Development of a Global Subjective Skin Aging Assessment score from the perspective of dermatologists

Punnapath Buranasirin1, Krit Pongpirul2,3* and Jitlada Meephansan1

Abstract

Objective: Several skin aging assessment scales exist but no standard scale is widely used. Dermatologists may be the most appropriate persons for skin assessment but their perceptions regarding the signs of skin aging are unexplored. To develop a simple global skin aging assessment score from the perspective of dermatologists, an online questionnaire, the Thai Dermatologist Survey of Skin Aging Assessment, was conducted from October to December, 2016. Twenty-nine signs with published evidence of their relevancy to skin aging were included. Certified dermatologists were asked to score each sign using a 5-point Likert scale. Descriptive statistics and exploratory factor analysis (EFA) were used for data analysis.

Results: Of 213 randomly selected dermatologists, 145 responded to the survey. EFA revealed 3 important factors related to skin aging: Factor 1 comprised 8 signs related to atrophy (deep/superficial wrinkles, eye bags, lax appearance, etc.); Factor 2 comprised 7 signs related to discoloration (freckles, lentigines, melasma, etc.); and Factor 3 comprised 3 malignant skin lesions. The Global Subjective Skin Aging Assessment (GS2A2) score is a simple numerical score that can be used to evaluate the anti-aging effects of a cosmetic product or dermatologic intervention. This score should be tested further for validity and reliability.

Keywords: Skin aging assessment, Dermatology, Attitude

Introduction

Nowadays, the aging society is increasing and concerns about aging sequence have started to receive special attention. The primary target of an aging study is to increase the quality of elderly life and to prevent age-related diseases. Especially, aged skin is more interesting because the skin is the most noticeable sign of the aging mechanism and demonstrative of human health, as it seems to prognosticate the systemic illness and prognosis.

Chronologic age and solar damage are the most important intrinsic and extrinsic factors associated with skin aging. Skin aging characteristics vary ethnically among races. No one is spared from skin wrinkling, sunspots, uneven skin color, vascular abnormality, benign tumor, and skin cancer; yet, these symptoms manifest differently according to one's ethnic origin. Among the races, Caucasians manifest signs of photo-damage earlier than other races because of their relatively low melanin content [1]. Furthermore, Caucasians manifest earlier onset of skin aging together with more obvious skin wrinkles and sagging signs than other races [2]. Additionally, the clinical manifestation of skin aging among Asians differs considerably compared to Caucasians. In Asians, pigmentation changes are the most frequent and significant manifestation of skin aging [3]. Despite this fact, the mechanisms responsible for the deleterious effects and cutaneous alterations are distinct, and general skin signs are essential for the clinical assessment of skin aging. An ideal skin aging assessment should include all signs of skin aging.

As features of skin aging depend on genetics, race, age, sex, and exposure to different external factors [4],
developing a global skin aging assessment scale has been difficult. Most skin aging scales have been used for research studies [5]. Currently, there are more than 100 skin aging assessment scales. They have been proposed and used in clinical practice and in research to compare treatment outcomes. Some assessment methods have used only selected signs, such as those from the Glogau [6] and Fitzpatrick classification systems [7], focused on specific areas (e.g., the face, neck, and chest) [8] or a specific ethnicity [9]. Others have applied multidimensional assessment of multiple skin signs but the content and measurement properties were sub-standard [10].

From 1978 to 2016, only 5 published multidimensional assessment scales were determined to have high methodological quality for all skin types according to the COnsensus-based Standards for the selection of health Measurement INstruments (COSMIN)—a group that developed a critical appraisal tool (checklist) containing the standards for evaluating the methodological quality of studies investigating the properties of health measurement instruments [10]. Of these, 5 scales are known as the Merz Aesthetics Scale (MAS) [11], SCore of INtrinsic and EXtrinsic skin Aging (SCINEXA) [12], Skin Aging Score (SAS) [13], and two unnamed scales (Table 1) [14, 15]. Additionally, there are some limitations: most scores focus on Caucasians, they do not have complete skin aging signs, they are not widely used, and they do not have clear terminology. All of these scales were developed based on the relationship between skin aging signs and many factors that involve aging. Dermatologists are important persons who diagnose any skin signs because of their expertise and experience. However, none of these scales were designed to perform an assessment from a dermatologist’s perspective.

