Characterization of a New Murine Model of Cancer Pain

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This paper describes a model of tumor-induced bone destruction and hyperalgesia produced by implantation of fibrosarcoma cells into the mouse calcaneal bone. Histological examination indicates that tumor cells adhere to the bone edge as early as post-implantation day (PID) 3, but osteolysis does not begin until PID 6, correlating with the development of hyperalgesia. C3H/He mice exhibit a reproducible hyperalgesia to mechanical and cold stimuli between PID 6 and 16. These behaviors are present but significantly reduced with subcutaneous implantation that does not involve bone. Systemic administration of morphine (ED50 9.0 mg/kg) dose-dependently attenuated the mechanical hyperalgesia. In contrast, bone destruction and hypersensitivity were not evident in mice implanted with melanoma tumors or a paraffin mass of similar size. A novel microperfusion technique was used to identify elevated levels of the putative algogen endothelin (ET) in per fusates collected from the tumor sites of hyperalgesic mice between PID 7 and 12. Increased ET was evident in microper fusates from fibrosarcoma tumor-implanted mice but not from melanoma tumor-implanted mice, which are not hyperalgesic. Intraplantar injection of ET-1 in naive and, to a greater extent, fibrosarcoma tumor-bearing mice produced spontaneous pain behaviors, suggesting that ET-1 activates primary afferent fibers. Intraplantar but not systemic injection of the ET-A receptor antagonist BQ-123 partially blocked tumor-associated mechanical hyperalgesia, indicating that ET-1 contributes to tumor-induced nociception. This model provides a unique approach for quantifying the behavioral, biochemical, and electrophysiological consequences of tumor–nerve interactions.

Key words: hyperalgesia; primary afferent fibers; tumor nociception; endothelin; cancer pain; tumor microperfusion

Pain is often the first indication of tumor presence or recurrence (Caraceni and Portenoy, 1999) and is present in 80% of cancer patients at death (Reale et al., 2001). Metastasis to bone, often with adjacent soft tissue involvement, is thought to be the most common cause of cancer-related pain (Banning et al., 1991; Mercadante, 1997; Reale et al., 2001). Clinical evidence indicates that tumors cause pain in bone and soft tissues via nerve compression, the release of chemical algogens, and pH changes, whereas pain from microfractures and stretching of the periosteum are unique to bone (Mercadante, 1997). Despite an extensive literature regarding cancer pain assessment and management, our understanding of the basic mechanisms that underlie the production of pain associated with malignancy is meager at best. An understanding of these basic mechanisms is essential for the development of better therapeutic approaches to cancer pain treatment, but efforts to gain such information are hampered by the lack of adequate animal tumor models. Recently, Schweig et al. (1999) described a femur model of bone cancer pain that allowed characterization of neurochemical changes in the spinal cord associated with development of tumor-induced nociception. Although the femur bone tumor model provides a valuable archetype to assess CNS changes, it is difficult to examine algogen release, quantify primary hyperalgesia, and perform electrophysiological analysis of primary afferent fibers innervating the tumor, bone, and other deep tissues in these mice. Here we report on the development of a hindpaw tumor model that incorporates both bone and adjacent soft tissue involvement to produce a localized cancer pain that lends itself more readily to behavioral, electrophysiological, and peripheral neurochemical analysis.

Tumor cells are known to secrete a variety of different substances (Hall, 1997; Chirgwin and Guise, 2000), many of which are potential algogens. One of the goals of the present study was to examine the putative role of one of these mediators, endothelin-1 (ET-1), in the induction of cancer pain in the hindpaw tumor model. Endothelin is a 21 amino acid peptide derived from a larger precursor, big-endothelin, by action of endothelin-converting enzyme; three isoforms of endothelin, named ET-1, ET-2, and ET-3, have been identified (Rubanyi and Polokoff, 1994). ET-1, generated by a number of cell types and a number of tumor cell lines, exerts various important biological actions mediated by two receptor subtypes, ET₁ and ETᵢ (Gandhi et al., 1994; Webb, 1997). Ferreira et al. (1989) were among the first to show that ET-1 participates in the production of inflammatory
pain. Subsequently all three isoforms of ET were shown to elicit an abdominal constriction response in mice, ET-2 being the most potent (Raffa and Jacoby, 1991). Since these initial studies, a role for ET in nociception has been well documented (Davar et al., 1998; De-Melo et al., 1998; Piovezan et al., 1998, 2000; Fareed et al., 2000), and both receptor subtypes have been shown to participate in ET-induced nociception (Raffa et al., 1996). The present study targets the peptide ET-1 and tests the hypothesis that it participates in the production of bone tumor pain.

**MATERIALS AND METHODS**

**Animals**

A total of 313 C3H/He and 42 B6C3fie/1 mice (National Cancer Institute) aged 8–10 weeks and weighing 24–28 gm were used in all fibrosarcoma or mixed melanoma/fibrosarcoma experiments, respectively. The inbred mouse strain C3H/He is syngeneic to the fibrosarcoma cells used in these experiments and allows these cells to grow tumors without rejection (Clohisy et al., 1996). The B6C3fie/1 mice, the F1 cross between C57BL/6 and C3H/He strains, readily accept both fibrosarcoma (C3H/He origin) and melanoma (C57BL/6 origin) cells, allowing direct comparison of the effects of these tumor types. Mice were housed in boxes of 8–10 in a temperature- and humidity-controlled environment and maintained on a 12 hr light/dark cycle with ad libitum access to mouse chow and water. All experimental protocols were approved by the Animal Care and Use Committee of the University of Minnesota.

