Postoperative epidural analgesia for patients undergoing pectus excavatum corrective surgery: a 10-year retrospective analysis

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Introduction: Managing postoperative pain in patients undergoing minimally invasive pectus excavatum repair (Nuss procedure) is challenging but essential in facilitating ambulation and minimizing the length of stay. Although multiple epidural regimens with varying opioids are presently used for pain management, there is currently no clinical consensus regarding which epidural regimen provides the best analgesia outcomes with the fewest side effects. This 10-year retrospective cohort study was performed to compare the quality of analgesia and the incidence of side effects associated with the three most common epidural regimens used at a tertiary care children’s hospital, in patients undergoing the Nuss procedure.

Methods: Seventy-two pediatric patients were identified as having been treated with one of three epidural regimens for postoperative pain management following the Nuss procedure: Group A (n=12) received 0.125% bupivacaine and 5 µg/mL fentanyl, Group B (n=21) received 0.125% bupivacaine and 10 µg/mL hydromorphone, and Group C (n=39) received 0.1% ropivacaine and 20 µg/mL hydromorphone. Our primary outcome was maximal daily pain scores (numerical rating scale 0–10), with an analytical focus on postoperative day 1 scores. The primary outcome was analyzed using linear regression. The secondary outcomes included the length of stay, side-effect profiles as reflected by the number of treatments for nausea and pruritus, pain scores according to epidural site insertion, occurrence of breakthrough pain, and presence of severe pain throughout their hospital stay. Secondary outcomes were analyzed using linear or logistic regression adjusted for pain scores at baseline. The criterion for statistical significance was set a priori at alpha = 0.05.

Results: Group A had significantly higher day-1 pain scores (score 5.42/10) than Group B (4.52/10; P=0.030) and Group C (4.49/10; P=0.015) after adjusting for baseline pain and age. No significant difference in maximum daily pain scores was found between groups during postoperative days 2–5. Among secondary outcomes, Group C had a significantly lower incidence of nausea/vomiting than Group B (P=0.003). There was also significantly more severe pain in Group A than in Group C (P=0.031). No significant difference was found between the three groups for the incidence of pruritus, critical events, breakthrough pain, or patient satisfaction.

Conclusion: There is no significant difference in managing postoperative pain overall between the three epidural regimens employed at our center. However, in managing day-1 postoperative pain and minimizing nausea/vomiting, our study suggests that a hydromorphone–ropivacaine epidural regimen appears to have more favorable results than a fentanyl–bupivacaine regimen or a hydromorphone–bupivacaine regimen.

Keywords: Nuss procedure, pain, opioids, pediatric patients, epidural
**Introduction**

Pectus excavatum (PE) is a common congenital anomaly of the anterior thorax characterized by sternal depression that results in a concave precordium. Occurring in 1:400 births, PE is six times more common than pectus carinatum and is cited as the most frequent congenital deformity of the anterior chest wall. Clinically, symptoms of easy fatigability, dyspnea on exertion, wheezing, anterior chest wall pain, palpitations, and recurrent lower respiratory tract infections may develop as a result of this, but these symptoms are not common; rather, most patients are healthy and asymptomatic. Despite this, the role of surgery in correcting PE is underscored by the findings that repair of PE (post bar removal) results in an increase in cardiopulmonary function. In addition, surgical intervention improves exercise tolerance, cosmesis, and self-image.

In 1997, Nuss et al described a minimally invasive technique where they preserved costal cartilages and elevated the sternum with a convex internal steel bar. At that time, it had been performed in 42 patients older than 10 years. The more recent 21-year report on the Nuss procedure shows that the procedure had been performed in 1,215 patients with 95.8% good-to-excellent anatomic result. Naturally, it is now becoming widely accepted as the primary surgical intervention for the correction of PE deformity due to its shorter operating duration, better cosmesis, and less blood loss in comparison to the more invasive Ravitch costal resection procedure.

