Intractable pain treated with intrathecal isotonic iced saline

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SUMMARY Hitchcock’s original method of hypothermic subarachnoid irrigation employed both temperature and osmolarity. Spinal cooling was then abandoned in favour of intrathecal injection of normothermic hypertonic saline. Modifications of the procedure that followed have continued to accept hyperosmolarity as the factor causing pain relief. Fifty patients were treated by a technique evolved to enhance the effect of hypothermia while avoiding the complications associated with hyperosmolar solutions. For the cases of terminal carcinoma and others considered to be poor surgical risks, the results have been quite satisfactory. For non-neoplastic painful syndromes, rapid perfusion cooling of the subarachnoid space offers an alternative therapeutic approach.

The effects of hypothermia on the nervous system (Denny-Brown, Adams, and Brenner, 1945) and perfusion of the subarachnoid space (Albin, White, and Donald, 1961) have been studied for many years. Hitchcock (1967a, b) introduced subarachnoid irrigation with cold solutions for the relief of intractable pain and subsequently inquired into the hyperosmolarity of his solution. Ensuing clinical research explored the use of anaesthesia (Nicholson and Roberts, 1968; Collins, Juras, and Van Houten, 1969), varying sites of injection (Hitchcock, 1969; Mathews, Ambruso, and Osterholm, 1970), prolonged irrigation (Tsubokawa, 1969a), increasingly hypertonic saline (Hitchcock, 1970), and elaborate perfusion apparatus (Negrin, 1970). To our knowledge, no previous investigators have reported the primary results of brief intrathecal injection of isotonic iced saline.

METHOD

The irrigating solution was prepared by keeping 500 ml. pour bottles of sodium chloride injection (U.S.P.) in the freezer of an ordinary refrigerator until ice cubes formed in a separate tray. During the procedure, a small bucket filled with crushed ice maintained the temperature in the bottle between 0° and −1° C ('hypothermic solution'). Analysis of our iced saline failed to reveal any increase in osmolality: 308 m-osmol/l., consisting of 154 m-equiv/l. sodium and 154 m-equiv/l. chloride.

The patient was placed on a tilting table in the lateral position for lumbar tap with a 16 gauge spinal needle. Cerebrospinal fluid was allowed to flow out spontaneously until subambient pressure was produced. Mechanical withdrawal by syringe was specifically avoided to prevent aspiration of a nerve root or alteration of the needle tip within the subarachnoid space.

When the pain was unilateral, the patient remained with the affected side down. When the pain was bilateral or midline, the patient was placed in the prone position. The table was then tilted from −60° to +60°, so that the appropriate area could be adequately irrigated.

A second 16 gauge needle was intermittently attached to a 12 ml. syringe for sterile aspiration of the hypothermic solution from the supply bottle. Repetitive injections were made as rapidly as possible to produce maximum cumulative cooling. Between injections, the spinal needle was left open; and the free back-flow allowed for continuous evaluation of needle placement as well as prevention of pressure build-up. After the final 12 ml. injection, back-flow was not permitted, and the needle was removed.

With the use only of local injection of lidocaine, there was severe experience of paraesthesias and fasciculations which interfered with rapid, repetitive irrigation and dissuaded patients from allowing subsequent cooling. A combination of fentanyl 0.1–
0.2 mg and droperidol 5–10 mg, followed by either sodium thiamylal 200–400 mg or diazepam 20–40 mg, given intravenously just before the period of actual perfusion, was found to effectively reduce the discomfort while allowing evaluation of regional muscular contractions. At this light plane of anaesthesia, a total of 72 ml iced saline could regularly be employed for lower quadrant pain, and 96 ml for upper quadrant pain.

Fasciculations were limited to the period of actual cooling, while dysesthesias continued in some of our patients for up to two hours and not infrequently required additional medication. Hypaesthesia was often noted in the affected area for periods of up to 48 hours. Headache and local discomfort from spinal tap occasionally necessitated administration of minor analgesics. Four patients with preexisting weakness of one extremity, secondary in two cases to known myelopathy and in two cases to suspected radiculopathy, experienced a transient increase in paresis for several days.

If the relief from pain was incomplete or less than 48 hours in duration, the entire procedure was repeated. If multiple areas of the body were involved with diffuse metastases, different levels were irritated serially. In two patients the entire spinal cord was cooled, by three separate procedures in one and four separate procedures in the other.

RESULTS

Eighty subarachnoid irrigations with isotonic iced saline were performed on 50 patients: 33 with metastatic neoplasm, two with causalgia, four with neuralgia, five with low back pain persisting after laminectomy for herniated nucleus pulposus and six miscellaneous painful syndromes. In effect, 36 cases of lower quadrant pain and 18 cases of upper quadrant pain were treated during a one year period. Eleven patients with carcinoma obtained complete relief from intractable pain until death eight days to eight months later; 10 others had almost total relief for eight weeks before return of the original pain, and the remaining 12 had an equivocal result. Both patients with causalgia were free of symptoms for one month before the return of some discomfort in the affected area; neither showed preference for repeat spinal cooling. Essentially no relief was produced in the two patients with post-herpetic neuralgia, the one patient with post-thoracotomy intercostal neuralgia, or the one with post-herniorrhaphy ilioin-

guinal neuralgia. Three patients with intractable low back pain after laminectomy and excision of herniated nucleus pulposus have improved up to 12 months to date, while the other two had an equivocal result. The six miscellaneous painful syndromes included intractable buttock pain after coecoygectomy, cervical radiculopathy persisting after laminectomy and fusion, superior vena caval syndrome, ulcerative colitis, Buerger’s disease, and phantom limb; three obtained partial relief, and three were unimproved in a one year follow-up.

