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Intrathecal Morphine Tolerance: Use of Intrathecal Clonidine, DADLE, and Intraventricular Morphine

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Continuous intrathecal morphine infusion provides a new approach to provide analgesia for the terminally ill patient in pain. Unfortunately, tolerance to morphine given intrathecally may predate death from the tumor. With such tolerance, the pain again can become unbearable unless permanent neurolysis or neuroablative surgery succeeds. 1

One approach to overcoming the tolerance is to "rest" the mu opioid receptors while activating alternate receptor systems known to mediate antinociception. Yaksh et al. 2 demonstrated both adrenergic and serotonergic descending inhibitory systems in the spinal cord. Activation of postsynaptic adrenergic receptors at the spinal level with either the lipophilic alpha agonist clonidine or its more polar analogue ST-91 yields a potent and prolonged antinociception as measured by the shock titration paradigm in monkeys. 3 Alternatively, activation of delta opioid receptors in the dorsal horn produces analgesia in various laboratory models. 4–6 A third approach to overcoming tolerance to narcotics given intrathecally is to administer morphine into the cerebral ventricles (ICV). Theoretically, this might increase morphine availability at opiate receptors modulating pain at higher levels of the cerebrospinal axis. Lobato et al. 7 and Leavens et al. 8 reported analgesia in humans with this route of morphine administration.

We report temporary control of cancer pain during chronic intrathecal infusion of clonidine in a patient with metastatic rectal adenocarcinoma, already tolerant to high doses of morphine administered systemically and intrathecally. We also report lack of analgesia following intrathecal administration of the specific delta agonist Da La 2 D-Leu 5 enkephalin (DADLE). Additionally, the ineffectiveness of high-dose ICV morphine was noted in this same patient. Finally, 3 months after intrathecal neurolysis, spinal opioid receptor sensitivity to low-dose intrathecal morphine reemerged.

Materials and Case Report

Drug Delivery System. The drug delivery system consisted of a subcutaneously implanted Infusaid® Model 400 reservoir pump (Infusaid Corp., Norwood, Massachusetts) with a constant flow rate (4 ml per day) connected in series with an L1–2 intrathecal Silastic® catheter. Detailed descriptions of this system are reported elsewhere. 9 Drug dosages were varied by changing the concentration of the drugs in the pump reservoir. If needed, bolus intrathecal doses were given via the auxiliary port of the reservoir.

Prior to human use, compatibility and stability of clonidine HCl were tested in vitro in a reservoir maintained at 37°C in a water bath for 56 days via serial
measurement of clonidine HCl levels and resultant flow rates (Data on file at Infusaid Corp., Norwood, Massachusetts). Clonidine concentrations were measured by a double isotope radioimmunoassay. Clonidine concentrations were observed to fluctuate less than 12% over the 56 days. Liability of serious neurotoxicity resulting from intraspinal clonidine was precluded by investigations in sheep prior to obtaining an investigational new drug application (IND), Institutional Review Board approval, or initiating clinical studies.\textsuperscript{††} DADLE was prepared in a saline solution from lyophilized powder (Peninsula Laboratories, Palo Alto, California) and sterilized by passage through a 0.22-μm filter. Once in solution, the DADLE was brought to the patient’s bedside and injected immediately through a 22-gauge lumbar spinal needle. Administration of drugs was performed under a protocol approved by the institutional review board; both patient and daughter gave written and oral informed consent.

This patient had originally met our study criteria for continuous intraspinal morphine infusion via an implantable reservoir device in November 1982.\textsuperscript{11} Demonstration of tolerance to high-dose intrathecal morphine (15 mg intrathecally daily) was required prior to admission to the bolus intrathecal clonidine study. Significant psychologic depression was excluded by clinical examination and a Zung self-rating depression score <0.65.\textsuperscript{12} A minimum acceptable therapeutic response to bolus intrathecal clonidine was required prior to initiation of continuous infusion. Minimal response was defined as at least a 50% reduction from the patient’s preclonidine pain report on visual pain analogue scale scores (VPASS),\textsuperscript{13} with the decrease sustained for at least 8 h.

\textbf{Patient 1.} In May 1979, the patient, a 52-year-old obese woman, underwent an abdominal perineal resection of a 2.0 × 1.5 cm Duke’s B1 adenocarcinoma. Postoperative course was complicated by a perineal body abscess and sigmoid colostomy stricture; and eventually constant left-sided sacral pain prevented her return to a desk job. Over the next 2 years, this pain was complicated by perineal body suture granuloma and flares of degenerative joint disease, primarily involving lumbosacral and knee joints. Neurologic evaluation in March 1981 demonstrated a polysegmental postoperative sacral neuropathy. During September 1981, intermittent urinary retention appeared. Evaluation for recurrent disease and bilateral leg and back pain demonstrated a presacral mass that was irradiated with 5,000 rads. Her daily analgesic intake increased to 20 mg po methadone. In February 1982, recurrent pain and sympathetic dystrophy in the extremities led to an additional 2,000 rads of pelvic irradiation and cesium inserts.