The Nuss procedure is performed as follows: 1) a small lateral subcostal incision is made on the right and left side for insertion of a convex steel bar under the sternum; 2) a separate, small lateral incision is made to allow for a thoracoscope for direct visualization as the bar is passed under the sternum; 3) thoracoscopy has been added to minimize the risk of mediastinal and liver injury during insertion of the subternal bar; 4) the convex bar is rotated to elevate the sternum, and is then fixed to the ribs on either side; 5) the incisions are closed and dressed; and 6) after a 2–4 year period, bar removal is performed as an outpatient procedure. In the past, bar displacement has been a major complication resulting in both recurrence of pectus excavatum and a need for reoperation. Since then, however, modifications to the procedure and restrictions on patient mobility have been made to minimize complications of bar displacement; bar stabilizers are now used and patients are restricted from flexion and rotation at the waist and hips during the perioperative period.

Despite its minimally invasive approach, it is well recognized that the Nuss procedure has a more painful postoperative course than the Ravitch procedure for reasons not entirely elucidated. It has been hypothesized that forcing the sternum into normal alignment causes stress on the ribs and sternum, as well as strain on the costal cartilage. This can eventually result in small fractures visible on ultrasound and nuclear medicine scans. Postoperative pain is furthermore associated with an increase in complication rates, including bar displacement. Therefore, this emphasizes the importance of determining the most effective pain management technique for postoperative care.

Studies have examined the difference in visual analog pain scores between patient-controlled analgesia (PCA) and epidural treatments for the postoperative care of patients undergoing the Nuss procedure. A study of 40 children randomized between morphine PCA and a fentanyl/ropivacaine epidural regimen demonstrated lower visual analog pain scores in the group receiving epidural pain management. Another study of 28 randomized patients showed that there was no difference in postoperative pain control using PCA with fentanyl in comparison to an epidural block with fentanyl/bupivacaine. The use of PCA versus epidural intervention for pain management is thus not yet definitive and has consequently been left to the discretion of the anesthesiologist.

Independently, the epidural technique has been found to be effective in postoperative pain management for PE repair. One study of 21 patients who underwent the Nuss procedure used one of two epidural regimens to relieve postoperative pain: fentanyl in bupivacaine and fentanyl in ropivacaine. Use of these regimens resulted in 57.1% of patients requiring no additional analgesic, 19.0% of patients required one additional dose of pain management with diclofenac sodium or pentazocine, and 23.9% of patients required two or more additional doses of pain management. This study illustrates the utility of epidural pain management postoperatively for the Nuss procedure but does not specifically compare pain outcomes between the two different epidural groups.

Similarly, in a study conducted by Densmore et al, different epidural regimens were tested and compared for postoperative pain management in the Nuss procedure. The study determined a success rate of 96.4% in terms of pain management in its patient’s post-Nuss procedure. The regimens utilized included an opioid (morphine, hydromorphone, or fentanyl), a local anesthetic (bupivacaine or ropivacaine), and in 20% of cases, clonidine. The study found that the type of opioid used did not change outcomes with regard to the pain scores. The study, however, did not report on the incidence of side effects that were associated with each epidural regimen used.
At McMaster University, patients undergoing the Nuss procedure have been followed closely by our Acute Pain Service (APS), and we have been collecting analgesia outcomes since we launched a quality assurance database (APS Manager; Adjuvant Informatics) in 2002. Over the course of 10 years, we have tried a number of epidural analgesia regimens for these patients, and it is not clear which is optimal and which has the fewest side effects. The purpose of this retrospective quality assurance study was to compare the quality of analgesia and the incidence of side effects associated with the three most common epidural regimens, over a 10-year period, used at our center for the treatment of patients undergoing the Nuss procedure.

Methods
This was a single-center, retrospective cohort study comparing postoperative pain management in patients undergoing the Nuss procedure between 2002 and 2012 at the McMaster University Medical Centre in Hamilton, Ontario. After obtaining Research Ethics Board approval from Hamilton Health Sciences Research Ethics Board, one surgeon (Peter Fitzgerald) provided a list of 153 patients originally referred for an assessment for the Nuss procedure. Written informed patient consent was not obtained as this was a retrospective chart review. Of these patients, 112 underwent the Nuss procedure and were identified for review. Eighteen patients were immediately excluded from the review based on the criteria that medical records could not be found (14 patients), redo pectus repair was necessary (two patients), or they were the sole recipients of a unique type of epidural regimen (two patients). Original medical files, microfiche, and computer-based copies of medical files were thus extracted for 94 patients for retrospective chart analysis. From this, 72 patients were identified as having received treatment with the three most common epidural regimens employed at our center for postoperative pain management of the Nuss procedure: Group A (n=12) received 0.125% bupivacaine and 5 µg/mL fentanyl, Group B (n=21) received 0.125% bupivacaine and 10 µg/mL of hydromorphone, and Group C (n=39) received 0.1% ropivacaine and 20 µg/mL hydromorphone (Table 1).

