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# Oral propranolol therapy in 23 infants with infantile hemangioma

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**Background** Infantile hemangioma (IH) is a common vascular tumor in pediatric patients, and is commonly treated with propranolol. We describe our experiences with dosage, response to treatment, and side effects in 23 IH patients treated with propranolol.

**Methods** For this nonrandomized comparative cohort study, the authors enrolled 23 patients treated with propranolol. Photographs were taken before propranolol administration and at 3, 6, 9, and 12 months after treatment. Treatment responses were objectively analyzed with a computer program.

**Results** There were three male and 20 female patients. Common tumor locations were the head and neck (13 cases, 56.5%), trunk (four cases, 17.4%), extremities (three cases, 13.0%), and combined locations (three cases, 13.0%). The response to propranolol was significantly lower in patients with two or more lesions than in patients with a single lesion in terms of both color fading (P<0.001) and size reduction (P<0.001). In male patients ( $42.2 \pm 3.9$ ), the change in a-values, indicating coloration, was higher than in female patients ( $19.8 \pm 13.8$ ) (P<0.001). In patients who started treatment before 6 months after birth, the size reduction was greater than in their counterparts (62.3%; range, 3.0%–93.0% vs. 15.8%; range, 1.0%–79.0%; P<0.001).

**Conclusions** Propranolol is an efficacious treatment with a good safety profile. In patients with a single lesion, the response to treatment was better in terms of color fading and size reduction. Furthermore, male patients responded better to propranolol treatment in terms of color fading than female patients, and starting treatment before 6 months after birth was more advantageous for size reduction.

Keywords Hemangioma / Propranolol / Therapeutics

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**INTRODUCTION** 

Infantile hemangioma (IH) is a common vascular tumor in children. Schupp et al. [1] reported an estimated incidence rate in Asia of one in 125 persons annually. Female infants are more frequently affected than are male infants [2]. The disease is caused by an abnormal proliferation of endothelial cells of the capillary vessels and typically occurs at 3 to 4 weeks of age, peaking at 6 to 7 months of age, in the proliferative phase. After this period, the size of the tumor rarely decreases naturally, al-though its color can suddenly become less livid [3]. Most IHs are diagnosed via patient history and physical examination. In addition, imaging studies, such as Doppler ultrasound and magnetic resonance imaging (MRI), can characterize the location

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and extent of cutaneous and extracutaneous hemangiomas. MRI can also help differentiate IHs from high-flow vascular lesions such as arteriovenous malformations. Ultrasonography is useful for differentiating hemangiomas from other deep dermal or subcutaneous structures, such as cysts, pilomatrixomas, and lymph nodes [4].

Although hemangiomas are not routinely biopsied because the diagnosis can generally be made clinically, specimens may be evaluated for tissue-specific immunohistochemical markers, such as GLUT-1, merosin, Fc-gamma-RII, and Lewis Y antigens. These markers can differentiate IHs (positive staining for all markers) from other vascular neoplasms and malformations [5,6].

Most IHs are small and self-limiting; however, complicated IHs with accompanying bleeding or infection may impact vital structures, such as the eye or airway, and medical or surgical treatment is therefore indicated in such cases [7]. Furthermore, the efficacies of existing treatments vary, and all have associated safety concerns [8-10]. Systemic glucocorticoids are the primary treatment for complicated IHs, with interferon- $\alpha$  and vincristine used for lesions that do not respond to glucocorticoid therapy [10,11].

In 2008, Leaute-Labreze et al. [12] reported the first case in which propranolol was administered for the treatment of obstructive hypertrophic myocardiopathy in a 1-month-old infant who had undergone treatment with steroids. Thereafter, treating IH with propranolol was deemed effective, with a negligible frequency of side effects. Propranolol is a nonselective betablocker that is used in patients of all ages with hypertension and various cardiac diseases. It is also known to be safe for administration in infants [13].

The proposed mechanisms of the effects of propranolol on IH include the inhibition of tumor growth and angiogenesis, as well as vasoconstriction [14]. Its known side effects include hypotension, bradycardia, bronchospasm, hypoglycemia, and diarrhea [12]. However, studies of the effects of propranolol on infant growth and development after its prolonged administration are limited, as are studies of the acute side effects mentioned above [15,16]. The optimal duration of propranolol treatment in cases of IH has not yet been established; its long-term administration (>1 year) may be required, depending on tumor size and treatment response. Many studies of IH have been published, but few have reported objective measures of treatment response. We thus measured the change of color and size of the lesions to evaluate their response to treatment. We also describe our experience with propranolol dosage and side effects in 23 IH patients who were treated with propranolol.

## **METHODS**

Research ethics approval was obtained from the Institutional Review Board at Severance Hospital (Approval No. 4-2017-1158). Informed consent was obtained from all patients or their representatives for inclusion in this study.

We evaluated 29 infants with newly diagnosed IH who were given propranolol at a dosage of 3 mg/kg/day, given as two divided doses, between 2015 and 2017, at a tertiary care hospital. Six patients were excluded from the final analysis because of a lack of follow-up visits.

In a retrospective study based on computerized medical records, the charts were reviewed for sex, age, location of the hemangioma, response to treatment as determined by the analysis of photos, age of the patient at the beginning of therapy, duration of therapy, complications, and adverse effects. We used propranolol after an evaluation of the cardiac status of each infant for the treatment of IH in an outpatient clinic.

