Photohyperthermal therapy using liposomally formulated indocyanine green for feline nasal lymphoma: A case report

MASAMICHI YAMASHITA1, MAYUMI MAYAMA2, AKIKO SUGANAMI3, KAZUO AZUMA1, TAKESHI TSUKA1, NORIHIKO ITO1, TOMOHIRO IMAGAWA1, YUTAKA TAMURA3 and YOSHIHARU OKAMOTO1

1Joint Department of Veterinary Clinical Medicine, Faculty of Agriculture, Tottori University, Tottori 680-8550; 2Mayama Animal Hospital, Hokkaido 068-0028; 3Department of Bioinformatics, Graduate School of Medicine, Chiba University, Chiba 260-8670, Japan

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Abstract. Our previous research has focused on the development of a novel cancer therapy by using photohyperthermal therapy (PHT) with indocyanine green (ICG) as an optical sensitizer. ICG-Lipo is a liposomally formulated ICG derivative in which ICG is tagged with an octadeca-alkyl chain to incorporate into liposome bilayers, and contains antitumor drugs such as carboplatin and paclitaxel within the inner membrane space. The present study reported a case of feline nasal lymphoma that was treated with combination therapy of PHT with ICG-Lipo. An antitumour effect was observed, and the patient entered remission. Complications from the radiation treatment included skin burns and bleeding from the irradiated hard palate. Serious side effects related to the drugs were not observed. This report suggested that PHT using ICG-Lipo enabled efficient and safe treatment of lymphoma, and that treatment with a liposomal drug delivery system was enhanced by PHT.

Introduction

Feline nasal lymphoma is the most common nasal tumour diagnosed in cats (1,2). It is a relatively treatable tumour, with remission rates of 65-75% observed when various chemotherapy protocols are used (3). The median survival time (MST) of feline nasal lymphoma is 473 days with combined radiation therapy and chemotherapy, and 320 days with chemotherapy alone (4). Chemotherapy and radiation therapy are effective for the treatment of feline nasal lymphoma, and surgery is not required (4). Despite the effectiveness in treating feline nasal lymphoma, chemotherapy and radiation have significant limitations; both have severe side effects, including bone marrow suppression and skin ulcers (5-8). Moreover, treatment is restricted by a repeated requirement for anaesthesia and high cost. The development of less burdensome and low-cost treatments would be beneficial in clinical settings.

Photohyperthermal therapy (PHT), combining photodynamic therapy (PDT) and photothermal therapy, has been investigated as safe and low-cost treatments for cancer (9-11). PHT is used in combination with near-infrared light (NIR) and a photosensitizer, such as indocyanine green (ICG) or aminolevulinic acid. The resulting activated oxygen has an anti-tumour effect (10,11). ICG has a peak spectral absorption of ~780 nm and peak fluorescence emission at ~820 nm. It induces heat and singlet oxygen formation in response to NIR light with a wavelength of 800 nm and is characterized by low general toxicity (10). The principal disadvantage of ICG is its rapid clearance from the body (plasmatic half-life of 2-4 min), limiting its accumulation within tumours (12). Several studies have reported the use of liposomally formulated ICG for optical imaging and cancer therapy, which improves its tumour-accumulating ability and stability (12-14). Liposomal drug delivery systems have enhanced permeability retention effects, which can increase the stability and accumulation within tumours of ICG (15,16). Suganami et al (13), designed and synthesised a novel NIR photoactivating probe, which is more hydrophobic compared with conventional ICG to promote liposome formation. Toyota et al (17), reported that a liposomally formulated ICG derivative (ICG-Lipo) yielded strong fluorescence images under an NIR-fluorescence imaging system. PDT using ICG-Lipo was reported to induce antitumour effects in vitro and in vivo (18,19). The present study describes a case of feline lymphoma that was treated with the combination therapy of PHT with ICG-Lipo.

Case report

A 10-year-old male cat (weight, 4.1 kg) presented with primary symptoms of sneezing, and nasal mucus and conjunctival injection in the right eye. The cat was initially diagnosed with an upper respiratory infection and prescribed an antibiotic. All the symptoms disappeared, with the exception of conjunctival hyperaemia. After 1 month, sneezing, nasal haemorrhage, protrusion of the right eye and facial swelling
were observed (Fig. 1A). Gingivitis was observed around the right upper premolar, which led to the diagnosis of a root abscess. Tooth extraction was performed with a routine course of post-operative antibiotics. However, the symptoms did not improve, and the facial swelling deteriorated. Cancer was suspected, and the owner was advised to seek treatment at a secondary hospital. The owner selected to have the cat treated at the primary hospital, where the combination therapy of PHT with ICG-Lipo was performed. This procedure was approved by the Organization for Research Initiative and Promotion in Tottori University (ethical approval no. H28-007).

