Photodynamic hyperthermal chemotherapy with indocyanine green: a novel cancer therapy for 16 cases of malignant soft tissue sarcoma

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Sixteen cases of malignant soft tissue sarcoma (STS; 10 canines and six felines) were treated with a novel triple therapy that combined photodynamic therapy, hyperthermia using indocyanine green with a broadband light source, and local chemotherapy after surgical tumor resection. This triple therapy was called photodynamic hyperthermal chemotherapy (PHCT). In all cases, the surgical margin was insufficient. In one feline case, PHCT was performed without surgical resection. PHCT was performed over an interval of 1 to 2 weeks and was repeated three to 21 times. No severe side effects, including severe skin burns, necrosis, or skin suture rupture, were observed in any of the animals. No disease recurrence was observed in seven out of 10 (70.0%) dogs and three out of six (50.0%) cats over the follow-up periods ranging from 238 to 1901 days. These results suggest that PHCT decreases the risk of STS recurrence. PHCT should therefore be considered an adjuvant therapy for treating companion animals with STS in veterinary medicine.

Keywords: cancer, chemotherapy, hyperthermia, PDT, soft tissue sarcoma

Introduction

Soft tissue sarcoma (STS) develops in a variety of mesenchymal tissues [20]. In general, local recurrence is common following conservative excision. Therefore, other therapies including radiation therapy and chemotherapy are used to prevent recurrence after performing a wide first excision [13,21,22,26,27]. Radiation therapy plays a particularly important role in the management of STS [9]. However, this type of therapy can only be performed in restricted facilities. Some patients cannot undergo radiation therapy due to financial difficulties or because the number of available facilities is limited. Therefore, it is necessary to develop novel and effective techniques to treat STS.

Indocyanine green (ICG) induces heat generation in response to light at a wavelength of 808 nm [5-7] and oxygen radicals upon exposure to light at wavelengths of 600 ∼ 800 nm [18]. Based on these evidences, we developed a novel cancer therapy using the properties of ICG and a broadband light source instead of a diode laser to establish a combination of photodynamic therapy and hyperthermia. This method is known as photodynamic hyperthermia (PHT) and can be combined with local chemotherapy to create a triple therapy strategy (photodynamic therapy, hyperthermia therapy, and chemotherapy) called photodynamic hyperthermal chemotherapy (PHCT). To date, no studies on the application of PHT and PHCT for the treatment of STS have been published. This is the first report of the use of PHCT to treat STS.

Materials and Methods

Animals

Table 1 presents a summary of the 10 canine and six feline cases. These animals were treated at Veterinary Teaching...
Table 1. Data for the clinical cases

| Species | Case number | Breed | Age | Sex | BW (kg) | Type of tumor       | Site          | Size (cm) | TNM stage |
|---------|-------------|-------|-----|-----|---------|---------------------|---------------|-----------|-----------|
| Dogs    | C01         | Mongrel | 10  | ♂   | 10.0    | Malignant schwannoma | L. forlimb    | 3.0 × 3.0 | T1N0M0    |
|         | C02         | L. retriever | 9   | ♂   | 29.0    | Malignant schwannoma | R. hindlimb   | 1.9 × 3.1 | T1N0M0    |
|         | C03         | Miniature schnauzer | 13  | ♀   | 7.9     | Malignant schwannoma | L. axilla    | 6.0 × 5.0 | T2N0M0    |
|         | C04         | G. retriever | 13  | ♂   | 22.7    | Hemangiopericytoma    | L. forlimb    | 5.0 × 8.0 | T2N0M0    |
|         | C05         | Welsh corgie   | 15  | ♂   | 11.0    | Hemangiopericytoma    | L. hindlimb   | 8.0 × 8.0 | T2N0M0    |
|         | C06         | French bulldog | 11  | ♀   | 3.0     | Liposarcoma          | L. hindlimb   | 3.0 × 3.0 | T1N0M0    |
|         | C07         | L. retriever | 7   | ♂   | 31.4    | Fibrosarcoma         | R. hindlimb   | 3.0 × 3.0 | T1N0M0    |
|         | C08         | G. retriever | 10  | ♂   | 33.0    | Undifferential soft tissue tumors | Perineum | 3.0 × 3.0 | T1N0M0    |
|         | C09         | Cocker spaniel | 10  | ♂   | 10.0    | Undifferential soft tissue tumors | R. forlimb  | 8.0 × 8.0 | T2N0M0    |
|         | C10         | Mongrel      | 8   | ♀   | 10.0    | Hemangiopericytoma    | L. forlimb    | 5.0 × 8.0 | T2N0M0    |
| Cats    | F01         | DSH         | 11  | ♂   | 6.7     | Malignant schwannoma | R. forlimb    | 2.0 × 2.0 | T1N0M0    |
|         | F02         | DSH         | 13  | ♂   | 3.5     | Fibrosarcoma         | Dorsal region | 2.7 × 3.1 | T1N0M0    |
|         | F03         | DSH         | 12  | ♂   | 3.2     | Fibrosarcoma         | R. hindlimb   | 3.0 × 4.0 | T1N0M0    |
|         | F04         | American short-hair | 4   | ♂   | 3.2     | Fibrosarcoma         | L. axilla    | 0.8 × 0.8 | T1N0M0    |
|         | F05         | DSH         | 11  | ♂   | 3.5     | Undifferential soft tissue tumors | R. forlimb  | 1.5 × 1.0 | T1N0M0    |
|         | F06         | DSH         | 7   | ♀   | 5.0     | Rhabdomyosarcoma     | R. forlimb   | 2.0 × 3.0 | T1N0M0    |

