Implant failure following pedicle based dynamic stabilization of the lumbar spine

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Pedicle-based dynamic stabilization (PBDS) devices such as Dynesys are promoted as an alternative and less invasive option for rigid stabilization of one and even more levels of the lumbar spine. Promising features of the Dynesys system, as well as shortcomings, became obvious in several clinical studies. Since 2012, we started using a new PBDS device as an alternative for the Dynesys, to avoid the screw loosening and the kyphosing effect.

The objective is to compare failure rates between the Dynesys and Balan-C type PBDS implant and factors affecting outcome.

In a retrospective study we investigated a total of 90 patients with lumbar pedicle screw dynamic stabilization (a group of 64 patients with Dynesys stabilization is compared to a group of 26 patients with Balan-C stabilization). Mean follow-up was 48 and 38 months, respectively. Using logistic regression analysis the impact of baseline characteristics such as gender, age, body mass index (BMI), indication for surgery, primary or revision surgery, single versus more level surgery, surgeon’s experience and type of the implant on implant failure was analyzed.

We found a statistically significant difference in failure rates between the two systems (13% in the Dynesys group versus 62% in the Balan-C group). In multivariate analysis, type of implant was associated with implant failure (odds ratio : 13).

Our current results call for an optimization of the pre-and post-marketing surveillance of pedicle-based dynamic stabilization.

Keywords: medical device; pedicle based dynamic stabilization; market authorization; failure; product liability.

INTRODUCTION

The last two decades, the range of lumbar spine pedicle screw stabilization techniques was expanded with more dynamic and hybrid techniques. The Dynesys system is promoted as a less invasive technique for stabilization of one and even more levels (1,2,3,4,5). This system offers the possibility to restore disc height and to preserve some mobility. The Dynesys system consists of polycarbonate urethane spacers to limit spinal extension and a polymer cord acting as a tension band to limit spinal flexion combined with a pedicle screw system. It also exists in a dynamic transition option (DTO) variant allowing a traditional rigid fixation at the lower level combined with a dynamic stabilization at the upper level. The Dynesys system was introduced in spine surgery by Dr. Gilles Dubois.
from Toulouse. This system can rely on a follow-up of 22 years of clinical experience (1,2,3,4,5). In 2012 we launched the Balan-C system in our department as a promising alternative for the Dynesys system to avoid the problem of screw loosening and the kyphosing effect (1,2,4). The Balan-C system consists of Polyether ether ketone (PEEK) rods with silicone bumpers, combined with a pedicle screw system. The PEEK-rod-silicone construct is acting as a hinge, allowing a limited motion of the functional spinal unit. This system also exists in different types of lordotic curves and in a DTO variant. To our knowledge, no long-term results are available about the Balan-C system (7). This study will focus on the outcome of the Balan-C system compared to the Dynesys implant.

**MATERIALS AND METHODS**

A total of 90 patients with lumbar pedicle screw dynamic stabilization surgery between January 1, 2006, and December 31, 2015, was investigated in our department of orthopedic surgery (a group of 64 patients with a stabilization of the Dynesys type compared to a group of 26 patients with a stabilization of the Balan-C type). All patients were operated by the same surgeon. We investigated the impact of baseline characteristics such as gender, patient age at the time of the intervention, BMI, indication for stabilization, primary or revision surgery, single versus more level construct and years of experience with stabilizing in case of failure. The aim of the present study was to compare the failure rate between the Dynesys group and the Balan-C group. Characteristics of patients in the Dynesys and Balan-C group were compared using the Kruskal-Wallis test for continuous and the Chi-Square test for categorical variables. Association of gender, age, BMI, indication for surgery, revision surgery, more level surgery, surgeon’s experience and type of implant with implant failure was analyzed through logistic regression analysis. All statistical analyses were performed using SPSS version 24.

**RESULTS**

The failure rate in the Dynesys group was 13%, in accordance with previous research (1,2,5). In the Dynesys group, 8 revisions (13%) were performed, including 7 for screw loosening and 1 for screw breakage following trauma, several years after surgery in a patient with a previously favorable result. The failures occurred all in more level constructs. In the Balan-C group, we found 16 revisions (62%). One revision involved the symptomatic aseptic inflammation around the implant in a hybrid construct. Laboratory research and PET-scan showed signs of infectious process on the implant. Intraoperative culture did not reveal any pathogens. Favorable MRI findings were found both at the fusion and transition level. Another revision related to screw loosening was done in a hybrid construct with satisfactory NMR findings at the fusion-level as well as on the transition level. In either case, the implant could be removed safely.

