Primary breast angiosarcoma with disseminated intravascular coagulation is successfully treated with self-subcutaneous unfractionated heparin calcium injection: A case report

TOSHINARI YAGI1, HARUMI NAKAMURA2, TORU WAKAMATSU3, YOSHINORI IMURA3, HIRONARI TAMIYA3, HIDEAKI SABE3, KATSUNARI YAMASHITA3, MAKIYO WATANABE3 and SATOSHI TAKENAKA3

Departments of 1Outpatient Chemotherapy, 2Diagnostic Pathology and Cytology, and 3Orthopedic Surgery, Osaka International Cancer Institute, Osaka 541-8567, Japan

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Abstract. Angiosarcoma is a rare sarcoma with a poor prognosis and is prone to disseminated intravascular coagulation (DIC), where DIC often interferes with chemotherapy. Primary angiosarcoma of the breast (PASB) is a type of angiosarcoma that is located in mammary parenchyma and is not associated with radiation exposure. The current study reported a 47-year-old female with DIC associated with PASB. The DIC of the patient relapsed during mono-chemotherapy with paclitaxel (PTX) after first-line anticoagulant therapy using thrombomodulin-α. The second-line danaparoid sodium therapy, followed by self-subcutaneous injection of unfractionated heparin calcium (UFH), resulted in long-term stabilization of DIC. Under this second-line anticoagulant therapy, the patient continued chemotherapy and chemoradiotherapy for >13 months in the outpatient setting without impairment of quality of life. The present case suggested that self-subcutaneous injections of UFH may be a useful therapeutic option for long-term control of DIC associated with PASB. However, further prospective clinical trials are needed to verify the efficacy of self-subcutaneous injection of UFH in similar settings.

Introduction

Disseminated intravascular coagulation (DIC) is a well-known hemostatic complication of solid tumors. It is usually identified at the time of presentation and can sometimes cause excessive bleeding or thromboembolic complication. In comparison, primary angiosarcoma of the breast (PASB) is a malignant primary endothelial neoplasm of mammary parenchyma that is not associated with radiation exposure. The median age of patients at diagnosis is 40 years, compared with a median age of 70 years for the secondary angiosarcoma (1). Adem et al (2) reported that angiosarcoma accounts for one-fourth of all breast sarcomas, excluding metaplastic carcinomas and phylloides tumors. D'Angelo et al (3) performed a retrospective analysis of 119 patients with metastatic angiosarcoma, 24% of which had radiation-associated angiosarcoma. The most frequent primary sites were the chest wall/breast (37.31%), viscera (22%), and head/neck (20%). The median overall survival of patients with metastatic angiosarcoma with origin at all sites was 12.1 months. Even if DIC is a common complication associated with angiosarcomas and often interferes with patients' chemotherapy (4), there have been few case reports on anticoagulant therapies for DIC associated with angiosarcoma because angiosarcoma is rare. In the current study, we present a rare case of a PASB with DIC that was controllable by a self-subcutaneous injection of unfractionated heparin calcium (UFH).

Case report

In November 2019, a 47-year-old woman with a breast tumor and multiple lung and bone metastases was referred to our hospital for further diagnosis and treatment. She had consulted a nearby hospital complaining of swelling of the left breast in July 2018. Imaging examinations revealed a giant mass in her breast. Core needle biopsy was performed, resulting in the histological diagnosis of hemangioma. Lumpectomy was performed in September 2018. In August 2019, her left nipple became swollen, and computed tomography (CT) revealed bilateral multiple lung and bone metastases (Fig. 1).
At this point, a malignant tumor was suspected and she was transferred to our hospital. She was admitted to our hospital in November 2019. Upon admission, a red mass that was 3 cm in diameter was observed on the left nipple during physical examination. Whole body magnetic resonance imaging revealed multiple bone metastases in the vertebra, sternum, ilium, bilateral humeri, and bilateral femurs. Laboratory tests yielded the following results (Table I): White blood cells=4.67x10⁹/l, hemoglobin=11.3 g/dl, platelet count=79x10⁹/l [normal range (NR): 158‑348x10⁹/l], total bilirubin=0.6 mg/dl, blood urea nitrogen=15.0 mg/dl, creatinine=0.64 mg/dl, total protein=7.9 g/dl, C-reactive protein=0.14 mg/dl, PT‑INR=1.11, APTT=31.8 sec, fibrinogen=127 mg/dl, and D‑dimer=112.9 µg/ml. Her laboratory data met the diagnostic criteria (6 points) for DIC from the Japanese Society on Thrombosis and Hemostasis (5). However, there were no signs of deep vein thrombosis or pulmonary embolism, or signs of bleeding tendency such as subcutaneous bleeding or mucosal bleeding.