**Cell culture and implantation**

National Collection of Type Cultures (NCTC) clone 2472 fibrosarcoma cells, originally derived from a connective tissue tumor in a C3H mouse, were obtained from the American Type Cell Culture Collection (Rockville, MD). G3.26 cells, a B6 subclone melanoma cell, originally derived from a C57BL/6 mouse, were obtained from Dr. Christopher W. Stackpole (New York Medical College, Valhalla, NY) (Stackpole et al., 1985). All cells were maintained as described previously (Clohisy et al., 1996). Just before implantation, cells were counted with a hemacytometer, pelleted, resuspended, rinsed in PBS, pelleted a second time, and then resuspended in PBS for implantation.

Mice were placed in an enclosed chamber and anesthetized with 2% halothane in preparation for cell implantation. When the animal demonstrated nonresponsiveness to paw pinch, it was removed from the chamber and fitted with a facemask that continuously delivered 2% halothane in an air/oxygen mixture throughout the procedure. Cells (2 x 10^5 fibrosarcoma or 1.5 x 10^5 melanoma) in a volume of 10 μl of PBS were injected unilaterally into the heel using a 29 gauge, sterile single-use needle attached to a 0.3 ml insulin syringe (Becton Dickenson) to manually bore through the calcaneus bone. Sham mice underwent the same procedures with a heated 27 gauge needle and syringe to produce a nonmalignant mass approximately the size of a post-implantation day (PID) 10 fibrosarcoma tumor.

**Tumor histology**

At days 3, 6, 9, and 12 after tumor implantation, mice were deeply anesthetized with 100 mg/kg sodium pentobarbital and transcardially perfused with 15 ml of cold PBS followed by 30 ml 4% paraformaldehyde. Both hindpaws were removed and post-fixed for 4 hr in 4% paraformaldehyde and then transferred to decalifying solution (0.002 M EDTA, 1.35N HCl) for 24 hr. The tissue was rinsed, dehydrated, and paraffin-embedded, cut into 5 μm cross sections using a rotary microtome, and stained with hematoxylin and eosin. Sections through both the ipsilateral hindpaw containing the tumor and the contralateral hindpaw were examined histologically under bright-field microscopy. Sections were examined for the presence and degree of bone destruction and for evidence of immune cell infiltration.

**Tumor size and calcaneus thickness**

Both the melanoma and fibrosarcoma were localized and formed a mass in the region of the injection site with the fibrosarcoma tumors forming a particularly even sphere around the heel. Therefore, the relative tumor (days 3–15) and paraffin bleb (days 1–7) sizes were determined by measurement of the heel width. Measurements were taken after behavioral testing in the following manner. Mice were held by the tail and allowed to grasp a wire mesh, leaving their hindpaws free while a micrometer was positioned over the heel and the diameter of the tumor was measured percutaneously. On PID 15, mice were killed; tumor/ connective tissue was carefully removed from the remaining calcaneus bone, and the maximum diameter of the remaining bone was measured with a micrometer.

**Behavioral methods**

**Mechanical hyperalgesia assay**

Withdrawal responses evoked by mechanical stimuli were obtained in tumor-bearing mice and compared with sham-treated animals at several time points after implantation as well as in a series of pharmacological experiments described below. Groups of mice were prescreened for hypersensitivity with a von Frey monofilament, 3.4 mN (C3H/He mice) or 1.6 mN (B6C3fie/1 mice) bending force, and high responders (responses of ≥50% before treatment) were removed from further experimentation (<5% of mice). The 3.4 mN monofilament was used in the C3H/He mice because their responses to this monofilament were reproducible and sufficiently large to detect dose-dependent attenuation by analogics or ET-1 antagonists (Wacnik et al., 2000).

Baseline values for mechanical sensitivity were determined for each animal 4 d before, 1 d before, and on the day of implantation (cell-implanted and sham groups) or immediately before analgesic/antagonist administration; testing was repeated throughout the time course of each study. Briefly, animals were placed on a wire mesh platform, covered with a hand-sized container, and allowed to acclimate to their surroundings for a minimum of 30 min before testing. The monofilament was applied to the point of bending six times on the plantar surface of each hindpaw for tumor time course studies. In pharmacological studies (morphine, cycloheximide, and ET-1 antagonists), the monofilament was applied 10 times on the paw ipsilateral to the tumor. The number of vigorous responses to the monofilament was counted and expressed as percentage of stimuli giving rise to a withdrawal response.

**Cold hyperalgesia assay**

The same groups of mice were tested for cold hyperalgesia after determining responses to mechanical stimuli using a constant temperature cold plate repeatedly over time as tumors grew. The cold plate consisted of an aluminum test surface (10 x 15 cm) enclosed in a clear Plexiglas container (20 cm high) maintained at 3°C by a thermostatically controlled water bath and circulating pump. After placing the mouse on the cold plate, the experimenter counted the frequency of withdrawal responses over a 4 min period. A withdrawal response included one or more of the following behaviors: hindpaw held above the cold plate (one response per second), shaking or licking a hindpaw (one response per a whole mouse movement or one response per leap). This cold hyperalgesia protocol is based on a rat model used to analyze hyper-responsiveness in inflammatory (complete Freund’s adjuvant) and neuropathic (chronic constriction injury) experimental hyperalgesia models (Jasmin et al., 1998).