BW: body weight, TNM: T, size of the primary tumor; N, condition of the regional lymph nodes; M, absence/presence of distant metastasis.

Hospital of Tottori University (six dogs and four cats; Tottori, Japan), Takayama Pet Clinic (two dogs; Osaka, Japan), Aino Animal Hospital (one dog and two cats; Shizuoka, Japan) and Tokyo Animal Medical Center (one dog; Tokyo, Japan), respectively. The animals were diagnosed with STS based on preoperative biopsies. The ages of the animals ranged from 4 to 15 years. The dogs included two Labrador retrievers, two golden retrievers, one cocker spaniel, one Welsh corgi, one miniature schnauzer, one French bulldog, and two mongrels. The cats included five domestic shorthair (DSH) cats and one American shorthair. We measured the original tumor size by caliper before surgery. All cases were classified according to TNM (T, size of the primary tumor; N, condition of the regional lymph nodes; M, absence/presence of distant metastasis) stage [21]. Tumors in five dogs and six cats were classified as T1 (≤ 5 cm in diameter at the greatest dimension), and T2 (> 5 cm in diameter at greatest dimension) in five dogs. Among the dogs, the tumor types included three cases of malignant schwannoma, three cases of hemangiopericytoma, one case of liposarcoma, one case of fibrosarcoma, and two cases of undifferentiated soft tissue tumors. In the cats, the tumor types included three cases of fibrosarcoma, one case of malignant schwannoma, one case of rhabdomyosarcoma, and one case of an undifferentiated soft tissue tumor. The tumor sites included the trunk in four cases (one in the dorsal region, two in the axilla, and one in the perineum) and the limbs in 12 cases. No lung metastasis was observed by radiography in any case.

We explained the risk of recurrence and treatment options, including surgery, radiation, and chemotherapy, to the animal owners. When tumors had developed in the limbs, we proposed amputation as the first choice of treatment to the owners. However, the owners did not desire amputation and radiation. We then proposed to all owners other treatments including the combination of PHCT and surgery. We explained that PHCT is an experimental therapy, and all owners of the pets enrolled in this clinical trial provided informed consent.

Surgical treatment
For tumors that had developed in the limbs, we maintained sufficient skin for skin closure with skin sutures after the tumors were removed because the owners did not want us to perform reconstructive surgery such as skin grafting. As a result, we could not obtain sufficient surgical margins. We removed the underlying fascia of the tumors. In some cases, we could not perform perfect skin closure due to the presence of skin ulcers (Fig. 1A). In some cases (cases C06, C09, and F04), we removed the tumors with an ultrasonic aspiration device (Qucer; M & M, Japan). For case F02, only PHCT was performed because the owner did not wish the animal to undergo surgery.

