Calcification in Werner syndrome associated with lymphatic vessels aging

Hideyuki Ogata¹, Shinsuke Akita¹, Sanae Ikehara²,³, Kazuhiko Azuma³, Takashi Yamaguchi³, Maihulan Maimaiti⁴, Yoshito Maezawa⁴,⁵, Yoshitaka Kubota¹, Koutaro Yokote⁴,⁵, Nobuyuki Mitsukawa⁵, Yuzuru Ikehara²,³

¹Department of Plastic, Reconstructive, and Aesthetic Surgery, Graduate School of Medicine, Chiba University, Chiba, Japan
²Department of Pathology, Graduate School of Medicine, Chiba University, Chiba, Japan
³Cellular and Molecular Biotechnology Research Institute, National Institute of Advanced Industrial Science and Technology (AIST), Tsukuba, Japan
⁴Department of Endocrinology, Hematology and Gerontology, Graduate School of Medicine, Chiba University, Chiba, Japan
⁵Division of Diabetes, Metabolism and Endocrinology, Chiba University Hospital, Chiba, Japan

Correspondence to: Shinsuke Akita, Yuzuru Ikehara; email: shinsukeakitarps@gmail.com, https://orcid.org/0000-0001-7075-6602; yuzuru-ikehara@chiba-u.jp

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ABSTRACT

In addition to the symptoms of aging, the main symptoms in Werner syndrome (WS), a hereditary premature aging disease, include calcification of subcutaneous tissue with solid pain and refractory skin ulcers. However, the mechanism of calcification in WS remains unclear. In this study, the histological analysis of the skin around the ulcer with calcification revealed an accumulation of calcium phosphate in the lymphatic vessels. Moreover, the morphological comparison with the lymphatic vessels in PAD patients with chronic skin ulcers demonstrated the ongoing lymphatic remodeling in WS patients because of the narrow luminal cross-sectional area (LA) of the lymphatic vessels but the increment of lymphatic microvessels density (MLVD). Additionally, fluorescence immunohistochemical analysis presented the cytoplasmic distribution and the accumulation of WRN proteins in endothelial cells on remodeling lymphatic vessels. In summary, these results point out a relationship between calcification in lymphatic vessels and the remodeling of lymphatic vessels and suggest the significance of the accumulation of WRN mutant proteins as an age-related change in WS patients. Thus, cytoplasmic accumulation of WRN protein can be an indicator of the decreasing drainage function of the lymphatic vessels and the increased risk of skin ulcers and calcification in the lymphatic vessels.

INTRODUCTION

Werner syndrome (WS) is a rare autosomal recessive disorder in which a variety of age-related diseases appear at an early age [1, 2]. The cause of this disease is a mutation of the WRN gene located in the WS locus at 8 p 12 [3]. WRN proteins have helicase and exonuclease activities [4–6] and appear to be involved in DNA repair [7].

Typical age-related disorders in WS include arteriosclerosis, malignant neoplasms, type 2 diabetes mellitus, dyslipidemia, osteoporosis, and ocular infections. In addition, calcification of the skin and soft tissues is associated with severe pain and skin ulceration, which is often challenging to treat and, therefore, impairs patients' quality of life [8]. It is noteworthy that calcification of the skin and soft tissues is a characteristic feature of WS: [1, 2, 9–12]
because it is often painful enough to lead to a physician's visit [9–13]. Moreover, calcification is used as an indicator for an early diagnosis of WS in less clear cases of age-related diseases [13] because calcification in WS presents on x-rays with characteristic flame-like shadows are pathognomonic findings and differ from the calcification patterns in other disorders [12].

Our previous study showed that overexpression of Na-Pi cotransporter (Pit-1) in dermal fibroblasts in WS contributed to deposit calcium in vitro as the cause of calcification [13], while the mechanism of calcification at the tissue level in vivo has not yet been shown despite the essential findings to diagnosis for WS in current guidelines.

In the present study, we examined the pathological features of skin ulcer lesions with the characteristic calcification of WS. We observed that the calcification in patients with WS spread along the lymphatic vessels on the boundary between the dermis and subcutaneous tissue. and that the proliferation of lymphatic vessels with a narrow lumen was prominent in the interstitial tissue with less calcification. Because these findings strongly suggested a relationship with the radiographic appearance of flame-like shadows, we examined the expression of WRN protein in lymphatic endothelial cells (LECs) by immunohistochemical staining to investigate the relationship between soft tissue calcification and lymphatic vessel function in WS.

