A preliminary study of a novel robotic system for pedicle screw fixation: A randomised controlled trial

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A R T I C L E   I N F O

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A B S T R A C T

Background and objective: Existing orthopaedic robotic systems are almost restricted to provide guidance for trajectory direction. In the present study, a novel spinal robotic system with automatic drilling power was introduced. The aim of this study is to evaluate the feasibility and safety in pedicle screw insertion of posterior lumbar interbody fusion assisted by this novel robotic system.

Methods and materials: A randomised controlled trial was conducted for 17 participants who were required posterior lumbar interbody fusion process. Seven (3 M/4 F) were randomly assigned to the robot-assisted group (RA group), and the other ten (4 M/6 F) were assigned to the conventional technique group (FH group). A novel robotic system was used in the RA group. All measurements were based on postoperative computed tomography (CT) data. Accuracy of screw insertion was determined using the Gertzbein and Robbins Scale. Precision was measured by the entry point deviation distance and the trajectory rotation. Other variables included operation time, radiation time, length of stay, and screw-related complications.

Result: A total of 82 pedicle screws were placed in the 17 participants. In the RA group, 90.6% of screws placed were Grade A, and 9.4% were Grade B. In the FH group, 78.0% of screws were Grade A, 20.0% were Grade B, and 2.0% were Grade C. No statistical difference was found in the operation time, radiation time per case, and length of stay between both groups. The radiation time per screw is significantly lower in the RA group. No screw-related complications or revision occurred in the present study.

Conclusion: The outcome of screw accuracy of this robotic system was comparable with that of experienced surgeons, and no screw-related complication was found in the RA group during hospitalisation. In addition, radiation time per screw in the robotic group was significantly lower than that in the conventional group, which shows the potential to reduce radiation exposure of pedicle screw fixation assisted by this robotic system.

Translational potential: Our study shows that pedicle screw fixation assisted by “Orthbot” system is accurate and safe. It is concluded that this novel robotic system offers a new option for internal implantation in spine surgery.

Background

Pedicle screw insertion is widely used in posterior lumbar fusion for spinal stabilisation. Conventional internal fixation procedure, with free-hand tools, is based on anatomical landmarks and intraoperative fluoroscopic images [1]. Although this method has been shown to be safe and used worldwide, screw misplacement rates vary greatly among previous research studies [2–6]. Malposition of the screw may result in severe surgical complications, such as vascular or visceral damage, neurological deficit, and dural tearing [3,7]. To acquire higher accuracy in pedicle screw placement, several techniques have been introduced. Computer-assisted system [8,9] or three-dimensional (3D) fluoroscopy–based system [10] seems to be effective to improve screw position in earlier literature. The robotic system, developed as a newer navigation technique with high intrinsic precision and stability, has obtained popularity in a multitude of spinal internal fixation.
procedures [11]. Current research tends to pay close attention to Renaissance (Mazor Robotics Ltd, Caesarea, Israel) [12], SpineAssist (Mazor Robotics Ltd, Caesarea, Israel) [13], ROSA spine (Zimmer Biomet, Warsaw, Indiana) [14], and the TiRobot system (TINAVI Medical Technologies Co. Ltd, Beijing, China) [15]. However, these robotic systems seem to be limited to provide the position of trajectory guiding surgeons to insert K-wires and screw manually. In the present study, a novel spinal robotic system with automatic drilling power is introduced. The aim of this study is to evaluate the feasibility and safety in the insertion of pedicle screws for posterior lumbar interbody fusion assisted by this robotic technique.

Methods and materials

Approved by the Southern Medical University Ethics Committee, the patient recruitment for this randomised controlled trial was performed from August 2018 to December 2018 in our single medical centre. Informed consent was obtained from each individual participant in our study. Those who do not have any mental disorders were enrolled, and all of them signed the informed consent form in a clear-conscious situation without any temptation. Patients who required lumbar interbody fusion and pedicle screw fixation treatments for degenerative lumbar disc disease were prospectively included in the study and randomly divided into two groups. Patients in the ‘Orthbot’ group were consecutively treated under robotic assistance (RA group), while the other patients in the conventional technique group were treated with free-hand approach assisted by intraoperative fluoroscopy equipment (FH group).