In November 1982, the pain was uncontrolled by 80 mg po methadone daily, and an Infusaid Pump\textsuperscript{®} was implanted for continuous infusion of intrathecal morphine. Pain control initially was achieved with 2 mg intrathecal morphine per day and prn po acetaminophen and oxycodone. Methadone withdrawal was averted with a 2-week course of oral clonidine.\textsuperscript{14} Two subsequent courses of cis-platinum,


\textbf{Figs. 1.} Combined intrathecal morphine and clonidine infusion over the initial 45-day trial with corresponding po methadone requirements. The corresponding analgesic response is shown in figure 2. The patient’s hydromorphone requests were minimal but are graphed as methadone equivalents (7.5 mg po hydromorphone = 20 mg po methadone).

BCNU, and DTIC, with weekly 5-FU had no effect on tumor size. Examination in July 1983 revealed sacral invasion by tumor including the caudal dural sac. Intrathecal morphine doses gradually were increased and methadone re instituted and increased until the combination of 30 mg intrathecal morphine with 120 mg po methadone daily failed to control her pain. She was bedridden and in constant pain for the week prior to readmission.

In late September 1983, the patient was placed on a double-blind protocol for bolus infusion of intrathecal clonidine. A 0.3 mg intrathecal clonidine bolus yielded more than 18 h analgesia, characterized by greater than 50% reduction in pain report (VPASS). In contrast, placebo (2 ml intrathecal normal saline) yielded no pain relief, while a 0.15 mg bolus of clonidine yielded intermediate relief. Side effects included 3–4 h of sedation and transient hypotension to 70 mmHg systolic, easily controlled with IV metaraminol injection. Bradycardia (48 bpm) responded to a single IV dose of 0.5 mg atropine. Thus, a trial of combined intrathecal infusion of 0.3 mg/day clonidine with 2 mg/day morphine was instituted. This was supplemented with a reduced po methadone maintenance schedule of 10 mg/day and by 2 mg po hydromorphone prn. Intrathecal and po analgesic use during this trial is shown in figure 1, while the resultant analgesia as measured by mean daily visual pain analogue scores during the 45 days on this regimen is shown in figure 2. On this regimen she again was able to ambulate. Despite good analgesia, orthostatic hypotension (systolic blood pressure 76–98 mmHg) unresponsive to ergonovine,\textsuperscript{15} phentolamine, and po ephedrine led us to taper the clonidine dose by day 27 (intrathecal clonidine was discontinued for two days). A concomitant escalation of narcotic requirement occurred. Pelvic computed axial tomography (CAT) scan revealed further marked increase in pelvic tumor size with intraspinal tumor extending to L3. Reinsti tution on day 30 of 0.2 mg/day intrathecal clonidine infusion failed to control pain, but
Intrathecal morphine substantially enhances the analgesic effect of systemic opiates.\textsuperscript{16,17} The magnitude of this effect suggests a multiplicative interaction between the spinal and supraspinal opiate receptor systems.\textsuperscript{18} However, as with tolerance in the centrally mediated opiate system, tolerance also develops in the spinal system.\textsuperscript{19,20} In terms of cancer pain control, intraspinal morphine is a useful adjunct with obvious limitations once increasingly high doses are ineffective.\textsuperscript{19,20}

Intrathecal morphine initially produces analgesia characterized by tachyphylaxis via its action primarily upon spinal mu receptors. However, at least two spinal opiate receptor populations, so-called mu and delta receptors, are believed to modulate nociceptive pain transmission in the dorsal horn via specific receptor agonists.\textsuperscript{5} Thus, at least two possible ways exist to circumvent the limitation imposed by spinal narcotic tolerance. Both attempt to put the mu receptors to “rest.” In the first case, analgesia is provided by intermittent or continuous regional anesthesia until the opiate receptors recover opiate sensitivity.\textsuperscript{21,22} Yaksh observed recovery in opiate-tolerant animals with 2 weeks opiate abstinence.\textsuperscript{23} Logistically, this method requires long periods of hospitalization with close monitoring and, in our experience, does not assure complete recovery of the mu receptor-mediated analgesia. Second, analgesia could be achieved through stimulation of an alternate nontolerant spinal receptor population subserving antinoception.