RESULTS

WS causes calcification in the lymphatic vessels of the skin around the ulcer

Radiographs of four patients with WS who had ulcers on the elbow revealed calcification in the subcutaneous tissue of the ulcerated elbow joint in all patients (Figure 1A, 1B). In skin tissue taken from the same ulcers, calcification was present in three of the four patients (Figure 1C, 1D). We observed the skin tissue of a patient with WS (WS2) with calcification of the skin by scanning electron microscopy. There was a deposition of crystalline substances in the luminal structure under the dermis (Figure 2A, 2B). Next, we performed energy dispersive X-ray (EDX) analysis of the crystalline substance (Figure 2C, 2D) and detected the characteristic X-rays of phosphorus and calcium with high counts suggestive that the crystalline substance was calcium phosphate. To determine whether tissue other than luminal structures contained calcium, we compared calcium content in luminal structures, dermal connective tissues, and areas that did not contain tissue by EDX. We measured three points in each area and performed similar measurements in three fields (Figure 2E). One-way analysis of variance (ANOVA) was performed for the three groups, followed by Tukey's test. The calcium content in the luminal structure was significantly higher than that in the dermal connective tissue (P<0.0001; mean difference 22101 keV; CI=-18844 to -25359) and the calcium content in the dermis connective tissue was not significantly different from that in the non-tissue region (P=0.0001; mean difference -1985 μm2; CI=-2514 to -1456) (Figure 2F). Immunohistochemical staining was performed using podoplanin, CD31, and αSMA antibodies to identify the luminal structure in which the calcified material was deposited. The luminal structure around the calcification was positive for the D2 -40 antibody, weakly positive for the CD 31 antibody, and negative for the αSMA antibody. These results indicate that this luminal structure was a LEC (Figure 3A–3D).

Increased lymphangiogenesis and impaired lymphatic vessel space development around ulcers in patients with WS

To clarify the histological features of lymphatic vessels in patients with WS, we compared immunohistochemical staining using the podoplanin antibody in the skin of the non-ulcerated area (non-ulcer WS group) and the skin around the ulcer (ulcer WS group) of four patients with WS and the skin around the foot ulcer (ulcer PAD group) of three patients with peripheral arterial disease (PAD). In the non-ulcer WS group, the lymphatic vessels were poorly endoluminal and sparse in density. In the ulcer WS group, lymphatic vessels with poorly developed lumens proliferated in an irregular arrangement. In the ulcer PAD group, the lymphatic lumina expanded and aligned linearly (Figure 4B–4D). We measured the intradermal luminal cross-sectional area (LA) of the lymphatic vessels in the three groups (Table 1). One-way analysis of variance (ANOVA) was performed for the three groups, followed by Tukey’s test. LA in the ulcer WS group was larger than in the non-ulcer WS group (P=0.0034; mean difference -730.1 μm2; CI=-1220 to -240.0) and smaller than in the ulcer PAD group (P<0.0001; mean difference -1985 μm2; CI=-2514 to -1456) (Figure 4E). Microlymphatic vessel density (MLVD) was measured in all three groups (Table 1). According to Tukey test, post one-way ANOVA, MLVD in the ulcer WS group was higher than in the non-ulcer WS (P<0.0001; mean difference -1.21x10^5/μm2; CI=1.67x10^5 to -7.60x10^6) and ulcer PAD groups (P<0.0001; mean difference 1.27x10^5/μm2; CI=7.80x10^6 to 1.766x10^5) (Figure 4F).
In LECs of WS, WRN protein accumulates in the cytoplasm.

We investigated the subcellular localization of WRN protein in lymphatic endothelial cells of ulcerated skin tissue of 2 WS patients (WS1, WS4) and 1 PAD patient (PAD1) using a confocal microscope (Figure 5). In PAD patients, WRN protein is mainly present in the perinuclear lesion and shows nucleolus-like staining in nuclear. Conversely, the expression in WS1 and WS4