PHCT
ICG (25 mg/vial, Giagnogreen; Daiich Sankyo, Japan) was dissolved in 9 mL of saline with an adjusted pH of 5.0. As an anti-tumor drug, 1 mL of bleomycin (1 mg/mL; Bulse;
A new adjuvant therapy for malignant soft tissue sarcoma

Fig. 1. (A) Skin incision (arrow) in case C07. (B) PHCT was performed after surgery. The arrow indicates the broadband light source while the arrowheads indicate the thermometer and thermometer sensor.

Nippon Kayaku, Japan) or carboplatin (10 mg/mL; Nippon Kayaku) was added to the ICG solution. For some cases, a small volume (0.1 or 0.2 mL) of paclitaxel (Bristol-Myers Squibb, USA.) was added to the ICG solution. For one case (case C04), no anti-tumor drugs were administered in accordance with the owner’s request. A broadband light source (Super Lizer 5000; Tokyo Iken, Japan) emitting a wavelength spectrum from 600 to 1,600 nm with a 5,000 mW maximum output power was used because ICG responses to light at wavelength of 600 ∼ 800 nm.

For each case, the tumors were resected and ICG solution was injected into the resected area 3-dimensionally, including the skin surgical margin (3 cm). One mL of the ICG solution was administered per cm² of the wound bed. Irradiation was administered at a distance of 10 cm from the resected area (irradiation area: 113 cm², 40 mW/cm²) for 20 min per 113 cm² (48 J/cm²; Fig. 1B) immediately after the ICG solution was injected. The temperature at the surface of the resected area was kept under 45°C by moving light source near and away from the skin surface and monitored with a thermometer. The first round of PHCT was performed immediately after skin suturing following surgery. The treatment interval between the second and fourth round of PHCT was generally 1 week, and then the treatment was performed at intervals of 2 to 4 weeks. At a minimum, treatment was continued for 3 months after surgery. At that point, we continued the treatment if the owner desired. For the second and subsequent rounds of PHCT, the treatments were performed with all animals under sedation. In some cases, local anesthesia was induced by Lidocaine of 15 ∼ 50 mg/head (Xylocaine; AstraZeneca, Japan).

Follow-up after PHCT

In all cases, follow-up examinations for STS recurrence and metastasis were performed at intervals of several months for 1 year after the first round of PHCT. Thereafter, follow-up examinations were performed once a year for 5 years.

Results

Table 2 presents a summary of the study results. PHCT was performed three to 21 times. The treatment frequency depended on the wishes of the owner. No severe side effects, including severe skin burns, necrosis, or rupture of skin sutures, were observed in any of the animals although skin redness and minor skin burns occurred. The overall canine survival time (ST) except for 1 case of amputation (case C06) ranged from 225 to 1,901 days (median survival time: 767 days). In seven out of 10 dogs (70.0%), no recurrence was observed during the follow-up periods ranging from 238 to 1,901 days. In five of these seven dogs, no recurrence was observed after more than 2 years. The remaining three dogs (two with undifferentiated soft tissue sarcomas and one with liposarcoma) experienced recurrence or metastasis over intervals ranging from 72 to 162 days after surgery. One case with liposarcoma (case C06) experienced local recurrence occurring within 1 month after the first surgery. In this animal, local recurrence was also observed 72 days after the first round of PHCT following the second surgery. The disease-free time (DFT) was prolonged by the use of PHCT. However, amputation was performed in accordance with the owner’s desire, and the dog is currently alive. One dog (case C08) of the remaining three dogs died due to tumor progression. In case 08, metastasis to the sublumbar lymph nodes was
Table 2. Summary of treatment outcomes

