Objective Spectrometric Measurement of Keloid Color in the East Asian Population: Pitfalls of Subjective Color Measurements

Masayo Aoki, Satoshi Akaishi, Junichi Nakao, Teruyuki Dohi, Hiko Hyakusoku and Rei Ogawa

Department of Plastic, Reconstructive and Aesthetic Surgery, Nippon Medical School, Tokyo, Japan

Background: Keloids are characterized by the formation of excessive scar tissue that extends beyond the area of the initial wound. Keloid redness is due to angiogenesis and chronic inflammation and is an important indicator of the severity of the lesion and the effectiveness of treatment.

Methods: The color of 33 untreated keloids from 30 patients was measured with a narrow-band reflectance colorimeter. The erythema and melanin levels in the keloids (Ek and Mk, respectively) were recorded with control data obtained from the flexor aspect of the forearm (Ec and Mc, respectively). The keloid color was also evaluated subjectively.

Results: The Ek or Mk values did not vary significantly according to symptom intensity, scar region, patient age, and patient sex. Younger patients (<40 years) and female patients had significantly higher Ek/Ec ratios than did older patients and male patients, respectively. Subjective keloid redness evaluations distinguished keloids with high Ek/Ec ratios from keloids with low Ek/Ec ratios (P<0.0001) but could not distinguish keloids with high Ek from keloids with low Ek.

Conclusions: Subjective evaluations of keloids in Japanese subjects reflected Ek/Ec ratios, which were strongly affected by variation in background skin color. The subjective assessment of the color of keloids or other skin disorders should be performed with caution in Asian populations.

Key words: keloid, scar assessment, skin color assessment, spectrometer, erythema

Introduction
Keloids are a result of abnormal wound healing that results in excessive scar tissue formation that extends beyond the area of the initial wound and does not regress spontaneously. These scars are generally difficult to treat, although there are many therapeutic options, including surgery, corticosteroid injections, cryotherapy, radiation, and antitumor/immunosuppressive agents. Determining the efficacy of these therapies is sometimes difficult. One possibility is to assess these scars’ color, which reflects hemoglobin and melanin concentrations. The redness of keloids and hypertrophic scars (HSs) appears to relate to angiogenesis and chronic inflammation, which suggests that the color of keloids is an important indicator of the severity of the lesion and the effectiveness of treatment.

Scars are rated with a variety of scales. The most well-known and widely used scale is the Vancouver Burn Scar Assessment Scale, which was established in 1995 and is usually referred to as the Vancouver Scar Scale. This scale also assesses color and thus can be used to assess the severity of keloids/HSs or the efficacy of a therapy on such scars. Other scales have also been used for this purpose, namely, the Manchester Scar Scale, a visual analog scale scar scoring, and the Japan Scar Workshop Scar Scale. However, these scales require that color is judged with the naked eye; thus, the measurement is subjective and may vary depending on the evaluator.

The color of scars and other skin disorders has been measured for more than 50 years with spectrophotometry. The ability of spectrophotometry to measure the color of different skin lesions (both pigmented and vas-
cular) is good\cite{9,10}. The measurement of scar color with either a tri-stimulus colorimeter (Minolta Chroma Meter\cite{11}) or a narrow-band reflectance meter (DermaSpectrometer\cite{12}); has yielded fairly good results\cite{11}. However, whether these methods can measure HS/keloid color in an objective and quantitative fashion has not been determined.

On the basis of our clinical experience, we have postulated that keloid erythema levels are higher in female patients than in male patients and higher in younger patients than in older patients. We have also hypothesized that keloid erythema and melanin levels correlate with symptom intensity and vary depending on the body region of the keloid. In the present study, keloid color was measured with a color measurement device. The relationships of these color measurements with the scar region, the intensity of the symptoms, and patient age, and sex were assessed.

Materials and Methods

Ethical Considerations

The research complied with the Declaration of Helsinki and was approved by the Ethical Committee of Nippon Medical School. Informed consent was obtained from all patients.

Instruments

The DermaSpectrometer\cite{12} (Cortex Technology, Hadsund, Denmark) narrow-band reflectance meter was used to measure the color of keloids. The light-emitting diodes of this instrument emit light at 2 defined wavelengths: 568 nm (green) and 655 nm (red). A photodetector measures the light reflected by the skin. Because hemoglobin strongly absorbs green light wavelengths (which is why blood is red), the DermaSpectrometer estimates erythema levels by measuring how much green light is absorbed. In contrast, melanin readily absorbs both red light and green light. The DermaSpectrometer can measure melanin levels by measuring the reflectance of red light. The area of skin that is measured is 6 mm in diameter (surface, 0.28 cm$^2$). The probe is applied to the skin surface (3.46 cm$^2$) with no more force than the weight of the instrument (640 g). Thus, the skin is placed under a pressure of 158 g/cm$^2$. The instrument is calibrated with a black and white calibration plate\cite{12}.