**In vitro and in vivo analysis of endothelin secretion**

As part of our investigation of endothelin and its potential role in tumor-induced pain, we conducted three experiments to measure (1) relative levels of ET secreted by tumor cells in vitro, (2) relative concentration of ET in homogenates of developing hindpaw tumors, and (3) relative concentration of ET secreted by tumor cells in vivo at various time points after implantation. The level of ET secreted by tumor cells in vitro was determined by growing cells in serum-free media for 24 hr and subsequently analyzing ET levels in the culture media (conditioned media). Whole-tumor levels were determined by harvesting and homogenizing tumors and then evaluating relative ET levels in homogenate supernatants, whereas the level of ET secreted into the extracellular fluid of the tumor in vivo was sampled by microperfusion, as described below. Homogenate samples were collected at PID 5–12, and microprobe samples were collected at PID 8–13. All tumor samples were taken from mice that exhibited mechanical hyperalgesia at the given time points.

**ET levels in tumor cell cultures**

Tumor cell cultures (cell lines 2472 and G3.26) were grown in serum-free media for 24 hr, at which time the cells were at equal con-
milliliters of conditioned media were collected and centrifuged for 20 min at 1000 rpm, and the supernatant was aliquoted and frozen at − 80°C until time of analysis. ET concentration was compared between conditioned and unconditioned media (media not incubated with cells).

**ET levels in tumor homogenates**

Hindpaw tumors were dissected away from the surrounding connective tissue. Normal tissue from a comparable area of the contralateral hindpaw was collected for comparison. Samples were placed in ice-cold buffer (PBS with 0.4% NaCl, 0.05% Tween 20, 0.5% NGS, 0.1 mm phenylmethylsulfonyl fluoride, 0.1 mm benzethonium chloride, 10 mm EDTA, and 1% protease inhibitor mixture) and finely minced with a scissors. The tissue suspension was ground with a disposable pestle (Fisher Scientific, Houston, TX) and then centrifuged at 15,000 rpm for 30 min at 4°C. The supernatant was aliquoted and frozen at − 80°C until the time of analysis.

**ET in tumor extracellular fluid**

**Microprobe design.** Microdialysis, traditionally used to sample low molecular weight substances from extracellular fluid, has limited ability to dialyze proteins and peptides. In practice, a membrane with a 20 kDa molecular weight cutoff (MWCO, Gambro Hospel Ltd., Huntingdon, UK) yields significant transport only at and below 5 kDa. Moreover, some proteins such as tumor necrosis factor α (TNFα) and nerve growth factor (NGF) adhere to the microdialysis membrane, further limiting their passage through the membrane. Therefore, we developed and applied a perfusion microprobe consisting of a 23 gauge stainless steel hypodermic needle with a 2 mm opening in the middle, extending a “push–pull” microperfusion design based on the procedure recently described by Patterson et al. (2001). This microprobe design allowed increased and reproducible peptide/protein yields and increased mechanical stability and durability.

**In vitro testing of the microprobe.** We compared in vitro recoveries of ET-1 between traditional microdialysis (20 kDa MWCO probe) and push–pull microperfusion techniques using a modified Ringer’s solution at a flow rate of 3 μl/min for dialysis (Renno et al., 1998) and 10 μl/min (each flow rate was confirmed for run time) for microperfusion (Patterson et al., 2001). Both techniques used peristaltic pumps (Rabbit Plus, Rainin Instrument Co.). In vitro microperfusion and microdialysis techniques were compared by determining the percentage recovery of ET-1 or bovine serum albumin (BSA) from a 100 μg/ml solution of these substances in a restricted region of a Petri dish delineated with a hydrophobic barrier (Pap Pen, Research Products International, Natick, MA).

**Microperfusion levels of ET in the extracellular fluid of the hindpaw fibrosarcoma.** We implanted a microprobe into the tumor site or into a comparable site in control animals. This procedure is terminal so that each mouse contributes data to a single time point. Preliminary studies showed no difference in ET release in naive versus sham-injected animals at all time points examined; therefore, naive animals were used as controls in the microperfusion experiments. A small amount of heparin (2 μl) prevented clotting in the microprobe. Mice were anesthetized initially with 0.6 mg acemephrame and 10 min later with 40 mg/kg Nembutal. When nociceptive withdrawal and eyeblink reflexes were absent, the microprobe was inserted through the center of the tumor site and secured to the skin with super glue. Two lengths of polyethylene tubing (PE-10, prefilled with heparin-nPBR, 30 μl/ml) connected the microprobe to the fluid swivel and two peristaltic pumps. The inlet perfusion pump pushed modified Ringer’s solution into the microprobe at a rate of 10 μl/min the outlet pump was set at a rate of 20 μl/min (as determined from in vitro experiments), which pulled at an effective rate of 10 μl/min in vivo. Preliminary experiments had determined that this setting on the outlet pump maintained a constant pressure at the microprobe opening and prevented clogging of the probe. After 45 min equilibration and awakening, samples were collected for a period of 3–4 hr, centrifuged to assess the amount of red blood cells (micrograms per milliliter), augumented with a protease inhibitor mixture (Sigma, St. Louis, MO; concentration = 0.00014%, based on manufacturer’s recommendation), and stored at −80°C for later analysis.

