Original article

DiscoGel® in patients with discal lumbosciatica. Retrospective results in 25 consecutive patients

A. Léglise a, *, J. Lombard b, A. Moufid a

a Service de chirurgie orthopédique et traumatologique, CHU La Milérié, 2, rue de La Milérié, 86000 Poitiers, France
b Service de chirurgie orthopédique et traumatologique, centre hospitalier Georges-Renon, 40, avenue Charles-de-Gaulle, 7900 Niort, France

A R T I C L E   I N F O

Article history:
Received 22 July 2014
Received in revised form 31 December 2014
Accepted 11 May 2015

Keywords:
Lumbosciatica
Percutaneous
DiscoGel®

A B S T R A C T

Introduction: Discogenic lumbosciatica is a common disorder in patients between 30 and 40 years old. Because of the frequency and socio-professional impact of this entity, it represents a real public health problem. DiscoGel® is a class III medical device used for nucleolysis to avoid discectomy. The goal of this study was to evaluate the effectiveness of this treatment in patients with discogenic lumbosciatica following unsuccessful conservative medical treatment.

Materials and methods: This is a retrospective, single-center study including 25 patients who were treated with DiscoGel® between 2010 and 2011 at Niort Hospital, France. The severity of lumbar and radicular pain was assessed by a verbal numeric scale (VNS) and patient satisfaction. Patients were classified as successes or failures.

Results: Treatment was found to reduce the severity of lumbar pain in 73% and of radicular pain in 80% of patients in the success group. Treatment was a failure in 64% of patients. A comparison of the two groups showed that a preoperative MODIC 2 MRI signal of the adjacent vertebrae end plate was significantly associated with treatment failure (Chi² = 8572, P < 0.01).

Discussion: The VNS for lumbar pain and radicular pain decreased in 42% and 50% of patients respectively after the use of DiscoGel®. In our series, DiscoGel® treatment was unsuccessful for discogenic lumbosciatica in 16 patients. These results do not support others in the literature. A lack of statistical power could partly explain these results. The most important result of this study is found in the subgroup analysis which suggests that indications for DiscoGel® treatment could be modified in the future in relation to preoperative imaging data.

Level of evidence: 4.

1. Introduction

Common discogenic lumbosciatica is frequent in patients between 30 and 40 years old. Between 13% and 40% of the general population will experience an episode of sciatica during his/her life due to discoradicular conflict. Because of its frequency and socio-professional impact of this entity, it is a public health issue.

Percutaneous intradiscal techniques have been developed to provide treatment for discogenic lumbosciatica that is more effective than epidural infiltrations and less invasive than surgery. These techniques result in nucleolysis with a physicochemical action and nucleotomy with a physical action [1–3].

DiscoGel® is a class III intradiscal medical device composed of gelified ethanol associated with tungsten in suspension for nucleolysis. It is injected into the nucleus pulposus to reduce intradiscal pressure. The presence of cellulose, which is a gelling agent, limits the risk of epidural leaks that may occur with pure ethanol. The tungsten particles are used to control progression of the gel in the disc and through any annular fissures by fluoroscopic control. The mechanism of action of DiscoGel® is based on several hypotheses:

- decreasing intradiscal pressure because of the dehydration caused by injection of the product to decrease pain;
- lytic action on new nerve growth in the disc [4];
- necrotic action of alcohol on the nucleus pulposus [5].

In 2007, Théron defined the indications for DiscoGel® use in his study of 202 cases [6]. For this author, DiscoGel® is an alternative to surgical treatment of radicular pain from contained or...
uncontained herniated discs at the cervical or lumbar levels, that have been present for more than 3 months, are resistant to medical treatment with no neurological complications, and which have been confirmed by imaging techniques.

Twenty-eight patients in our center were treated with DiscoGel®. The goal was to evaluate the efficacy of treatment of lumbar radicular pain as well as to compare the success and failure treatment groups to identify any possible factors of failure.

2. Materials and methods

Twenty-eight patients, aged 19 to 58 were treated with DiscoGel® between 2010 and 2011 by the same operator in the same center (Niort Hospital, France).

The initial population included 10 women and 18 men presenting with clinical lumbariscatica due to median, para-median or foraminal ligamentous herniated discs on imaging.

Inclusion criteria were a clinical picture of discal lumbariscatica that was resistant to appropriate conservative medical treatment and confirmed by imaging. Exclusion criteria were a history of surgery of the spine at the involved levels and/or imaging results that did not support the clinical results, the presence of a herniated disc excluded on imaging, involvement of several levels and/or incomplete data collection.

The severity of lumbar and radicular pain was evaluated by a verbal numeric scale (VNS, 0 = no pain; 10 = maximum pain) before treatment and at the postoperative consultation between D30–D45 after injection of DiscoGel® to define the variation in the VNS as a result of treatment.

Patients were contacted by telephone to determine their level of satisfaction with treatment. The patient was asked to rate the outcome of treatment as poor/mediocre/satisfactory/very satisfactory/excellent; and in case of failure to know if additional treatment had been necessary. Seven of the 28 patients could not be reached by telephone, and these data were obtained from the elements in the medical file during the different consultations. Three files were excluded from analysis due to lack of data.

The standardized DRAD questionnaire (Dallas pain self-evaluation [7]) was used to evaluate the repercussions of lumbariscatica on daily and professional activities before and after DiscoGel® treatment.

The characteristics of the study population are presented in Table 1.