To assess cardiac status for pediatric cardiac outpatient care, electrocardiography and echocardiography were conducted. Additional imaging studies (ultrasonography or MRI) were conducted if necessary.

The oral administration of propranolol was begun during the next outpatient clinic visit after the evaluation of cardiac status and imaging studies were complete. Patients' caregivers were instructed to cease administering propranolol if severe coughing, dyspnea, vomiting, or diarrhea developed during treatment. Follow-up visits were initially scheduled 2 weeks after the initiation of therapy and then monthly thereafter. Visits were also encouraged in case of any complications. We determined that the duration of treatment would be dependent on the response to treatment.

Patients were evaluated regularly in the outpatient clinic. Patients were photographed prior to the administration of propranolol and at 3, 6, 9, and 12 months after treatment, with a commercial digital camera and a digital camera equipped with the Antera 3D system (Miravex, Dublin, Ireland), which enables analyses of size and color from the chromatic aberration values (L, a, and b values). These values express color as three numerical values: L for lightness and a and b for the green-red and blueyellow color components, respectively [17]. The a-value was measured in our study for assessing the redness of the lesions. To assess general aesthetic outcomes and the overall improvement in the appearance of the hemangioma after treatment, we measured and compared the initial and final a-values and the size of the hemangiomas.

We further analyzed changes in color and size using four variables: sex, number of hemangiomas, location of the hemangioma, and the age of patients at the start of treatment. The starting age was divided into two groups: before and after 6 months after birth. The location of hemangiomas was also divided into two groups: the head and neck group and the non-head and neck group. Furthermore, patients were divided into two groups by the number of hemangiomas: group 1 included patients with only a single hemangioma, whereas group 2 included patients with two or more hemangiomas.

Statistical tests were used to analyze the results. The normality of the distribution of the groups was assessed using the Shapiro-Wilk test. Statistical analyses were performed using the t-test for independent samples and the Mann-Whitney U-test. A P-value less than 0.05 was considered to indicate statistical significance, and statistical analyses were performed using SPSS 15.0.1 for Windows (SPSS Inc., Chicago, IL, USA).

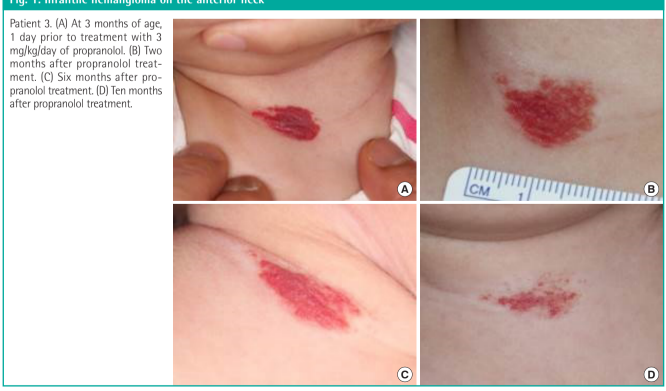
## RESULTS

In the present study, we included 23 children who were diagnosed as having IH and who were referred for propranolol treatment. The mean age of the patients at the initiation of treatment was  $6.5 \pm 3.6$  months. The average follow-up period was  $7.0 \pm$ 3.2 months. There were three male and 20 female patients. Common locations were the head and neck (13 cases, 56.5%), trunk (four cases, 17.4%), extremities (three cases, 13.0%), and combined locations (three cases, 13.0%). We divided patients into two groups based on the number of hemangiomas. Group 1, which included patients with only a single hemangioma, comprised 19 patients, whereas group 2, which included patients with two or more hemangiomas, was composed of four patients (Figs. 1 and 2).

The response to treatment was assessed in terms of color and size using the Antera 3D system (Miravex). At the time of the initial diagnosis, the average measured a-value was  $39.7 \pm 7.9$ . The average final a-value was  $17.0 \pm 14.5$ . The difference between the two values was calculated to assess the response of treatment in terms of color fading. The mean change of the avalue was  $22.7 \pm 15.0$ . Overall, the a-value decreased.

The Shapiro-Wilk test was performed to test the normality of the size-related data. The P-value was less than 0.05, so we concluded that the data were not normally distributed. The median initial size of the IHs was 115.0 mm<sup>2</sup> (range, 4.0–1,517.0 mm<sup>2</sup>) and the median final size was 56.0 mm<sup>2</sup> (range, 1.0-891.0 mm<sup>2</sup>). The percent change in size was calculated as the difference between the initial size and the final size divided by the initial size. The median percent change in size was 32.3% (range, 1.1%-92.6%) (Table 1).

When color fading was analyzed by sex, no significant difference was found in the initial average between male patients  $(46.1 \pm 1.1)$  and female patients  $(38.8 \pm 8.1)$  (P > 0.05). How-



#### Fig. 1. Infantile hemangioma on the anterior neck

#### Fig. 2. Infantile hemangioma on the right cheek



Patient 4. (A) At 3 weeks of age. (B) Three months after the initiation of therapy with 3 mg/kg/day of propranolol. (C) Six months of age. (D) Residual hemangioma at 11 months of age.

ever, there was a statistically significant difference according to sex in the final a-value, indicating color, and the color change. In male patients, the final a-value was lower (P < 0.001) and the color change was greater than in female patients (P < 0.001). Similar results were found for the number of hemangiomas; group 1 showed a lower final a-value (P < 0.001) and a greater color change than group 2 (P < 0.001). No other statistically significant differences were found in the analyses according to the location of the hemangioma and the age of patients at the start of treatment (Table 2).