**Quantification of ET using microbead immunosorbent assay**

The following microbead immunosorbent assay (MBISA) was used to measure ET levels in all release and homogenate studies. The use of flow cytometry has recently been adapted to evaluate the presence of solubilized proteins that have been immobilized on polystyrene beads (Curt-singer et al., 1997), and we have adapted this method to determine relative levels of individual proteins in a mixture. The adapted procedure is cost effective and efficient, requires very little sample, and allows analysis of multiple proteins from single samples. Briefly, beads (4 μm sulfate polystyrene; Interfacial Dynamics, Portland, OR) are coated with proteins and peptides contained in a sample. The protein of interest is labeled with a fluorophore, and the beads are subjected to flow cytometry. Previous experiments indicated a linear relationship between the amount of a known protein bound to the bead and the fluorescence intensity of the beads (data not shown).

Analysis was begun by incubating 1 μl of conditioned cell culture medium in 9 μl of Dulbecco’s PBS or 10 μl of perfusate (average total protein of 3 μg) with 107 (1 μl) beads overnight at 4°C. To ensure equal loading of protein onto beads between tumor homogenates, a total protein concentration was determined (Coomassie Plus, Pierce, Rockford, IL), and 10 μg of protein in a volume of 10 μl was incubated with 107 beads for 2 hr at room temperature. After the sample incubation period, the beads with bound protein from conditioned medium, perfusate samples, and homogenates were treated identically. Nonspecific binding to the beads was blocked with PBS containing 1% normal goat serum and 5 mm sodium azide (FACS wash buffer) for 30 min at room temperature. Beads were washed 5 times at 14,000 rpm. Protein adsorption was discarded, and the bead pellet was resuspended in 20 μl of FACS wash buffer. Ten microliters of resuspended beads were added to rabbit anti-mouse ET serum (1:1000; Sigma), and 10 μl was added to control serum (1:1000 normal rabbit serum) and incubated at room temperature for 1 hr. The primary antibody used in this study recognizes all three forms of ET (ET-1, ET-2, and ET-3) and an alpase antibody that is specific to ET-3 to determine whether ET-3 levels are specifically elevated in tumor microperfusates.

Beads were pelleted and washed with FACS wash buffer, resuspended, and incubated with FITC-conjugated goat-antirabbit Ig (1:100; Jackson ImmunoResearch Laboratories, West Grove, PA) for 1 hr at room temperature. Beads were pelleted, washed, and resuspended in FACS wash for analysis. For these assays, the mean fluorescence intensity (MFI) of 5000 beads was determined using a flow cytometer (FACS-Calibur, Becton Dickinson, Mountain View, CA). The net mean fluorescence was determined by subtracting the MFI of antibody isotype control-incubated beads from the MFI of beads incubated with ET-specific antibodies. Because previous experiments (data not shown) have demonstrated that the resulting net MFI is linearly correlated with the ET concentration, MFI was used for statistical analyses.

**ET-1-induced noicception**

Mice were placed on a wire mesh platform, covered with a hand-sized container, and allowed to acclimate to their surroundings for a minimum of 30 min before testing. ET-1 (4.0 pmol–1.2 nmol/30 μl; American Peptide Co., Sunnyvale, CA) or vehicle was injected into the tumor site or subcutaneously into the heel of control (naïve) mice, and the cumulative licking responses directed to the hindpaw were counted over a 10 min post-injection period. An injection was assumed that both the tumor and surrounding tissue were bathed in the drug or vehicle. The duration of observation of licking time was chosen on the basis of preliminary studies showing that the majority (64%) of nocifensive behaviors occurred in the first 10 min of a 20 min test period. ET-1 was dissolved at 1 mg/ml in 1% NaHCO3, and further dilutions to appropriate concentrations were made in sterile saline so that vehicle for control experiments ranged from 0.001 to 0.5% NaHCO3 in sterile saline (ET-1, 4.0 pmol–1.2 nmol/30 μl).

**Inhibition of tumor-induced nociception**

**Withdrawal responses after mechanical stimuli**

Mechanical hyperalgesia was used to evaluate tumor-induced nociception (pre-drug baseline) and to measure the anesthetic effect of selected inhibitors. The advantage of testing mechanical sensitivity with von Frey monofilaments is that it allows several post-drug measurements to be made without handling the mouse, permitting time-dependent and dose-dependent analysis in tumor-implanted mice. When tumor-induced mechanical hyperalgesia was evident, each C3H/He mouse was tested before and after drug administration with the 3.4 mN von Frey monofilament. Times on the plantar surface of the ipsilateral paw with responses >50% and with no signs of skin lesions were included in the analgesic/antagonist studies. The degree of drug-induced inhibition of mechanical hypersensitivity was determined relative to the pre-drug baseline. Percentage inhibition was calculated using the following for-
mula: % inhibition = (% response pre-drug - % response post-drug) \times 100/\% response pre-drug

The ED_{50} values and 95% confidence limits were calculated according to the method of Tallarida and Murray (1987). This protocol and analysis have been reproduced in multiple experiments to measure dose-dependent attenuation of tumor-induced mechanical hyperalgesia (Wacnik et al., 2000) by the analgesics morphine and clonidine, as reported previously (Fairbanks et al., 2000).