Figure 1. Clinical images of patients. A series of photographs (A) of elbow ulcers in four patients with Werner’s syndrome (WS1-4), radiographs (B) of the same area, and HE stained tissue (C, D) of the skin around the ulcers. WS1: 62-year-old male, WS2: 48-year-old male, WS3: 50-year-old female, WS4: 57-year-old male. (A) The skin around the ulcer is sclerotic and atrophic. (B) Various degrees of calcification were observed in the soft tissues around the elbow joint. In WS2 and WS3, it becomes a shadow of the flame state. (C, D) Strong fibrosis of the dermis (all cases) and calcification within the luminal structures (WS2, 3, 4). Scale bars are 700 μm (C) and 200 μm (D).
Figure 2. Scanning electron microscopy images and EDX analysis. Scanning electron micrograph of skin around an ulcer on a patient’s elbow with WS (A: x 60, B: x 200). Crystalline substance in luminal structures under the dermis (red squares). This crystalline substance was examined with energy dispersive X-ray (EDX) (C, D). Each color and element combination corresponds to C for yellow, O for orange, Na for light blue, Mg for pink, Si for green, P for blue, and Ca for red. Using EDX, we measured the calcium content of the luminal structure, dermal connective tissue, and non-tissue areas at three points in each area (yellow circles), and performed similar measurements in three fields (E). Comparison of the calcium content of the luminal structure, dermal connective tissue, and non-tissue areas (F). Data are expressed as mean ± standard error. One-way ANOVA followed by Tukey test were performed (* p < 0.01, **** p < 0.0001).
diffusely increased and distributed through the cytoplasm of the podoplanin positive cells identified as LECs. LECs with the increased WRN protein in the cytoplasm appeared in the regenerative LECs consisting of the lymphatic vessels with structural abnormalities. Moreover, WS4 showed the increased WRN protein in the cytoplasm and the nucleus. In conclusion, mutated WRN protein accumulation in the cytoplasm may

Figure 3. Immunohistochemical staining image of the patient's skin. Various stained tissue images of calcified tissues from a patient with WS. (A) HE. (B) Immunohistochemical staining with podoplanin antibody. Cells around the calcification were positive. (C) Immunohistochemical staining with αSMA antibody. Cells around the calcification were negative. (D) Immunohistochemical staining with CD31 antibody. The cells around the calcification were weakly positive. All scale bars are 200 μm.
Figure 4. Image analysis of lymphatic vessels in the skin of patients. (A) The area of skin tissue sections was measured by tracing the skin margins using an Aperio Image Scope. (B) A representative example of skin histology taken from an ulcer-free area in a patient with Werner syndrome. Scale bar is 300 μm. The lumen is narrow and small in number. (C) A representative example of skin tissue taken from the ulcer circumference of a patient with PAD. Scale bar is 300 μm. The lumen is dilated and lined up. (D) A representative example of skin tissue taken from the ulcer circumference of a Werner syndrome patient. Scale bar is 300 μm. The lumen is indistinct and randomly proliferating. (E) Comparison of luminal cross-sectional area (LA) of the lymphatic vessels. In four patients with Werner syndrome, two different sites were measured in the non-ulcerated area and in the skin surrounding the ulcer. Three patients with PAD underwent measurements at two different sites of skin around the ulcer. Data are expressed as mean ± standard error. One-way ANOVA followed by Tukey test were performed (* * p < 0.001, **** p < 0.0001). (F) Comparison of lymphatic vessel density (MLVD). In four patients with Werner syndrome, two different sites were measured in the non-ulcerated area and in the ulcer’s skin. Three patients with PAD underwent measurements at two different sites of skin around the ulcer. Data are expressed as mean ± standard error. One-way ANOVA followed by Tukey test were performed (n.s. No significant difference, **** p < 0.0001).
Table 1. LA and MVLD.

| Group           | N | LA (μm²) | MVLD (/μm²) |
|-----------------|---|----------|-------------|
|                 |   | Mean SE  | Mean SE     |
| non-ulcer WS    | 8 | 434.7 40.25 | 7.24x10⁻⁶ 8.15x10⁻⁷ |
| ulcer WS        | 8 | 1165 139.3 | 1.94x10⁻⁵ 1.91x10⁻⁶ |
| ulcer PAD       | 6 | 3150 234.5 | 6.68x10⁻⁶ 3.04x10⁻⁷ |

LA, luminal cross-sectional area; MLVD, lymphatic microvessel density; WS, Werner syndrome; SEM, standard error of the mean.

associate with the developing abnormalities in the lymphatic vessels. One of the plausible mechanisms is that the increased accumulation of WRN in the cytoplasm causes the degeneration and the abnormal remodeling of lymphatic vessels with aging in WS patients, resulting in abnormalities in the lymphatic vessels and the developed luminal crystalloids through the impaired drainage function.