In an analysis of the response to treatment based on size change according to the number of hemangiomas, group 1 showed a significantly greater size change (median, 49.5%; range, 1.1%-92.6%) than group 2 (median, 12.1%; range, 4.0%-16.0%) Additionally, the change in size in patients who started treatment before 6 months of age (median, 62.3%; range 3.0%-93.0%) was greater than in their counterparts (median, 15.8%; range, 1.0%-79.0%; P < 0.001). No other statistically significant differences were found in the analyses according to sex and the location of the hemangioma (Table 3).

No severe adverse events were noted in any patient. We observed mild adverse effects (diarrhea or mild fever) that were tolerated without the need to discontinue treatment (Table 4).

## DISCUSSION

C

Propranolol is widely used for the effective treatment of IH, with a good response rate compared with prednisolone, which is the conventional treatment [18]; in our experience, it also has a good safety profile. Propranolol was the first β-adrenergic receptor antagonist introduced for the treatment of angina. It is known to reversibly bind  $\beta$ 1 and  $\beta$ 2 receptors, reducing heart rate and cardiac output, and causing peripheral vasoconstriction soon after administration [19]. Although the mechanism of the effects of propranolol on IH remain unclear, some studies have reported that vasoconstriction plays a major role in lesion improvement [20], appearing to soften lesions and make them less livid. Moreover, the late effects of propranolol, which are evident several days after administration, include lesion improvement via the inhibition of proangiogenic signals, such as vascular endothelial growth factor and basic fibroblast growth factor, and the stimulation of microvascular endothelial cell apoptosis [21].

The determination of the therapeutic dose of propranolol appears to differ for each hospital. According to Kim et al. [22], treatment with propranolol generally begins at admission, with 0.5 mg/kg/day administered on the first day of treatment, and 1 mg/kg/day given on the second day. If no side effects are observed on the third day, the final treatment dose is increased to 2