Today, there are many techniques and instruments to detect skin aging for research and treatment, including the 3-dimensional camera [16], dermoscopy [17], a physical sample analysis of color [18], and measurement of elasticity [19]. However, these include only some of the signs of skin aging. Some cosmetic products can improve skin elasticity, but changes in other clinical symptoms may be unclear. In addition, pigmentation and malignancy may require a dermatologist to identify signs of aging.

This is the first study that aimed to develop a simple global skin aging assessment score based on the perceptions of certified dermatologists.

**Main text**

**Materials and methods**

**Design and participants**

This study was conducted between October 1 and December 31, 2016. An internet-based survey using Google form was delivered to randomly selected dermatologists certified by the Dermatology Society of Thailand, including those with board certification, Doctor of Philosophy in Dermatology, Master of Science in Dermatology, and Fellowship and Diploma in Dermatology.

**Questionnaire**

The questionnaire comprised three main sections. The first section asked about respondents’ demographic characteristics and clinical training and experience with no identifiable data. In the second section, all skin signs of five COSMIN-qualified checklists were reviewed. To ensure that all possible and commonly known signs were included, two major textbooks [20, 21] that have been widely used for dermatology training in Thailand were also reviewed. To ensure scale simplicity, skin signs that required facial expression or were localized to specific facial areas were excluded. Therefore, 29 signs with published evidence of their relevancy to skin aging were included (Table 2). Respondents were asked whether each skin sign was essential for a skin aging diagnosis using the question, “I think that each of the following signs or diseases is essential for skin aging diagnosis” Their responses were based on a 5-point Likert scale (1, strongly disagree; 2, disagree; 3, neither agree nor disagree; 4, agree; 5, strongly agree) (Additional file 1). The third section included the question, “I am familiar with the following scale” to evaluate the familiarity and popularity of currently available, well-developed skin aging assessment scales among Thai dermatologists.

**Statistical analysis**

The Pearson Chi-square and Student t tests were used to analyze categorical and continuous variables,
respectively, where appropriate. Exploratory factor analysis (EFA) was performed to analyze the responses to explore the essential signs of skin aging based on the knowledge and experience of Thai dermatologists. EFA was chosen because the aim of this process was to identify linear group combinations of items that identify the optimal number of factors. The Kaiser-Guttman criterion [22] and scree test [23] were used to explore the optimal number of factors, which were used as an input for repeated EFA. Then, the factors were rotated to spread variability more evenly, so that all solutions were relatively the same. Items with high uniqueness were removed, whereas the remaining items had high factor loading, which should be at least 0.70, as this corresponds to approximately half of the variance of the variable being explained by the factor of interest. The retained items were grouped into factors, each of which was named based on the member items. Orthogonal (varimax) and oblique (promax) rotation provided similar results, but we decided to proceed with the latter because of the potential non-independent nature of the factors. An initial reliability test and item-based statistical analysis also were performed in conjunction with EFA [24]. Stata/SE version 15 (StataCorp) was used for all statistical calculations. p values < 0.05 were considered statistically significant.