Revision surgery was needed for rod breakage in 14 cases. In all cases, dynamic stabilization was converted to conventional fusion. Average hospital stay was 2 days, considering it concerned only a partial replacement of the dynamic implant by a rigid rod. Two revisions for breakage were performed for acute back pain after a fall and following a car accident. In both cases, breakage of the connector was diagnosed on CT imaging with reconstruction in 3 planes (3D CT-scan). Before the causative trauma, there was no clinical or radiological evidence of implant failure.

The implant breakage occurred always in the connecting rod section of the Balan-C implant. Breakage was bilateral in 13 of the 14 (93%) cases and 8 times (30%) at the rod portion, and in 19 times (70%) at the dynamic bumper portion. Breakage was found in 3 (21%) single and 11 (79%) double level constructs. The Balan-C system led to a failure rate of 62% (Fig. 1: breakage of a Balan-C implant through the bumper portion; Fig. 2: breakage through the rod portion).

Statistical analysis of baseline characteristics could not reveal a significant difference between the two groups except for surgical experience. This finding can be explained through the history of both series: the Balan-C started since 2012 as an alternative to the Dynesys technique (table 1).

Using logistic regression analysis, the type of the implant was the only parameter that showed
Of the 8 failures in the Dynesys group, 3 (38%) failures were confirmed by X-ray and 4 (50%) by CT-scan. Implant failure was not visible on conventional imaging in only 1 patient (12%) though failure was suspected due to a huge recurring herniated disc on the stabilized segment. The failure was per-operatively discovered as being caused by loosening of the implant.

We had to rely on more targeted medical imaging techniques to demonstrate implant breakage.

Table I. — Baseline Characteristics

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Dynesys</th>
<th>Balan-c</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (mean, yr)</td>
<td>48</td>
<td>39</td>
<td>0.105</td>
</tr>
<tr>
<td>Gender male</td>
<td>35 (55%)</td>
<td>13 (50%)</td>
<td>0.686</td>
</tr>
<tr>
<td>BMI mean</td>
<td>44</td>
<td>48</td>
<td>0.553</td>
</tr>
<tr>
<td>Indication: disc hernia/disc degeneration: dh</td>
<td>34 (53%)</td>
<td>9 (35%)</td>
<td>0.111</td>
</tr>
<tr>
<td>Primary/revision: primary</td>
<td>50 (78%)</td>
<td>22 (85%)</td>
<td>0.485</td>
</tr>
<tr>
<td>Single or more level surg.: more level</td>
<td>50 (78%)</td>
<td>21 (81%)</td>
<td>0.781</td>
</tr>
<tr>
<td>Experience: (mths) (mean)</td>
<td>35</td>
<td>71</td>
<td>0.000</td>
</tr>
<tr>
<td>Follow-up: (mths) (mean)</td>
<td>48</td>
<td>38</td>
<td>0.097</td>
</tr>
<tr>
<td>Failed or not failed: failed</td>
<td>8 (13%)</td>
<td>16 (62%)</td>
<td>0.000</td>
</tr>
</tbody>
</table>

Table II. — Logistic regression model with 8 variables correlated to failure

<table>
<thead>
<tr>
<th>Variables</th>
<th>Odds ratio</th>
<th>95% C.I. lower</th>
<th>95% C.I. higher</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gender</td>
<td>0.679</td>
<td>0.190</td>
<td>2.433</td>
</tr>
<tr>
<td>Age</td>
<td>1.000</td>
<td>0.948</td>
<td>1.055</td>
</tr>
<tr>
<td>BMI</td>
<td>1.030</td>
<td>0.900</td>
<td>1.178</td>
</tr>
<tr>
<td>Indication: DH/DD</td>
<td>3.208</td>
<td>0.843</td>
<td>12.206</td>
</tr>
<tr>
<td>Primary/revision</td>
<td>0.684</td>
<td>0.175</td>
<td>2.990</td>
</tr>
<tr>
<td>One or more level surg.</td>
<td>1.546</td>
<td>0.259</td>
<td>9.238</td>
</tr>
<tr>
<td>Experience (Y)</td>
<td>0.966</td>
<td>0.662</td>
<td>1.409</td>
</tr>
<tr>
<td>Type of surgery</td>
<td>12.825</td>
<td>2.310</td>
<td>71.212</td>
</tr>
</tbody>
</table>
study with the Balan-C system, no screw loosening or breakage was reported (7). As opposed to our current practice, postoperative imaging was limited to conventional. In our series, breakage was not visible on conventional X-ray in 10 of the 14 (71%) cases and the mean time to breakage was 30 months, which may explain the difference in outcome. This finding underscores the importance of a regular follow-up until one year postoperatively followed by a yearly follow-up. In cases with an innovative implant we propose a 5-year follow-up period. Our results are in disagreement with present scientific evidence about PBDS-systems (1,2,3,5).