		lable 1. Course of propranolol treatment in patients with	ith infantile hemangioma (n=23)	gioma (n=23)							
Vertation of hemaly limitsNon of all all all all all all all all all al					Response	to treatmen	nt (color)	Response	to treatmo	ent (size)	
Turk Grup1 38.7 27.1 11.6 2.4 210 10.3   Head & neck Group1 41.9 39.0 2.9 17.4 172 1.1   Head & neck Group1 41.9 36.2 4.7 32 2.7 15.6   Head & neck Group1 41.1 15.3 25.6 1.208 89 92.6   Head & neck Group1 41.1 15.3 25.6 1.208 89 92.6   Head & neck Group1 41.1 15.3 25.6 1.208 89 32.3   Lower extremity Group1 41.4 2.6 44.8 30.9 24.7   Head & neck Group1 45.4 2.6 45.8 32.3 32.3   Upper extremity Group1 45.4 7.0 14.8 53.6 44.4   Head & neck Group1 45.4 7.0 14.8 53.6 54.9   Head & neck Group2 47.7 47.0	Duration of follow-up (mo)	Propranolol dose (mg/kg/day)	Location of hemangioma	No. of hemangiomas	Initial test (a-value)	Final test (a-value)	Final- initial test (a-value)	Initial test (mm²)	Final test (mm <sup>2</sup> )	Final- initial/ initial test (%)	Complication
Head & neck Group1 419 39.0 2.9 174 172 11   Head & neck Group2 40.9 36.2 4.7 32 27 15.6   Head & neck Group1 37.1 9.1 28.0 111 56 49.5   Tunk Group1 41.1 15.3 25.8 1,208 89 26.0   Head & neck Group1 47.4 2.6 42.8 189 17 91.0   Head & neck Group1 21.8 7.0 14.8 53 37 29.7   Lower extremity Group1 21.8 7.0 14.8 33 32.3   Lower extremity Group1 47.4 2.6 44.8 33 33   Lower extremity Group1 45.4 7.0 14.8 36 33   Lower extremity Group1 45.4 7.0 14.8 36 33   Head & neck Group1 45.4 7.6 16		ę	Trunk	Group 1	38.7	27.1	11.6	234	210	10.3	No
Head & neck Goup2 409 36.2 4.7 32 27 15.6   Head & neck Goup1 37.1 9.1 28.0 111 56 49.5   Tunk Goup1 41.1 15.3 25.8 1,208 89 92.6   Head & neck Goup1 41.4 21.6 42.8 130 89 23.3   Head & neck Goup1 21.8 7.0 14.8 53 37 29.7   Lower extremity Goup1 21.8 7.0 14.8 53 37 30.9   Lower extremity Goup1 41.4 26 44.8 33 32.3   Upper extremity Goup1 43.4 7.0 14.8 33.9 33.9   Head & neck Goup1 43.4 7.6 35.8 33.7 11 35.7   Head & neck Goup1 45.5 65 39.0 25.6 17.7 16.8 35.8   Head & neck Goup1		S	Head & neck	Group 1	41.9	39.0	2.9	174	172	1.1	No
Haad & neck Group1 37.1 9.1 2.80 111 5.6 49.5   Turrk Group1 41.1 15.3 25.8 1.208 89 92.6   Haad & neck Group1 41.1 15.3 25.8 1.208 89 92.6   Haad & neck Group1 21.8 7.0 14.8 53 37 29.7   Haad & neck Group1 21.8 7.0 14.8 53 37 29.7   Upper extremity Group1 45.2 46.1 0.1 115 35   Head & neck Group1 45.2 46.1 0.1 116 35   Head & neck Group1 45.2 46.1 0.1 116 35   Head & neck Group2 47.7 47.0 0.7 117 162 85   Head & neck Group2 41.3 7.6 117 162 85   Head & neck Group1 45.7 37.9 177		ç	Head & neck	Group 2	40.9	36.2	4.7	32	27	15.6	Diarrhea
TurkGoup141.115.325.81,2088992.6Head & neckGoup145.42.64.281891791.0Head & neckGroup121.87.014.8533729.7Lower externityGroup121.87.014.8533729.7Lower externityGroup147.42.644.81308832.3Upper externityGroup138.89.329.54931.920.9Upper externityGroup145.56.539.071185.7Head & neckGroup145.56.539.0229.99.9Head & neckGroup247.747.00.790864.4Head & neckGroup247.747.00.790864.4Head & neckGroup247.747.00.790864.4Head & neckGroup247.747.00.790864.4Head & neckGroup137.411.525.01578575.5Head & neckGroup137.411.525.61.5136775.5Head & neckGroup137.411.525.61.5137.676.5Head & neckGroup147.326.61.5137.776.576.5Head & neckGroup147.326.61.5137.676.576.5Head & ne		c	Head & neck	Group 1	37.1	9.1	28.0	111	56	49.5	No
Head & neck Goup1 45.4 2.6 4.28 189 17 91.0   Head & neck Group1 21.8 7.0 14.8 5.3 37 29.7   Lower extremity Group1 21.8 7.0 14.8 53 37 29.7   Lower extremity Group1 38.8 9.3 29.5 49 36 30   Upper extremity Group1 46.2 46.1 0.1 115 35.8   Head & neck, trunk Group1 45.5 56.5 35.8 7 1 56.7   Head & neck, trunk Group1 45.5 56.5 35.8 7 1 56.7   Head & neck, trunk Group2 47.7 7.6 35.8 7 1 56.7   Head & neck, trunk Group1 45.6 55.6 57.8 56.7 56.5   Head & neck, trunk Group1 37.4 11.5 56.7 56.5 56.5 56.5   Head & neck, trunk <td></td> <td>ę</td> <td>Trunk</td> <td>Group 1</td> <td>41.1</td> <td>15.3</td> <td>25.8</td> <td>1,208</td> <td>89</td> <td>92.6</td> <td>No</td>		ę	Trunk	Group 1	41.1	15.3	25.8	1,208	89	92.6	No
Head & neck Goup1 21.8 7.0 148 53 37 29.7   Lower axtremity Group1 47.4 2.6 44.8 130 88 32.3   Head & neck Group1 38.8 9.3 29.5 49 34 30.9   Upper extremity Group1 46.2 46.1 0.1 115 111 35   Head & neck, trunk Group1 45.5 65.5 35.8 7 1 85.7   Head & neck, trunk Group2 47.7 7.6 35.8 7 1 15 35.9   Head & neck, trunk Group2 47.7 7.6 35.9 7 1 35   Head & neck, trunk Group2 47.7 7.6 37 1 1 35   Head & neck, trunk Group1 37.4 17 162 8.5 1   Head & neck, trunk Group2 31.9 23.1 1 1 2 1   Head & n		c	Head & neck	Group 1	45.4	2.6	42.8	189	17	91.0	Diarrhea
Lower extramity Group 1 47.4 2.6 44.8 130 88 2.3   Head & neck Group 1 38.8 9.3 29.5 49 34 309   Upper extramity Group 1 38.8 9.3 29.5 49 34 309   Upper extramity Group 1 45.5 6.5 39.0 7 11 35   Head & neck Group 2 47.7 47.0 0.7 90 86 44   Head & neck, trunk Group 2 43.5 53.93 32.9 72 90 90   Head & neck, trunk Group 2 43.5 53.93 4.2 177 162 8.5   Head & neck, trunk Group 2 31.9 23.1 9.2 177 162 8.5   Head & neck, trunk Group 2 31.9 23.1 8.8 157 158   Head & neck, trunk Group 2 11.5 25.5 1517 357 76.5   Head & neck		с	Head & neck	Group 1	21.8	7.0	14.8	53	37	29.7	No
Head & neck Goup1 388 9.3 29.5 49 34 309   Upper extremity Goup1 46.2 46.1 0.1 115 111 35   Head & neck Group1 45.5 6.5 35.8 7 1 85.7   Head & neck Group1 45.5 6.5 39.0 22 2 909   Head & neck, tunk Group2 47.7 47.0 0.7 90 86 4.4   Head & neck, tunk Group2 43.5 3.3.3 4.2 177 162 8.5   Head & neck, tunk Group2 31.9 23.1 8.4 177 162 8.5   Head & neck, tunk Group1 37.4 11.5 25.6 1,517 15.8 15.8   Head & neck Group1 37.4 11.5 25.6 1,517 357 76.5   Head & neck Group1 37.4 11.5 25.6 1,517 357 76.5		c	Lower extremity	Group 1	47.4	2.6	44.8	130	88	32.3	No
