THE USE OF NITINOL RODS IN SURGICAL TREATMENT OF DEGENERATIVE SCOLIOSIS. 2.5-YEAR FOLLOW-UP

ABSTRACT

Objectives: To compare the outcomes of surgical treatment with lumbar fixation using nitinol rods without fusion and with standard lumbar fixation with titanium rods and interbody fusion. Methods: Treatment results of 70 patients with degenerative lumbar scoliosis aged 40 to 82 were analyzed. In all cases pedicle screws and nitinol rods with a diameter of 5.5 mm were used. Thirty patients underwent fixation at L1-S1 and 40 patients underwent fixation at L1-L5. Spinal fusion was not performed. All patients had radiography, CT and MRI performed. The results were assessed according to the Oswestry scale, SRS 22, SF 36 and VAS. The minimum follow-up period for all patients was 2.5 years. For the control group, consisting of 72 patients, pedicle fixation with titanium rods and interbody fusion in the lumbosacral region were performed. Results: The average level of deformity correction equaled 25° (10° - 38°). The analysis of X-ray and CT-scans revealed a single patient with implant instability, two patients with bone resorption around the screws and one patient with rod fractures. Functional radiography 2.5 years after surgery showed an average mobility of the lumbar spine of 21° (15° - 30°). There were no problems at the adjacent levels. Conclusions: The use of nitinol rods in spinal deformity surgery is promising. This technology is an alternative to rigid fixation. Continued gathering of clinical data and its further evaluation is necessary.

Keywords: Scoliosis; Internal fixators; Orthopedic fixation devices.

ORIGINAL ARTICLE/ARTIGO ORIGINAL/ARTÍCULO ORIGINAL

USO DE HASTES DE NITINOL NO TRATAMENTO CIRÚRGICO DE ESCOLIOSE DEGENERATIVA. ACOMPANHAMENTO DE DOIS ANOS E MEIO

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ABSTRACT

Objetivos: Comparar os resultados de tratamento cirúrgico com fixação lombar usando hastas de nitinol sem arthrodesis e com fixação lombar padrão com hasites de titânio e fusão intersomática. Métodos: Foram analisados os resultados do tratamento em 70 pacientes com escoliose lombar degenerativa com idades entre 40 e 82 anos. Em todos os casos, foram usados parafusos pediculares e hastas de nitinol com diâmetro de 5,5 mm. Trinta pacientes foram submetidos à fixação em L1-S1 e 40 pacientes tiveram fixação em L1-L5. Não foi realizada artrodesis da coluna. Todos os pacientes fizeram radiografias, TC e RM. Os resultados foram avaliados de acordo com a escala de Oswestry, com o SRS 22, o SF 36 e o VAS. O período mínimo de acompanhamento para todos os pacientes foi de 2,5 anos. No grupo controle, com 72 pacientes, realizou-se a fixação do pedículo com hastas de titânio e fusão intersomática no nível lombossacral. Resultados: O nível médio de correção da deformidade correspondeu a 25° (10°-38°). A análise das radiografias e das TC revelou um único paciente com instabilidade, dois pacientes com reabsorção ósea ao redor dos parafusos e um paciente apresentou fraturas da haste. A radiografia funcional 2,5 anos após a cirurgia mostrou mobilidade média da coluna lombar de 21° (15°-30°). Não foram encontrados problemas nos níveis adjacentes. Conclusões: O uso de hastes de nitinol na cirurgia de deformidades da coluna é promissor. Essa tecnologia é uma alternativa à fixação rígida. É preciso manter a coleta contínua de dados clínicos e sua posterior avaliação.

Descritores: Escoliose; Fixadores internos. Dispositivos de fixação ortopédica.
Degenerative scoliosis is a disease that is more common among people in the later stages of life. Its clinical course involves severe pain, radicular damage and loss of sagittal and frontal balance of the body. Due to lateral listhesis of the lumbar vertebrae, spinal stenosis develops at multiple levels. Intermittent claudication may also develop in patients. In addition, patients may also have osteoporosis, which exacerbates the progression and creates treatment difficulties. Conservative treatment is not very effective, so the method of choice is surgical treatment. During surgical treatment, a multilevel transpedicular fixation from the lower thoracic or upper lumbar to the L5 or S1 vertebrae is carried out. When performing fixation below L5, it is recommended to combine sacroc-pelvic fixation with an interbody fusion at L5-S1 and L4-L5.1,2 When conducting a standard rigid fixation, a titanium rod of 5.5 mm or 6 mm in diameter is used. As a result of the fixation and deformity correction, bending decreases, and sagittal and frontal balance improve. Decompression of the spinal canal and the nerve roots is also performed. In most cases, surgery improves the patients’ quality of life. However, when performing surgery for spinal deformity in adults, the percentage of complications can be as high as 53%, with “serious” complications as high as 8.4%. The most common complications of degenerative scoliosis surgery are proximal junctional kyphosis (PJK) (28%), infection (46%), and pseudoarthrosis (10%). According to the literature, 44% of patients will require revision surgery within 5 years.3,4 Furthermore, rigid fixation of the lumbar spine significantly violates its biomechanics, especially if the fixation area includes the sacrum. Because of the rigidity of the titanium rods in the absence of fusion, the implants become loose in the bone, hence the search for new solutions in spinal fixation.