We scrutinized in a report of the Belgian Health Care Knowledge Centre on the Dynesys system. Promising features of the Dynesys system as prevention of adjacent-level-degeneration have been critically considered. One 4-year follow-up study failed to demonstrate the property of the Dynesys system to prevent adjacent level degeneration. Complications included a revision rate of 21% and a screw loosening rate of 11% (5). These findings are in accordance with Schroeder et al (3). In a recent review failure and iterative surgeries rates following a Dynesys implant were similar to conventional rigid pedicle-screw systems. Apparently, the rate of adjacent segment degeneration (ASD) appeared to be lower supporting the theory of reduced biomechanical stress. The screw loosening rate was 12%. Implant failure, such as spacer and screw breakage was rare (1).

A recent systematic literature review about PBDS-systems, in general, found adjacent segment degeneration in 0 to 30%. Revision surgery for breakage was performed in 9%. Technical failures were design related but also linked with patient-related properties such as poor bone quality. Surgeon’s adherence to strict indications as moderate degeneration, mild degenerative facet arthritis, low-grade spondylolisthesis, segmental instability and dynamic stenosis is of paramount importance. The sagittal balance of the vertebral column and the degree of instability deserve special emphasis (2,4,8). The posterior shift of the center of rotation is an important feature of dynamic stabilization. PBDS-systems must be able to withstand shear forces to prevent failure (2,9,10,11).

Fig. 4. — Implant failure (see arrow) documented on 3D CT-scan

Dynesys screws. In the Balan-C group failure arising from rod breakage was confirmed by X-ray in 4 (29%) patients and by CT-scan in another 4 (29%) cases. All cases needed a 3D CT-scan to confirm the broken Balan-C rod component. Breakage was not visible on X-ray or 3D CT-scan in 6 patients (42%) where it was clinically suspected and confirmed per-operatively. (Fig. 3: implant failure documented on conventional X-ray image by the asymmetric aspect of the left-sided marker in the bumper portion; Fig. 4: implant failure (see arrow) documented on 3D CT-scan).

DISCUSSION

Our results showed a worse result for the Balan-C system. To our knowledge, a failure rate of that size has never been reported before in literature. Only one recent study described the radiological features of a new pedicle based dynamic stabilization (PBDS) system and reports implant breakage in 27% of the cases which is considered as striking failure (10 out of 37 interventions) and lead to retrieval of the implant from the market. The PBDS technique concerned the CD-Horizon Agile system, which has some similar features with the Balan-C system. Consistent with our findings, the authors reported a breakage in the dynamic spacer section of the implant (6). In a recent 2-year follow-up
The literature about PEEK-rods is extremely scarce. PEEK-rods proved both an advantage and a drawback to spinal stabilizing. The advantages include better load-sharing with the lateral column, the slightly greater flexibility of the PEEK-rod biomechanically beneficial in terms of fusion and lower risk of screw breakage at the expense of an increased risk of PEEK-rod breakage (2,7,12,13). In the Balan-C, there is a combination of a PEEK-rod with a PEEK-silicone bumper. In our series, we noted 8 (30%) breakages of the rod and 19 (70%) of the bumper. There is nothing in the literature up-to-date about this. The composite nature of the Balan-C implant does not allow to draw any conclusion on PEEK rods in general.

Though our study was retrospective, the strength of our study includes the large sample size and that the intervention was performed by the same surgeon. Given the young average age of the Balan-C group (39 years), we did not investigate bone quality.

Our study has some policy implications:

The regulatory framework for obtaining market access for high-risk medical implants in Europe differs strongly to the United States where this Balan-C system was never allowed. In the United States, RCT’s are a necessary tool in obtaining market access. To gain market access in Europe, studies that prove safety and performance are mandatory and RCT’s are not necessary (14,15,16). Safety studies are carried out by notified bodies selected by the manufacturer which establishes a relationship of dependency. European healthcare providers are not always aware of the weakness in the pre-clinical evaluation of these implantable medical devices not providing the same level of safety as drugs (16). Testing of new innovative devices should be entrusted to high volume centers with sufficient expertise. Reinforcing post-market surveillance is equally necessary. Council Directive 93/42/EEC of 14 June 1993 on medical devices is a European harmonization law that is transposed into Belgium legislation. Pursuant to article 11 of The Royal Decree of March 18, 1999, the manufacturer, the distributor, the notified body and (para)-medical practitioners have the duty to inform the Federal Agency for Medicines and Health Products (FAMHP) in case of incidents (7,16).
important burden on healthcare in terms of hospital stay and absenteeism. In addition, implant breakage in road accident victims is likely to prove contentious in view of causation analysis. We conclude with following concerns: Breakage of PBDS implants is a main issue for spine surgeons. A regular follow-up during the first postoperative year, followed by a yearly checkup in the next 5 years is recommended. Unexplained pain following PBDS-stabilization requires 3D CT scan for further exploration.

REFERENCES