Upper extremityGroup 1 $46.2$ $46.1$ $0.1$ $115$ $111$ $35$ Head & neckGroup 1 $43.4$ $7.6$ $35.8$ $7$ $1$ $85.7$ Head & neckGroup 2 $47.7$ $47.0$ $0.7$ $90$ $86$ $4.4$ Head & neck, tunkGroup 2 $47.7$ $47.0$ $0.7$ $90$ $86$ $4.4$ Head & neck, tunkGroup 2 $43.5$ $39.3$ $4.2$ $177$ $162$ $8.5$ Head & neck, tunkGroup 2 $31.9$ $23.1$ $8.8$ $152$ $128$ $4.6$ Head & neckGroup 1 $37.4$ $11.5$ $25.6$ $1,517$ $357$ $6.5$ Head & neckGroup 1 $45.4$ $21.6$ $1,208$ $891$ $26.2$ Head & neckGroup 1 $45.4$ $26.6$ $1,208$ $891$ $26.2$ Head & neckGroup 1 $45.4$ $26.6$ $4.2$ $891$ $79$ $76.5$ Head & neckGroup 1 $25.8$ $9.1$ $11.8$ $53$ $381$ $76$ $78$ Head & neckGroup 1 $26.9$ $9.1$ $11.8$ $53$ $381$ $76$ $78$ Head & neckGroup 1 $25.8$ $9.1$ $11.8$ $79$ $79$ $76.5$ Head & neckGroup 1 $25.8$ $9.1$ $11.8$ $79$ $79$ $78$ Head & neckGroup 1 $25.8$ $9.1$ $11.8$ $79$ $79$ $78$ Head & neckGroup 1 $25.8$ $9.$		с	Head & neck	Group 1	38.8	9.3	29.5	49	34	30.9	No
Head & neck Group 1 43.4 7.6 35.8 7 1 85.7   Head & neck Group 1 45.5 6.5 39.0 22 2 90.9   Head & neck, trunk Group 2 47.7 47.0 0.7 90 86 4.4   Head & neck, trunk Group 2 43.5 39.3 4.2 177 162 8.5   Head & neck, trunk Group 2 31.9 23.1 8.3 4.2 177 162 8.5   Head & neck Group 1 37.4 11.5 25.9 1.517 357 76.5   Head & neck Group 1 40.9 15.3 25.6 1.517 357 76.5   Head & neck Group 1 40.9 15.3 25.6 1.517 357 76.5   Head & neck Group 1 19.9 8.1 1.51 357 76.5   Itunk Group 1 25.6 1.50 1.50 37.4 1.5 1.5		S	Upper extremity	Group 1	46.2	46.1	0.1	115	111	3.5	Diarrhea
Head & neck Group 1 45.5 6.5 39.0 22 2 90.9   Head & neck, trunk Group 2 47.7 47.0 0.7 90 86 4.4   Head & neck, trunk Group 2 43.5 39.3 4.2 177 162 8.5   Head & neck, trunk Group 2 31.9 23.1 8.4 152 15.8 8.5   Head & neck Group 1 37.4 11.5 25.6 1,517 357 76.5   Head & neck Group 1 40.9 15.3 25.6 1,208 891 26.2   Head & neck Group 1 19.9 81 11.8 25.6 1,517 357 76.5   Head & neck Group 1 19.9 26.6 1,517 357 76.5   Tunk Group 1 19.9 27.6 1,517 357 76.5   Head & neck Group 1 19.9 26.6 1,517 357 76.5   Tunk Gr		ო	Head & neck	Group 1	43.4	7.6	35.8	7	-	85.7	No
Head & neck, tunk Group 2 47.7 47.0 0.7 90 86 4.4   Head & neck, tower Group 2 43.5 39.3 4.2 177 162 8.5   wattemity Group 2 43.5 39.3 4.2 177 162 8.5   Head & neck, tounk, upper extremity Group 1 37.4 11.5 28.1 18.8 157 15.8   Head & neck Group 1 37.4 11.5 25.6 1,517 357 76.5   Head & neck Group 1 40.9 15.3 25.6 1,208 891 26.2   Head & neck Group 1 40.9 15.3 25.6 1,208 891 26.2   Hunk Group 1 19.9 8.1 11.8 537 36.2   Head & neck Group 1 25.8 9.1 11.8 537 36.2   Hunk Group 1 25.8 9.1 11.6 79 78 78   Inunk <td< td=""><td></td><td>ę</td><td>Head &amp; neck</td><td>Group 1</td><td>45.5</td><td>6.5</td><td>39.0</td><td>22</td><td>2</td><td>90.9</td><td>No</td></td<>		ę	Head & neck	Group 1	45.5	6.5	39.0	22	2	90.9	No
Head &neck, lower Group 2 4.3 39.3 4.2 177 162 8.5   wattermity Group 2 31.9 23.1 8.8 152 128 15.8   Head & neck, tunk, upper extremity Group 1 37.4 11.5 25.9 1,517 357 76.5   Head & neck Group 1 37.4 11.5 25.6 1,208 891 26.2   Head & neck Group 1 40.9 15.3 25.6 1,208 891 26.2   Itunk Group 1 19.9 8.1 11.8 53 38 27.8   Itunk Group 1 25.8 9.1 16.7 37 76.5   Itunk Group 1 25.8 9.1 16.7 37 76.5   Itunk Group 1 25.8 9.1 16.7 76.5 76.5   Itunk Group 1 25.8 9.1 16.7 76.5 76.5   Itunk Group 1 25.8 9.1		S	Head & neck, trunk	Group 2	47.7	47.0	0.7	06	86	4.4	No
Head & neck, tunk, upper externity Group 2 31.9 23.1 8.8 152 128 15.8   Head & neck Group 1 37.4 11.5 25.9 1,517 357 76.5   Head & neck Group 1 40.9 15.3 25.6 1,208 891 26.2   Head & neck Group 1 45.4 2.6 42.8 189 17 91.0   Itunk Group 1 19.9 8.1 11.8 53 382 27.8   Head & neck Group 1 25.8 9.1 16.7 130 79 27.8   Itunk Group 1 25.8 9.1 16.7 130 79 39.2   Itunk Group 1 25.8 9.1 16.7 130 79 27.8   Itunk Group 1 25.8 39.9 37.4 16.7 76 76.5   Itunk Group 1 45.5 5.9 37.4 1 76 76 76		с С	Head &neck, lower extremity	Group 2	43.5	39.3	4.2	177	162	8.5	No
Head & neck Group 1 37.4 11.5 25.0 1,517 357 76.5   Head & neck Group 1 40.9 15.3 25.6 1,208 891 26.2   Head & neck Group 1 45.4 2.6 42.8 189 17 91.0   Tunk Group 1 19.9 8.1 11.8 53 38 27.8   Head & neck Group 1 25.8 9.1 16.7 130 79 39.2   Inurk Group 1 25.8 9.1 16.7 130 79 39.2   Inurk Group 1 25.8 9.1 16.7 130 79 39.2   Inurk Group 1 47.3 9.9 37.4 4 1 75.0   Head & neck Group 1 45.5 6.5 39.0 42 9 76 76.0   Upper extremity Group 1 40.2 11.1 29.1 111 76 76 76		ო	Head & neck, trunk, upper extremity	Group 2	31.9	23.1	8.8	152	128	15.8	Fever
Head & neck Group 1 40.9 15.3 25.6 1,208 891 26.2   Head & neck Group 1 45.4 2.6 42.8 189 17 91.0   Trunk Group 1 19.9 8.1 11.8 53 38 27.8   Head & neck Group 1 25.8 9.1 16.7 130 79 39.2   Trunk Group 1 25.8 9.1 16.7 130 79 39.2   Trunk Group 1 47.3 9.9 37.4 4 1 75.0   Head & neck Group 1 45.5 6.5 39.0 42 9 78.6   Upper extremity Group 1 40.2 11.1 29.1 111 56 49.5		ო	Head & neck	Group 1	37.4	11.5	25.9	1,517	357	76.5	No
Head & neck Group 1 45.4 2.6 42.8 189 17 91.0   Tunk Group 1 19.9 8.1 11.8 53 38 27.8   Head & neck Group 1 25.8 9.1 16.7 130 79 39.2   Tunk Group 1 25.8 9.1 16.7 130 79 39.2   Head & neck Group 1 47.3 9.9 37.4 4 1 75.0   Head & neck Group 1 45.5 6.5 39.0 42 9 76.0   Upper extremity Group 1 40.2 11.1 29.1 111 56 49.5		с	Head & neck	Group 1	40.9	15.3	25.6	1,208	891	26.2	No
Tunk Group 1 19.9 8.1 11.8 53 38 27.8   Head & neck Group 1 25.8 9.1 16.7 130 79 39.2   Tunk Group 1 25.8 9.1 16.7 130 79 39.2   Head & neck Group 1 47.3 9.9 37.4 4 1 75.0   Head & neck Group 1 45.5 6.5 39.0 42 9 76.0   Upper extremity Group 1 40.2 11.1 29.1 111 56 49.5		S	Head & neck	Group 1	45.4	2.6	42.8	189	17	91.0	No
Head & neck Group 1 25.8 9.1 16.7 130 79 39.2   Tunk Group 1 47.3 9.9 37.4 4 1 75.0   Head & neck Group 1 45.5 6.5 39.0 42 9 78.6   Upper extremity Group 1 40.2 11.1 29.1 111 56 49.5		3	Trunk	Group 1	19.9	8.1	11.8	53	38	27.8	No
Trunk Group 1 47.3 9.9 37.4 4 1 75.0   Head & neck Group 1 45.5 6.5 39.0 42 9 78.6   Upper extremity Group 1 40.2 11.1 29.1 111 56 49.5		33	Head & neck	Group 1	25.8	9.1	16.7	130	79	39.2	No
Head & neck Group 1 45.5 6.5 39.0 42 9 78.6   Upper extremity Group 1 40.2 11.1 29.1 111 56 49.5		33	Trunk	Group 1	47.3	9.9	37.4	4	-	75.0	No
Upper extremity Group 1 40.2 11.1 29.1 111 56 49.5		ю	Head & neck	Group 1	45.5	6.5	39.0	42	6	78.6	No
		3	Upper extremity	Group 1	40.2	11.1	29.1	111	56	49.5	No