Patients

Inclusion criteria were as follows: (1) age between 18 and 65 years; (2) indication of posterior fusion for degenerative lumbar disc disease or lumbar spinal stenosis; (3) signed informed consent to accept the subsequent randomised trial process. Exclusion criteria were as follows: (1) patients with severe degenerative spinal deformity; (2) patients with a previous surgery on the lumbar spine; (3) patients with coagulopathy or metal allergy; (4) pregnant or lactating women; (5) patients who participated in relevant clinical trials of other drugs or medical devices in the past 3 months. We prospectively enrolled 17 participants, 7 males and 10 females, aged 23–63 years. Seven (3 males and 4 females) were randomly assigned to the RA group, and the other 10 (4 males and 6 females) were assigned to the FH group.

Robotic procedure

Based on machine vision tracking algorithm, the “Orthbot” system (Fig. 1), which is comprised of a robotic workstation, coordinate position plate, tracking camera (Fig. 2a) and a 6-degree of freedom (6-DOF) robotic arm with an automatic bone drill (Fig. 2b), can finish inserting a K-wire autonomously by following the preoperative planning of screw trajectory. The working procedure is as follows.

Preoperative planning

A thin-cut (1 mm) computed tomography (CT) scan was performed for each patient before surgery. The CT images were saved in the digital imaging and communications in medicine (DICOM) file format and transferred to the “Orthbot” system’s proprietary software in the planning workstation. The workstation was used to integrate preoperative planning, image acquisition and registration, kinematic calculations, and supervisory control. After the reconstruction of CT data into a 3D model, surgeons could formulate and modify the preoperative plan with appropriate screw sizes and optimal trajectory position in coronal, sagittal, and axial planes (Fig. 3).

Registration

Patients, after receiving general anaesthesia, were placed in prone position. Surgeons assembled the coordinate position plate and set it up above the surgical area. Anteroposterior and lateral fluoroscopic images were required by the C-arm to match with the preoperative CT data for registration. Surgeons’ inspection for registration outcomes was of utmost importance to ensure that the surgical plan had been precisely imported into the robotic system.

Robotic position and drilling procedure

In the RA group, first, the “Orthbot” system was passively positioned alongside the surgical table. The tracking camera then guided the robotic arm into the surgical site by recognising the marker sign set on the coordinate position plate. The automatic bone drill thereafter began to make a channel into the target pedicle, and a K-wire was placed subsequently by following the trajectory in the surgical planning. After all K-wires were inserted, the surgical arm is cleared off the region. Surgeons finally finished inserting screws over each K-wire. The primary surgical procedures in the RA group are shown in Fig. 4.

Conventional surgical procedure

In the conventional technique group, experienced surgeons insert K-wires and screws according to routine processes by a fluoroscopy-guided
technique.

Evaluation and data collection

All patients required postoperative CT scans. The CT reconstructed images with actual screws position were also loaded on to the workstation to compare the deviation from the planned trajectories. Precision assessment of the “Orthbot” system depended on two parameters—the deviation distance of the entry point and the rotation angle of the trajectory. Based on the 3D surgical space coordinate system constructed in the robotic workstation, entry point position and trajectory location were transformed to numerical parameters with X-value, Y-value, and Z-value. Three-dimensional measurements of these two parameters were also performed in the proprietary software (Fig. 5). Meanwhile, each screw position was appraised by the Gertzbein and Robbins Classification Scale to obtain the evaluation of accuracy. In the Gertzbein and Robbins Scale [16], Grade A indicated no perforation of the pedicle; Grade B indicated < 2 mm of perforation of the pedicle; Grade C indicated 2–4 mm of perforation of the pedicle; and Grade D indicated > 4 mm of perforation. Grades A and B were considered clinically satisfactory. All of the evaluation was assessed by an independent author who was blinded to patients’ allocation after receiving the postoperative CT data. Other perioperative variables collected included operation time, blood loss, radiation time (radiation time per case and per screw), length of stay, and screw-related complications (e.g., neurologic deficit, vital vascular injury, and revision surgery).