Yaksh and Reddy successfully have demonstrated the intrathecal antinoceptive potential of the alpha agonist clonidine in macaque monkeys.\textsuperscript{2} Convincing evidence exists that this effect is mediated through a post-synaptic receptor of a descending inhibitory adrenergic neuron\textsuperscript{2,24-27} with synapses in the dorsal horn. In contrast to morphine, clonidine analgesia is only mediated spinal,\textsuperscript{28} since it is not influenced by high thoracic spinal cord transection. This clonidine mediated analgesia may be achieved best by intraspinal administration, since at brain stem levels the net effect of such alpha agonists may be antalgic.\textsuperscript{17,26,28,29}

In this patient we observed a marked analgesic effect during initial intrathecal clonidine infusion documented by decreased pain reports and increased mobility, despite a marked reduction in oral narcotic use and a tenfold
reduction in intrathecal morphine use (fig. 1). Analgesia and decreased narcotic use were sustained until clonidine was tapered. During intrathecal clonidine reinstitution, however, analgesia was not obtained at the previously effective morphine and clonidine doses. This may represent adrenergic-opiate cross tolerance. However, differentiation between narcotic tolerance and increasing pain from tumor progression was difficult in part because substantially higher concentrations of an intrathecal clonidine preparation were not available for use. Alternative explanations for failure of the previously effective clonidine–morphine combination exist. Higher doses of analgesics often are needed to overcome pain that is out of control. Also, abandonment of clonidine may have induced a psychologic “setback.” The earlier increased mobility of the patient may have increased painful afferent input. Narcotic withdrawal is an unlikely explanation, since no classic symptoms were observed and both methadone and intrathecal morphine later were stopped acutely following neurolysis without withdrawal symptoms.

Speculatively, this patient might have benefited from higher concentrations of intrathecal clonidine. Yaksh et al., using shock titration in monkeys tolerant to intrathecal morphine injections, observed no waning in analgesia with a daily intrathecal bolus of both clonidine and morphine through 21 days of study. In contrast, Paalzow noted that rats made tolerant to intrathecal clonidine demonstrated no analgesic response to intrathecal morphine. From this it is logical to conclude that the spinal alpha receptors are “downstream” from the opiate receptor; further, a similar biochemical common denominator effectively must yield tolerance to agonists of both receptors, if tolerance to the alpha agonist is present. We had hoped that intrathecal clonidine might allow the mu receptors to “rest” long enough to recover sensitivity to low-dose intrathecal morphine. Though a substantial reduction in oral and intrathecal narcotic was accomplished temporarily, complete opiate abstinence was not achieved, while subsequent narcotic escalation was rapid. To truly test this hypothesis in future studies, intrathecal morphine will have to be halted prior to intrathecal clonidine initiation.

During studies of bolus clonidine infusion in our patient, hypotension and bradycardia were pronounced and required alpha-adrenergic and chronotropic support. During infusion of lower daily doses (0.3–0.4 mg/day) of the drug, resting bradycardia and hypotension were noted but were absent in the higher dose range (0.4–0.6 mg/day). This hypotension is consistent with the observations of Yasuoka and Yaksh in cats given bolus intrathecal clonidine. Stable blood pressure at higher infusion rates perhaps results from peripheral alpha receptor effects of clonidine, a partial agonist at both presynaptic and postsynaptic alpha receptors. At higher intrathecal doses increasing concentrations of clonidine may gain access to the systemic circulation, with vasoconstriction predominating over presynaptic adrenergic effects; a similar phenomenon was observed by one of the authors (DWC) during chronic intraspinal clonidine infusions in sheep. An alternative to adrenergic receptor mediated antinociception in the spinal cord is delta opiate receptor mediated analgesia. Several authors have reported lack of cross-tolerance between spinal mu and delta opiate receptors, suggesting each receptor system, in the presence of specific agonists, might exert an independent analgesic effect. Therefore, we gave intrathecal DADLE, a specific delta agonist, at a dose exceeding that shown to be analgesic in both animals and humans. We observed no analgesia. One explanation for this finding is partial receptor cross-tolerance between DADELLE and morphine. Teng has shown that rats tolerant to intrathecal morphine (tail-flick latency test) demonstrate no cross-tolerance with intrathecal DADLE. However, rats tolerant to intrathecal DADLE are also tolerant to intrathecal morphine. Therefore, DADLE also must activate the mu receptor. In our patient the degree and duration of mu receptor activation by morphine was of a greater magnitude than in these animal models. Further, the patient also had received intrathecal clonidine. Unfortunately, absence of analgesia with intrathecal DADLE in the setting of advanced tolerance to intrathecal morphine and concomitant clonidine suggests that exposure to clonidine may render both DADLE and morphine ineffective as spinal analgesics. The clinical utility of chronically alternating mu, delta, and adrenergic spinal receptor agonists thus becomes suspect. An alternative explanation for lack of analgesia, namely, that DADLE did not gain access to the delta receptor population in the spinal cord, seems unlikely given the period of observation.