Table 2. Response to treatment (color)	eatment (co	olor)											
	Totol		Sex		No. 0	No. of hemangiomas	mas	Locatio	Location of hemangioma	gioma	Age of	Age of patients at start	art
Category	101al (n = 23)	Male (n = 3)	Female (n = 20)	P-value	Group 1 (n = 19)	Group 2 (n = 4)	P-value	H&N (n = 13)	Non-H&E (n = 1)	P-value	Before 6 mo After 6 mo $(n = 14)$ $(n = 9)$	After 6 mo $(n = 9)$	P-value
Response to treatment (color)													
Mean initial color (a-value)	$39.7 \pm 7.9$	39.7 ± 7.9 46.1 ± 1.1	$38.8 \pm 8.1$	0.137	$39.5\pm8.3$	$41 \pm 6.7$	0.732	$39.2 \pm 7.5$	39.2 ± 7.5 40.4 ± 8.8	0.733	40.3±8.2	$38.9 \pm 7.9$	0.694
Mean final color (a-value)	17.0 ± 14.5	$3.9 \pm 2.3$	$19.0 \pm 14.6$	< 0.001	$13.0 \pm 11.9$	13.0 ± 11.9 36.4 ± 10.0	< 0.001	$12.5 \pm 11.6$	$23.0 \pm 16.3$	0.087	13.0±12.7	$23.3 \pm 15.7$	0.100
Mean color change (a-value)	22.7 ± 15.0 42.2 ± 3.9	$42.2 \pm 3.9$	$19.8 \pm 13.8$	< 0.001	$26.5 \pm 13.6$	$4.6 \pm 3.3$	< 0.001	$26.7 \pm 13.6$	26.7 ± 13.6 17.4 ± 15.8	0.144	27.2 ± 14.4	$15.6 \pm 13.7$	0.069
Values are presented as mean ± standard deviation. Group 1, 1 hemangioma lesion; Group 2, 2 or more hemangioma lesions; H&N, head and neck; Before 6 mo, the patients who were treated before 6 months after birth; After 6 mo, the patients who were treated after 6 months after birth.	± standard devi Group 2, 2 or r	ation. more hemangic	oma lesions; H&N	N, head and n€	sck; Before 6 mc	), the patients $v$	who were treate	sd before 6 mo	nths after birth; /	After 6 mo, the	: patients who wer	e treated after 6 r	nonths after
Table 3. Response to treatment (size)	eatment (siz	ze)											
	Totol		Sex		No. of	No. of hemangiomas	las	Locatio	Location of hemangiomas	iomas	Age of	Age of patients at start	art
Category	nulai (n = 23)	Male (n = 3)	Female (n = 20) (P	MWU (P-value)	Group 1 (n = 19)	Group 2 (n = 4) (	MWU (P-value)	H&N (n=13)	Non-H&N MWU $(n = 10)$ (P-value)	MWU (P-value)	Before 6 mo After 6 mo $(n = 14)$ $(n = 9)$		MWU (P-value)
Doctorion to troatmost (ciro)													