### Table 2 List of 29 signs and relevant sources

| Skin aging signs          | Skin aging scales | Authors: Guinot et al. | Vierkotter et al. | Rzany et al. | Bazin et al. | Allehand et al. | N/A | N/A | Textbooks |
|---------------------------|-------------------|-------------------------|-------------------|--------------|--------------|-----------------|-----|-----|-----------|
| Wrinkles-superficial      | 0                 |                         |                   |              |              |                 | 0   |     | Yaar et al. |
| Wrinkles-deep             | 0                 |                         |                   |              |              |                 |     |     |           |
| Wrinkles-criscross        |                   |                         |                   |              |              |                 |     |     |           |
| Reduce fat tissue         |                   |                         |                   |              |              |                 |     |     |           |
| Nasolabial folds          | 0                 |                         |                   |              |              |                 |     |     |           |
| Eye bags                  | 0                 |                         |                   |              |              |                 |     |     |           |
| Ptosis of eyelids         |                   |                         |                   |              |              |                 |     |     |           |
| Yellowish discoloration   |                   |                         |                   |              |              |                 |     |     |           |
| Lax appearance/tissue slacking | 0             |                         |                   |              |              |                 |     |     |           |
| Solar elastosis           |                   |                         |                   |              |              |                 |     |     |           |
| Pseudoscar                |                   |                         |                   |              |              |                 |     |     |           |
| Cutis rhomboidalis nuchae |                   |                         |                   |              |              |                 |     |     |           |
| Freckles                  |                   |                         |                   |              |              |                 |     |     |           |
| Solar lentigines          |                   |                         |                   |              |              |                 |     |     |           |
| Melasma                   |                   |                         |                   |              |              |                 |     |     |           |
| Uneven pigment/pigment spot | 0               |                         |                   |              |              |                 |     |     |           |
| Guttate hypomelanosis     |                   |                         |                   |              |              |                 |     |     |           |
| Venous lakes              |                   |                         |                   |              |              |                 |     |     |           |
| Senile purpura            |                   |                         |                   |              |              |                 |     |     |           |
| Telangiectasias           |                   |                         |                   |              |              |                 |     |     |           |
| Milia                     |                   |                         |                   |              |              |                 |     |     |           |
| Sebaceous hyperplasia     |                   |                         |                   |              |              |                 |     |     |           |
| Senile comedone           | 0                 |                         |                   |              |              |                 |     |     |           |
| Favre-Racouchot syndrome  |                   |                         |                   |              |              |                 |     |     |           |
| Actinic keratosis         |                   |                         |                   |              |              |                 |     |     |           |
| Xerosis                   |                   |                         |                   |              |              |                 |     |     |           |
| Squamous cell carcinoma   |                   |                         |                   |              |              |                 |     |     |           |
| Basal cell carcinoma      |                   |                         |                   |              |              |                 |     |     |           |
| Malignant melanoma        |                   |                         |                   |              |              |                 |     |     |           |

COSMIN CONsensus-based Standards for the selection of health Measurement INstruments, SAS Skin Aging Score, SCINEXA SCore of INtrinsic and EXtrinsic skin Aging, MAS Merz Aesthetics Scale, N/A not applicable
Results
Of 213 randomly selected dermatologists, 145 responded to the survey (response rate, 68.1%). Seventy-five percent of respondents were women, and their mean age was 35.2 years. Most respondents worked at a private clinic (46.2%), followed by a medical school (31.7%), public hospital (29.7%), and private hospital (22.8%). Mean work experience was 7.7 years, and they attended approximately 3 international conferences annually on average.

Both the Kaiser–Guttman criterion and scree plot suggested that 3 factors were optimal in our EFA (Table 3). Factor 1 comprised 8 signs of atrophy (wrinkles-superficial, wrinkles-deep, wrinkles-crisscross, reduced fat tissue, nasolabial folds, eye bags, lax appearance, and solar elastosis), which reflected dermis and soft tissue lesions. Factor 2 comprised 7 signs of discoloration (freckles, lentigines, melasma, venous lakes, purpura, telangiectasias, and milia). Factor 3 comprised of 3 malignant lesions.

One-fifth (21.6%) of respondents were familiar with the SAS, compared with 11.2% and 9.4% who were familiar with MAS and SCINEXA, respectively. Average Likert scores were 2.54, 2.26, and 2.24 for the SAS, MAS, and SCINEXA, respectively. SAS was significantly better known than MAS (p < 0.001) and SCINEXA (p < 0.001); there was no difference between MAS and SCINEXA (p = 0.59).

Discussion
Although more than 100 skin aging assessment scales have been introduced, none has gained sufficient popularity to become widely used. This is supported by our finding that a minimal number of Thai dermatologists were familiar with even the methodologically robust scales. Many reasons for this finding are possible. First, the terminology may be unclear and the assessment results may be complicated, requiring competent assessors with special training. For example, the term inability to redden [13], pigment spot [15], and benign tumor [12] could be interpreted in many different ways, making the scale unreliable. Second, some scales may be too comprehensive for real-life clinical practice. For instance, SCINEXA covers almost all signs of aging, and of the 13 studies that used this scale, only two completed the full assessment [25, 26]. Potential reasons were its complexity and dependency on competent assessors with special training [12]. Additionally, some scales required permission before use, whereas other scales leave out key skin aging signs, such as pigmentation and benign tumors.

We propose a simplified Global Subjective Skin Aging Assessment (GS2A2) score comprising 3 factors, which are based on empirical evidence from dermatologists in practice and representative of the pathophysiologic changes that occur during the skin aging process. The term “atrophy” comprises reduction of elastic fibers as well as changes in collagen components and subcutaneous tissues (fat, muscle, and bone), leading to skin signs of wrinkles, solar elastosis, reduced fat tissue, nasolabial folds, etc. The term “discoloration” includes melanin, changes in vascular pigmentation, and keratin-filled cysts. All common skin aging malignancies are included in factor 3. Based on these 3 factors, skin aging is multidimensional, and one skin sign is insufficient to express the skin aging process in general.