Re-examination of the resected left breast tumor by our hospital pathologists resulted in the diagnosis of high‑grade angiosarcoma, as the mass was mainly composed of atypically anastomosing vessels that spread invasively, and dense solid foci containing oval or spindle cells with nuclear enlargement and hyperchromasia (Fig. 2). Tumor cells were immunohistochemically positive for cluster of differentiation (CD)34, CD31, and erythroblast transformation‑specific [ETS]‑related gene (ERG). The immunohistochemistry was performed using monoclonal antibodies for CD34 (clone QBEnd/10, cat. no. 518‑102418, Roche), CD31 (clone JC70, cat. no. 518‑103231, Roche), and ERG (clone EPR3864, cat. no. 518‑110819, Roche). The immunostaining was performed using a Ventana BenchMark ULTRA IHC/ISH Staining Module®. Antigen retrieval was performed by placing tissue sections in ULTRA CC1 buffer (pH 9.0) for the optimum amount of time (32‑64 min). Subsequently, slides were incubated at 37˚C for varying amounts of time: 8 min for CD34, 32 min for CD31, and 24 min for ERG. The OptiVIEW DAB Detection Kit (Ventana Medical Systems, Inc.) was used to detect the mouse IgG, mouse IgM, and rabbit primary antibodies. Slides were then counterstained with hematoxylin. Thrombomodulin‑α (380 U/kg) was started on the day of admission and lasted for 7 days. Weekly PTX infusion (100 mg/m²) was started on the second day. Her hemostasis test improved and she was discharged 13 days after admission. However, she was hospitalized a second time 3 days after discharge because her DIC relapsed.

Table I. Summary of laboratory data.

| Parameter                        | Normal range | 1st Admission | 2nd Admission |
|----------------------------------|--------------|---------------|---------------|
| Complete blood count             |              |               |               |
| White blood cells (x10⁹/l)       | 3.30‑8.60    | 4.67          | 3.35          |
| Hemoglobin (g/dl)                | 11.6‑14.8    | 11.3          | 9.3           |
| Platelets (x10⁹/l)               | 158‑348      | 79            | 52            |
| Biochemistry                     |              |               |               |
| Aspartate aminotransferase (U/l) | 13‑30        | 17            | 23            |
| Alanine aminotransferase (U/l)   | 7‑23         | 9             | 22            |
| Lactate dehydrogenase (U/l)      | 124‑222      | 323           | 433           |
| Total bilirubin (mg/dl)          | 0.4‑1.5      | 0.6           | 0.4           |
| Blood urea nitrogen (mg/dl)      | 8.0‑20.0     | 15.0          | 12.0          |
| Creatinine (mg/dl)               | 0.47‑0.79    | 0.64          | 0.51          |
| Total protein (g/dl)             | 6.6‑8.1      | 7.9           | 6.7           |
| C‑reactive protein (mg/dl)       | <0.14        | 0.14          | 0.09          |
| Coagulation                      |              |               |               |
| PT‑INR                           | 0.85‑1.20    | 1.12          | 1.22          |
| APTT (sec)                       | 24.0‑39.0    | 31.8          | 30.9          |
| AT‑3 (%)                         | 75‑130       | 100           | 100           |
| Fibrinogen (mg/dl)               | 200‑400      | 127           | 84            |
| FDP (µg/ml)                      | <5.0         | 297.9         | No data       |
| D‑dimer (µg/ml)                  | <1.0         | 112.9         | 215.8         |

PT‑INR, prothrombin time‑international normalized ratio; APTT, activated partial thromboplastin time; AT‑3, antithrombin‑3; FDP, fibrinogen/fibrin degradation product.
of PTX. Therefore, to control DIC, danaparoid sodium (1,250 units, every 12 h) was infused for 13 days, and chemotherapy was switched from PTX to eribulin mesylate (1.4 mg/m²; infusions performed on days 1 and 8, of every 21-day cycle). After treatment with danaparoid sodium was started, her DIC had improved. For outpatient treatment, infusion of danaparoid sodium was switched to self-subcutaneous injection of UFH (5,000 units, once every 12 h). After this anticoagulant change, her DIC remained controllable, and eribulin mesylate was continued for 2 months (Fig. 3).