**DISCUSSION**

This study reports that the subcutaneous calcification of WS grows in lymphatic vessels, and the crystalloids materials in luminal grow with the continuing disorders on lymphatic vessels related to aging. Moreover, it reveals the characteristic features of the subcutaneous calcification of WS patients that the lymphatic endothelial cells stored the WRN mutant protein in the...
cytoplasm and formed narrower vessels than them in the control group. Furthermore, it is noteworthy that lymphatic vessels are of the mesoderm origin. Because age-related disorders of WS generally occur predominantly on mesodermal organs such as bones and arteries, the subcutaneous calcification of WS can be regarded as the age-related changes in WS patients [14].

First, we performed a histological and ultrastructural analysis of soft tissue calcification appearing in WS and observed that crystalline substances consisting mainly of calcium phosphate were distributed exclusively in expanded lymphatic vessels located deep in the subcutaneous tissue. The crystalline material was deposited between the dermis and subcutaneous tissue, with prominent lymphangiogenesis of the small luminal structures. A previous report indicated that cultured dermal fibroblasts from WS deposit calcium in vitro [13]. Similarly, it is inferred that calcium phosphate is overproduced by WS fibroblasts, even in vivo.

In this study, we revealed that the site of subcutaneous calcification in WS patients is lymphatic vessels. Conversely, it has been considered that fibroblast ossification forms subcutaneous calcification of WS patients. Because calcification occurred on the in vitro cultured skin fibroblasts from WS patients, and the cells exhibited a higher bone-inducing ability in vitro assays [13, 15]. In contrast, this study demonstrated that the crystalloids materials exclusively exist in lymphatic vessels using quantitative elemental analysis for the subcutaneous calcification from WS patients. Moreover, we detected no significant increase in the amount of calcium element in the stromal tissue surrounding the lymphatic vessels. Therefore, generally, we concluded that lymphatic vessels are a place where calcified crystalloids grow in WS patients.

PAD is a specific disease that forms ulcers in the extremities of older adults and has characteristics of arteriosclerotic calcification. Therefore, we used PAD as a control group to clarify the characteristics of WS. There was no evidence of lymphatic calcification in the ulcerated area of PAD, suggesting that lymphatic calcification may be involved in the specific mechanism of WS.

The other characteristic findings in subcutaneous calcification of WS patients is the decreased diameter of lymphatic vessels, which can refer to abnormal aging in WS. In support of this hypothesis, it has been pointed out that decreased diameter and drainage function in lymphatic vessels are typical age-related changes [16, 17]. In this respect, WS needs to link with lymphatic changes as an age-related disease, whereas PAD must correlate with arteriosclerotic calcification.

It is noteworthy that calcification in the lymphatic vessels appears in only the ulcerative tissue and that the density was increased. This is not observed in non-ulcerated areas and is a distinct feature of typical age-related changes, suggesting the presence of unique factors in WS ulcerated areas. In this regard, studies comparing skin fibroblasts from the trunk of WS patients with those from lower extremity limbs have shown that lower extremity fibroblasts strongly express genes related to bone formation. Differences in preload of calcium phosphate to lymphatic vessels may account for differences in findings by site [15]. Furthermore, not all tissues are generally aged in WS. Responses to inflammatory lymphangiogenesis signals may be preserved compared with ordinary older adults, and further studies are needed to clarify differences in lymphatic vessel density.

This study found that LECs of WS patients showed accumulation of WRN protein diffusively distributed in the cytoplasm by confocal microscopic analysis. Moreover, we raised the possibility that mineralization in the lymphatic vessels links with abnormality in lymphatic endothelial cells with WRN accumulation. In this respect, we consider that the cytoplasmic accumulation of WRN proteins is one of the factors as aging in LECs, which causes the impaired functions of lymphatic vessels through the increasing cellular stresses, either supplying or draining interstitial fluid and metabolites.

Although, to date, various types of mutations in the WRN gene are in WS patients, while mutated proteins generally lack nuclear localization signals [2, 18–20]. Supporting this hypothesis, earlier studies using knock-in mice have shown that WRN mutant proteins accumulated in the cytoplasm of WRN mutant mice cause oxidative stress in the cells, resulting in various symptoms associated with aging [21–23]. However, the phenotype in WS patients does not appear in the knock-out mice lacking the WRN gene.