The GS2A2 score includes factors relevant to skin aging from the perspective of dermatologists, and it is easily calculated as a 3-factor score. Although each item of the score is assessed independently, the final summative number is used for representing the overall skin aging of a person as perceived by the dermatologist. This approach mimics a real-life situation, in which a layperson would justify the skin aging condition of another person from overall appearance; rather than from area-by-area assessment usually performed in a research study. Given a 5-point Likert scale for each item (from 1—Strongly disagree to 5—Strongly agree), the total score for this 18-item scale may range from 18 to 90. While the change of the total score can be a simple quantification of skin aging

| Table 3 Factors related to skin aging | Loading |
|--------------------------------------|---------|
| Factor 1—atrophy                      |         |
| Wrinkles-superficial                 | 0.8587  |
| Wrinkles-deep                        | 0.8999  |
| Wrinkles-crisscross                  | 0.7437  |
| Reduced fat tissue                   | 0.8359  |
| Nasolabial folds                     | 0.7898  |
| Eye bags                             | 0.7696  |
| Lax appearance                       | 0.8027  |
| Solar elastosis                      | 0.7991  |
| Factor 2—discoloration               |         |
| Freckles                             | 0.8519  |
| Lentigines                           | 0.7946  |
| Melasma                              | 0.7565  |
| Venous lakes                         | 0.7056  |
| Purpura                              | 0.7346  |
| Telangiectasias                      | 0.7749  |
| Milia                                | 0.8056  |
| Factor 3—malignancy                  |         |
| Squamous cell carcinoma              | 0.8555  |
| Basal cell carcinoma                 | 0.8620  |
| Malignant melanoma                   | 0.8120  |
improvement of a subject, the score of each factor could help identify specific needs for a further correction in order to look younger. Nonetheless, as this is still an initial phase of scale development, the score should be further tested for validity and reliability before it can be used by a dermatologist to perform a visual assessment of overall skin aging condition of a subject in a simple summative number. The summary score may be used for assessing a cosmetic product that claims to have a global anti-aging effect, whereas a factor-based score could be used to differentiate whether the change in overall clinical outcomes was because of atrophy, discoloration, or malignancy. Moreover, standard terminology is used for each factor, resulting in better inter-rater reliability, which should be proven by further study. These results would support the GS2A2 score as a simple and informative tool. This tool is particularly relevant for Asian skin. However, it should be applied to non-Asian skin and/or non-Thai dermatologists. This score can also be used to measure overall anti-aging effects before and after dermatologic treatment, cosmetic products, or non-surgical skin rejuvenation procedures, such as laser therapy or microdermabrasion that claim to minimize overall skin aging. For example, the clinical effects of a multi-component nutritional supplement for photo-aged skin could be evaluated using the GS2A2 score instead of the Glogau classification system, which reflects only anti-wrinkle effects and not overall anti-aging effects. However, as the subjectivity of this score could be affected by many factors, the GS2A2 score should be used as a before-and-after comparison rather than as an average across individuals. Because this score was developed based on inputs from a representative group of Thai dermatologists who have experience with Asian patients, GS2A2 score is particularly relevant for Asians.

Conclusions
The GS2A2 score was empirically generated based on multiple signs of skin aging and the perceptions of certified dermatologists. It is a simple numerical score that can be used to evaluate the anti-aging effects of a cosmetic product or dermatologic intervention. This score can be helpful for dermatologists to evaluate skin aging in clinical practice and research to compare treatment outcomes. It should be tested further for validity and reliability.

Limitations
This study presented only the initial phase of scale development. As suggested by COSMIN, the GS2A2 score should be tested further for validity and reliability. Moreover, it should be conducted in other ethnicities to improve its generalizability.

Additional file

Additional file 1. Global Subjective Skin Aging Assessment (GS2A2).

Abbreviations
COSMIN: COnsensus-based Standards for the selection of health Measurement Instruments; EFA: exploratory factor analysis; GS2A2: Global Subjective Skin Aging Assessment; MAS: Merz Aesthetics Scale; SCINEXA: SCOrer of INtrinsic and ETrinsic skin Aging; SAS: Skin Aging Score.

Acknowledgements
The conduct of this study would not be possible without the dermatologists who participated in the survey.