| Species | Case number | Treatment time | Anti-cancer drug | DFT | ST | Update   | Comments                                                                 |
|---------|-------------|----------------|------------------|-----|----|----------|--------------------------------------------------------------------------|
| Dogs    | C01         | 7              | CA               | 1041| 1041| Alive    |                                                                          |
|         | C02         | 3              | CA               | 1185| 1185| Alive    |                                                                          |
|         | C03         | 12             | BL               | 238 | 238 | Alive    |                                                                          |
|         | C04         | 8              | No use           | 1774| 1774| Alive    |                                                                          |
|         | C05         | 8              | CA, PA           | 767 | 767 | Alive    |                                                                          |
|         | C06         | 8              | CA, PA           | 72  | 453 | Alive    | Tumor removal using ultrasonic device. Underwent amputation               |
|         | C07         | 4              | CA               | 1901| 1901| Alive    | Died due to tumor progression.                                            |
|         | C08         | 11             | CA               | 151 | 269 | Deceased | Tumor removal using ultrasonic device. Died due to cardiac failure.       |
|         | C09         | 5              | BL               | 162 | 225 | Deceased |                                                                          |
| Cats    | F01         | 5              | CA, PA           | 1344| 1344| Alive    |                                                                          |
|         | F02         | 17             | BL, CA           | 1521| 1521| Alive    | No surgery                                                               |
|         | F03         | 13             | BL               | 1515| 1515| Alive    |                                                                          |
|         | F04         | 21             | CA               | 82  | 1254| Alive    | Tumor removal using ultrasonic device. Underwent amputation.              |
|         | F05         | 15             | CA, PA           | 175 | 383 | Deceased | Died due to tumor progression.                                            |
|         | F06         | 6              | BL, CA           | 20  | 1879| Deceased | Underwent amputation.                                                     |

DFT: disease-free time, ST: survival time, CA: carboplatin, BL: bleomycin, PA: paclitaxel.

observed. In that animal, no local recurrence was found. In case C09, local recurrence was observed 162 days after the first round of PHCT. PHCT was repeated for this animal five times at intervals of 1 to 2 weeks. Thereafter, the dog was only followed up. Sixty-seven days after the fifth round of PHCT, local recurrence was observed. The dog eventually died of cardiac failure 225 days after the first round of PHCT.

The overall feline ST, excluding two cases of amputation (cases F04 and F06), ranged from 383 to 1,521 days (median survival time: 1,344 days). In three out of the six cats (50.0%), no recurrence was observed over the follow-up periods ranging from 1,344 to 1,521 days. The remaining three cats (one case of fibrosarcoma, one case of rhabdomyosarcoma, and one case of undifferentiated STS) experienced recurrence over intervals ranging from 20 to 175 days after surgery. In two (cases F04 and F06) of the remaining three cases, amputation was performed on one cat (case F04) that is currently alive and one cat (case F06) that died due to unknown cause. The remaining cat died due to tumor progression. In case F02 for which no surgery was performed, we periodically assessed tumor status using fine needle aspiration (FNA). No living tumor cells were observed by FNA during the eighth round of PHCT. We diagnosed the tumor as degenerative following PHCT, and the degenerative tumor tissue was removed using an ultrasonic aspiration device (Sono Cure; Tokyo Iken, Japan). Thereafter, PHCT was performed for 3 months. This cat had no recurrence for about 5 years.

Discussion

Malignant STSs rarely metastasize; however, they are locally invasive [3]. If the surgical margin is insufficient, the rate of recurrence is 10 times of that when the surgical margin is sufficient [20]. Therefore, complete resection with a sufficient surgical margin is necessary. In general, adjuvant therapies including radiation therapy and chemotherapy are administered to cases of insufficient surgical margins [13,21,22,27]. In particular, radiation therapy plays an important role in the management of STS. In the present study, the surgical margins were insufficient in all cases. However, 70.0% (7/10) of the dogs and 50.0% (3/6) of the cats did not experience recurrence over follow-up periods ranging from 238 to 1,901 days. These results suggest that PHCT decreases the risk of recurrence.

As an alternative to radiation therapy, we developed a new therapy, PHT, by combining Photodynamic therapy (PDT) and hyperthermia with ICG using a broadband light source. Furthermore, we administered PHT in combination with local chemotherapy and designated this technique “PHCT”. No previous studies have demonstrated the efficacy of the combination of ICG and use of a broadband light source in
cancer therapy.

It has been suggested that fibrosarcomas, malignant schwannomas, and hemangiopericytomas are sensitive to PHCT considering the relationship between tumor type and PHCT efficacy. The local control rate of fibrosarcoma, malignant schwannoma, and hemangiopericytoma was 100% in the present study. However, PHCT is not effective for other tumor types including liposarcoma, rhabdomyosarcoma, and undifferentiated STSs occurring in the limbs. In these cases, the tumors recurred ranging from 20 to 175 days after the first PHCT. These results suggest that the effects of PHCT might differ depending on tumor type. Further investigation into this possibility is necessary. In the present study, no relationship between treatment frequency and recurrence was observed. In one case (case C09), local recurrence was observed 63 days after the final round of PHCT. This phenomenon suggests that it is important to periodically continue treatment with PHCT. Additionally, no relationship was observed between the use of anti-cancer drugs and outcomes.