Statistical analysis

Quantitative data were expressed as mean ± standard deviation (SD). The two-sample t test was used to compare continuous variables between the two groups, and the Fisher exact test was used for ranked variables. Alpha was set at 0.05, while P-values < 0.05 was considered statistically significant.

Result

There were 17 participants enrolled in the randomised controlled trial for treatment by posterior lumbar interbody fusion. The median age of these patients was 50.0 years, and the confidence interval was 48.9 ± 11.4 years (ranging from 23 to 63 years). The median body mass index was 24.3 kg/m², and the confidence interval was 24.5 ± 2.3 kg/m² (ranging from 20.7 to 30.5 kg/m²). Demographic data collected are shown in Table 1. A total of 82 pedicle screws were placed for the two groups. In the RA group, according to the Gertzbein and Robbins Scale, 29 of 32 screws placed were Grade A, and 3 were Grade B. In the FH group, 39 of 50 screws were Grade A, 10 were Grade B, and 1 was Grade C (p > 0.05) (Tables 2 and 3). A comparison of the screw position according to the

Figure 2. Photographs of the (A) tracking camera and (B) the automatic bone drill set on the robotic arm.

Figure 3. Screw planning performed in the workstation based on the preoperative CT reconstruction data. CT = computed tomography.
Gertzbein and Robbins Classification Scale in each group is presented in Fig. 6. However, the deviation distance of entry point \( (p < 0.001) \) and the rotation angle of the trajectory \( (p < 0.001) \) were both significantly lower in the RA group (Table 4). There were no statistical differences in the operation time \( (p = 0.620) \), blood loss \( (p = 0.878) \), radiation time per operation \( (p = 0.291) \), and length of stay \( (p = 0.811) \) between both groups. But radiation time per screw is significantly lower in the RA group \( (0.56 \text{ s in the RA group and 1.04 s in the FH group, } p < 0.05) \) (Table 5). No screw-related complications occurred, and no intraoperative or postoperative revision was necessary for either group.

**Discussion**

Commonly, procedures of spinal surgery require meticulous manipulation to avoid complications due to screw misplacement. Spinal surgery involves complex anatomical structures, and precise screw placement is crucial for optimal surgical outcomes. The use of robotic assistance in spinal surgery, as demonstrated in this study, offers several potential advantages over traditional free-hand techniques.

**Figure 4.** (A) The coordinate position plate was assembled by fellows and set up above the surgical area for registration; (B) the robotic arm of the “Orthbot” system was passively positioned alongside the surgical table; (C) supervised by surgeons, k-wires were placed by the “Orthbot” system automatically during the operation; (D) K-wires were finished inserting, and the robotic system was cleared off the surgical field.

**Figure 5.** Screw accuracy and precision (based on the entry point and trajectory) were evaluated on the postoperative CT images (one of the robot-assisted groups). CT = computed tomography.

| Table 1 | Demographic data of the study cohort. |
|---------|--------------------------------------|
| Characteristic | RAG \( (n = 7) \) | FHG \( (n = 10) \) | P-value |
| Gender (male/female) | 3/4 | 4/6 | 1.000 |
| Age (median; M ± SD) (y) | 48.0; 47.4 ± 12.9 | 51.5; 49.9 ± 10.9 | 0.675 |
| BMI (median; M ± SD) (kg/m²) | 24.3; 24.3 ± 1.8 | 24.0; 24.6 ± 2.6 | 0.773 |
| Diagnosis (no. patients, percentage) | Degenerative disc disease 5 (71.4%) | 6 (60.0%) |
| | Lumbar spinal stenosis 2 (28.6%) | 4 (40.0%) |