**Morphine**

The activity of morphine as an analgesic was tested in this model to support the idea that a hyperalgesic condition underlies the behavior. Hyperalgesic mice (PID 15) were administered morphine systemically (3–30 mg/kg, i.p.) and tested again 30 min after administration, and the percentage inhibition was calculated. This morphine dose range was based on previous work (Wacnik et al., 2000) in which mechanical hyperalgesia produced by fibrosarcoma tumors in the femur was attenuated.

**Cycloheximide**

To test for inhibition of mechanical hypersensitivity induced by putative peptidergic algogens secreted at the tumor site, the protein synthesis inhibitor cycloheximide (150 µg/10 µl PBS) was injected into the tumor site of PID 8 hyperalgesic mice. Mice were tested again 0.5, 2, 4, 6, 8, 12, and 24 hr after cycloheximide administration, and the percentage inhibition was calculated.

**ET receptor antagonists**

To determine whether ET contributes to tumor-evoked mechanical hyperalgesia and to determine the receptor types responsible, the ET-A receptor antagonist BQ-123 [0.16–16 nmol/30 µl PBS, c(Trp-D-Asp-Pro-D-Val-Leu)] and the ET-B receptor antagonist BQ-788 (0.16–48 nmol/30 µl PBS, N-citr-2,6-dimethylpyrrolidinocarbonyl-L-γ-methylleucyld-1-methoxybenzyltripropanyl-D-Nle; American Peptide Co.) or saline was administered into the tumor site of PID 8 hyperalgesic mice. Mice were tested again 0.5, 2, 4, 6, 8, 12, and 24 hr after ET-1 antagonist administration, and the percentage inhibition was calculated. Injecting a volume of 30 µl assured that the tumor and the surrounding paw were bathed in drug or vehicle. Antagonist testing was conducted at a time after implantation when both microperfusion and homogenate levels of ET where found to be elevated (PID 10).

**Statistical analysis**

Mean withdrawal response frequency to mechanical stimuli, cold-plate responses, and ET-induced nociceptive behaviors as well as mean fluorescence intensity data were analyzed by repeated-measures ANOVA; Bonferroni or Fisher’s PSDL post hoc comparisons were used for behavioral time course and fluorescence intensity data as appropriate to determine significance across the tumor time course analysis using StatView 5.0 (SAS Institute). Data are presented as mean and SEM for treatment groups. Statistical significance is reported for p < 0.05 except as noted.

**RESULTS**

**Tumor morphology**

Histological examination of fibrosarcoma tumors revealed a non-encapsulated tumor mass consisting of spindle-shaped cells (Fig. 1B), characteristic of a fibrosarcoma. As early as PID 3, tumor cells adhered to the bone edge, but osteolysis was not evident until PID 6 (Fig. 1). Bone destruction progressed through PID 12, as indicated by increasing irregularity of the bone edge as well as decreasing bone thickness and eventual breakthrough. At PID 6, 9, and 12, nerves within the tumor mass could be identified, and at these time points there was no evidence of either nerve degeneration or invasion by tumor cells. At all time points examined, the skin overlying the tumor had normal morphology and was not invaded by tumor cells. Comparison of fibrosarcoma and melanoma tumors on PID 9 revealed that although the fibrosarcoma tumors showed evidence of osteolysis, melanoma tumors did not (Fig. 2). The melanoma tumor was separated from bone matrix by an intact layer of periosteum, and at no point did the bone edge become irregular, thus indicating a lack of bone invasion by the melanoma tumor. Histological examination of the fibrosarcoma and melanoma tumors revealed very little inflammatory cell (neutrophil and lymphocyte) infiltration of the tumor site (tumor, bone, muscle, and surrounding connective tissue). These results corroborate the histological findings of Clohisy et al. (1996) after implantation of these fibrosarcoma cells into the medullary space of the femur.

**Tumor-induced behavioral changes**

Behaviorally, there was neither evidence of ongoing pain (guarding of the hindpaw) nor any signs of evoked pain during palpation of the hindpaw in animals receiving either sham injection or melanoma tumor cell injection into the calcaneus bone. In contrast, animals injected with fibrosarcoma cells into the calcaneus showed pronounced curling of the toes, cupping and guarding of the ipsilateral paw, and a distinct preference for weight bearing on the contralateral hindpaw during normal ambulation on a wire mesh surface. Curling of the toes and cupping and guarding of the paw were also evident while the animals were being handled. Furthermore, palpation of the tumor-bearing heel from PID 8 to 12 elicited a withdrawal response.