ICG-Lipo, comprised of 2.25 mg ICG-C18 (ICG derivative in which ICG is tagged with an octadeca-alkyl chain), 10 mg carboplatin (Nichi-Iko Pharmaceutical Co., Ltd) and 0.6 mg paclitaxel (Bristol-Myers Squibb), was diluted with 50 ml PBS (pH 7.4) at room temperature and administered intravenously at a rate of 50 ml/h. Light irradiation was performed using a basic semiconductor laser (DVL-15; Asuka Medical, Inc.) five times a week for 2 weeks (Fig. 2). The total dose of light that the cat was subjected to per irradiation was 6,000 J; light was irradiated at a dose rate of 5-10 W. Adjuvant therapy for the tumour was not administered. Fluid infusion was administered only when the cat exhibited a poor appetite. Five courses of combination therapy of PHT with ICG-Lipo were administered at 0, 25, 43, 60 and 74 days. Progressive improvement and deterioration of symptoms was observed between the first and fourth course of treatment. After the fifth course, facial swelling and nasal congestion showed dramatic improvement (Fig. 1B). Remission was confirmed and combination therapy was completed at the end of the fifth course. The tumour remained in remission for ~3 months. It is proposed that continuous treatment is required for long-term remission.

**Discussion**

The survival time (401 days) in the present case was longer than the MST (320 days) observed with chemotherapy alone (4). The increased survival time compared with the MST suggested that the combination therapy of PHT with ICG-Lipo is potentially effective for treating feline nasal lymphoma. The effects of the combination therapy were not observed until 2 months after the commencement of treatment, at the fifth course, when there was a dramatic decrease in the tumour. The effect of the treatment continued, and the remission phase lasted for ~3 months. It is proposed that continuous treatment is required for long-term remission.
It was recently reported that the antitumour effect of the PHT/ICG-Lipo combination therapy was associated with an immunological mechanism (20). According to a report into using immunotherapy with programmed death (PD)-ligand 1 to treat a tumour, the tumour size temporarily increased and subsequently decreased (21). The tumour volume may have increased due to an inflammatory response. Alternatively, there may be a time-lag between the initiation of treatment and observation of the effect. In the present case, improvement and deterioration of the symptoms were observed throughout the course of treatment. The patient's response may have been a result of an increased immune response induced by PHT with ICG-Lipo.

However, significant treatment effects were not observed after relapse. The tumour cells may have become drug-resistant. If the antitumour effect was related to an immunological mechanism, the tumour may have expressed immune checkpoint molecules (such as PD-1 or cytotoxic T-lymphocyte associated protein 4). However, a biopsy could not be performed, thus it was not possible to compare molecular expressions between the primary and recurrent tumour. The immune effects of PHT with ICG-Lipo may be demonstrated via immunohistochemical examination for immune checkpoint molecules.

This is the first report of PHT with ICG-Lipo containing an antitumour drug. In this case, ICG-Lipo contained carboplatin and paclitaxel, and their dose was ~10% of the standard administration protocol (22-24). Notable therapeutic effects were observed without the usual side effects of antitumour drugs. The encapsulation capacities of carboplatin and paclitaxel in liposomal pharmaceuticals were reported as 12% and 28%, respectively (25). Antitumour effects were demonstrated in a previous study (25); however, the compound ratio of ICG-Lipo was not validated in this study. It is proposed that the compound ratio is similar to that utilised in the previous report (25), thus an antitumour effect was predicted.

In human medicine, nasal lymphoma is known as extranodal natural killer/T-cell lymphoma, nasal type (26). The lymphoma originates from either natural killer cells or γδ T-cells, both of which express CD56 (26). Conversely, a B-cell phenotype comprises 40% of feline nasal lymphomas, and a T-cell phenotype comprises 47% (27). Moreover, the pathological features of human and feline nasal lymphomas are different. Thus, the findings from the present study may not contribute directly to human medicine. However, the treatment strategies for feline lymphoma are relatively similar to that for human lymphoma. It is proposed that PHT with ICG-Lipo has the potential to be effective in treating human lymphomas.

In conclusion, the present case report described the first treatment of nasal feline lymphoma using combination therapy of PHT with ICG-Lipo. The present case showed a satisfactory outcome compared with chemotherapy alone. PHT with ICG-Lipo is easy to perform and the side effects are less severe. However, a single case is insufficient to prove the exact treatment effects of PHT with ICG-Lipo. A clinical trial with a large study population is required to prove the efficacy of PHT with ICG-Lipo for treating nasal feline lymphoma.

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Availability of data and materials
All data generated or analyzed during this study are included in this published article.

Authors' contributions
MY was involved in designing the treatment protocol of PHT with ICG-Lipo, evaluating the treatment effect and preparing the manuscript draft. MM oversaw all aspects of treatment and drafted a report on the case. AS and YT synthesised ICG-Lipo. KA and TT were involved in designing the fluid therapy and reviewing the manuscript. NI and TI were involved in analysing haematological data and reviewing the manuscript. YT and YO designed the treatment plan, and reviewed and edited the manuscript. All authors read and approved the final manuscript.

Ethics approval and consent to participate
The treatment protocol was approved by the Organization for Research Initiative and Promotion in Tottori University (ethical approval no. H28-007).

