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## **FAST** TRACK

**Cryotherapy** is better than heat for initial treatment of acute muscle strain

# Does heat or cold work better for acute muscle strain?

### **Evidence-based answer**

Cryotherapy is better than heat for treating acute muscle strain (strength of recommendation [SOR]: C, consensus, usual practice, and expert opinion).

Insufficient patient-oriented evidence exists regarding use of heat to treat acute softtissue injuries.

# Evidence summary

A comprehensive review of the literature revealed no studies that compare heat and cryotherapy to treat acute softtissue injury. Well-designed human trials of general management of acute softtissue injury are rare.1

Cryotherapy has been the recommended initial treatment for muscle strain for more than 30 years, based generally on expert opinion and physiological models, not clinical trials.2 Theoretically, cryotherapy controls hemorrhage and tissue edema, whereas heat enhances the inflammatory response.<sup>2</sup>

## One human RCT and animal studies find benefits from cold

A 2007 review evaluated 66 publications and found only 1 randomized controlled trial conducted on humans.3 The intervention in this trial involved applying cold gel 4 times a day for the first 14 days after the injury. The control group received a room-temperature gel application; neither group was aware of the temperature differential.

The study found significant reduction in pain at rest, pain with movement, and functional disability at intervals of 7, 14,

and 28 days postinjury (P<.001) among patients receiving cold-gel applications. Patients receiving cold-gel treatment also reported increased satisfaction with treatment compared with the controls. At 28 days, cold-gel treatment patients scored 71 on a 100-point satisfaction scale compared with 44 for controls (P<.001).3 Inconclusive results or significant design flaws limited the validity of all other trials cited in this review.3

> Laboratory studies on rats have also demonstrated beneficial effects of cryotherapy after simulated soft-tissue injuries.4,5 One study cited a significant reduction in inflammatory cells, based on histologic examination, in 43 rats between 6 and 24 hours after trauma.4 A second study of 21 rats showed improvement in associated physiological components with cryotherapy, but no statistically significant improvement in edema.<sup>5</sup>

#### How cold is too cold?

Most authorities recommend empiric treatment with cryotherapy during the acute inflammatory phase—the first 24 to 48 hours after injury.6 Although not rigorously studied, some sources recommend applying cold to the involved muscle for NSAIDS, Aspirin, and Warfarin)- Serotonin release by platelets plays an important role in hemostasis. Epidemiological studies of case-control and cohort design have demonstrated an association between use of psychotropic drugs that interfere with serotonin reuptake and the occurrence of upper gastrointestinal bleeding. These studies have also shown that concurrent use of an NSAID or aspirin may potentiate this risk of bleeding. Altered anticoagulant effects, including increased bleeding, have been reported when SSRIs and SNRIs are coadministered with warfarin, Patients receiving warfarin therapy should be carefully monitored when Pristiq is initiated or discontinued. Ethanol - A clinical study has shown that deswenlafaxine does not increase the impairment of mental and motor skills caused by ethanol. However, as with all CNS-active drugs, patients should be advised to avoid alcohol consumption while taking Pristiq. Potential for Other Drugs to Affect Desvenlafaxine-Inhibitors of CYP3A4 (ketoconazole)- CYP3A4 is a minor pathway for the metabolism of Pristiq. Concomitant use of Pristiq with potent inhibitors of CYP3A4 and presult in higher concentrations of Pristiq. Inhibitors of other CYP enzymes. Based on in wirro data, drugs that inhibit CYP isozymes 1A1, 1A2, 2A6, 2D6, 2C6, 2C9, 2C19, and 2E1 are not expected to have significant impact on the pharmacokinetic profile of Pristiq. Potential for Desvenlafaxine to Affect Other Drugs- Drugs metabolized by CYP2D6 (dispramine)- Mro studies showed minimal inhibitory effect of desvenlafaxine on CYP2D6. Clinical studies have shown that drug. Drugs metabolized by CYP3A4 (and path of the CYP3A4 isozyme. Concomitant use of Pristiq with a drug metabolized by CYP2D6 (are result in higher concentrations of Mrd drug. Drugs metabolized by CYP3A4 (and path of the CYP1A2, 2A6, 2C8, 2C9 and 2C19 in witz, desvenlafaxine loses not inhibit CYP1A2, 2A6, 2C8, 2C9 and 2C19 in witz, desvenlafaxine is not a substate or an inhibitor for the P-glycoprotein transporter. The pharmacokinetics

OVERDOSAGE: Human Experience with Overdosage—There is limited clinical experience with desvenlafaxine succinate overdosage in humans. In premarketing clinical studies, no cases of fatal acute overdose desvenlafaxine were reported. The adverse reactions reported within 5 days of an overdose > 600 mg that were possibly related to Pristig included headache, vomiting, agitation, dizziness, nausea, constipation, diarrhea, dry mouth, paresthesia, and tachycardia. Desvenlafaxine (Pristig) is the major active metabolite of venlafaxine. Overdose experience reported with venlafaxine (the parent drug of Pristig) is presented below; the identical information can be found in the \*Overdosage\* section of the venlafaxine package insert. In postmarketing experience, overdose with venlafaxine (the parent drug of Pristig) has occurred predominantly in combination with alcohol and/or other drugs. The most commonly reported events in overdosage include tachycardia, changes in level of consciousness (ranging from somnolence to coma), mydriasis, seizures, and vomiting, Electrocardiogram changes (e.g., prolongation of OT interval, bundle branch block, QRS prolongation), sinus and ventricular tachycardia, bradycardia, hypotension, rhabdomyolysis, vertigo, liver necrosis; serotonin syndrome and death have been reported. Published retrospective studies report that venlafaxine overdosage may be associated with an increased risk of fatal outcomes compared to that observed with SSRI antidepressand products, but lower than that for tricyclic antidepressants. Epidemiological studies have shown that venlafaxine-treated patients have a higher pre-existing burden of suicide risk factors than SSRI-treated patients. The extent to which the finding of an increased risk of fatal outcomes can be attributed to the toxicity of venlafaxine in overdosage, as opposed to some characteristic(s) of venlafaxine-treated patients, is not clear. Prescriptions for Pristig should be written for the smallest quantity of caspulses consistent with good patient manage

This brief summary is based on Pristiq Prescribing Information W10529C002, revised April 2008.

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the first 4 hours after injury at intervals of 10 to 20 minutes every 30 to 60 minutes.<sup>6</sup>

The literature focuses more on the optimal temperature for cryotherapy than on the duration and frequency of therapy.<sup>7</sup> Temperatures below 15° to 25°C may actually result in vasodilatation rather than vasoconstriction.<sup>7</sup>

#### **Evidence for heat is limited**

A 2006 Cochrane review that addressed treatment of lower back muscular strain, not soft-tissue injuries in general, found moderate evidence that heat therapy reduces pain by 17% and disability in the acute setting (*P*=.001).<sup>8</sup> The review also cited 2 head-to-head trials that compared heat and cryotherapy; however, the study designs were poor and the results were contradictory.<sup>8</sup>

#### **Recommendations**

Authoritative textbooks consistently recommend applying ice for initial treatment of musculoskeletal and soft-tissue strains.<sup>9</sup>

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