**Fibrosarcoma-induced hyperalgesic behaviors**

Fibrosarcoma implantation into and around the heel of C3H/He mice induced hyperalgesia to mechanical and cold stimuli when compared with naive and sham-implanted controls (Fig. 3). Mechanical hyperalgesia was evident in response to stimulation of the hindpaw with a normally non-noxious von Frey monofilament (3.4 mN bending force) (Fig. 3A) as early as PID 3. In addition, progressive cold hyperalgesia was observed as a significant increase in the number of nocifensive behaviors over a 4 min period on the 2–4°C cold plate beginning by PID 10 (Fig. 3B). Tumor cell implantation into the calcaneus bone produced a greater hypersensitivity with an earlier onset of increased responsiveness (Fig. 3) compared with subcutaneous implantation into the heel not involving the bone, which yielded 34.3 ± 2.7% response to mechanical stimuli [area under the curve (AUC), 6–17 PID] and 10.8 ± 1.3 nociceptive behaviors on the cold plate (AUC, 7–16 PID).

Tumor size in C3H/He mice, measured over the time course as the width of the heel, showed continuous progression with PID 10 measurements of 4.6 ± 0.2 mm compared with 3.3 ± 0.06 mm for the naive group (n = 10). C3H/He mice, injected with paraffin wax of size similar to the fibrosarcoma in the heel as a non-tumor mass control, were tested across days 1–7; observed heel width of 4.5 ± 0.08 mm was not accompanied by hyperalgesia (22.3 ± 5.5% response; n = 5). Behavioral testing was concluded at PID 15 because the incidence of skin lesions in a small number of animals might have confounded sensory testing results. Bone loss was evident at necropsy in mice that exhibited osteolysis, but not in the sham-treated mice or mice with tumors in subcutaneous tissue only (see below).

**Fibrosarcoma but not melanoma tumors of similar size induce mechanical hyperalgesia**

Nonosteolytic melanoma tumors of size similar to the fibrosarcoma did not produce hyperalgesia when implanted in the heel of B6C3H/er mice. Figure 4 presents the correspondence between increasing tumor/heel size (bars) and hyperalgesia (lines): fibrosarcoma growth was more localized to the heel than was melanoma through day 15 after implantation, and this was accompa-
nied by more intense hyperalgesia in the fibrosarcoma-implanted mice. Postmortem dissection of the tumor-implanted paws showed that the melanoma and fibrosarcoma tumors had equal mass on PID 15 but that fibrosarcoma tumors adhered to and invaded the calcaneus bone and caused its destruction. Dissection and subsequent measurement of the calcaneus bone from the fibrosarcoma-injected heel revealed a significant reduction in calcaneus thickness: 0.7 ± 0.1 versus 1.2 ± 0.03 mm on the contralateral side compared with 1.1 ± 0.03 mm in the melanoma-bearing heel.

**Systemic morphine dose-dependently attenuates mechanical hyperalgesia**

Systemic morphine injected (3–30 mg/kg, i.p) on PID 15 attenuated the fibrosarcoma-induced mechanical hyperalgesia in calcaneus-implanted C3H/He mice in a dose-dependent manner with an ED_{50} of 9.0 mg/kg (95% confidence interval, 6.8–11.7) and a maximum inhibition of 87% at 30 mg/kg (n = 12). In addition, morphine was found to be effective in attenuating tumor-induced cold sensitivity (data not shown). Neither sedation

**Figure 1.** Photomicrographs of hematoxylin and eosin-stained fibrosarcoma tumor sections at different PID time points. A, PID 3 fibrosarcoma: tumor cells are closely adhered to bone surface, but the bone edge is intact. B, PID 6 fibrosarcoma: bone edge is irregular, indicative of osteolysis. Arrows indicate individual spindle-shaped fibrosarcoma cells. C, Skin overlying tumor at PID 9; note the lack of skin invasion by tumor cells and normal skin morphology. D, PID 12 fibrosarcoma: intact nerve bundle surrounded by tumor cells. There is no evidence of nerve invasion by tumor cells or of nerve degeneration at this time point. Scale bars (shown in A): A, B, D, 20 μm; C, 60 μm.
nor motor impairment were observed during the post-drug testing period after morphine administration, although some hyperactivity was evident at the high dose (30 mg/kg). Analgesic attenuation of hyperalgesia without sedation or motor impairment validates this model of hyperalgesia.

**Measurement of ET in fibrosarcoma cells and tumors**

*In vitro* testing of the microprobe indicated that recoveries of ET-1 and BSA were 68 and 62%, respectively, as compared with a microdialysis probe recovery of 4.2% for ET-1 and 0% for BSA.

Figure 2. Photomicrographs of hematoxylin and eosin-stained sections of a normal and tumor-bearing mouse heel. A, Cross section of normal mouse heel. B, Cross section of a comparable area at PID 9 of fibrosarcoma cells. C–F, Comparison of fibrosarcoma tumor and control melanoma tumor morphology. C, PID 9 melanoma tumor. D, Enlargement of boxed area in C; note the regular bone edge and layer of periosteum (arrow) separating bone and tumor cells. E, PID 9 fibrosarcoma tumor. F, Enlargement of boxed area in E; note irregular bone edge and invasion of tumor cells into bone, indicating osteolysis. Scale bars: (shown in B) A, B, 500 μm; (shown in C) C, E, 100 μm; (shown in D) D, F, 30 μm.
Similar in vitro microperfusion recovery rates were found when testing NGF and TNFα (data not shown). Labeling of known amounts of ET adsorbed to beads with antibody and analysis by flow cytometry indicated that MFI increases linearly with the concentration of ET-1.