Patients

The color of 33 untreated keloids from 30 patients (12 men and 18 women) was measured with the spectrophotometer. All patients first presented at the outpatient department of our hospital in 2009 or 2010.

Assessment of Symptom Intensity

Before keloid color was measured, the patients were asked to subjectively evaluate the intensity of their keloid symptoms by scoring the overall pain, itch, and/or subjective redness of their keloid with a 4-point system: 1, slight; 2, mild; 3, strong; and 4, very strong.

Measurement of Keloid Color with Spectrophotometry

The erythema and melanin levels in the keloids (designated as Ek and Mk, respectively) were recorded with control data taken from the flexor aspect of the forearm (Ec and Mc, respectively). Each measurement was performed 3 times, and the mean was calculated. The Ek/Ec and Mk/Mc ratios were then calculated.

Subjective Evaluations of Keloid Redness

The redness of 8 of the keloids was evaluated subjectively by 6 plastic surgeons with a 10-point system. To ensure that this evaluation was not affected by the age or sex of the patient or by the keloid region, the plastic surgeons were given digital photos of the keloid lesions. The redness was evaluated by comparing the keloid to the color of the surrounding skin.

Statistics

The relationships of Ek, Mk, Ek/Ec, and Mk/Mc to the scar region, symptom intensity, patient age, and patient sex were assessed with the Tukey-Kramer method or Student’s $t$-test. Two-factor repeated measures ANOVA was used to assess the accuracy of the subjective evaluations of keloid redness relative to the DermaSpectrometer measurements. The Statcel2 software program (OMS Ltd., Saitama, Japan) was used for statistical analyses. Differences were considered to be statistically significant if $P$ was $<0.05$.

Results

Patient Characteristics

Of the 30 patients, 12 were men (40%) and 18 were women (60%; Table 1). The ages of the patients ranged from 18 to 64 years, and the mean age was 33.09 ± 12.79 years. In total, 23 patients (76.7%) were younger than 40 years, and 7 (23.3%) were 40 years and older. All patients were Japanese. Of the 33 keloids, 14 (42.4%) were on the anterior chest, 5 (15.2%) were on an ear lobe, 4 (12.1%) were on a shoulder, and 3 (9.1%) were on the lower abdomen. The remaining 7 keloids were on the mandible, lateral chest (2 each, 6.1%), dorsum, upper limb, and axilla (1 each, 3.0%). Of the patients, 2 (6.1%), 11 (33.3%), 15 (45.5%), and 5 (15.2%) rated the intensity of the symptoms of the keloid(s) as slight, mild, strong, and very
Table 1 Characteristics of the patients in this study

| Variable            | Number (% relative to the total number of patients or keloids) |
|---------------------|---------------------------------------------------------------|
| Patients            | 30                                                            |
| Keloids             | 33                                                            |
| Sex                 |                                                              |
| Male                | 12 (40.0%)                                                    |
| Female              | 18 (60.0%)                                                    |
| Age                 |                                                              |
| 18–39 years         | 23 (76.7%)                                                    |
| 40–64 years         | 7 (23.3%)                                                     |
| Mean (SD)           | 33.09 (12.79)                                                 |
| Location of keloids |                                                              |
| Anterior chest      | 14 (42.4%)                                                    |
| Ear lobe            | 5 (15.2%)                                                     |
| Shoulder            | 4 (12.1%)                                                     |
| Lower abdomen       | 3 (9.1%)                                                      |
| Mandible            | 2 (6.1%)                                                      |
| Lateral chest       | 2 (6.1%)                                                      |
| Dorsal              | 1 (3.0%)                                                      |
| Upper limb          | 1 (3.0%)                                                      |
| Axilla              | 1 (3.0%)                                                      |
| Intensity of symptoms|                                                              |
| 1: Slight           |                                                            |
| 2: Mild             | 11 (33.3%)                                                    |
| 3: Strong           | 15 (45.5%)                                                    |
| 4: Very strong      | 5 (15.2%)                                                     |

Table 2 Keloid color measured in all patients (n=30) by using a color measurement device

| Subject                              | Mean of measured values (SD) |
|--------------------------------------|-----------------------------|
| Erythema level in keloid (Ek)        | 18.29 (4.27)                |
| Erythema level in forearm control (Ec)| 9.76 (2.93)                |
| Ek/Ec ratios                         | 1.96 (0.72)                 |
| Melanin level in keloid (Mk)         | 42.70 (6.44)                |
| Melanin level in forearm control (Mc) | 33.30 (2.65)               |
| Mk/Mc ratios                         | 1.32 (0.22)                 |

strong, respectively (Table 1).

