Baseline Pain and Disability in the Investigational Vertebralplasty Efficacy and Safety Trial

BACKGROUND AND PURPOSE: Multiple case series of vertebroplasty outcomes have been published, though no large, placebo controlled trial has yet been performed. Our aim was to report baseline characteristics for the Investigational Vertebralplasty Efficacy and Safety Trial (INVEST), a randomized blinded controlled study of vertebroplasty.

MATERIALS AND METHODS: We compared baseline demographics, pain scores, and scores on the modified Roland-Morris Disability Scale (RMDS), a back pain-specific metric, between 2 groups. One group included subjects enrolled at the lead INVEST site (n = 27 to date). The second group consisted of eligible patients seen concurrently at the lead INVEST site, who declined enrollment (n = 70). Comparisons were made by using 2-sample t tests.

RESULTS: Mean ages were similar between groups, averaging approximately 74 years among study participants and 77 years among nonenrolled eligible patients (P = .17). Approximately 75% of subjects were female in both groups. RMDS scores of enrolled patients at the lead site (18.0 ± 4.2) were not statistically different from those of eligible nonenrolled patients at the lead site (18.6 ± 3.6, P = .49). Pain scores in the enrolled subjects were measured as “average intensity over the prior 24 hours” with mean scores of 7.6 ± 2.1 among enrolled patients at the lead site. Pain scores in eligible nonenrolled patients were measured as “pain at rest,” with mean score of 3.4 ± 3.3, and “pain with activity,” with mean score of 8.5 ± 2.0.

CONCLUSIONS: Patient demographics among subjects enrolled in the INVEST are similar to those in a cohort of eligible nonenrolled patients. Back pain-specific disability was similar between subjects enrolled in the INVEST study and eligible nonenrolled patients at the lead site.

Vertebroplasty is a widely applied procedure for palliation of pain from osteoporotic and neoplastic vertebral compression fractures. The procedure has been evaluated in numerous studies, including a large number of retrospective case series and a relatively small number of nonrandomized controlled studies. These controlled studies typically focused on patients who underwent vertebroplasty or conservative management, usually with the potential for crossover in the short term. Only 1 randomized trial comparing vertebroplasty with medical therapy has been published. Almost without exception, these studies found large treatment effects from the procedure, though the treatment effect in the randomized trial was more modest than that in other reports.

There is, to date, no blinded trial comparing vertebroplasty with a sham or placebo intervention. Such a study is needed to understand better the extent to which pain relief following vertebroplasty is due to a true treatment effect of the cement compared with other factors such as regression toward the mean, effects of local anesthesia, or nonspecific treatment effects, including patient and clinician expectations of pain relief.

The Investigational Vertebralplasty Efficacy and Safety Trial (INVEST) is a National Institutes of Health-sponsored international multicenter prospective randomized blinded trial comparing vertebroplasty with a “control intervention.” The control intervention consisted of a simulated vertebroplasty comprising all aspects of the typical vertebroplasty procedure except placement of the biopsy needle and cement. The study was designed to assess the true treatment effect of the cement in the vertebroplasty procedure compared with the other factors listed above.

The results of any clinical trial need to be interpreted in light of the study sample. Patient characteristics may influence response to any treatment and findings from a single clinical trial may or may not generalize to populations of patients with different demographic or clinical characteristics. Furthermore, patients who enroll in a randomized trial of a treatment may differ in important ways from patients who receive the treatment outside a research study. Therefore, the purpose of the present report was to describe the baseline demographic and clinical characteristics of patients who enrolled in the INVEST study and to compare them with those of patients who were eligible for the study but declined to enroll. This information will assist practitioners, researchers, and payers in interpreting vertebroplasty efficacy and safety results in the INVEST study when they are available.

Materials and Methods

Study Inclusion and Exclusion. Protocol details for INVEST are published in detail elsewhere. Institutional review board approval was obtained for this study, and all participants provided informed consent. Eligible patients are those with ≤3 osteoporotic compression fractures of <1-year duration. MR imaging or bone scanning was used to characterize fractures in cases without serial plain radiographs.
to document fracture chronicity. Patients with fractures from neoplasms, including multiple myeloma and metastases, were excluded as were patients maintained on anticoagulation.

**Procedural Details.** Subjects were randomized 1:1 to either a regular vertebroplasty or a “control intervention.” Vertebroplasty was performed in standard fashion, typically with unipediculate needle placement and deposition of barium-opacified polymethylmethacrylate, following skin and subcutaneous tissue infiltration with 1% lidocaine and infiltration of the periosteum of the target pedicle or pedicles with 0.25% bupivicaine. The control intervention comprised skin and subcutaneous tissue infiltration with 1% lidocaine and infiltration of the periosteum of the target pedicle or pedicles with 0.25% bupivicaine, similar to that administered during routine vertebroplasty. Pressure was applied to the back to simulate manipulation from vertebroplasty needles, and the methacrylate monomer was opened in the procedure room to simulate a vertebroplasty.

**Follow-Up.** Subjects and study coordinators performing follow-up interviews remained blinded to the procedure type. Subjects completed the study measures at predetermined time points up to 1 year. The primary outcomes were a numeric rating of pain on a scale from zero = no pain to 10 = pain as bad as could be and the modified Roland-Morris Disability Questionnaire, a measure of back pain–related physical disability at 30 days. Subjects were allowed to cross over to the other procedure after 30 days but remained blinded to procedure type for 1 year.

**Study Groups.** In the current report, we describe baseline characteristics of INVEST participants enrolled to date at the lead site. To better understand how eligible patients who enrolled versus those who did not enroll might differ, we compared the subgroup of patients who enrolled at the lead site with patients at the lead site who were eligible but who did not enroll. Such data about eligible nonenrolled patients were not available at the other sites. Because we did not perform prospective data collection in nonenrolled patients in the exact same fashion as was done for INVEST, data were typically limited to pain severity and scores on the modified Roland-Morris Disability Scale (RMDS).