FHG = free-hand technique group (conventional group); M = mean; RAG = “Orthbot”-assisted group; SD = standard deviation.
Table 2
Grading assessment for screw position of each patient.

| Patient | Group | Screw (n) | Grade of screw |
|---------|-------|-----------|----------------|
|         |       |           | A   | B   | C   |
| 1       | FHG   | 4         | 4   | 0   | 0   |
| 2       | FHG   | 6         | 4   | 2   | 0   |
| 3       | RAG   | 4         | 3   | 1   | 0   |
| 4       | RAG   | 6         | 6   | 0   | 0   |
| 5       | RAG   | 4         | 3   | 1   | 0   |
| 6       | FHG   | 4         | 4   | 0   | 0   |
| 7       | FHG   | 6         | 5   | 1   | 0   |
| 8       | FHG   | 4         | 2   | 0   | 0   |
| 9       | FHG   | 4         | 3   | 1   | 0   |
| 10      | RAG   | 4         | 4   | 0   | 0   |
| 11      | RAG   | 4         | 4   | 0   | 0   |
| 12      | FHG   | 4         | 1   | 2   | 1   |
| 13      | RAG   | 4         | 3   | 1   | 0   |
| 14      | RAG   | 6         | 6   | 0   | 0   |
| 15      | FHG   | 4         | 3   | 1   | 0   |
| 16      | FHG   | 8         | 8   | 0   | 0   |
| 17      | FHG   | 6         | 5   | 1   | 0   |

RAG = “Orthbot”-assisted group; FHG = free-hand technique group (conventional group).
No Grade D was evaluated in both the two groups.

Table 3
Grading assessment for screw position of each group.

| Grade | RAG (n = 32) | FHG (n = 50) | Total (n = 82) | P-value |
|-------|--------------|--------------|----------------|---------|
| A     | 29 (90.6%)   | 39 (78.0%)   | 68 (82.9%)     | 0.138   |
| B     | 3 (9.4%)     | 10 (20.0%)   | 13 (15.9%)     | 0.199   |
| A + B | 32 (100.0%)  | 49 (98.0%)   | 81 (98.8%)     | 1.000   |
| C     | 0            | 1 (2.0%)     | 1 (1.2%)       | 1.000   |

RAG = “Orthbot”-assisted group; FHG = free-hand technique group (conventional group).
No Grade D was evaluated in both the two groups.

Figure 6. A bar graph of comparison of the percentage for the accuracy on screw insertion between two groups in the study. Grades were determined by the Gertzbein and Robbins Classification Scale.

Table 4
Precision parameters measurement of each study group.

| Parameters | RAG (M ± SD) | FHG (M ± SD) | P-value |
|-----------|-------------|--------------|---------|
| EDP (M ± SD) (mm) | 0.70 ± 0.33 | 4.16 ± 2.21 | 0.000   |
| TRA (M ± SD) (°)   | 1.3 ± 0.8  | 11.9 ± 5.6  | 0.000   |

EDP = entry point deviation; FHG = free-hand technique group (conventional group); M = mean; RAG = “Orthbot”-assisted group; SD = standard deviation; TRA = trajectory rotation angle.
and the trajectory rotation angle. Comparing with the FH group, the RA group showed no statistical difference in the rate of acceptable trajectories (Grade A and B). But we achieved a significant reduction on the entry point deviation ($p < 0.001$) and the trajectory rotation angle ($p < 0.001$) in the RA group. It indicated that the “Orthbot” system could definitely follow the preoperative plan to drill k-wires precisely for assisting pedicle screw fixation. Meanwhile, without using the axial and sagittal views, it is possible to measure precision parameters on the 3D surgical space coordinate system constructed by the “Orthbot” system, which is a newer concept we came up with in the present study.

