PALLIATIVE JAW-SPARING TREATMENT OF A NON-RESECTABLE CANINE ORAL FIBROSARCOMA USING COMBINATION OF ELECTROCHEMOTHERAPY WITH BLEOMYCIN AND IL-12 GENE ELECTROTRANSFER

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Abstract: A 15-year-old male castrated English setter was presented for evaluation of a rapidly growing oral mass. Patient’s history was otherwise unremarkable, except of moderate proteinuria of 5 years duration. Clinical examination findings were within normal limits, except of an ulcerated lesion located at the left mandibular canine tooth, which was histologically confirmed as a high grade infiltrative fibrosarcoma with high mitotic index (61/10 HPF) and multifocal necrotic areas. The client declined full staging, so only hematological and biochemistry examinations of blood were performed, which were within normal limits. Furthermore a fine needle aspiration biopsy of regional lymph nodes was performed, which revealed reactive lymphadenopathy without signs of metastases. After declining other more invasive therapeutic procedures, the clients elected treatment with combination of electrochemotherapy and IL-12 electrogene therapy. Four consecutive treatment sessions were performed, resulting not only in complete response of the primary tumor, but also in regression of untreated distant metastases, which were diagnosed approximately one month after the initial examination. Furthermore, the percentage of circulating CD8+ cells was increased after each therapy session, indicating possible systemic induction of immune response by IL-12 gene therapy. This case shows that this type of therapy can represent an alternative type of both local and systemic treatment in selected tumor cases, where clients seek a less invasive nonsurgical treatment.

Key words: dog; fibrosarcoma; electroporation; electrochemotherapy; electrogene therapy; interleukin-12

Introduction

Malignant neoplasms of the oral cavity represent approximately 6% of all canine tumors, with fibrosarcoma (FSA) being one of the three most common malignant oral tumors in dogs (1). The most common site of presentation is gingiva, followed by lip, cheek and tongue (1). FSAs are reported to be primarily locally aggressive with a low potential for distant metastases. Gingival tumors can invade the bone and spread to the palate. Early diagnosed tumors (i.e., smaller tumors) are mostly treated by curative-intent surgery (2), which can be combined with radiotherapy in selected cases. However, oral tumors often develop unnoticed until the tumor reaches an advanced stage, when it is often inoperable without causing significant dysfunction and cosmetic changes to the animal. In such cases, different palliative therapeutic approaches may be used, depending on the sensitivity of the tumor. Outcome of any such therapy is usually short-lived, with reportedly high recurrence rates, even over 50% and short survival times (3). Wide margin surgical excision of oral FSA as a sole therapy results in mean survival time (MST) of 12-24 months (3, 4).

Electrochemotherapy (ECT) and gene electrotransfer are techniques, where electric pulses are
used to increase cell membrane permeability to enhance entry of the cytotoxic agents (i.e., electrochemotherapy, ECT) and plasmid DNA (i.e., gene electrotransfer or electrogene therapy, EGT) into the cells. ECT combined with EGT with plasmid encoding interleukin-12 (IL-12) has already been described for the treatment of a small number of naturally occurring spontaneous canine oral neoplasms with encouraging results (5, 6).

Important feature of ECT and EGT with plasmid encoding IL-12 is the elicitation of the immune response, which is believed to have systemic effects. IL-12 stimulates cytotoxic T lymphocytes, which are crucial in cancer elimination (7) as they target the tumor cells and eliminate them by the action of perforin and granzyme B (8). Perforin is involved in formation of pores on the membrane of tumor cells, thus enabling the granzyme B to enter the tumor cells where it causes apoptotic tumor cell death (9). Such induction of immune response can be detected using different techniques, including flow cytometry, which utilizes gating different types of peripheral blood mononuclear cells using specific antibodies.

The purpose of this case report is to describe a case of a large grade II mandibular gingival FSA in a 15-year-old dog successfully treated with ECT and EGT with plasmid encoding IL-12.