Therefore, it would be necessary for aging research to study the mechanisms between the lymphatic vessels and the cellular stress derived from the cytoplasmic accumulation of the mutated WRN protein. As there was no difference in clinical symptoms between patients with/without nuclear localization of WRN (WS1 and WS4), the prognostic roles on the diminished WRN protein in nuclear need to be elucidated in further study.

There are some limitations to this study. First, we did not use normal adult skin as a control instead, we initially used skin tissue discarded during surgery to treat ulcers. However, it is expected that there is a
difference even among healthy individuals and WS patients because there was a significant difference in comparison with the PAD patient who did not have WS.

Next, this study clarified the pathology of skin calcification in WS histopathologically, and we have not yet elucidated its etiology. To explain the mechanism by which the accumulation of intracellular WRN mutant protein brings about the change of the lymph duct and what kind of difference of the clinical manifestation the degree of the accumulation brings about, further research in the future is necessary. Finally, the sample size in this study was small. Because Werner syndrome is a rare disorder, removing skin tissue for therapeutic purposes is even rarer, it is not easy to conduct extensive observational studies. Most of the research reports on Werner syndrome so far have only a few cases. We analyzed specimens from valuable surgical cases and identified cutaneous soft tissue calcification features, a specific symptom in 80% of Werner's syndrome. Whether or not the findings obtained in this study are universally present needs to be widely verified and confirmed in future studies.

In this study, we observed that immature lymphatic capillaries proliferated in the skin around ulcers in WS, and calcium phosphate accumulated in the lumen. In the cytoplasm of lymphatic endothelial cells of patients with WS, WRN mutant proteins accumulated at high concentrations. These results suggest that the accumulation of mutant proteins damages lymphatic endothelial cells, causing changes in lymphatic vessels similar to aging, resulting in reduced drainage function of lymphatic vessels in WS. This decreased lymphatic drainage function is thought to be one of the causes of subcutaneous tissue calcification, prolonged inflammation, and delayed wound healing. Our new findings have revealed part of the calcification, ulceration, and other conditions that plague many WS patients and could be a new step toward further research.

MATERIALS AND METHODS

Patient and control individuals

This study was approved by the Institutional Review Board of the Graduate School of Medicine, Chiba University (approval number: 973). The participants provided written informed consent before participating in the study.

We analyzed the skin tissues of patients with WS who developed painful ulcers at radiographically calcified sites (elbows and ankles) and required surgical treatment. The control group consisted of skin tissue from patients with chronic skin ulcers of the lower extremities associated with PAD and chronic renal failure. The profiles of the included patients are presented in Table 2. The types of WRN mutations in each patient with WS are listed in Table 3. These mutations resulted in the loss of the C-terminus after 1369-1402aa, where a nuclear localization signal was present. Mutation c. 3139 -1 G > C of WS2, 3, and 4 is the most common mutation in Japan [24].

Tissue sampling

In patients with WS, we sampled sclerotic skin around the ulcer that was removed simultaneously with debridement and normal skin that was used as a donor site for skin grafting. Resected specimens were fixed in 10% buffered formalin. In the PAD patient group, we sampled the skin at the boundary between the necrotic tissue and normal tissue from the amputated forefoot. All specimens were fixed in 10% buffered formalin and embedded in paraffin (FFPE tissues).

Detection of crystalline substance using scanning electron microscopy

Scanning electron microscopy was used to map the distribution of crystalline substance in the tissue sections using an energy dispersive X-ray analyzer (SEM-EDX, TM4000Plus, Hitachi High-Technologies, Tokyo, Japan). Briefly, the 4 μm thick FFPE tissue sections were deparaffinized with xylene and washed well with 100% ethanol to eliminate wax and xylene remaining on the sections and dried. Then, well-dried sections were coated at 3 nm thickness with osmium using an osmium coater Neoc (Meiwafosis Co., Ltd., Japan) instrument and subjected to SEM-EDX analysis.

Histological analysis

Histological analysis was performed on hematoxylin-eosin (H-E) stained 4um sections prepared from FFPE surgically resected skin tissues as previously described.