The anesthetic records, pain management flow sheets, medication charts, and admission data were extracted for these 72 patients. Patients’ sex, age, weight, American Society of Anesthesiologists (ASA) classification, duration of hospital stay, satisfaction, site of insertion of epidural catheter, and time of starting the epidural infusion were recorded. Maximal daily pain scores (numerical rating scale: 0–10), breakthrough analgesia requirements, and the presence of severe pain, critical events, and side effects (nausea and pruritus) were extracted from a computerized acute pain database from the APS. We considered the maximum pain score postoperative day 0 to be considered the patient’s baseline pain score. A critical event was defined as a medication error, respiratory depression, respiratory or cardiac arrest, severe hypotension, prolonged motor blockade, or epidural hematoma or abscess.

All patients except one had a thoracic epidural catheter inserted preoperatively and epidural infusion either commenced or continued postoperatively. Epidural catheter insertions were considered optimal if inserted at or higher than T6/7 and were considered suboptimal if inserted below T6/7. Patients were discharged from the postsurgery care unit when their analgesia was adequate, and epidural infusions were maintained between 0.1 mL/kg/h and 0.4 mL/kg/h. The epidural regimen employed was left to the APS team and the regimen used for these cases changed over

### Table 1 Comparison of demographic characteristics between Groups A, B, and C

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Levels</th>
<th>Group A (0.125% bupivacaine/5 µg/mL fentanyl) (N=12)</th>
<th>Group B (0.125% bupivacaine/10 µg/mL hydromorphone) (N=21)</th>
<th>Group C (0.01% ropivacaine/20 µg/mL hydromorphone) (N=39)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years), mean (SD)</td>
<td>15.2 (6.1) (n=12)</td>
<td>15.9 (4.5) (n=21)</td>
<td>18.5 (4.1) (n=39)</td>
<td>20.2 (5.3) (n=39)</td>
<td>0.030*</td>
</tr>
<tr>
<td>Weight (kg), mean (SD)</td>
<td>60.9 (22.5) (n=10)</td>
<td>56.4 (16.7) (n=21)</td>
<td>64.8 (13.6) (n=39)</td>
<td>69.4 (16.7) (n=39)</td>
<td>0.156</td>
</tr>
<tr>
<td>Pain score at baseline, mean (SD)</td>
<td>2.1 (2.7) (n=11)</td>
<td>4.2 (2.2) (n=20)</td>
<td>4.5 (2.6) (n=39)</td>
<td>4.7 (2.4) (n=39)</td>
<td>0.026*</td>
</tr>
<tr>
<td>Sex (male), n (%)</td>
<td>11/12 (91.7%)</td>
<td>20/21 (95.2%)</td>
<td>13/39 (97.4%)</td>
<td>14/39 (35.9%)</td>
<td>0.673</td>
</tr>
<tr>
<td>ASA class, n (%)</td>
<td>1</td>
<td>7/10 (70.0%)</td>
<td>12/21 (57.1%)</td>
<td>13/39 (33.3%)</td>
<td>0.177</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>3/10 (30.0%)</td>
<td>9/21 (42.9%)</td>
<td>25/39 (64.1%)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>0/10 (0.0%)</td>
<td>0/21 (0.0%)</td>
<td>1/39 (1.6%)</td>
<td></td>
</tr>
<tr>
<td>Epidural level, n (%)</td>
<td>Optimal</td>
<td>5/11 (41.7%)</td>
<td>11/21 (52.4%)</td>
<td>25/39 (64.1%)</td>
<td>0.344</td>
</tr>
<tr>
<td></td>
<td>Suboptimal</td>
<td>7/11 (58.3%)</td>
<td>10/21 (47.6%)</td>
<td>14/39 (35.9%)</td>
<td></td>
</tr>
</tbody>
</table>

Notes: For continuous variables, the P-values are obtained by one-way ANOVA. For categorical variables, the P-values are obtained by Pearson chi-squared test. ANOVA tests whether there are any differences between the groups with a single probability for continuous outcomes, which is used when the number of groups is more than two. Pearson chi-squared test is used for categorical outcomes to investigate whether there are any differences on the distribution between groups. *Statistically significant: P<0.05.