Fibrosarcoma tumor homogenates contain increased levels of ET

MBISA of homogenates of whole tumors on PID 5, 7, 10, and 12 (Fig. 5) showed that ET MFI was increased on PID 7, 10, and 12 (19.7 ± 0.9, 26.7 ± 4.7, and 17.6 ± 5.2) compared with control MFI from the contralateral paw at PID 5 (5.6 ± 0.35). The inset in Figure 5 shows representative flow cytometry histograms for naïve and tumor-implanted mice at PID 7, 10, and 12. The production of ET by the fibrosarcoma tumor increased by PID 7 and peaked around PID 10.

Fibrosarcoma tumor microperfusates contain increased levels of ET

In vivo microperfusion of fibrosarcoma tumors in awake, freely moving mice between PID 8 and 13 evaluated the time course of ET release into the extracellular fluid of the tumor site (Fig. 6).
MBISA of the perfusates showed that ET MFI increased on PID 9, 10, and 11 (39.7 ± 6.1, 41.9 ± 4.7, and 53.0 ± 8.4 compared with naive mice, 20.8 ± 7.1). Replication of this experiment in B6C3fe/1 mice and comparison of mice with fibrosarcoma and melanoma tumors yielded consistent results: MBISA showed higher ET MFI in mice with fibrosarcoma tumors compared with mice implanted with melanoma tumors or compared with naive controls (57.9 ± 9.2 vs 28.3 ± 5.9 and 23.2 ± 8.1, respectively) (Fig. 7). As indicated in Materials and Methods, the ET antibody used recognizes all three forms of ET; absence of increased ET-3-immunoreactivity in the fibrosarcoma microperfusates (data not shown) suggests that the tumor releases either ET-1 or ET-2.

In vitro microperfusion

MBISA on media conditioned for 24 hr with fibrosarcoma cells yielded higher ET MFI than did MBISA of melanoma-conditioned and cell-free control media (50.1 ± 8.3 compared with 6.3 ± 1.2 for melanoma-conditioned and 11.2 ± 3.0 for cell-free control media; \( p < 0.0005 \)) (Fig. 7). This result demonstrates that the fibrosarcoma cell line, but not the melanoma cell line, produces and secretes ET, indicating that the fibrosarcoma cells contribute to the release of the ET measured in tumor homogenates and tumor microperfusates.

**Algogenic activity of ET-1**

Injection of ET-1 (4.0 pmol–1.2 nmol/30 μl) into the ipsilateral hindpaw of mice bearing fibrosarcoma tumors at PID 10 (when ET perfusate yield peaked) produced dose-related licking of and attending to the injected paw for 10 min when compared with vehicle-injected and naive controls (Fig. 8). This experiment shows that tumor-bearing mice manifest higher sensitivity to ET-1 than naive mice, supporting a local pronociceptive action of ET-1.

**Cycloheximide injected into the tumor site attenuates mechanical hyperalgesia**

Injection of the protein synthesis inhibitor cycloheximide (150 μg in 10 μl PBS) into the calcaneus tumor site of hyperalgesic mice (PID 8; \( n = 5 \)) attenuated responses to 40 ± 9.2% relative to preinjection baseline (82 ± 4.6%) more than PBS sham injection (67 ± 8.8% response) between 4 and 12 hr after injection.

**ET-A receptor antagonist BQ-123 attenuates ET-1- and tumor-induced hyperalgesia**

Intraplantar injection of the ET-A receptor antagonist BQ-123 (1.6 or 16 nmol/30 μl) in naive C3H/He mice before ET-1 (400 pmol/30 μl) injection reduced the time spent licking to control...
levels. Pretreatment with saline or the ET-B receptor antagonist BQ-788 (0.1, 1.0, and 10 nmol/30 μl) was without effect. Intraplantar injection of BQ-123 (16 nmol) had no effect on nociceptive mechanical sensitivity (von Frey monofilament, 12.1 mN bending force) in naive mice (n = 7; data not shown).

To test for ET receptor participation in tumor-induced hyperalgesia, BQ-123 (0.16–16 nmol/30 μl), BQ-788 (0.16–48 nmol/30 μl), or saline (30 μl) was injected into the tumor site of PID 10–12 fibrosarcoma-implanted C3H/He mice showing >60% responsiveness to von Frey stimulation (3.4 mN). Preliminary experiments had resolved the time course of BQ-123 effects (peak 45 min, duration until 180 min after injection). Figure 9 shows that BQ-123 reduced hyperalgesia 45 min after injection, reaching a maximum inhibition of 43 ± 11.6% at 1.6 nmol. BQ-788 was inactive at doses below 16 nmol as was saline, but 16 nmol of BQ-788 reduced hyperalgesia slightly (15 ± 11.6%). Systemic (intraperitoneal) administration of BQ-123 (16–48 nmol/30 μl) in tumor-bearing mice was inactive at any time point tested. These results suggest that ET-1 contributes to the mechanical hyperalgesia in tumor-bearing mice by activating ET-A receptors.