Lategory	(n=23)	Male (n = 3)	Female (n = 20)	MWU (P-value)	Group 1 (n = 19)	Group 2 (n = 4)	MWU (P-value)	H&N (n = 13)	Non-H&N MWU (n=10) (P-value)	MWU (P-value)	Before 6 mo (n = 14)	Before 6 mo After 6 mo $(n = 14)$ $(n = 9)$	MWU (P-value)
Response to treatment (size)													
Initial size (mm²)	115.0 (4.0–1,517.0)	115.0 129.9 113.0   (4.0-1,517.0) (22.0-189.0) (4.0-1,517.0)	113.0 (4.0–1,517.0)	28.5 (0.898)	115.0 (4.0–1,517.0)	121.0 (32.0–199.0)	35.0 (0.845)	110.8 (7.0–1,517.0)	122.45 (4.0–1,208.0)	58.0 (0.692)	122.5 (4.0–1,517)	110.0 (42.0–234.0)	61.5 (0.926)
Final size (mm²)	56.0 (1.0–891.0)	17.0 (2.0–88.0)	67.5 (1.0–891.0)	17.5 (0.268)	56.0 (1.0–891.0)	107.0 (27.0–199.0)	28.0 (0.456)	34.0 (1.0–891.0)	88.5 (1.0–210.0)	42.0 (0.166)	47.0 (1.0–891.0)	86.0 (9.0–210.0)	46.5 (0.305)
Size difference/initial size (%)	32.3 (1.1–92.6)	90.9 30.3 (1.1–92.6) (1.1–92.6)	30.3 (1.1–92.6)	12.5 (0.115)	49.5 (1.1–92.6)	12.1 (4.0–16.0)	10.0 (<0.001)	49.5 (1.0–91.0)	21.8 (3.5–92.6)	40.5 (0.131)	62.3 (3.0–93.0)	15.8 (1.0–79.0)	28.5 (<0.001)
Values are presented as median (range). A Mann-Whitney U-test indicated that group 1 who has 1 hemangioma lesion showed a significantly greater size change (median, 49.5%) than group 2 (median, 12.1%; P < 0.001). Additionally, the change in size in patients who started treatment before 6 months of age (median, 62.3%) was greater than in their counterparts (median, 15.8%; P < 0.001). MWU, Mann-Whitney U-test; Group 1, 1 hemangioma lesion; Group 2, 2 or more hemangioma lesions; H&N, head and neck; Before 6 mo, the patients who were treated before 6 months after birth; After 6 mo, the patients who were treated before 6 months after birth.	iedian (range). A ize in patients wh ;: Group 1, 1 hem birth.	Mann-Whitney o started treatn angioma lesion	/ U-test indication ment before 6 π η; Group 2, 2 or	ed that group 1 nonths of age (π more hemangic	who has 1 he ledian, 62.3%) \ ima lesions; H&	mangioma lesi vas greater tha N, head and ne	ion showed a s n in their counti sck; Before 6 m	ignificantly gree erparts (median, o, the patients w	ater size change 15.8%; P < 0.0 ho were treateo	e (median, 49.5 101). 1 before 6 montl	5%) than group 2 hs after birth; After	(median, 12.1%) r 6 mo, the patier	; P < 0.001). Its who were