Case presentation

A 15-year-old male castrated English setter, weighing 22 kg, was presented to the Small animal clinic of Veterinary faculty Ljubljana for evaluation of a rapidly growing oral mass. History of the patient was unremarkable, except persistent moderate proteinuria of 5 years duration, which has been well controlled with appropriate diet and enalapril. Clinical examination findings were within normal limits, apart the large mass at the left mandibular canine tooth. The client declined full staging including head CT scan and diagnostic imaging for possible distant metastases. Therefore only partial staging was performed, including complete blood count with white blood cell count (performed using an automated laser hematology analyzer with species-specific software Advia 120, Siemens, Munich, Germany) and detailed biochemistry panel (performed using automated chemistry analyzer RX-Daytona, Randox, Crumlin, UK). Biochemistry panel consisted of serum concentrations of glucose, urea, creatinine, Na, K, Cl, Ca, total proteins, albumin and serum activity of alkaline phosphatase, alanine aminotransferase and creatine kinase. All parameters were within normal limits except moderate, clinically

Figure 1: The figure is showing regression of the tumor mass at different time points. (A) tumor mass at the time of the first therapy. (B) clinical regression of the tumor at 4 weeks after the first therapy and before the second therapy. (C) four months after the first of the tumor therapy the size of the tumor significantly decreased. (D) seven months after initial therapy complete regression could be seen.
irrelevant thrombocytosis (platelets 575 x 10^-9/L, reference value 143-400 x 10^-9/L).

A detailed oral examination and dental radiographs with the dog under general anesthesia were performed, revealing a 4 cm x 3 cm x 3 cm partly ulcerated proliferative mass at the left mandibular canine tooth (Figure 1). Geographic bone loss at the left mandibular incisor and canine teeth was visible on dental radiographs, and permeative pattern of bone loss in the symphyseal region suggested bony involvement of both rostral mandibles (T3b) (Figure 2). At the same time incisional biopsy was performed, and the mass was histologically confirmed as a high grade infiltrative FSA with high mitotic index (61/10 HPF) and multifocal necrotic areas. Fine needle aspiration biopsy of regional lymph nodes was performed, revealing reactive lymph nodes.

After discussing possible treatment options, including bilateral rostral mandibulectomy in combination with radiotherapy, the client elected ECT and EGT. During the next four months, four therapy sessions were performed (Graph 1). Each session was performed with the dog under short (approximately 30 min) general anesthesia, starting with ECT using intravenous application of bleomycin (Blenoxane, Bristol-Myers, Princeton, USA; 3 mg/ml) at the dose 0.3 mg/kg, followed by delivery of electric pulses with electric pulses generator Cliniporator™ (IGEA s.r.l., Carpi, Italy). Train of 8 pulses was applied, each pulse of 100 µs duration and amplitude to electric distance ratio of 1300 V/cm and frequency of repetition 1 Hz, using two parallel stainless steel plate electrodes with 6 mm distance between them. This procedure was followed by combination radiotherapy, starting with ECT using intravenous application of bleomycin and radiotherapy.

Figure 2: (A) Dental radiograph, occlusal view of the rostral mandibles at initial presentation. Geographic bone loss at the left mandibular incisor and canine teeth was visible on dental radiographs, and permeative pattern of bone loss in the symphyseal region suggested bony involvement of both rostral mandibles (T3b) (Figure 2). Clinical assessment of radiographically abnormal (fractures or abrasion) left mandibular incisor teeth was impossible as they were embedded in the tumor mass. Right mandibular third incisor tooth is missing. (B) Dental radiograph, occlusal view of the rostral mandibles at 4 weeks. Radiographically visible severe progression of the osteolysis. Left mandibular second and third incisor teeth had exfoliated since the last visit. There is a total loss of attachment at the left mandibular first incisor tooth and near total loss of attachment at the left mandibular canine tooth, therefore these two teeth were removed (C). The round poorly mineralized structure remained, as it was hidden in the soft tissues of the tumor. (D) Dental radiograph, occlusal view of the rostral mandibles at 10 weeks. Radiographically subjectively decreased osteolysis progression.
by EGT with peritumoral application of 2 mg of plasmid encoding canine IL-12 (pCMVcaIL-12) into the mucosa. The same generator of electric pulses was used to deliver electric pulses for gene delivery. Two sets of pulses were used, one high- and four low-voltage. High-voltage pulses consisted of 100 µs duration and amplitude to electric distance ratio of 600 V/cm and low-voltage pulses consisted of 100 ms and amplitude to distance ratio of 200 V/cm. Plate electrodes with 6 mm distance between them were used.