CT in February 2020 revealed an increase in the left breast mass, whereas lung metastatic lesions became reduced. Eribulin mesylate was stopped and chemoradiotherapy was performed, which consisted of intensity-modulated radiotherapy (total 60 Gy/30 fraction) and concurrent weekly PTX infusions (63 mg/m²), as well as self-subcutaneous injections of UFH. PTX was re-started because a previous report suggested that chemoradiotherapy with taxane and maintenance chemotherapy in cutaneous angiosarcoma were efficacious (6), but there have been no reports of eribulin mesylate with radiotherapy.

After the radiotherapy, PTX (63 mg/m²; infusions performed on days 1, 8, and 15 of every 28-day cycle) was continued as a maintenance therapy for 6 months. In November 2020, multiple liver metastases were diagnosed by follow-up CT, but her DIC remained stable, and PTX was replaced by eribulin mesylate (1.1 mg/m²; infusions performed on days 1, day 8, of every 21-day cycle) again. At the time of reporting, the patient has been continuing the self-subcutaneous injections of UFH as well as eribulin mesylate infusions. No adverse events associated with UFH use have been observed. Her PASB remained in stable disease, whereas DIC became controllable.

**Discussion**

DIC is a common complication associated with angiosarcoma and often interferes with a patient’s chemotherapy. Farid et al (4) retrospectively reviewed case records of 42 patients diagnosed with angiosarcoma at Mount Sinai Hospital, wherein 7 patients (17%) met clinical criteria for DIC. In their report, all patients who received systemic antineoplastic therapy with resultant disease response or stability had their DIC resolved in tandem with clinical improvement. DIC also recurred at the time of disease progression in all cases. Only a few case reports mention anticoagulant therapies for DIC associated with angiosarcoma. Honda et al (7) reported the first case of a patient having primary cardiac angiosarcoma with coexisting DIC who was successfully treated with nab-PTX. This patient’s DIC, treated with transfusion of fresh-frozen plasma and recombinant thrombomodulin-α (used for 6 days), improved shortly after the initiation of chemotherapy. Rosen et al (8) reported a case of hepatic angiosarcoma with a primary focus on the management of the patient’s DIC. In their report, the patient’s DIC became controlled after initial treatment with unfractionated heparin and subsequent enoxaparin treatment.

In our study, the patient’s DIC improved through chemotherapy with PTX and recombinant thrombomodulin-α. However, it re-exacerbated soon after recombinant thrombomodulin-α was completed. As a second treatment for DIC, danaparoid sodium was used and her DIC improved again. To switch from inpatient treatment to outpatient treatment, infusion of danaparoid sodium was switched to self-subcutaneous injections of UFH. This UFH resulted in long-term stabilization of DIC (>13 months) and made the continuity of chemotherapy and chemoradiotherapy possible as an outpatient treatment.

Based on the degree of fibrinolytic activation, there are diverse subtypes in DIC, such as suppressed-fibrinolytic-type, balanced-fibrinolytic-type, and enhanced-fibrinolytic-type (9). Our patient’s laboratory findings, that was measured under the treatment of UFT, showed a marked elevation in both thrombin-antithrombin complex (more than 120.0 ng/ml: NR<4 ng/ml) and plasmin-α 2 plasmin complex...
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We speculated that our patient had enhanced-fibrinolytic-type DIC. For this type of DIC, combination therapy with heparin and tranexamic acid has been reported to be very effective (9). In our case, addition of tranexamic acid was not considered because there was no bleeding symptom, and her DIC was in a stable condition at

Figure 2. Microscopic features of the angiosarcoma of the left breast resected at the previous hospital (Osaka Breast Clinic). (A) The mass was mainly composed of atypically anastomosing vessels that spread invasively (white arrow; hematoxylin and eosin staining; magnification, ×100) and dense solid foci containing oval or spindle cells with nuclear enlargement and hyperchromasia (black arrow). Tumor cells were immunohistochemically positive for (B) CD34 (magnification, ×100), (C) CD31 (magnification, ×100) and (D) ERG (magnification, ×100). ERG, erythroblast transformation-specific (ETS)-related gene.