**DISCUSSION**

The present paper, together with its companion paper (Cain et al., 2001b), defines a new model of cancer pain in which an osteolytic tumor grows locally and around the calcaneus bone of the mouse hindpaw. Together, these two studies define the histological, behavioral, neurochemical, and neuroanatomical, and neurophysiological characteristics of the tumor model. The model is distinguished by the presence of spontaneously active and hyper-responsive C-fibers and the development of morphine-sensitive mechanical and cold hyperalgesia. The tumor-induced nociception appears to result from the release of algogenic mediators, the participation of one of which, endothelin, is documented in this paper. The joint conclusion of these two papers is that early hyperalgesia appears to be tumor induced and nociceptive in nature, but that neuropathic components may develop later in the course of the growth of the tumors.

Beyond assertions of tumor infiltration around bone or nerve and the utility of morphine in treating cancer pain, a lack of knowledge surrounds the mechanistic basis of cancer pain, primarily because animal models have only recently been described in mouse femur (Schwei et al., 1999; Wacnik et al., 2000) and humerus (Wacnik et al., 2000). These initial models are limited by the difficulty of electrophysiological and neurochemical evaluation of deep peripheral hyperalgesia. The hindpaw, by contrast, is a standard site for the development and study of many acute and chronic pain models in mice (Mansikka et al., 1999; Mogil et al., 1999; Mogil et al., 1999; Fairbanks et al., 2000) and rats (Stein et al., 1988; Schadrack et al., 1999; Jinks and Carstens, 2000; Du et al., 2001; Zheng and Chen, 2001) because it provides ready access for testing primary and secondary hyperalgesia and allows for electrophysiological recording (Cain et al., 2001a,b), local drug delivery, and fluid collection. This new tumor pain model applies fibrosarcoma cells into and around the hindpaw calcaneus bone to mimic the painful condition observed in human calcaneus bone cancer (Sarlak et al., 2000) and characterizes behavioral, morphological, neurochemical, and algogenic sequelae. The companion paper (Cain et al., 2001b) characterizes electrophysiological and neuroanatomical sequelae to tumor growth.
The NCTC 2472 fibrosarcoma cell line activates osteoclasts and promotes bone destruction when injected into the medullary cavity (Clohisy et al., 1996), and tumors resulting from implantation of cells in the femur produce spontaneous pain behavior (Schwei et al., 1999; Honore et al., 2000a) and secondary cutaneous hyperalgesia (Wacnik et al., 2000; Honore et al., 2000b). The present study shows that cancerous implantation involves both bone and soft tissue, enhances responses to mechanical and cold stimuli at the tumor site, and elicits spontaneous nociceptive behaviors (favoring, cupping, and guarding the affected paw) reminiscent of cancer pain symptoms involving the tibia or calcaneus (Caraceni and Portenoy, 2000). Morphine attenuation of hyperalgesia seen in this hindpaw model both confirms its usefulness as a model of cancer pain and indicates its susceptibility to opioids. Furthermore, it is notable that the systemic potency of morphine in this chronic pain model (ED50 ~10 mg/kg) is comparable to its potency in the mouse strain using common acute thermal nociceptive tests (ED50 3–15 mg/kg) and lower than its potency in tonic

**Analgesics and cancer pain**

Opioids remain the key treatment for chronic cancer pain (Portenoy, 2000). Morphine attenuation of hyperalgesia seen in this hindpaw model both confirms its usefulness as a model of cancer pain and indicates its susceptibility to opioids. Furthermore, it is notable that the systemic potency of morphine in this chronic tumor pain model (ED50 ~10 mg/kg) is comparable to its potency in the mouse strain using common acute thermal nociceptive tests (ED50 3–15 mg/kg) and lower than its potency in tonic

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**Figure 9.** The ET-A receptor antagonist BQ-123 (0.16–16 nmol/30 µl) injected into the tumor site of hyperalgesic CSH/He mice with fibrosarcoma tumors on PID 10 produced dose-dependent attenuation of mechanical hyperalgesia (45 min post-drug; n ≥ 9; filled squares). Injection of the ET-B receptor antagonist BQ-788 (0.16–48 nmol/30 µl) on PID 12 attenuated mechanical hyperalgesia only at the highest dose tested (45 min post-drug; n ≥ 9; shaded diamonds). Saline, similarly injected into the tumor site of PID 10 and 12 hyperalgesic CSH/He mice, was inactive (n = 20; open circles). Data are presented as mean and SEM, analyzed by ANOVA, and further tested for differences from saline–vehicle control (•) with the Fisher’s post hoc test; • indicates statistical significance; p < 0.05.
chemical nociceptive tests (ED_{50} ≤ 1 mg/kg) (Jacob et al., 1983; Elmer et al., 1998). This relatively low potency echoes the low potency observed in clinical cancer pain therapy (Portenoy, 2000). In the femur tumor model in which NCTC 2472 cells are injected into the medullary cavity (Clohisy et al., 1996), morphine attenuates the secondary mechanical hyperalgesia (measured on the plantar surface of the ipsilateral hindpaw) with similar potency and efficacy (Honore et al., 2000b; Wacnik et al., 2000). In contrast, implantation of fibrosarcoma cells bilaterally into the humeri induces a movement-related hyperalgesia that is only partially attenuated by morphine (P. W. Wacnik, L. J. Kehl, and G. L. Wilcox, unpublished observations). The combined application of these models would undoubtedly be useful for further characterizing the effectiveness of opiate analgesics in cancer pain but more importantly for the development of other nonopioid analgesics, such as ET receptor antagonists, based on novel mechanistic knowledge of cancer pain gained through the application of these and future animal models.

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