Abbreviations: SD, standard deviation; ANOVA, analysis of variance; ASA, American Society of Anesthesiologists.
The results from the linear and mixed-effects models are adjusted for pain scores at baseline and for age. Logistic regression. All linear and logistic regression analyses are illustrated by a box plot.

The primary outcome was maximal daily pain scores (numerical rating scale 0–10) as determined by the ward nurse for each postoperative day. Our primary time period of focus was day 1 pain scores as pain is likely most severe immediately postoperatively. Our secondary time period of focus was the pain scores for the remaining days. Daily morning pain scores assessed by the APS nurses were also extracted, and any pain problems found on follow-up outpatient assessments were recorded. The secondary outcomes included the length of stay (LOS) of the patients, the side-effect profiles as reflected by the number of treatments for nausea and pruritus, the occurrence of breakthrough pain, and the presence of severe pain throughout the course of their hospital stay. Another secondary outcome also included daily pain scores according to epidural site insertion point (optimal vs suboptimal).

Statistical analysis
The demographic and anesthetic factors of the patients were analyzed by group using descriptive statistics reported as a mean (SD) for continuous variables and count (percentage) for categorical variables. The primary outcome (daily pain scores after treatment) is illustrated by box plot and analyzed using linear regression. They were adjusted for age and baseline pain score. A generalized estimating equation (GEE) model was also used to analyze the pain scores and provide an overall pain score for each group, adjusting for repeated measurements acquired for each patient on a day-to-day basis. Daily pain scores were also analyzed by epidural catheter location (optimal vs suboptimal) using linear regression. They are illustrated by a box plot.

The secondary outcomes include LOS in hospital, requirement of breakthrough analgesia, severe pain, critical events, pruritus, nausea/vomiting, and satisfaction. The LOS is analyzed using linear regression. Other secondary outcomes are illustrated by bar plots and analyzed using logistic regression. All linear and logistic regression analyses are adjusted for pain scores at baseline and for age. The results from the linear and mixed-effects models are reported as adjusted difference between the treatment groups, corresponding two-sided 95% confidence intervals (CIs), and associated P-values. Results from logistic regression are reported as odds ratio, 95% CI, and associated P-values.

All statistical tests are performed using two-sided tests at the 0.05 level of significance. P-values are reported to three decimal places with values <0.001 reported as <0.001. All the analyses were performed using SAS 9.2 (SAS Institute Inc., Cary, NC, USA).

Results
Of the 72 patients, the most common epidural regimen was 0.1% ropivacaine/20 µg/mL hydromorphone (Group C, N=39), the next most frequent regimen was 0.125% bupivacaine/10 µg/mL hydromorphone (Group B, N=21), followed by the 0.125% bupivacaine/5 µg/mL fentanyl (Group A, N=12). The groups were similar with respect to sex, ASA status, weight, LOS, and site of epidural catheter insertion (Table 1). Group C had a significantly higher mean age (18.5) than did Group A (15.2) and Group B (15.9, P=0.030). One patient was considered a fail of epidural technique and was switched to PCA on postoperative day 2, when his pain control was considered inadequate. With respect to pain scores at baseline, Group A (2.1/10) was found to have a statistically significant (but clinically modest) lower pain score than both Group B (4.2/10) and Group C (4.5/10, P=0.026).

Primary outcome – maximal daily pain scores
On postoperative day 1, Group A was found to have a significantly higher amount of pain (score 5.42/10) than both Group B (4.52/10, P=0.030, 95% CI [-3.81, -0.21]) and Group C (4.49/10; P=0.015, 95% CI [-3.91, -0.43]) after adjusting for baseline pain between the groups. No significant difference in maximum daily pain scores was found between groups during the postoperative days 2, 3, 4, or 5 (Table 2 and Figure 1). When using the GEE method, no significant differences were found in overall pain scores between any of the groups.