No permanent sensory deficits, bladder dysfunctions, or other complications directly attributable to hypothermia have been observed in our series of 80 spinal cord coolings. Nevertheless, complications have been reported by other researchers, especially with the use of hypertonic saline. Transient changes in blood pressure (Hitchcock, 1970) and cardiac arrhythmias (Hitchcock and McKean, 1968) have been attributed to direct stimulation of sympathetic nerve fibres by lumbar injection or of parasympathetic nerve fibres by cisternal injection. O’Higgins, Padfield, and Clapp (1970) published an account of one complication of infarction in the territory of the left middle cerebral artery. Hitchcock and McKean (1968) described one permanent cauda equina lesion.

DISCUSSION

Considerable evidence has been presented in favour of the concept that sense organs of the skin with unmyelinated C nerve fibres have widely varying functional characteristics (Christensen and Perl, 1969; Bessou and Perl, 1970; Kumazawa, Perl, Burgess, and Whitehorn, 1971). In the mammal, a large proportion of such sensory units were excited by gentle mechanical stimulation and by transient cooling; a second group responded best to small temperature changes; and a third group had elevated thresholds for various or all stimuli. The intensity of the liminal stimulus and unique ability to provide signals marking the difference between innocuous, threatening, or damaging stimuli strongly indicated a nociceptive function for the high-threshold unmyelinated sensory units. The sensitization of polymodal nociceptors after noxious heat simulated certain kinds of hyper-
algesia. Furthermore, the pericorneal cells of Rexed’s lamina 1 could be fired antidromically by stimulation of the opposite anterior lateral funiculus in the cervical area, demonstrating the absence of internuncial cells and the dispensability of the modulation concept inherent in the gate control theory (Melzack and Wall, 1965).

Studies on the effects of temperature on neural conduction and the relative susceptibility of myelinated and unmyelinated nerve fibres have been contradictory. Von Euler (1947) suggested that unmyelinated C fibres were more susceptible to cooling than thickly myelinated fibres, which might even be stimulated. Lundberg (1948) reported conversely that the conduction of A fibres ceased at 7°C to 15°C in the cat, whereas conduction of splenic C fibres continued at 0°C C in the cat and cow. Albin, White, and Locke (1967) confirmed the earlier observations of Brooks, Koizumi, and Malcolm (1955) and Suda, Koizumi, and Brooks (1957) that the action of cold on nerve conduction was opposite to that of local anaesthetic agents—the larger the cross-sectional diameter of the fibre, the more it was sensitive to hypothermia.

Laboratory research into the mechanism of relief of pain has been extensive but inconclusive; no animal model yet designed adequately simulates the complex human experience of intractable pain. The effects which have been documented are either transient phenomena or the results of extremes of temperature (Denny-Brown et al., 1945) and osmolarity (Robertson, 1958; Jewett and King, 1971) not attained in the clinical situation. Only Tsubokawa (1969b) has made a comparative study of conduction block of A-delta and C fibres in the cat with appropriate cooling and without appreciable alteration of osmolarity: six hours after a 15 minute irrigation at spinal level L6–7 with 80 ml iced saline (314 to 376 m-osmol/l.), action potentials did not reappear in the A-delta or C fibres, while A-alpha and A-beta fibres were essentially unchanged; six days after two separate spinal coolings, 60% of the prepared animals failed to exhibit evoked potentials in the mesencephalic reticular formation or the centromedian nucleus of the thalamus after noxious stimulation of the sciatic nerve.

Nerve conduction studies were performed on a limited number of our patients. No change was demonstrated in the conduction velocity of normal peripheral nerves immediately after brief perfusion cooling of the subarachnoid space or at intervals up to six months. In one patient with radiculopathies involving spinal roots C6, 7, and 8 the right ulnar nerve tests were within normal limits, while the left ulnar nerve exhibited slowed conduction; after two separate irrigations, the left ulnar nerve remained electrically silent to testing.

Postmortem examination of one patient in our series failed to reveal any demonstrable neurolysis of the spinal cord or nerve roots. Mathews et al. (1970) also reported no histological changes in the medulla, spinal cord, or nerve roots of one patient who came to necropsy. Hitchcock (1967a), on the other hand, reported widespread vacuolization and pallor of myelin staining in one patient who came to necropsy.

Patients with successful relief of their intractable pain after spinal cord cooling still have appropriate perception of pin prick, skin pinching, and deep pressure and can identify correctly a hot or cold stimulus. These observations indicate that at least the crude reaction to noxious stimuli or temperature is present. Careful testing for threshold differences has not been carried out, yet the alleviation of intractable pain is in clear contrast to the apparent preservation of response to painful test stimuli.

Initial fasciculations and paraesthesias may be explained by transient activation of the larger A fibres in ventral and dorsal roots; subsequent hypaesthesia may be caused by inactivation of the sodium pump for a limited period of time. Sustained relief of pain may be attributed to cold trauma to a critical number of A-delta and C fibres or to pericorneal cells and their ascending axons. Alternatively, gate modulation could be rendered inoperative by damage to a still undemonstrated anatomical substrate of interneurones. The exact role of spinal cord ischaemia secondary to vasospasm of the intrinsic vasculature remains undetermined (Meyer and Hunter, 1957; Douglas and Ritchie, 1957). Further experiments are necessary to ascertain whether the alleviation of pain is due to deactivation of peripheral fibres or to prolonged quiescence of central internuncial neurones.

The authors are indebted to Dr. Kuo, Department...
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*J Neurol Neurosurg Psychiatry* 1973 36: 417-420
doi: 10.1136/jnnp.36.3.417

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