Patient consent for publication
The owner of the pet that was the subject of this case report provided written consent for the publication of the case report.

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

References
1. Henderson SM, Bradley K, Day MJ, Tasker S, Caney SM, Hotston Moore A and Gruffydd-Jones TJ: Investigation of nasal disease in the cat—a retrospective study of 77 cases. J Feline Med Surg 6: 245-257, 2004.
2. Mukaratirwa S, van der Linde-Sipman JS and Gruys E: Feline nasal and paranasal sinus tumours: Clinicopathological study, histomorphological description and diagnostic immunohistochemistry of 123 cases. J Feline Med Surg 3: 235-245, 2001.
3. Couto CG: Chapter 80: Lymphoma in the cat and dog. In: Nelson RW and Couto CG (eds.). Small animal internal medicine, 4th edition. St. Louis, Elsevier, pp1174-1186, 2009.
4. Haney SM, Beaver L, Turrel J, Clifford CA, Klein MK, Crawford S, Poulson JM and Azuma C: Survival analysis of 97 cats with nasal lymphoma: A multi-institutional retrospective study (1986-2006). J Vet Intern Med 23: 287-294, 2009.
5. Pinard CL, Mutsaers AJ, Mayer MN and Woods JP: Retrospective study and review of ocular radiation side effects following external-beam Cobalt-60 radiation therapy in 37 dogs and 12 cats. Can Vet J 53: 1301-1307, 2012.
6. Knapp DW, Richardson RC, Bonney PL and Hahn K: Cisplatin therapy in 41 dogs with malignant tumors. J Vet Intern Med 2: 41-46, 1988.
7. Machado MC, da Costa-Neto JM, Portela RD, D'Assis MJMH, Martins-Filho OA, Barrouín-Melo SM, Borges NF, Silva FL and Estrela-Lima A: The effect of nalbuphine as a carboplatin chemotherapy-associated drug on the immune response, quality of life and survival of dogs with mammary carcinoma. PLoS One 13: e0204830, 2018.

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8. Oun R, Moussa YE and Wheate NJ: The side effects of platinum-based chemotherapy drugs: A review for chemists. Dalton Trans 47: 6645-6653, 2018.

9. O’Reilly S, Rowinsky E, Slichenmyer W, Donehower RC, Forastiere A, Ettinger D, Chen TL, Sartorius S, Bowling K, Smith J, et al: Phase I and pharmacologic studies of topotecan in patients with impaired hepatic function. J Natl Cancer Inst 88: 817-824, 1996.

10. Radzi R, Osaki T, Tsuka T, Imagawa T, Minami S, Nakayama Y and Okamoto Y: Photodynamic hyperthermal therapy with indocyanine green (ICG) induces apoptosis and cell cycle arrest in B16F10 murine melanoma cells. J Vet Med Sci 74: 545-551, 2012.

11. Urbanska K, Romanowska-Dixon B, Matuszak Z, Oszajca J, Nowak-Sliwinska P and Stochel G: Indocyanine green as a prospective sensitizer for photodynamic therapy of melanomas. Acta Biochim Pol 49: 387-391, 2002.

12. Porcu EP, Salis A, Gavini E, Rassu G, Maestri M and Giunchedi P: Indocyanine green delivery systems for tumour detection and treatments. Biotechnol Adv 34: 768-789, 2016.

13. Suganami A, Toyota T, Okazaki S, Saito K, Miyamoto K, Akutsu Y, Kawahira H, Aoki A, Muraki Y, Madono T, et al: Preparation and characterization of phospholipid-conjugated indocyanine green as a near-infrared probe. Bioorg Med Chem Lett 22: 7481-7485, 2012.

14. Xue X, Fang T, Yin L, Jiang J, He Y, Dai Y and Wang D: Multistage delivery of CDs-DOX/ICG-loaded liposome for highly penetrating and effective chemo-photothermal combination therapy. Drug Deliv 25: 1826-1839, 2018.

15. Fang J, Nakamura H and Maeda H: The EPR effect: Unique features of tumor blood vessels for drug delivery, factors involved, and limitations and augmentation of the effect. Adv Drug Deliv Rev 63: 136-151, 2011.

16. Matsumura Y and Maeda H: A new concept for macromolecular therapeutics in cancer chemotherapy: Mechanism of tumor-tropic accumulation of proteins and the antitumor agent smancs. Cancer Res 46: 6387-6392, 1986.

17. Toyota T, Fujito H, Suganami A, Ouchi T, Ooishi A, Aoki A, Onoue K, Muraki Y, Madono T, Fujinami M, et al: Near-infrared-fluorescence imaging of lymph nodes by using liposomally formulated indocyanine green derivatives. Bioorg Med Chem 32: 721-727, 2014.

18. Suganami A, Iwadate Y, Shibata S, Yamashita M, Tanaka T, Shinozaki N, Aoki I, Saeki N, Shirasawa H, Okamoto Y and Tamura Y: Liposomally formulated phospholipid-conjugated indocyanine green for intra-operative brain tumor detection and resection. Int J Pharm 496: 401-406, 2015.