Tumor regression was observed 4 weeks after the first therapy (Figure 1, Graph 1), although dental radiographs revealed progression of the osteolysis, requiring removal of the left mandibular incisor and canine teeth. After the second session, subjective decrease in osteolysis progression was also noted on dental radiographs. However, one month after the first therapy a subcutaneous metastases in the intermandibular region was diagnosed by cytopathologic examination of fine needle aspirate. In the course of the next three months tumor burden steadily decreased (Graph 1) and re-check at four months after the initial therapy revealed only ongoing mild-moderate necrosis of the treated area. Client reported minimal side effects and an improved quality of life of the dog. At this time point complete remission of both intraoral tumor as well as subcutaneous metastases was reached. The dog was euthanized 8 months after the initial combined therapy due to causes unrelated to the oral tumor and at that time point the dog was without any macroscopic evidence of either oral tumor or metastases in surrounding tissue. Necropsy was declined by the client.

At different time points following each therapy (1, 2, 4 weeks and then monthly after remission was achieved), detailed bloodwork was performed, including the same hematological and biochemical parameters as at the initial staging. In addition, flow cytometry was performed on frozen whole blood collected at each visit. The lymphocytes were gated for CD45 (leukocytes), CD3 (lymphocytes T), CD4 (helper T lymphocytes) and CD8 (cytotoxic T lymphocytes) markers and the percentage of each cell population was calculated. The percentage of CD8+ cells temporarily increased after each therapy and decreased in the following weeks (Graph 1). We did not observe any change in the percentage of other cell populations. Furthermore, we used quantitative polymerase chain reaction (qPCR) with specific primers for the pCMVcaIL-12 plasmid to detect the presence of plasmid DNA in the urine, stool and oral mucosa swab samples in order to determine possible shedding of plasmid into the environment. Urine and stool samples were collected one week after each therapy and no plasmid DNA was detected in these samples. Oral mucosa swabs from the area of plasmid injection were collected immediately after the therapy and at different time points thereafter (1, 2 and 4 weeks). In the first sample maximal concentration of IL-12 plasmid detected at the site of injection was 100 ng/mL of plasmid DNA. After one week, the concentration already dropped to 0.5 pg/mL and at later time points no quantity of plasmid could be detected.

Graph 1: Percentage of CD8+ cells (●) detected by flow cytometry and tumor growth rate normalised on tumor size at first visit. (▲) The percentage of CD8+ cells temporarily increased after each therapy and decreased in the following weeks after EGT therapy.

Discussion

In this case report we describe effective use of ECT with bleomycin and EGT with canine IL-12 in the treatment of a canine oral FSA with subcutaneous metastases, resulting in complete regression of primary treated tumor as well as untreated distant metastases, which enabled marked prolongation and improved quality of the dog’s life. Oral FSA in dogs are usually fast growing neoplasms, prone to aggressive infiltrative growth. Therefore, the treatment of choice is surgical excision. However, in cases of advanced oral tumors, wide excision may cause significant cosmetic changes and, more importantly, impaired
function in the treated animal (e.g., mandibular drift after a segmental or total mandibulectomy with glossoptosis and drooling) (2). Prolonged recovery after large surgical resections, combined with usually high costs and poor outcome (high recurrence rate due to infiltrative tumor growth), makes the owners often reluctant to elect such invasive procedures, especially in older animals, as was also the case in the presented case. Based on our previous experience with treating oral and superficial cutaneous tumors in both dogs (10, 11) and cats (12) with either ECT or EGT or combined therapy, we offered this treatment combination as an alternative treatment option.

The dog received four therapy sessions in the course of 4 months and was euthanized 8 months after the first session due to tumor unrelated causes. At that time, there was no evidence of local tumor growth (oral cavity, intermandibular area, regional lymph nodes). Given the rapid tumor growth upon presentation (the mass approximately doubled in size in the course of one week before initiation of the treatment), the combined treatment resulted in slower progression of the tumor, with the first evidence of tumor regression 4 weeks after instituting ECT and EGT. With subsequent treatments we were able to induce complete regression of primary lesion, as well as subcutaneous metastases, which resulted in highly improved quality of the remaining life of the animal.