Immunohistochemical staining was performed using the Ventana XT system BenchMark (Ventana Medical Systems, Tucson, AZ, USA). Briefly, the tissue sections were automatically treated with an antigen retrieval solution (Ventana) and heated on a slide heater at 100° C for 30 min. Endogenous peroxidase activity was halted by immersion in 3% hydrogen peroxide for 4 min. The sections were then incubated with primary antibodies for 30 min at 37° C. According to the manufacturer's instructions, the LSAB Ventana Iview DAB detection system (Ventana) detects the immune complex and hematoxylin counterstained nuclei of 4um sections prepared from FFPE surgically resected tissues. Antibodies against podoplanin (D2-40, Ventana),
Table 2. The profiles of the patients.

| Patient | Age | Sex | Location | Comorbidity                      |
|---------|-----|-----|----------|----------------------------------|
| WS1     | 62  | M   | elbow    | DM, dyslipidemia, osteoporosis, CKD |
| WS2     | 48  | M   | elbow    | DM, dyslipidemia, osteoporosis   |
| WS3     | 50  | F   | elbow    | DM, dyslipidemia, osteoporosis   |
| WS4     | 57  | M   | elbow    | DM, dyslipidemia, osteoporosis   |
| PAD1    | 56  | M   | foot     | DM, CKD                         |
| PAD2    | 71  | M   | foot     | DM, CKD                         |
| PAD3    | 71  | M   | foot     | CKD                             |

WS, Werner syndrome; PAD, peripheral arterial disease; DM, diabetes mellitus; CKD, chronic kidney failure.

Table 3. WRN mutations in the patients.

| Patient | Nucleotide notation | Resulted codon | Resulted proteins (aa) |
|---------|---------------------|----------------|------------------------|
| WS1     | homo c.1105C>T      | Nonsense mutation | 368                   |
| WS2     | homo c.3139-1G>C     | Frame shift/Nonsense mutation | 1060 |
|         | c.3913C>T          | Nonsense mutation | 1304                  |
| WS3     | c.3139-1 G>C         | Frame shift/Nonsense mutation | 1060 |
| WS4     | c.3139-1 G>C         | Frame shift/Nonsense mutation | 1060 |
|         | c.3446 del T         | Frame shift/Nonsense mutation | 1160                  |

WS, Werner syndrome; aa, amino acid.

smooth muscle actin (1A4, Ventana), and CD31(JC70, Ventana), respectively, were used to detect cytoplasmic expression in blood and lymphatic vessels, as previously reported [25].

**Image analysis**

All image analyses were performed on digital slides (Leica Biosystems Aperio ImageScope v 12.4.7018). Microlymphatic vessel density (MLVD) in skin tissue was calculated by counting all D2-40 positive lymphatic vessels and dividing by the skin area. The skin area and luminal cross-sectional area (LA) of the lymphatic vessels were measured by tracing the lymphatic vessels on a digital slide (Figure 4A). LA was calculated as the mean of the largest to the 10th largest. To eliminate data variability, we performed measurements on two samples per case.

**Histological analysis through immunofluorescence detection**

Monoclonal antibodies against α-WRN antibody (ab200, Abcam) were used to detect Werner's syndrome helicase protein expression. This antibody binds to the N-terminus of the WRN protein. In addition, both hoechst33342 (ThermoFisher Scientific) and anti-podoplanin labeled with Alexa Fluor 594 (BioLegend, CA, USA) were used to evaluate the cellular localization of Werner's syndrome helicase protein expression in lymphatic vessels with a confocal microscope (LSM710, Carl Zeiss).

**Statistical analyses**

Data are presented as means ± SEM as indicated. One-way ANOVA followed by Tukey’s test was performed to compare the groups. Differences between groups were considered significant at p < 0.05. All tests were performed using GraphPad Prism 9.10 software (San Diego, CA, USA).

**Abbreviations**

WS: Werner syndrome; LA: luminal cross-sectional area; MLVD: lymphatic microvessel density; LECs: lymphatic endothelial cells; PAD: peripheral arterial diseases; ANOVA: analysis of variance; SEM: standard error of the mean.

**AUTHOR CONTRIBUTIONS**

HO, SA, and YI designed the study, analyzed the data, and wrote the manuscript; HO, SI, KA, TY, and MM carried out the experiments and made the figures; HO and
YK conducted sampling from patients; KY and YM contributed to the recruitment of the subject and the analysis of the gene mutation; YK and NM discussed the data; all authors provided critical feedback and helped shape the research, analysis, and manuscript.

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CONFLICTS OF INTEREST

The authors declare that they have no conflicts of interest.

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