Secondary outcomes
With respect to side effects, 67% of patients from Group A, 81% from Group B, and 34% from Group C experienced nausea/vomiting, requiring treatment throughout their stay. A significant difference was found between Group B and Group C (P=0.003, 95% CI [0.04, 0.52]). No significant difference was found for nausea/vomiting between the...
other groups. In addition, 67% of patients from Group A, 52% of patients from Group B, and 49% of patients from Group C experienced severe pain throughout their stay. Among these, there was a significant difference between Group C and Group A (P=0.031, 95% CI [0.03, 0.84]). No significant difference was found between the three groups for the incidence of pruritus, incidence of critical events, occurrence of breakthrough pain, or satisfaction with their pain management throughout their stay (Table 3 and Figure 2).

In our study, two critical events occurred: one patient in Group B had a medication error wherein inappropriate sedatives were prescribed by a non-APS physician, and another patient in Group C was found to have a respiratory rate of eight with no sequelae.

### Pain scores based on epidural catheter location

In total, 41 patients (57%) were found to have optimal epidural catheter location (inserted above T6/7), and 31 patients (43%) had suboptimal epidural catheter placement (insertion below T6/7). The groups were similar with regard to all baseline characteristics, including age, weight, pain scores at baseline, sex, and ASA class (Table 4).

A significant difference in pain scores between groups was found only on the fourth postoperative day (P=0.037, CI −3.25, −0.11). No significant differences were found on any of the remaining daily pain scores or using the GEE method for analysis (Table 5).

### Pain after discharge from hospital

All patients were discharged with oral analgesia – acetaminophen and ibuprofen plus codeine or oxycodone or MS Contin (morphine sulfate). Only one patient did not require opioid discharge medication. All patients were seen by the

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**Table 2 Comparison of maximum daily pain scores between Groups A, B, and C**

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Group A (N=12) Mean (SD) (n)</th>
<th>Group B (N=21) Mean (SD) (n)</th>
<th>Group C (N=39) Mean (SD) (n)</th>
<th>Comparing difference between groups</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Pain score at day 1</td>
<td>Pain score at day 2</td>
<td>Pain score at day 3</td>
<td>Pain score at day 4</td>
</tr>
<tr>
<td></td>
<td>5.42 (3.14) (n=12)</td>
<td>4.55 (2.38) (n=11)</td>
<td>4.00 (2.58) (n=10)</td>
<td>4.33 (1.97) (n=6)</td>
</tr>
<tr>
<td></td>
<td>Pain score at day 2</td>
<td>5.05 (2.09) (n=20)</td>
<td>4.08 (1.72) (n=20)</td>
<td>3.13 (1.88) (n=15)</td>
</tr>
<tr>
<td></td>
<td>4.49 (2.17) (n=39)</td>
<td>4.51 (2.24) (n=37)</td>
<td>4.09 (2.27) (n=35)</td>
<td>3.48 (3.09) (n=21)</td>
</tr>
<tr>
<td></td>
<td>Pain score at day 3</td>
<td>2.33 (1.97) (n=6)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Notes:** In the analysis of pain score in each day, age and pain score at baseline were adjusted as confounders. This means that the impacts of age and pain score at baseline have been adjusted when estimating the pain scores. In the analysis of pain score using GEE method, the repeated measurements of pain score for each patient were considered. GEE model is a popular choice when analyzing repeated measurements. When a patient was measured multiple times over a certain period, the data collected with same person were correlated. The estimates from GEE model have been adjusted for this type of correlation. Group A (0.125% bupivacaine/5 µg/mL fentanyl); Group B (0.125% bupivacaine/10 µg/mL hydromorphone); Group C (0.1% ropivacaine/20 µg/mL hydromorphone). *Statistically significant: P<0.05.

**Abbreviations:** SD, standard deviation; CI, confidence interval; GEE, generalized estimating equation.

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**Figure 1** Comparison of maximum daily pain scores between Groups A, B, and C.