The pronounced cytoreductive effect of the combined treatment resulting in remission of described FSA can mainly be attributed to the ECT, since previous studies showed excellent local antitumor effect of this type of therapy, resulting in even up to 100% of complete response (CR) rates in certain human tumor types (13). In veterinary medicine, a number of studies employed ECT as an antitumor treatment, in which CR rates around 70-80% were accomplished (10, 11, 14). On the other hand, experience with IL-12 EGT as a single treatment in veterinary patients is lacking. The only published study (15) reported less pronounced cytoreductive effect. In contrast to preclinical studies on laboratory animals in which IL-12 EGT as a single therapy resulted in even 100% eradication of different tumor types (16, 17), study in canine mast cell tumors resulted in only 36% of CR rate (15).

It should be emphasized that the presented case exhibited one important distinction to other reports of ECT efficacy, namely distant effect on subcutaneous untreated metastases, which, according to the current knowledge, cannot be attributed to any antitumor effect of ECT. Although it is known, that ECT causes tumor antigen shedding into surrounding tissue and blood due to ECT-induced immunogenic cell death, this is considered inadequate for prevention of the growth of distant tumors, thus resulting in antitumor effect only at local level (18, 19, 20). Therefore the distant effect of the combined therapy in our case can be attributed to immunotherapeutic effects of the procedure. It was already established, both on preclinical level as well as human clinical study, that IL-12 EGT exhibits antitumor effect on distant untreated tumors, distant metastases and even elicits long-term resistance to regrowth of tumors (17, 21). Similar distant effect on lymph node metastases was also seen in other studies, employing IL-12 EGT with or without ECT component (5, 15, 22).

Immunological response to therapy was followed by flow cytometry, measuring different population of T lymphocytes. It was shown that dogs with tumors have decreased percentage of circulating CD8+ cells in comparison with healthy dogs (23). In our case the percentage of circulating CD8+ cells was increased after each session, which lasted up to 2 weeks. Therefore, the fluctuation of CD8+ cells in blood could serve as a guideline for timing repetition of the therapeutic procedure. In other studies, CD8+ cells were measured only in tumors, and the results showed the correlation between the increase of these cells in the tumor and better response of these tumors to the treatment in comparison to tumors without the increased infiltration of CD8+ cells (24, 25). It is proposed that intratumoral cytotoxic T lymphocytes migrate into the circulation reaching distant metastases where they can exert their immunological mediated tumor cell death (20). Therefore, the systemic increase of CD8+ lymphocytes could have abscopal effect on distant metastases, which was observed in our case. Therefore, the increase of CD8+ cells in blood could serve also as a predictive factor for the abscopal effect of the therapy.

One of the most important aspects of any gene therapy is environmental safety of the procedure. Namely, therapeutic plasmid contains antibiotic resistance gene, which can be horizontally transferred into commensal bacteria present on either treated patient’s skin or in gastrointestinal tract when treating oral tumors (26). To our best knowledge, only our research group addressed this aspect of safety of IL-12 EGT in clinical setting (11). In the presented case, persistence of plasmid DNA was...
monitored, at the site of injection (mucosa) as well as in the feces and urine. Similar to the larger study (11), majority of the plasmid in oral mucosa swabs was detected immediately after application, with abrupt decline of its quantity in the first week and complete disappearance in the course of the second week after the procedure. Furthermore, no plasmid was detected in feces and urine, further confirming the results of our previous study (11), that the IL-12 plasmid cannot be horizontally transferred into the culturable bacteria from the patients, therefore the possibility of any environmental shedding of antibiotic resistance gene is negligible.

**Conclusion**

In conclusion, the combination of ECT with bleomycin and EGT with IL-12 plasmid in the presented case of canine oral FSA exhibited very good local and systemic antitumor effect and was safe for the treated patient. Furthermore, potential for any environmental hazard of this type of gene therapy is negligible. Therefore, this treatment modality can represent an alternative type of therapy in selected cases, where clients seek a less invasive nonsurgical treatment of these tumors.

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