**Erythema and Melanin Levels in the Keloids, as Measured with the Color Measurement Device**

The Ek and Ec means (SD) were 18.29 (4.27) and 9.76 (2.93), respectively, and the Mk and Mc means (SD) were 42.70 (6.44) and 33.30 (2.65), respectively (Table 2). The Ek and Mk values were significantly higher than the Ec ($P<1\times10^{-11}$) and Mc ($P<1\times10^{-9}$) values, respectively. The Ek/Ec and Mk/Mc ratios were 1.96 (0.72) and 1.32 (0.22), respectively (Table 2).

**Relationship of Ek and Mk to Scar Region, Severity of Symptoms, Age, and Sex**

Student’s t-test revealed that patients older than 40 years did not differ from patients younger than 40 years in terms of Ek and Mk values. Men and women also had similar Ek and Mk values. The Tukey-Kramer method revealed that these values of keloids on the lower abdomen did not differ significantly from those of keloids on the chest or shoulder. Finally, Ek or Mk values of keloids that were rated as being slightly or mildly symptomatic did not differ from those of keloids that were rated as being strongly or very strongly symptomatic (Fig. 1).

**Relationship between Ek/Ec and Mk/Mc Ratios and Scar Region, Severity of Symptoms, Age, and Sex**

Patients younger than 40 years had significantly higher Ek/Ec ratios than did patients older than 40 years. Furthermore, women had significantly higher Ek/Ec ratios than did men. Both of these observations reflected the fact that Ec values were significantly lower in both younger patients than in older patients ($P=0.01$) and in women than in men ($P=0.0003$). In other words, in Japanese subjects, there is a stronger color contrast between
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Fig. 1  Association of keloid erythema (Ek) or keloid melatonin (Mk) with symptom intensity, body area, age, or sex. None of these factors associated significantly with Ek (above) or Mk (below).

The keloid and normal skin when the patient is younger or a woman than when the patient is older or a man, even though the actual erythema levels of the keloids are equivalent. The patients whose keloids were slightly or mildly symptomatic did not have Ek/Ec ratios that differed significantly from those of patients who had strongly or very strongly symptomatic keloids. Similarly, Ek/Ec ratios of keloids on the lower abdomen did not differ from those of keloids on the shoulder or chest.

The Mk/Mc ratios of keloids on the lower abdomen were significantly higher than those of keloids of the chest or shoulder (Fig. 2). However, the Mk/Mc ratios of keloids that were rated as slightly or mildly symptomatic did not differ from those of keloids that were rated as strongly or very strongly symptomatic. The Mk/Mc ratios was also not affected by the patients’ age or sex (Fig. 2).

Ability of Subjective Evaluations to Distinguish between Keloids with High and Low Ek Values and between Keloids with High and Low Ek/Ec Values

We did not observe that the objectively measured Ek values of the younger or female patients differed significantly from those of older or male patients, respectively. Instead, we found that younger or female patients had significantly lower Ec values than did older or male patients, respectively, which in turn led to significant but clinically irrelevant age and sex associations with Ek/Ec ratios. This result suggested that subjective evaluations of keloid redness, which require that the keloid is compared to the surrounding skin, might overestimate the redness of keloids in younger or female patients.

To assess this possibility, 8 patients were selected on the basis of spectrophotometrically measured Ek and Ek/Ec values, as follows. The Ek values <16.5 and >20.0 were considered to be low and high, respectively, and the Ek/Ec ratios <1.65 and ≥1.65 were considered to be low and high, respectively. Keloids with Ek and Ek/Ec values that were both low (n=2), both high (n=2), low and high, respectively (n=2), and high and low, respectively (n=2), were selected. Figure 3 shows 1 patient from each of the 4 groups. Photos of the 8 selected keloids were given to 6 plastic surgeons, who were asked to evaluate the keloid redness subjectively with a 10-point system. The normal skin around the keloids served for comparison.

