 (e) in young subject 1. Average intensity of *m/z* 284.33 (f), *m/z* 340.39 (g), *m/z* 488.39 (h), *m/z* 628.37 (i), and *m/z* 863.65 (j) in young
subject 2. Average intensity of *m/z* 284.33 (k), *m/z* 340.39 (l), *m/z* 488.39 (m), *m/z* 628.37
(n), and *m/z* 863.65 (o) in young subject 5. Average intensity of *m/z* 284.33 (p), *m/z* 340.39
(q), *m/z* 488.39 (r), *m/z* 628.37 (s), and *m/z* 863.65 (t) in young subject 6.

- 374 Table S1. Constituents of the cleanser used for the face washing before stratum corneum
- 375 sample collection.

Cleanser constituents
Water (aqua)
Mineral oil (paraffinum liquidum)
Butylene glycol
Glycerin
Alcohol
Polysorbate 80
Glyceryl stearate
Ascorbyl tetraisopalmitate
Lactobacillus/Soybean extract ferment filtrate
Tocopherol
Behenyl alcohol
Carbomer
Cetearyl alcohol
Dimethicone
Lauroyl lysine
PEG-20 glyceryl triisostearate
Potassium cocoyl glycinate
Potassium hydroxide
Silk powder (Serica powder)
Sodium lauroyl glutamate
Stearic acid
Ethylparaben
Methylparaben
Phenoxyethanol