In two out of three canine and two out of three feline recurrence cases, the tumor cells had infiltrated deeply. We speculate that the light used for PHCT did not reach the tumor cells located in deep tissues. In a preliminary experiment, we found that the temperature at a depth of 2 cm from the skin surface did not reach 40°C (data not shown), indicating that PHCT is not effective for treating tissues beyond this depth. Additionally, the ICG solution was not equally distributed. These drawbacks will be investigated in the future.

Hyperthermia is not a common treatment in veterinary medicine because the required device is expensive and few fundamental data related to the efficacy of hyperthermia are available. On the other hand, there are reports of PDT being used to treat spontaneous tumors in canines and felines, and the efficiency of this technique has been recognized [11,28-30,33-35]. However, the photosensitizer for PDT is very expensive and a special diode laser is needed. Therefore, PDT is also rarely administered by veterinarians. ICG was developed as a drug to promote liver and bile duct function, and it has been used medically since 1956. This drug is safe for both human and animals [8]. In particular, ICG is widely used in ophthalmology for hyperthermia therapy with an 808-nm diode laser to treat chorioretinopathy [12]. ICG is also used for sentinel biopsies [17,19].

Some reports have suggested that ICG induces the formation of oxygen radicals [1,2,4,10,23]. Most of these findings are from in vitro studies of different human cell lines. Bozkulak et al. [4] reported that ICG with near-infrared light (809 nm, 60 mW/cm², 24 J/cm²) is very effective for eliminating human breast cancer cells. In this study, the temperature of the medium was controlled to ensure that it remained at 37°C. The authors concluded that ICG acts as a new photosensitizer. Hirano et al. [18] was the first to demonstrate that ICG induces the formation of oxygen radicals in response to irradiation with light at a wavelength of 600 – 800 nm. The concentration of oxygen radicals produced by ICG dissolved in ethanol is similar to that generated by other photosensitizers. Oxygen radicals are formed even if ICG is dissolved in water although the concentration is lower. The total volume of oxygen radicals produced is thought to result from the accumulation of oxygen radicals produced with each wavelength. Therefore, a broadband light emitting light over a range of 600 – 800 nm would induce the formation of more oxygen radicals than a diode laser. Additionally, a broadband light that has an emission range of 600 – 800 nm would stimulate ICG to simultaneously induce heat and oxygen radical production in tumor tissues.

We previously investigated the effects of PHT on B16F10 murine melanoma cells in vitro [31,32]. Our results demonstrated that PHT induces early morphological changes in tumor cells that promote more cell death than hyperthermia alone. Furthermore, PHT induces early apoptosis and cell arrest. These in vitro data support the present in vivo results.

Tumor cells are more sensitive to heat under acidic conditions [14]. Therefore, saline with a pH adjusted to 5.0 with acetic acid was used as a solvent for ICG. In a preliminary experiment, we found that saline with a pH of 4.0 induced tissue inflammation (data not shown).

The effects of anti-tumor drugs are enhanced by heat [15,16,24,25]. Furthermore, tumor tissues have a tendency to be acidic [36]. Therefore, we selected bleomycin as an anti-tumor drug because it is effective in acidic environments [15]. Platinum-based drugs such as cisplatin and carboplatin are also useful anti-tumor reagents. In the present study, we were concerned that PHCT would affect the stability of the anti-tumor drug. However, our preliminary data showed that the anti-tumor effects of PHCT were enhanced by the addition of an anti-tumor drug (data not shown). This means that PHCT does not only affect the stability of the anti-tumor drug but also enhances its effects.

In conclusion, PHCT is a simple procedure, is not associated with any severe side effects, and requires no special facilities. However, further investigation is necessary to establish PHCT as a therapeutic technique due to some associated problems involving treatment times and intervals, and selection of anti-cancer drugs. Nevertheless, this modality is expected to become a useful alternative to radiation therapy for treating superficial tumors such as STS in companion animals.

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