**Note:** Group A (0.125% bupivacaine/5 µg/mL fentanyl); Group B (0.125% bupivacaine/10 µg/mL hydromorphone); Group C (0.1% ropivacaine/20 µg/mL hydromorphone). *Statistically significant: P<0.05.

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A significant difference in pain scores between groups was found only on the fourth postoperative day (P=0.037, CI −3.25, −0.11). No significant differences were found on any of the remaining daily pain scores or using the GEE method for analysis (Table 5).
surgeon in a clinic postoperatively, and none of the patients were using the analgesics at the follow-up appointment or experienced any pain problems.

**Discussion**

This study found no significant differences in pain scores between the three epidural regimens overall. However, this study demonstrated that patients who received fentanyl in their epidural regimen (Group A) had significantly higher pain scores on day 1 than did those who received hydromorphone (Groups B and C) when adjusted for age and baseline pain score. This finding, better analgesia with epidural hydromorphone than fentanyl, has not yet been reported in the literature. The analgesia results of this study are consistent with the literature demonstrating the efficacy of epidural regimens for postoperative pain control in patients undergoing the Nuss procedure. Several studies have reported the need for supplemental intravenous morphine in order to achieve adequate pain control in their

Table 3 Comparison of secondary outcomes between groups A, B, and C

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Group A (N=12) Mean (SD) (n)</th>
<th>Group B (N=21) Mean (SD) (n)</th>
<th>Group C (N=29) Mean (SD) (n)</th>
<th>Comparing difference between groups</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Requirement of breakthrough analgesia</td>
<td>Severe pain</td>
<td>Critical events</td>
<td>Pruritis</td>
</tr>
<tr>
<td>Event/n (%)</td>
<td>Event/n (%)</td>
<td>Event/n (%)</td>
<td>Event/n (%)</td>
<td>Event/n (%)</td>
</tr>
<tr>
<td></td>
<td>10/12 (83.33)</td>
<td>14/21 (66.67)</td>
<td>26/39 (66.67)</td>
<td>4/11 (36.36)</td>
</tr>
<tr>
<td></td>
<td>8/12 (66.67)</td>
<td>11/21 (52.38)</td>
<td>19/39 (48.72)</td>
<td>1/2 (4.76)</td>
</tr>
</tbody>
</table>

**Notes:** For all the above analyses, both age and pain score at baseline were adjusted as confounders. OR ≤ 1 means the odds from two groups are equal, OR > 1 means higher risk, OR < 1 means lower risk. For example, when looking at severe pain between Group A and Group C (C/A), OR = 0.15 (0.03, 0.84), P=0.031, this result tells us that the odds of patients from Group C having severe pain are significantly lower than the patients in Group A. Group A (0.125% bupivacaine/5 μg/mL fentanyl); Group B (0.1% ropivacaine/20 μg/mL hydromorphone); Group C (0.1% ropivacaine/20 μg/mL hydromorphone). *Statistically significant.

**Abbreviations:** SD, standard deviation; CI, confidence interval; OR, odds ratio.
Figure 2 Comparison of secondary outcomes between Groups A, B, and C.

Notes: Group A (0.125% bupivacaine/5 µg/mL fentanyl); Group B (0.125% bupivacaine/10 µg/mL hydromorphone); Group C (0.1% ropivacaine/20 µg/mL hydromorphone). No significant difference was found between the three groups for the incidence of pruritus, incidence of critical events, occurrence of breakthrough pain, or satisfaction with their pain management throughout their stay.

Table 4 Comparison of demographic characteristics between optimal and suboptimal epidural placement groups