Nitinol, a nickel (55%) and titanium (45%) alloy, is a unique material with shape memory. According to its physical characteristics, its plasticity is eight times greater than titanium. The crystalline structure of nitinol is more durable and has better resistance to cyclic loads. However, there are no analytical articles devoted to its use in the treatment of spinal deformity. Our early surgical experience of using nitinol rods should prove quite informative for spinal surgeons. The main purpose of our work was to determine the effectiveness of the use of nitinol rods for deformity correction and fixation of the lumbar spine region in degenerative scoliosis patients.

Fixation of the lumbar spine was performed without fusion. We then sought to determine whether the motion of the fixed segment is preserved; how common bone resorption around the implant is; what kinds of complications are possible and how often they occur; and how the results of treatment differ between the control group with standard fixation and the group with the use of nitinol.

MATERIALS AND METHODS

Investigation approved by the Roszdravnadzor Ethics Committee (approval number: 18194533). The study was conducted using two groups of patients, all of whom signed an informed consent form. The first group included 70 patients with degenerative lumbar scoliosis, who were operated on using 5.5 mm rods made of nitinol and standard polyaxial pedicle screws without using bone grafts. The second group included 72 patients with the same condition, who were operated on according to the standard procedures. Rigid titanium rods of 5.5 mm in diameter and polyaxial pedicle screws were used. When the level of fixation involved the S1 vertebra, an additional L5-S1 interbody fusion was carried out.

All procedures were carried out between 2011 and 2014 by two surgeons. The first group consisted of patients aged 40 to 82 years. The patients in the second group were aged 39 to 84 years. In the first group, the gender ratio was 37 female to 33 male patients. All patients complained of pain in the lower back, leg pain, and radicular disorders (reduced sensitivity). Some patients had symptoms of myelogenous intermittent claudication (15 patients in group one and 18 in group two). All patients were examined using plain radiography, CT and MRI. All patients were operated on at the Department of Spinal Pathology, CITO named after N.N. Priorov (Moscow, Russia).

Clinical evaluation

Before and after surgery, all patients were tested using several questionnaires: VAS, Oswestry, SF36 and SRS22. The testing was conducted at 3 months, 6 months, 1.5 years and 2.5 years after surgery.

Radiographic picture

In the evaluation of the radiographs, the Cobb angle of deformity and lumbar lordosis were measured, and the global sagittal and frontal balances before and after surgery were evaluated. Functional radiographs were performed 2.5 years after surgery to evaluate the range of motion of the lumbar spine. The Cobb angle was evaluated using radiographs in flexion and extension of the top and bottom vertebrae involved in the fixation. In addition, the range of flexion-extension movements in each fixed segment was measured using the Cobb angle technique. Two and a half years after surgery, CT studies were performed on all patients, to evaluated the presence of bone resorption around the pedicle screws.

Surgical technique

In the first group, the standard posterior approach was carried out. When the pedicle screws were inserted, the intervertebral joints were kept intact. After installing the standard polyaxial screws, decompression of the spinal canal by hemi- and full laminectomy was performed. Next, two nitinol rods, precooled to the activation temperature, were installed. The rods were modeled according to the lordosis of lumbar lordosis (35°–40°). Spinal fusion and bone grafting were not carried out. Fixation was performed between levels L1 and S1 in 30 patients, and between L1 and L5 in 40 patients. In the second group, the standard posterior approach was also carried out. Polyaxial pedicle screws were installed. The spinal canal was decompressed by hemi- and full laminectomy. When involving the S1 vertebra, the interbody fusion was carried out at L5-S1 by the means of posterior lumbar interbody fusion (PLIF) or transforaminal lumbar interbody fusion (TLIF) using cages. The rods were then installed. Additionally, bone grafting was performed on transverse processes and posterior elements of the vertebrae using autografts. Fixation from L1 to S1 was performed in 30 patients, and from L1 to L5 in 42 patients.

Statistical analysis

For statistical analysis IBM SPSS Statistics version 23.0.0.0 was used.