Authors’ contributions
PB and KP initiate the idea. PB and JM collected data. PB, JM, and KP analyzed the data and prepare the manuscript. All authors read and approved the final manuscript.

Funding
This study did not receive any specific grant from funding agencies in the public, commercial, or not-for-profit sectors.

Availability of data and materials
All data generated or analysed during this study are included in this published article.

Ethics approval and consent to participate
Ethical approval was obtained from the Human Research Ethics Committee of Thammasat University (Institutional Review Board no. 045/2560). The committee waived the need for written informed consent because responding to this anonymized questionnaire contained no more than minimal risk and consent by action was used.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

Author details
1 Division of Dermatology, Department of Medicine, Chulabhorn International College of Medicine, Thammasat University, Pathum Thani, Thailand. 2 Department of Preventive and Social Medicine, Faculty of Medicine, Chulalongkorn University, Bangkok 10330, Thailand. 3 Department of International Health, Johns Hopkins Bloomberg School of Public Health, Baltimore, MD, USA.

Received: 7 July 2018 Accepted: 25 June 2019

Published online: 28 June 2019

References
1. Vashi NA, de Castro Maymone MB, Kundu RV. Aging differences in ethnic skin. J Clin Aesthet Dermatol. 2016;9:31–8.
2. Vierkotter A, Krutmann J. Environmental influences on skin aging and ethnic-specific manifestations. Dermatoendocrinol. 2012;4:227–31.
3. Tschachler E, Morozov F. Ethnic differences in skin aging. In: Gilchrest BA, Krutmann J, editors. Skin aging. Berlin: Springer; 2006. p. 23–31.
4. Farage MA, Miller KW, Elsner P, Malbach Hl. Characteristics of the aging skin. Adv Wound Care (New Rochelle). 2013;2:5–10.
5. Dobos G, Lichterfeld A, Blume-Peytavi U, Kottriner J. Evaluation of skin ageing: a systematic review of clinical scales. Br J Dermatol. 2015;172:1249–61.
6. Glogau RG. Aesthetic and anatomic analysis of the aging skin. Semin Cutan Med Surg. 1996;15:134–8.
7. Fitzpatrick RE, Goldman MP, Satur NM, Tope WD. Pulsed carbon dioxide laser resurfacing of photo-aged facial skin. Arch Dermatol. 1996;132:395–402.
8. Sattler G, Carruthers A, Carruthers J, Flynn TC, Geister TL, Gortelmeyer R, Hardas B, Himmrich S, Jones D, Kerscher M, et al. Validated assessment scale for neck volume. Dermatol Surg. 2012;38:343–50.

9. La Padula S, Hersant B, SiddAhmed M, Noddam J, Meningaud JP. Objective estimation of patient age through a new composite scale for facial aging assessment: the face—objective assessment scale. J Craniofac Surg. 2016;44:775–82.

10. Mokkink LB, Terwee CB, Patrick DL, Alonso J, Stratford PW, Knol DL, Bouter LM, de Vet HC. The COSMIN checklist for assessing the methodological quality of studies on measurement properties of health status measurement instruments: an international Delphi study. Qual Life Res. 2010;19:539–49.

11. Rzany B, Carruthers A, Carruthers J, Flynn TC, Geister TL, Gortelmeyer R, Hardas B, Himmrich S, Jones D, de Maio M, et al. Validated composite assessment scales for the global face. Dermatol Surg. 2012;38:294–308.

12. Vierkotter A, Ranft U, Kramer U, Sugir D, Reimann V, Krutmann J. The SCINEXA: a novel validated score to simultaneously assess and differentiate between intrinsic and extrinsic skin aging. J Dermatol Sci. 2009;53:207–11.

13. Guinot C, Malvy DJ, Ambroisine L, Latrelle J, Mauger E, Tenenhous M, Morzit F, Lopez S, Le Fur I, Tschachler E. Relative contribution of intrinsic vs extrinsic factors to skin aging as determined by a validated skin age score. Arch Dermatol. 2002;138:1454–60.

14. Cattell R. The screen test for the number of factors. Multivar Behav Res. 1966;1:245–76.

15. Cinotti E, Perrot JL, Labeille B, Biron AC, Vierkotter A, Heusele C, Nizard C, Schnebert S, Barthelemy JC, Cambazard F. Skin tumours and skin aging in 209 French elderly people: the PROOF study. Eur J Dermatol. 2016;26(5):470–6.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.