Figure 3. Clinical course of the patient. TM, thrombomodulin-α; DP, danaparoid sodium; HP, unfractionated heparin sodium; PTX, paclitaxel; EB, eribulin mesylate; RT, radiotherapy; PASB, primary angiosarcoma of the breast; PLT, platelet count.

(8.5 µg/ml: NR 0-0.8 µg/ml). We speculated that our patient had enhanced-fibrinolytic-type DIC. For this type of DIC, combination therapy with heparin and tranexamic acid has been reported to be very effective (9). In our case, addition of tranexamic acid was not considered because there was no bleeding symptom, and her DIC was in a stable condition at
the time of UFH start. The mean survival time in metastatic disease was 6.4 months (range: 1-11 months) in 7 cases of angiosarcoma with DIC reported by Farid et al (4). Therefore, we speculate that the use of UFH helped prolong our patient's life. Mono-chemotherapy with PTX after first-line anticoagulant therapy with recombinant thrombomodulin-α could not stabilize DIC, but combination therapy of PTX and UFT after chemoradiotherapy could. So, we considered that UFH, rather than PTX, contributed to the stabilization of her DIC.

Self-subcutaneous injection of heparin is effective for chronic DIC associated with aortic aneurysm and vascular malformations, and is sometimes indicated for these patients (10,11). However, there have been few case reports where it was used to treat DIC with malignancy. Some studies on cancers other than angiosarcoma reported results similar to ours. Makari et al (12) reported a case of gastric cancer complicated by DIC. The DIC relapsed after the first line danaparoid sodium followed by self-subcutaneous injections of heparin.

In Japan, many oncologists tend to try first-line chemotherapy with anticoagulant therapy for solid tumors with DIC. However, once the patient's DIC relapses, if there is no second-line chemotherapy with a high response rate, they often decide that the tumors, as well as DIC, are beyond control and hence give up on aggressive treatments. Our case suggests that if self-subcutaneous injections of UFH are used, some patients with PASB complicated by DIC can continue to receive chemotherapies or chemoradiotherapies in the outpatient setting without impairing quality of life. Nevertheless, we would like to acknowledge a limitation in this case report. Including our case, there have been few case reports on DIC associated with malignancies treated by self-subcutaneous injection of UFH. DIC associated with more aggressive PASB may not be controllable by self-subcutaneous injections of UFH. This study is based on only one case report; therefore, further prospective investigations are needed to elucidate the clinical efficacy of self-subcutaneous injections of UFH for the treatment of PASB with DIC.

In summary, we reported the case of a woman with PASB complicated by DIC. The DIC relapsed after the first line anticoagulant therapy of thrombomodulin-α, but the second line danaparoid sodium followed by self-subcutaneous injection of UFH resulted in long-term stabilization of the DIC. Our case suggests that if self-subcutaneous injections of UFH are used, some patients diagnosed with PASB with DIC can continue to receive chemotherapies or chemoradiotherapies in the outpatient setting without having their quality of life impaired. Nevertheless, further prospective clinical trials are needed to verify the efficacy of self-subcutaneous injection of UFH in similar settings.

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Availability of data and materials
The datasets used and/or analyzed during the present study are available from the corresponding author on reasonable request.

Authors' contributions
TY conceived the present study, HN pathologically diagnosed the patient and revised the manuscript. TW, YI, HT, HS, KY, MW and ST collected clinical data. All authors read and approved the final manuscript prior to submission. TY and ST confirm the authenticity of all the raw data.

Ethics approval and consent to participate
The soft tissue mass of the patients was collected after written informed consent was obtained from the patient according to the protocol approved by the Osaka International Cancer Institute (Osaka, Japan).

Patient consent for publication
Written informed consent was obtained from the patient for the publication of patient data and associated images.

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

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