Patients were classified into 2 groups:

- in the success group if the VNS decreased by >50% and the patient was satisfied;
- in the failure group if the VNS decreased by \( \leq 50\% \) and/or the patient was dissatisfied.

Table 1

<table>
<thead>
<tr>
<th>Traitement</th>
<th>Médian</th>
<th>10–90%</th>
<th>% des patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>DRAD</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lumbociatica</td>
<td>4,12</td>
<td>2,78–8,85</td>
<td>38,6%</td>
</tr>
<tr>
<td>Radiculaire</td>
<td>4,08</td>
<td>2,46–7,73</td>
<td>37,9%</td>
</tr>
<tr>
<td>DRAD Lumbociatica</td>
<td>4,12</td>
<td>2,78–8,85</td>
<td>38,6%</td>
</tr>
<tr>
<td>DRAD Radiculaire</td>
<td>4,08</td>
<td>2,46–7,73</td>
<td>37,9%</td>
</tr>
</tbody>
</table>

Satisfaction with DiscoGel® treatment was classified as:

- poor;
- mediocre;
- average;
- good;
- very good;
- excellent.

Patients who responded poor, mediocre or average were considered to be dissatisfied with treatment while those who responded good, very good or excellent were considered to be satisfied.

Treatment was considered to have failed in the following situations:

- decrease in VNS < 50%);
- patients who underwent a secondary procedure (infiltration or surgery);
- patients who classified the results of treatment as poor, mediocre or average even if they did not undergo a secondary procedure.

The study population was divided into two groups based on the results: patients for whom the DiscoGel® procedure was successful and those for whom it was a failure. These two groups were analyzed and compared to identify any prognostic factors of failure.

The DiscoGel® injection protocol was the following (Figs. 1 and 2):

- outpatient procedure;
- in the surgical block under strictly aseptic conditions;
- under local and neuroleptic anesthesia;
- with the patient lying on the stomach;
- under fluoroscopic control;
- 18G needle;
- slow percutaneous injection of DiscoGel® 0.1 mL/30 seconds;
- progression of the product controlled by fluoroscopy;
- a single operator;
- with three hours of postoperative surveillance.

3. Results

There were no complications during the procedure, and no leaking of the product. Fifteen of the 25 patients (60%) seen at follow up were on sick leave at the first postoperative consultation due to lumbosciatica pain.

Treatment with DiscoGel® was considered to be a failure in 16 patients, or 64% of the population. Fourteen of these 16 failures underwent secondary treatment. A total of 56% of the population
underwent either foraminal infiltration, a cure for a herniated disc or arthrodesis. Two out of 16 failures did not undergo secondary treatment but rated the DiscoGel® treatment as mediocre or poor, respectively.

The VNS for lumbar pain in the 25 patients who were evaluated in the study went from 5.17 to 3.00 or a decrease of 42%. The VNS for radicular pain went from 6.87 to 3.43 or a decrease of 50%. Nevertheless, the decrease in VNS was still ≤50%, the threshold chosen for the treatment to be considered a success (Figs. 3 and 4).

Two groups can be identified in this study:

- a group of 9 patients in whom DiscoGel® was a success because the lumbar and radicular VNS decreased by >50% AND the patient was satisfied with treatment;

- a group of 16 patients in whom DiscoGel® treatment was a failure because the lumbar and radicular VNS decreased by ≤50% and/or the patient stated that s/he was dissatisfied.

In the group in which treatment failed 12/16 patients underwent secondary surgery (Fig. 5.pdf). The characteristics of the two groups are presented in Table 2.

In the satisfied group, the decrease in VNS for lumbar pain was 73% and for radicular pain was 80% compared to the dissatisfied group in which the lumbar VNS decreased by 21% and the radicular VNA by 26% (Fig. 6).

There were more men in the success group, but this difference was not significant. On the other hand, a significant number of patients had a MODIC 2 sign in the failure group (Chi² = 8.572, P < 0.01). In other words, a disc with fatty degeneration was associated with failure of DiscoGel® treatment. We did not find any other significant factors to explain our failures.

4. Discussion

DiscoGel® is a new minimally invasive therapeutic technique for the treatment of discogenic lumbar sciatica and an alternative to invasive surgery in case of unsuccessful medical treatment.

![Fig. 2. (Ph Brunner): mode of injection.](image)

![Fig. 3. Results of treatment in the study population.](image)

![Fig. 4. Variation in lumbar and radicular VNS before and after treatment for the entire study population.](image)

![Fig. 5. Secondary treatments in the failure group.](image)

![Table 2. Characteristics of 2 subgroups.](table)

Data in bold correspond to significant differences.
signal and treatment efficacy. In our study, the presence of fatty disc degeneration, presenting as a preoperative MODIC 2 MRI signal was significantly associated with failure. None of the existing studies have taken into account this factor, which our results seem to indicate is a potential prognostic parameter of treatment failure. Treatment with DiscoGel® is being evaluated and is an alternative to invasive procedures, which is why it is so important to determine its specific field of action.

5. Conclusion

DiscoGel® is an effective treatment for discogenic lumbosciatica. The high rate of failures in our study was significantly related to the presence of peridiscal fatty degeneration (MODIC 2). Further studies are therefore needed to confirm this hypothesis in a larger group and to confirm this feature as a potential prognostic factor of treatment failure.

Disclosure of interest

The authors declare that they have no conflicts of interest concerning this article.

References