At each study site, enrolled patients were interviewed with study case report forms by a research coordinator. At the lead site, nonenrolled patients were interviewed as part of clinical practice by a nurse dedicated to the vertebroplasty practice. In clinical practice at the lead INVEST site, pain questions included queries about “pain at rest” and “pain with activity,” whereas in the INVEST trial, patients were asked to verbally rate their average pain intensity during the preceding 24 hours on a scale of zero = no pain to 10 = pain as bad as could be.

Differences in pain questionnaires between the INVEST trial and the routine clinical practice arose because the study was designed before the principal investigator relocated to the current lead site. We, therefore, summarized the pain ratings made by study participants and those made by patients who did not enroll in the study, but we did not make any formal statistical comparison. Other outcomes measures, such as the SF-36, were not in routine clinical use at the lead site during the time of the INVEST study, so they were not available in the chart review.

**Statistical Analyses**

We calculated means and SDs for age, RMDS, and pain scores and compared study participants with eligible nonenrolled patients on each measure (except for pain scores) by using 2-sample t tests, assuming unequal variances. We presented and compared the proportion of women for both groups by using χ² tests. All statistical analyses were conducted by using STATA/IC 10.0 (StataCorp, College Station, Tex).

**Results**

Of the 972 patients screened at the lead site, 12% (n = 119) were eligible for enrollment, with the most common reasons for exclusion being presence of tumor, infection, coagulopathy, and the absence of documented osteoporosis. Of the 119 meeting eligibility criteria, 27 (23%) patients approached for study participation enrolled in the INVEST study. Of the 92 eligible patients who declined enrollment at the lead site, we had demographic, RMDS scores, and pain information on 70 patients. To date, we have enrolled 125 patients in INVEST at 10 sites in 4 countries.

Mean ages were similar across both comparison groups, averaging 73.7 years among enrolled participants and 76.6 years among nonenrolled patients (P = .17, Table). Approximately 75% of patients were women in both groups, with no significant difference between groups (P = .64). Enrolled patients at the lead site did not differ from eligible nonenrolled patients at the lead site in RMDS scores (mean = 18.0 ± 4.2 versus 18.6 ± 3.7, P = .51). Among the enrolled patients, the mean “average pain intensity during the prior 24 hours” was 7.6 ± 2.1 among patients enrolled at the lead site. Among eligible nonenrolled patients at the lead site, the mean pain at rest was 3.4 ± 3.3 and the mean pain with activity was 8.5 ± 2.0.
Discussion

Our data indicate that age, distribution of men versus women, and back pain–related disability, as measured by the RMDS, were similar between enrolled and nonenrolled patients at the lead site in INVEST. Mean RMDS scores did not differ between groups, as judged by either a conventional test of statistical significance or by currently accepted definitions of clinically meaningful differences.16 These data suggest that outcomes from INVEST, yet to be reported, likely will be generalizable to typical vertebroplasty patients who have baseline characteristics that would have allowed enrollment in INVEST.

The modified RMDS was initially developed for the study of patients with low-back pain and sciatica. The original report of the RMDS compared pain severity descriptors with scores on this scale.15 Pain that was “almost unbearable” or “very bad” was associated with RMDS mean scores of 14–15, with 95% confidence intervals ranging from 14 to 19. The mean score for enrolled patients in the INVEST trial currently is approximately 17–18, indicating both severe functional disability and pain levels.

Unfortunately, we are unable to compare directly baseline pain scores across groups. Patients enrolled in the INVEST trial responded to an ordinal 0–10 pain question describing “average pain over the past 24 hours,” whereas nonenrolled patients responded to a question posed as “pain at rest” and “pain with activity.” This disparity concerning how pain questions are posed to vertebroplasty patients has received little, if any, attention previously. Indeed, apart from studies published by the lead INVEST site in which specific pain questions about rest and activity are reported,14,17 we were unable to find any previous vertebroplasty publication that describes exactly how pain-related questions are asked. Previous studies have shown that simply asking for pain at rest versus with activity will change the mean baseline pain by 4–5 points on a 10-point scale.14,17 Further modifiers such as “average pain over the past 24 hours” as in the INVEST trial likely will modulate the reported pain level, compared with series that ask for worst pain. Unfortunately, given the dearth of detailed information concerning the pain measures used in previous publications, we cannot directly compare the INVEST baseline pain data with pain data in other studies. We surmise, however, that an “average” pain of approximately 7 of 10 indicates pain severity comparable with the level of 7–9 of 10 in prior studies.1-9

We have previously reported baseline RMDS data from a consecutive group of vertebroplasty patients treated at the lead site for INVEST but entirely separate from the INVEST trial.14 The mean RMDS score for this previously reported cohort was approximately 18, similar to that of the overall enrolled cohort and nearly exactly the same as that of enrolled patients at the lead site. This finding lends further credence to the idea that patients enrolled in INVEST are at least as disabled by their back pain as most typical vertebroplasty patients.

This study has several limitations. Although the comparison group of nonenrolled patients was treated at the lead INVEST site concurrent with the enrolled cohort, data collection schemes for the enrolled–versus–nonenrolled patients differed. Enrolled patients were interviewed with dedicated case report forms by a research coordinator. Nonenrolled patients were interviewed as part of clinical practice by a nurse dedicated to the vertebroplasty practice, with different questions used to assess pain. Another limitation is that RMDS scores were available for eligible nonenrolled patients only at the lead site. Despite these limitations, the data presented here indicate that the pain and disability experienced by enrolled INVEST patients mimicked that of nonenrolled eligible patients.

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