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Fig. 1. T1-weighted sagittal midline magnetic resonance images show the spinal column of a 38-yr-old, 170-cm, 67-kg man (top) and a 19-yr-old, 155-cm, 54-kg woman (bottom). The examinations were performed in the volunteers in the supine position with legs extended. The dural sac lies within the spinal canal and terminates at S2. The level of termination of the spinal cord is seen at L1–L2. In both images, L3 is located at a position higher than L3, whereas T8 is located at a position lower than T6.

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EMLA Cream in the Treatment of Causalgie Pain

To the Editor:—The management of causalgie pain includes sympathetic blockade, transcutaneous electrical nerve stimulation (TENS),1 vasodilators,2 carbamazepine,3 calcitonin,4 and intravenous local anesthetics.5 Some success also has been reported with topical creams such as prostaglandin E1 and 50% dimethylsulphoxide, a free radical scavenger.6 EMLA cream (a 1:1 oil and water eutectic mixture of 2.5% lidocaine and 2.5% prilocaine)7 has been used successfully in the management of herpetic neuralgia.8,9 We would like to report

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A patient in whom EMLA cream produced marked and sustained alleviation of severe sciatic nerve causalgic pain.

After a parachuting accident, a 24-yr-old man, who sustained multiple lower limb and pelvic fractures, required internal fixation of the right femur. An epidural infusion of bupivacaine and fentanyl was commenced at the time of operation, and excellent analgesia was achieved in the postoperative period. After removal of the catheter on day 7, weakness of the right extensor hallucis longus and absent ankle dorsiflexion was evident. The patient also complained of constant and severe burning pain of the dorsum of the foot, with excruciating pain on light touch. Clinical examination showed hyperesthesia and allodynia in an L5 dermatomal distribution, which was attributed to sciatic nerve injury at the greater sciatic notch due to fracture of the iliac wing. Increasing doses of parenteral opioids and a course of carbamazepine failed to achieve pain control.

Pain clinic review was requested on day 20. Because of suspicion that the first interdigital cleft was spared, a superficial peroneal nerve block was performed. Pain in the dorsum of the foot was controlled and intense interdigital hyperesthesia unmasked, confirming the L5 pattern. A second epidural infusion was commenced, supplemented by TENS and 75 mg dextepine. Satisfactory analgesia was achieved, and the infusion was gradually decreased and stopped after 7 days. Causalalgic pain initially was much improved and the catheter was removed. Severe nocturnal spasms of hyperesthesia recurred 24 h after catheter removal, and pain, assessed using a visual analog 10-point scale, increased from 4 to 10. Treatment options, including a third epidural, chemical sympathectomy, and intravenous regional anesthesia, were declined by the patient. It was decided to trial EMLA cream to the affected area.

The cream (5 g) was thickly applied for 18 h/day, between 6 pm and midnight. Dramatic relief was obtained after the first application with a decrease in pain score from 8 to 4. The patient stopped TENS after 24 h and dextepine after 48 h. After 5 days, he mobilized the limb for the first time and, although altered sensation was evident in the absence of EMLA cream, causalalgic pain had ceased and light touch was easily tolerated. He was discharged 6 days after commencing EMLA cream. At the 2-week followup, there was no recurrence of causalalgic symptoms, and the patient had ceased to use EMLA cream after 8 days. There were no trophic or residual vascular changes in skin at this time.

The characteristic spontaneous burning pain, allodynia, and hyperesthesia of causalalgic pain was described by de Takats in 1945. The association with pelvic fracture is well recognized. The modern theory about the pathogenesis of causalalgic pain includes both peripheral and central causes. Peripheral causes include spontaneous discharge from neural sprouts, with an increase in their sensitivity to adrenergic compounds and increased firing in dorsal root ganglia. Central causes include spontaneous activity of deafferented nerves in the dorsal horn and development of responses to new receptive fields. If untreated, the abnormality becomes self-sustaining and independent of peripheral input. The self-perpetuating cycle of pain can be broken more easily in the early phase by sympathetic blockade achieved by epidural or subarachnoid anesthesia, lumbar sympathectomy, or a guanethidine block. Although not measured, it is unlikely that circulating levels of local anesthetic were responsible for the patient's improvement. Stowe et al. reported that plasma concentrations of lidocaine and prilocaine were well below toxic levels when 5–10 g was applied for 24 h in patients with postherpetic neuralgia. Methemoglobinemia was not noted in our patient. It is possible that the EMLA cream may help break the cycle of pain by either local sympathetic inhibition or preferential binding to traumatized tissue. The use of EMLA cream in the management of causalalgic pain is worth further investigation.

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