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Levels</th>
<th>Optimal group (n=41)</th>
<th>Suboptimal group (n=31)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years), mean (SD)</td>
<td>16.85 (4.78) (n=41)</td>
<td>17.68 (4.75) (n=31)</td>
<td>0.470</td>
<td></td>
</tr>
<tr>
<td>Weight (kg), mean (SD)</td>
<td>67.72 (14.93) (n=39)</td>
<td>61.71 (17.95) (n=31)</td>
<td>0.998</td>
<td></td>
</tr>
<tr>
<td>Pain score at baseline, mean (SD)</td>
<td>3.94 (2.91) (n=39)</td>
<td>4.13 (2.17) (n=31)</td>
<td>0.759</td>
<td></td>
</tr>
<tr>
<td>Sex (male), n (%)</td>
<td>39/41 (95.12)</td>
<td>30/31 (96.77)</td>
<td>0.728</td>
<td></td>
</tr>
<tr>
<td>ASA class</td>
<td>1, n/N (%)</td>
<td>17/39 (43.59)</td>
<td>15/31 (48.39)</td>
<td>0.638</td>
</tr>
<tr>
<td></td>
<td>2, n/N (%)</td>
<td>21/39 (53.85)</td>
<td>16/31 (51.61)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>3, n/N (%)</td>
<td>1/39 (2.56)</td>
<td>0/31 (0.00)</td>
<td></td>
</tr>
</tbody>
</table>

Note: Optimal = epidural inserted above T6/7, Sub-optimal = epidural inserted below T6/7.

Abbreviations: SD, standard deviation; ASA, American Society of Anesthesiologists.

patients; of these, one study\textsuperscript{15} did not report pain scores, and the other\textsuperscript{16} found that IV (intravenous) morphine was needed in 47\% of their patients. Another case series reported excellent epidural analgesia without IV opioids, but they required regular, large doses of epidural opioid in order to achieve this – 40 µg/kg morphine Q8H.\textsuperscript{17} In either case, both of these studies corroborate with our finding that severe pain occurs in patients despite using a multimodal analgesic approach. A larger retrospective chart review using morphine, hydromorphone, or fentanyl as the opioid accompanying the local anesthetic in the epidural catheter found that IV ketorolac was used in 69\% of epidural patients to provide further pain relief.\textsuperscript{14} Further statistical analysis failed to find a significant difference between those who received it and those who did not.\textsuperscript{14} When actually comparing the types of opioids used in the epidural catheters, their study’s findings were similar to our own in that no significant difference in pain scores was found between groups.\textsuperscript{14} Thus, our paper supports the current evidence that the choice of epidural opioid does not have a significant impact on postoperative pain scores in patients undergoing the Nuss procedure.

The use of postoperative daily maximal pain scores may not reflect epidural efficacy if a brief moment of severe pain (commonly seen during ambulation) occurred for only a short duration of time. However, it still reflects the lack of adequate...
analgesia. Although a record of the duration of maximal pain score may better reflect the efficacy of the epidural, this was not possible at our center given the variations of the recording process on the ward in the 10-year period and also given that the pain nurse only visited the patient to record the pain scores once daily. Thus, a prospective randomized trial with hourly pain score recordings might better reflect the ability of the epidural regimen to control pain.

The chronological progression from Group A to B to C at our center occurred due to several perceived benefits as per the anesthesiology team. Switching from a fentanyl-based regimen (Group A, 2002–2003) to a hydromorphone-based regimen (Group B, 2004–2005) was thought to decrease the opioid-related side effects associated with fentanyl. Furthermore, the rationale from switching the local anesthetic from bupivacaine (Group B) to ropivacaine (Group C, 2005–2012) was to allow for an increase in the amount of hydromorphone that could be administered in order to improve overall analgesia. Severe pain was found to be improved in Group C (49%) when compared to Group A (67%), which may suggest that a higher dose of hydromorphone in a ropivacaine regimen may better control severe pain than a bupivacaine–fentanyl-based regimen. With regard to side effects, no significant difference was seen in nausea/vomiting once the opioid was changed from fentanyl to hydromorphone. This was contrary to the expectations of the anesthesia team. Surprisingly, however, a decrease in nausea/vomiting was seen when the hydromorphone dose was doubled and the local anesthetic was changed from bupivacaine to ropivacaine. Although there was not a statistically significant difference in the proportion of optimal epidural insertions between the treatment groups, there was a trend for more optimal placement over time (from 42% in Group A to 64% in Group C) and this may reflect an improvement in anesthesia practice and regional anesthesia training in recent years. There was no significant difference in pruritus between any of these groups. We cannot corroborate these findings with the available literature, as this is the first study to evaluate the side-effect profiles of various epidural regimens for postoperative pain control for the Nuss procedure.