**Operation time**

Operation time is an important variable to assess the effectiveness of a novel surgical technique. Theoretically, the robotic system is able to remarkably shorten the duration of the operation because it can directly finish a point-to-point task without overmuch adjustment. Schatto et al. [22], however, found equal operation time between the robot-assisted group and the fluoroscopy-guided group, and Khan et al. [26] obtained the consistent finding when comparing robotic assistance and 3D CT navigation. Similar to several previous research studies, this study also detected a similar outcome in this part for these two groups during the experiment. In our opinion, the lack of a record on time-per-screw placement in both two groups.

**Radiation**

It is widely concerned that radiation exposure is an exact hazard for the patient and the operating room personnel, especially orthopaedists [27]. Some articles confirmed the use of the robotic technique seems to reduce trajectory verification by fluoroscopic images after screw fixation [28,29]. In the present study, average radiation time was 7.29 s per case in the robot-assisted group compared with 9.00 s per operation in the conventional technique group. But we detected average radiation time per screw was significantly lower in the robotic group ($p < 0.05$). The main reason might be that overmuch C-arm exposure was used to inspect the accuracy of registration in the RA group because of inexperience in this new technique. In general, the radiation time per screw in the robotic group was significantly lower, which shows the potential to reduce radiation exposure of pedicle screw fixation assisted by this robotic system when more robot-assisted surgeries were conducted.

**Complication**

Pedicle screw malposition always increases the rate of complication. The major screw-related complications including vascular or visceral damages, neurologic deficits, and cerebrospinal fluid leakage are identified [7]. More attention had been paid on neurologic deficits in previous experiments of free-hand screw insertion technique. Davne and Myers [30] reported the rate of neural injury reached 1.1% in lumbar spine fusion, and Faraj and Webb [31] found a similar result (1.09%).

**Limitation and expectation**

There are some inherent limitations to our study. First, it is not enough to obtain generalizability of the results of our study with a small sample size. A primary reason is that we restrict indications involved in our study to obtain a preliminary evaluation of this robotic system. Deep research with a wider range of indications including cervical spondylosis, thoracic disease, and scoliosis will be required. Furthermore, time-per-screw placement was not documented, which made us fail to accurately estimate the influence on the operation time of this robotic platform. The study was a preliminary assessment of this novel robotic system which performs the pedicle screw insertion with high precision and stability. To our knowledge, this was the first clinical assessment about the accuracy and safety of this novel robotic technique. The unique automatic bone drilling system exhibits a new technology improving the operative technique compared with the conventional robotic system, which brought a new concept for surgical robotic technology changing it to an operative technique more than a guidance machine. Further studies with more cases performed are required to verify the accuracy of this novel system. With new applications or capabilities updated, in the future, the use of this system in spinal surgery clearly offers great promise.

**Conclusion**

We concluded that pedicle screw fixation procedure assisted by the “Orthbot” system was accurate and safe. All screws in the robotic group, with no complication, were determined to be of a clinically acceptable level, and screw accuracy of this robotic system was comparable with the outcomes intervened by experienced surgeons. Furthermore, the execution of the “Orthbot” system following the preoperative planning was more precise than surgeons, and we also found this robotic system has potential to reduce radiation exposure of pedicle screw fixation.

**Ethical approval**

All procedures performed in studies involving human participants were in accordance with the 1964 Helsinki declaration, and the protocol was approved by the Institutional Review Board of Nanfang Hospital, Southern Medical University (IRB no. NFEC-2018-082).

**Conflict of interest**

The study was conducted under the cooperation of the Department of Spinal Surgery, Nanfang Hospital, and Xinjunte Smart Medical Equipment Co. Ltd, Shenzhen. All authors declared that they have no conflict of interests.

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