Lobato et al. have advocated ICV narcotic administration as a means of overcoming po and intrathecal narcotic tolerance. In their series of patients, 0.5–2 mg ICV morphine, given up to once every 3 h, provided marked analgesia. The bolus dosage of intraventricular morphine administered to our patient exceeded the highest daily ICV morphine dose (16 mg) in the Lobato et al. series. We noted no observable analgesia, respiratory suppression, sedation, or seizure activity. Lobato et al. did not define prior narcotic use in their report; thus, it is tempting to conclude that marked opiate tolerance did not exist in their patients. The same is

likely true of the successful results in the series reported by Leavens et al.8 The tolerance observed in our patient perhaps approaches record levels; presumably, to achieve similar CSF morphine concentrations to that resulting during delivery of 75 mg/day intrathecal morphine and 17.5 mg ICV morphine truly would require astronomic parenetal morphine doses. Under conditions in which both high-dose methadone and intrathecal morphine have been administered, intuitively, a point must be reached when no multiplicative interaction can be obtained between spinal and supraspinal opiate receptors.17

There is a natural reluctance to use percutaneously exposed intrathecal catheters for fear of infection. In dealing with pain control in terminal phases of malignancy, however, the hazard of meningitis may be of less importance than the expense and inconvenience of implanted systems. Under such circumstances, a chronic lumbar CSF drain may offer a better conduit for analgesic delivery, even when local anesthetic delivery is to be part of the plan. Though the limits of such an approach are unknown, perhaps this route should be considered more frequently for terminal therapy of short or intermediate duration.

In summary, in this complex case we observed continuous intrathecal clonidine infusion to yield temporary pain control, despite advanced opiate tolerance. Limited availability of clonidine led to a trial of both intrathecal DADLE and high-dose ICV morphine; both failed to decrease pain. Following temporary pain control with intrathecal neurolytics (3 months), responsiveness to intrathecal morphine returned. This case suggests that cross-tolerance exists between spinal opiate and adrenergic systems after chronic exposure. Further study of intraspinal delta and alpha-adrenergic agonists as pain relievers are underway to define their analgesic role in this and other settings.

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Extracorporeal Circulation in a Patient with Heparin-induced Thrombocytopenia

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Heparin-induced thrombocytopenia may occur after two to 15 days of heparin therapy. It has been reported after iv and subcutaneous administration with various heparin preparations and is not dose dependent. Effective management requires cessation of heparin therapy, administration of platelet antiaggregating drugs, and, when indicated, the use of another form of anticoagulation. When patients with heparin-induced thrombocytopenia are rechallenged with heparin within 12 months of the initial event, recurrent thrombocytopenia and/or thrombosis may occur.1–3 Alternatively, reinstitution of heparin therapy within 4 months after the initial event has been performed without a decrease in platelets.3 We report a patient with heparin-induced thrombocy-

topenia, documented by aggregation testing,4 who required myocardial revascularization. The patient received heparin during coronary artery bypass 14 days after the initial thrombocytopenia had resolved and while aggregation tests indicated the persistence of a heparin-induced antiplatelet antibody.

REPORT OF A CASE

A 57-year-old man was referred for evaluation of unstable angina pectoris. On admission he was given beta-adrenergic receptor blockers, calcium channel blockers, nitrates, and prophylactic subcutaneous beef lung heparin (Upjohn) 5,000 units every 8 h. Platelet count on admission was 300,000/mm3. Prothrombin time and partial thromboplastin time were normal, and there was no splenomegaly. On the fourth hospital day, cardiac catheterization documented severe triple-vessel coronary artery disease. Subcutaneous heparin therapy was continued.

On the eighth hospital day, an acute inferior myocardial infarction was documented, intracoronary balloon pump (IABP) assist was begun, and a temporary transvenous pacemaker inserted to manage an associated atrioventricular conduction abnormality. Heparin anticoagulation was increased with beef lung heparin (Upjohn), 7,600 units iv, followed by a continuous iv infusion of 600 units of heparin per hour. The infusion rate subsequently was increased to 1,000 units per hour due to inadequate anticoagulation. A platelet count of 6.5 x 10^11 after starting iv heparin was 56,000/mm3, and heparin was discontinued. Warfarin then was used for anticoagulation. The platelet count decreased to 32,000/mm3 14 h after the heparin was discontinued. During this time the fibrinogen was 404 mg/dl, and fibrin degradation products were increased at 50 units. Platelet aggregation testing was positive with both beef and porcine heparin. Heparin was deleted from all solutions used to irrigate vascular cannuli.