One of the biggest strengths of this review is that it is the first retrospective cohort study that evaluates epidural regimens for both their efficacy in postoperative pain control and side-effect profiles for patients undergoing the Nuss procedure. Its sample size also reflects all the patients who underwent the Nuss procedure at a large tertiary center over a significant period of time (10 years). The current literature does not have similar studies examining patients over this period of time. Finally, the pain measurements and data were collected systematically and prospectively using dedicated APS nurses and a quality assurance database for the duration of the review.

The weaknesses of the study include its retrospective nature, lack of randomization, and limited ability to detect significant differences between treatment groups due to its seemingly small sample size. Other confounders not controlled for in this retrospective design include the era of the surgery, the severity of the deformity, length of surgery, and the degree of surgical manipulation required. However, given that our Nuss procedure is relatively uncommon and given that our data comprise all the patients who underwent minimally invasive repair over an entire decade, we argue that our sample size is acceptable. It should also be noted that our statistical analysis adjusted for the difference in age and baseline pain scores between the three groups in order to prevent confounding from these variables.

**Conclusion**

Our study demonstrates that there is no significant difference between the three epidural regimens in managing overall postoperative pain in patients undergoing the Nuss procedure. However, in managing day-1 postoperative pain and minimizing nausea/vomiting, our study suggests that a

### Table 5 Comparison of maximum pain scores between optimal and suboptimal epidural placement groups

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Optimal group (n=21) Mean (SD) (n)</th>
<th>Suboptimal group (n=51) Mean (SD) (n)</th>
<th>Comparing difference between groups</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pain score at day 1</td>
<td>4.30 (2.59)</td>
<td>5.11 (2.26)</td>
<td>Suboptimal–optimal 0.81 (0.36, 1.97)</td>
</tr>
<tr>
<td>Pain score at day 2</td>
<td>4.84 (2.27)</td>
<td>4.47 (2.13)</td>
<td>Suboptimal–optimal −0.38 (−1.45, 0.70)</td>
</tr>
<tr>
<td>Pain score at day 3</td>
<td>4.15 (2.19)</td>
<td>3.96 (2.08)</td>
<td>Suboptimal–optimal −0.18 (−1.26, 0.89)</td>
</tr>
<tr>
<td>Pain score at day 4</td>
<td>4.12 (2.61)</td>
<td>3.96 (2.08)</td>
<td>Suboptimal–optimal −1.68 (−3.25, 0.11)</td>
</tr>
<tr>
<td>Pain score at day 5</td>
<td>2.4 (2.61)</td>
<td>1.41 (1.41)</td>
<td>Suboptimal–optimal −1.4 (−5.83, 3.03)</td>
</tr>
<tr>
<td>Pain score (using GEE method)</td>
<td>2.4 (2.61)</td>
<td>1.41 (1.41)</td>
<td>Suboptimal–optimal −0.10 (0.76, 0.56)</td>
</tr>
</tbody>
</table>

**Notes:** Optimal = epidural inserted above T6/7, Sub-optimal = epidural inserted below T6/7. *Statistically significant.

**Abbreviations:** SD, standard deviation; CI, confidence interval; GEE, generalized estimating equation.
hydromorphone–ropivacaine epidural regimen appears to have more favorable results than a fentanyl–bupivacaine regimen or a hydromorphone–bupivacaine regimen. Although pain is problematic for several weeks to months after surgery, the incidence of chronic pain following the Nuss procedure has yet to be reported and could be an area of future research. A prospective randomized trial with a larger sample size that recorded the duration of maximal pain scores may better reflect the ability of the regimen to control pain, along with the prevalence of side effects.

**Author contributions**

Asad Siddiqui, Andrew Tse, and James E Paul, helped design the study, conduct the study, reviewed the analysis of the data, wrote and approved the final manuscript; Peter Fitzgerald, helped to design the study and analyze the data; Bernice Teh, helped to design the study and approved the final manuscript. All authors contributed toward data analysis, drafting and critically revising the paper and agree to be accountable for all aspects of the work.

**Disclosure**

The authors report no conflicts of interest in this work.

**References**