l adverse effects	No. (%)	0	0	0	3 (13.0)	s 0	1 (4.3)	
Table 4. Complications and adverse effects	Observed adverse effects	Symptomatic hypoglycemia	Hypotension or bradycardia	Cold hands or feet	Gastrointestinal problems	Aggravation of respiratory symptoms	Fever	

mg/kg/day (oral administration 3 times per day every 8 hours), and the patient is discharged. Schupp et al. [1] reported that patients admitted to the hospital were given a dose of 1 mg/kg/ day for 3 days. If no specific adverse events were observed, the dose was increased to 2 mg/kg/day, and the patient was discharged with this final dose. In other words, it is common to begin with a small dose of the drug initially and to gradually increase the dose over time. Leaute-Labreze et al. [12] reported that their most effective treatment involved a 6-month administration of propranolol at a dose of 3 mg/kg/day; thus, we prescribed 3 mg/kg/per day as our initial dose. In our experience, there were no major side effects in the 23 patients, except for 3 cases with minor problems such as diarrhea and fever.

In a large study of 1,264 IHs by Wedgeworth et al. [23], electrocardiography and echocardiography were not necessary for the detection of cardiovascular side effects during treatment with propranolol. Leaute-Labreze et al. [12] reported that adverse events related to propranolol (hypoglycemia, hypotension, bradycardia, and bronchospasm) did not occur frequently, with no significant differences observed between the placebo and treatment groups. However, Holland et al. [16] reported a case of hypoglycemia and recommended the discontinuation of propranolol therapy when accompanied by reduced dietary intake.

Noticeably, previous studies used subjective methods to assess the perceptible aesthetic treatment results. The reliability of subjective assessments is affected by several factors, such as the evaluator's prejudices, variation in lighting conditions, and even the emotional status of the infant. Thus, a subjective analysis can produce only partially reliable clinical implications. In our series, we used the objectively measured parameters of color difference and size, which were calculated with a digital camera and the Antera 3D system.

Interestingly, the response to propranolol was significantly lower in patients with a larger number of lesions than in patients with a smaller number of lesions in terms of color and size change. This finding suggests that propranolol does not effectively inhibit abnormal endothelial cell proliferation and angiogenesis, especially in patients with a large number of hemangiomas.

Male patients showed a greater color change (P < 0.001), whereas there was no sex difference in size changes (P < 0.001). The age at the start of treatment did not show a significant relationship with color change (P < 0.001), but patients who were treated before 6 months after birth showed a significantly greater size change than their counterparts (P < 0.001).

Corticosteroids were initially discovered to induce the accelerated regression of proliferating IHs in the 1960s [10,24] and have been considered the first-line treatment of IH for over 40 years [25]. However, because of recent clinical successes using the beta-blocker propranolol, the recommended first-line treatment for these lesions is changing.

Izadpanah et al. [18] compared propranolol and corticosteroids in terms of their treatment response rates, demonstrating that the response to propranolol was significantly better (97.3% vs. 71.0%, respectively; P < 0.001), consistent with our study findings that 21 out of 23 patients responded to the treatment (91.3%). In their meta-analysis of results related to corticosteroid use, the most common side effects reported were altered growth (6%) and development of a moon face (5%). Other reported side effects included osteoporosis, fungal infections, and hypertension. Side effects were encountered in 475 of 2,697 patients (17.6%). Meanwhile, 96 of 699 patients (13.7%) experienced side effects after propranolol therapy. The complication rate of steroid therapy was thus higher than that associated with propranolol [18]. The side effects observed at our institution were primarily gastrointestinal problems and fever (17.3%); although this rate was high, these complications may not have been directly related to propranolol treatment.

This study has limitations in that the number of patients in each group was relatively small; the inclusion of additional cases would improve the significance of our findings. In addition, further large-scale studies are needed to clarify the effects of propranolol on infant growth and development after long-term propranolol treatment. Another limitation is a lack of randomized controlled trials comparing propranolol therapy for IH with a more conservative treatment. Randomized controlled trials with a set dosage, duration of therapy, and appropriate follow-up are necessary to validate the observed therapeutic benefits of propranolol. The examined factors of the follow-up period, age at the initiation of treatment, treatment duration, and patient weight could influence the response to propranolol therapy. It is thus necessary to control for these factors in future studies.

In conclusion, propranolol is efficacious and has a good safety profile for the treatment of IH. A short duration of treatment ( < 6 months) at a dose of 3 mg/kg/day is able to stabilize and induce regression in certain IH patients. In patients with a single lesion, the response to treatment was better in terms of color fading and size reduction. Furthermore, male patients responded better to propranolol treatment in terms of color fading than female patients, and starting treatment before 6 months after birth was more advantageous for size reduction.

#### NOTES

#### **Conflict of interest**

No potential conflict of interest relevant to this article was reported.

## **Ethical approval**

The study was approved by the Institutional Review Board of Severance Hospital (IRB No. 4-2017-1158) and performed in accordance with the principles of the Declaration of Helsinki. Written informed consents were obtained.

#### Patient consent

The patients provided written informed consent for the publication and the use of their images.

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