RESULTS

During the analysis of the treatment results, the following data was obtained. The degree of curvature correction of the lumbar spine was similar in both groups. Before the operation, the angle of scoliotic curvature was an average of 35° (± 5°) in group one, and an average of 40° (± 5°) in group two. After the operation, this averaged was 14° (± 5°) in group 1, and 13° (± 5°) in group 2. Analyzing the sagittal profile of the deformation before surgery, lumbar lordosis was an average of 5° (± 3°) in group 1, and an average of 7° (± 3°) in group 2. Lumbar lordosis after surgery was 25° (± 3°) in group 1, and 32° (± 3°) in group 2. Global frontal balance in group 1 was 3 cm (± 1 cm) before surgery, and 0.5 cm (± 1 cm) after surgery, while in group 2 it was 3.5 cm (± 1 cm) before surgery and 0.3 cm (± 1 cm) after surgery. The global sagittal balance in group 1 was 2.5 cm (± 1 cm) before surgery, and ± 0.2 cm (± 1 cm) after surgery, while in group 2 it was 2.7 cm (± 1 cm) before surgery, and 0.3 cm (± 1 cm) after surgery. During the assessment of mobility of the lumbar spine region, the radiographs in flexion and extension showed the following results: lumbar motion from L1 to S1 averaged 50° (± 3°) in group 1, and 58° (± 3°) in group 2. After surgery, motion of the lumbar spine region fixed by nitinol rods averaged 21° (± 3°) in group 1, while in the second group, no motion was observed in the fixed segments. In the functional radiographs, motion of 11° (± 1°) was observed in segment L5-S1 only if it was not included in the zone of fixation.
When analyzing the duration of the surgery in the first group, the average operating time was 185 minutes (± 15 min). In the second group it was more than 243 minutes (± 15 min). This difference in time was due to time-consuming interbody fusion. The average amount of blood lost in group 1 was 400 cc. (± 50 cc.), and in group 2, 700 cc. (± 50 cc.). This difference is due to interbody fusion using the PLIF and TLIF methods, as well as decortication of posterior elements while performing posterior fusion. While carrying out these phases of the operation, an additional loss of blood occurred, usually from the epidural veins and the bone.

The results of the surveys according to different questionnaires were as follows. In group 1, the SF36 physical health questionnaire before surgery produced on average 32.4 points and after surgery, 63.5 points (p<0.05). The mental health questionnaire before surgery produced an average of 42.5 points, and after surgery, 76.3 points (p<0.05). (Figure 1) SRS 22 in the first group showed the following results: Table 1 Oswestry questionnaire data: 64.6 points before surgery and 17.8 (p<0.05) after surgery. (Figure 2) Average VAS before surgery was 9.3, while after surgery, average VAS was 2.4 (p<0.05). (Figure 3, 4, 5 e 6).

In group 2, the average SF 36 index of physical health before surgery was 33.4 points, while after surgery, the average SF was 58.5 points (p<0.05). The average score on the mental health index before surgery was 41.4 points, while after surgery was 72.2 points (p<0.05). The results of the SRS 22 index in the second group were as follows. (Table 1). Oswestry questionnaire data: 64.0 points before surgery, 17.7 points (p<0.05) after surgery. Preoperative VAS score: average of 9.4 before surgery, 2.1 (p<0.05) after surgery.

It is clear that according to various questionnaires, the outcomes 2.5 years after surgery are better in group 1 where nitinol rods were used. Analyzing complications, the following data was obtained. In the first group of patients, 2.5 years after surgery the following complications were recorded: implant instability – 1, bone resorption around the screws – 2, rod fractures - 1. Revision surgery was required in all cases. Two patients had infection-related complications (one superficial and one deep suppuration). In both cases, open wound drainage was used, followed by secondary suturing. Removal of the instrumentation was not required. In the second group of patients, with rigid fixation, the number of complications was higher. Suppuration was observed in one patient. Open wound drainage was used, followed by secondary suturing. Pseudarthrosis was found in 6 patients. Proximal kyphosis was found in 2 patients, and rod fractures in 3 patients. Revision surgery was performed in 12 patients. Thus, in the group in which nitinol rods were used, the results 2.5 years after treatment were better in terms of the basic parameters: blood loss, operative time, number of complications, and quality of life assessment according to various questionnaires.

Table 1. Comparison of preoperative and postoperative (2.5 years) SRS-22 scores for each of the four categories.

<table>
<thead>
<tr>
<th>Group</th>
<th>Preoperative score (mean)</th>
<th>Postoperative score at 2.5 years (mean)</th>
<th>P</th>
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<tr>
<td>Pain</td>
<td>2.4</td>
<td>4.2</td>
<td>&lt;0.05</td>
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<td>2</td>
<td>5</td>
<td>&lt;0.05</td>
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<tr>
<td>Function</td>
<td>2.2</td>
<td>4.6</td>
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<tr>
<td>Mental health</td>
<td>3</td>
<td>4.4</td>
<td>&lt;0.05</td>
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CONCLUSIONS

The use of nitinol rods without fusion in the surgical treatment of degenerative scoliosis showed good results 2.5 years after surgery. Their use preserves motion in the fixed spine, thereby reducing the number of complications commonly seen in rigid fixation. However, we must continue to gather clinical data and study the long-term results.

All authors declare no potential conflict of interest concerning this article.

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