Epidural steroid injections are not effective for patients with lumbar spinal stenosis

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Context
Degenerative lumbar spinal stenosis is a common cause of low back and leg pain in older individuals. The evidence supporting non-surgical treatments, including oral medications, physical treatments and spinal injections, is limited.1 Despite the lack of strong evidence supporting the use of epidural steroid injections (ESIs) for lumbar spinal stenosis, their use has increased dramatically since 2000.2 This multicentre, randomised trial compared the effectiveness of ESIs with lidocaine to epidural injections with lidocaine only, with potential for a repeat study injection at 3 weeks. Twenty-six anaesthesiologists, physiatrists and radiologists experienced in epidural injections from 16 US sites performed the procedures. The primary outcomes assessed at 6 weeks were an average pain rating for buttock, hip or leg pain in the past week and functional impairment measured by the Roland–Morris Disability Questionnaire (RMDQ). Patients, treating physicians and research staff performing baseline and outcome assessments were blinded to treatment assignment.

Methods
Patients were at least 50-years-old with pain and functional impairment attributable to lumbar spinal stenosis of the central canal, with no prior lumbar surgery or ESI within the past 6 months. Four hundred patients were randomly assigned to either a standard ESI (glucocorticoids) with lidocaine, or an injection of lidocaine alone, with potential for a repeat study injection at 3 weeks. Twenty-six anaesthesiologists, physiatrists and radiologists experienced in epidural injections from 16 US sites performed the procedures. The primary outcomes assessed at 6 weeks were an average pain rating for buttock, hip or leg pain in the past week and functional impairment measured by the Roland–Morris Disability Questionnaire (RMDQ). Patients, treating physicians and research staff performing baseline and outcome assessments were blinded to treatment assignment.

Findings
Baseline patient characteristics were similar among groups, and they had the same 6-week follow-up rate of 96.5%. Clinically and statistically significant decreases in pain and RMDQ scores were seen in both groups. There were no significant differences between groups in the average pain (adjusted difference −0.2 points, 95% CI −0.8 to 0.4) or RMDQ scores (adjusted difference −1.0 points; 95% CI −2.1 to 0.1). A prespecified subgroup analysis of interlaminar (n=282) and transformaminal injection revealed a small but statistically significant benefit to RMDQ score for interlaminar injection in the ESI with lidocaine group (adjusted difference −1.4 points; 95% CI −2.8 to −0.1, p<0.05), but no benefit to average pain. There was no difference in primary outcomes for the subgroup who received transformaminal injections. Among secondary outcomes, most showed no group differences at 6 weeks, but there were small, statistically significant improvements in pain and RMDQ in the ESI with lidocaine group at 3 weeks. Patients who received an ESI with lidocaine were more satisfied with their treatment than those who received lidocaine alone (67% vs 54%, p=0.01).

Commentary
This study adds to a growing body of research demonstrating the lack of important benefit of ESIs in patients with lumbar spinal stenosis. Overall, these results raise serious concerns about the overuse of ESIs in patients with symptomatic central canal stenosis. Although this study showed a small, statistically significant effect favouring the ESI group at 3 weeks, this benefit was no longer apparent at 6 weeks. Moreover, this statistical difference among treatment groups at 3 weeks would not be considered a clinically meaningful effect. Other secondary pain and functional status outcomes also showed no significant benefit at 6 weeks.

Should this study change current practice? Patients who remain symptomatic and functionally impaired despite initial empiric therapy with oral medications, physical treatments and time, should be engaged by providers to explore options and risks/benefits of various treatments with a shared decision-making and educational approach.4 Given the findings of short-term benefit, one could argue this could include ESIs.6 As an example, a highly symptomatic patient unable to actively participate in physical therapy due to pain and functional impairment could be considered for a single ESI as a way to engage her/him in exercise-based treatments. However, the lack of longer term benefit and the potential risks of ESIs suggest that using a series of injections to manage patients over time cannot be endorsed in the absence of more compelling evidence.4

This study leaves several unanswered questions. By focusing on individuals with central canal stenosis, it does not address patients undergoing injections for primarily foraminal and lateral recess stenosis. The subgroup analysis suggesting greater effectiveness for interlaminar techniques requires further investigation. Finally, one may question whether the lidocaine only control group represents a true placebo arm. Although it is hard to understand the physiological basis for a short-acting anaesthetic providing prolonged benefit, the current results argue for increased use of saline controls. Regardless of these limitations, the current study provides strong evidence that ESIs for patients with symptoms due to central canal lumbar spinal stenosis do not offer prolonged benefit compared to lidocaine only epidural injections.

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References