Division of the keloids into high and low Ek groups revealed that while the high-Ek keloids did tend to be subjectively evaluated as being somewhat redder than the low-Ek keloids, a statistically significant difference with two-factor repeated measures ANOVA was not observed (P=0.094). In contrast, when the keloids were di-
vided into low and high Ek/Ec ratio groups, the high Ek/Ec ratio keloids were evaluated as being highly significantly redder than the low Ek/Ec ratio keloids ($P<0.0001$; Fig. 4).

**Discussion**

Pathological cutaneous scars, such as keloids and HSs, are characterized by a diffuse redness that is caused by chronic inflammation-driven overgrowth of capillary vessels\(^1\). Hyperpigmentation is also commonly attributed to the inflammatory response. The mechanisms involved might be related to the activation of melanocytes by inflammatory mediators or reactive oxidative species that are released by the damaged skin\(^1\). Additional melanin is produced in response to the injury, and this production is engulfed by macrophages, which are also involved in the inflammatory response. The macrophages then retain the melanin in the dermis until the cells and melanin are degraded; in this way, the pigment can persist in the dermis for an extended period after the injury\(^1\). Notably, topical intraleosional steroid injections have been shown to suppress the expression of vascular endothelial growth factor (VEGF) in keloid tissue and induce its regression \(\textit{in vivo}\)\(^5\). Thus, improving the inflammation in keloids might also cause the redness or pigmentation to subside. This knowledge, in turn, suggests that redness and hyperpigmentation might be important indicators of the intensity of inflammation or the effect of topical intraleosional steroid injections on aberrant scars.

However, the present study has shown that the intensity of symptoms, the body region, patient age, and patient sex were not associated with keloid colors (Ek and Mk) when they were measured objectively. Thus, at least in Japanese patients, the severity of keloids/HSs cannot be evaluated on the basis of redness only, even if it is measured objectively. However, because lesion color might change during the course of treatment and, thus, can be used to estimate the therapeutic effect, the accurate evaluation of color might be useful for both the treatment and study of aberrant scarring.

Several scar assessment scales include a color assessment component. However, the present study was, to the best of our knowledge, the first to suggest that subjective evaluations of lesion color in Japanese patients are inaccurate: although young or female patients were found to have Ek/Ec ratios significantly higher than those of older...
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or male patients, this increase was not because their lesions had more erythema; rather, it was because they had significantly lower erythema levels in the forearm control which resulted in a more marked contrast between the colors of the keloid and normal skin. Thus, in subjective evaluations of skin color in Japanese patients and, perhaps, also in other East Asian patients, skin lesions might be evaluated as being redder in patients with fair skin than in patients with darker skin. Indeed, this subjective error led to our now-disproven hypothesis in the present study that younger or female patients are more likely to have greater erythema (strong redness) than are older or male patients. Thus, subjective evaluations of lesion color in Japanese patients (and probably most other East Asian patients) should be approached with great caution.

The present study has shown that, although the severity of the color of keloids/HSs was difficult to judge with the naked eye, skin color can be characterized and changes in small skin color can be quantified with the DermaSpectrometer. Such instruments have been used for many years to quantitatively estimate the results of the treatment of pigmented lesions. These instruments are useful for evaluating the effectiveness of laser treatment, for quantitatively assessing the skin erythema that is due to radiotherapy, and for evaluating scars. Thus, a DermaSpectrometer or other color measurement devices can be used to estimate the color of red scars accurately and in an objective and quantitative manner. These instruments for measuring color might also be useful for evaluating the effects of therapies on keloids/HSs. When a scar is assessed, a high priority should not be placed on the subjective evaluation of color. Furthermore, the objective and accurate evaluation with color-assessment devices might be useful for monitoring HS/
keloid color in patients.

Conclusions

When keloid color was objectively and accurately measured with a color measurement device, the intensity of the symptoms, the body region, patient age, and patient sex were not associated with either erythema or melanin levels in the keloids. However, young or female patients were found to have statistically significantly higher Ek/Ec ratios than did older or male patients. Subjective evaluations of Ek were then shown to actually reflect these ratios rather than Ek itself. As a result, in Japanese patients and, perhaps, other East Asian patients, the lesions of patients who are young or female may be erroneously considered to be redder than the lesions of patients who are older or male when relying on subjective skin color evaluations. In contrast, a DermaSpectrometer or other color measurement devices can be used to estimate the color of red scars accurately and in an objective and quantitative manner and, thus, might also be useful for evaluating the effects of therapies on keloids/HSs. Thus, subjective evaluations of the color of keloids or other skin disorders in Japanese patients and, probably, most other East Asian patients should be approached with great caution.

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