

**Change in patients' psychophysical performance following lumbar discectomy
relative to the postoperative rehabilitation programme**

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42 **Abstract**

43 **Background.** The aim of this study was to assess the change in psychophysical performance of
44 patients after lumbar discectomy in relation to the postoperative rehabilitation programme.

45 **Material and Methods.** The study involved 60 participants randomly divided into two groups of
46 30 individuals. Both groups participated in a basic version of the rehabilitation programme, and
47 individuals in the study group additionally received manual therapy. The evaluation was performed
48 twice, before the start and after the completion of the 3-month rehabilitation programme. The tests
49 were carried out to measure static balance, functional status using Oswestry Disability Index (ODI)
50 and Roland-Morris Disability Questionnaire (RMDQ), lumbar spine range of motion using the
51 original Schober's test and the intensity of pain using the Visual Analogue Scale (VAS).

52 **Results.** Both groups showed significant improvement in most of the psychophysical parameters
53 assessed (study versus control $p<0.01;p<0.05$), except for the parameters of balance. In the study
54 group, significant changes occurred in all parameters except X average, Area circular and average
55 velocity in trials with eyes closed and left leg stance after rehabilitation ($p<0.05$). The findings
56 showed significant differences in the reduced pain intensity on the VAS ($p=0.0001$), improved
57 functional status in ODI and improved static balance ($p<0.01$), in favour of the study group.

58 **Conclusion.** The protocol which additionally included manual therapy was found to be more
59 effective than the basic programme. Its superiority was reflected by greater pain reduction, more
60 visibly improved functional status as well as improved static balance.

61 **Keywords:** Lumbar, rehabilitation, therapy, pain
62
63

64 **Introduction**

65 Lumbar pain is one of the most common problems leading to reduced quality of life. Its conse-
66 quences include decrease in physical activity, which adversely affects the production of hor-mones
67 in the adrenal cortex and contributes to poor mood, possibly even leading to depression [22]. In
68 many of these patients, a herniated nucleus pulposus (HNP) is the underlying cause of pain in the
69 sacrum area. Conservative treatment is effective in a vast majority of patients with HNP. A surgical
70 procedure is necessary for a much smaller percentage of patients, although in absolute numbers
71 this is still a large group of patients. The most typical surgery in the case of HNP is lumbar
72 _ discectomy which is one of the most commonly performed procedures in neuro-surgery and
73 spine surgery in general [22]. It is highly effective in reducing HNP related pain, known as sciatica.
74 Approximately 30,000 procedures are performed annually in Poland, com-pared to 90,000 in South
75 Korea [13]. Revision procedures four years after discectomy are per-formed at a rate of 7.1-12%
76 [13, 24].

77 Functional outcomes of the surgery can be improved by well-designed postoperative exercise, as
78 shown by a systematic review listed in the Cochrane Database [28]. However, many patients in
79 Poland and in other countries receive little or no formal rehabilitation after the surgery [36].
80 Moreover, the problems faced by the surgeons performing the procedure and the physiotherapists
81 taking care of the patients after the surgery are associated with the lack of standards or recom-
82 mendations for postoperative rehabilitation. Additionally, inadequate postoperative rehabilitation
83 can thwart the positive effect of the surgical procedure. In clinical practice, there are no evi-dence-
84 based protocols, available to surgeons or physiotherapists, for rehabilitation after spinal surgery.
85 The scientific literature in this area is scarce [1, 2, 6, 9, 11, 14, 32, 26, 27]. There are a few
86 randomised clinical studies investigating the effects of rehabilitation on the functional performance
87 of patients fol-lowing discectomy as well as their return to work [1, 2, 6, 9, 11, 14, 32, 26, 27].
88 However, no research reports assess the effects of kinesiotherapy combined with elements of
89 manual therapy and physical therapy which are investigated in the present study.
90 The aim of this study was to assess the change in psychophysical performance of patients after
91 lumbar discectomy in relation to the postoperative rehabilitation programme.

92

93 **Materials & Methods**

94 **Participants and setting**

95 The study was conducted in the Neurosurgery Clinical Hospital of the University of Rzeszów, St.
96 Luke's Regional Hospital in Tarnów and in Reha Medica Rehabilitation Centre in Tarnów, Poland.
97 Sixty participants enrolled for the study were randomly divided into two groups, each with 30
98 individuals. Both groups participated in the basic version of the rehabilitation programme, and
99 individuals in the study group additionally received manual therapy.

100 The following inclusion criteria were applied: diagnosed intervertebral disc injury in L-4-L5 or
101 L5-S1 segments, no previous surgical intervention, age in the range of 18-65 years, no
102 comorbidities or neurological deficits, no active cardiovascular disease, and no contraindications
103 to administration of physical medicine or manual therapy. The exclusion criteria were defined as
104 follows: lack of informed consent to participate in the study, complications following the surgery,
105 poor exercise tolerance (dizziness, nausea, reported decline of daily functioning), postoperative
106 infections, and damaged nerves of the central nervous system.

107 **Ethics**

108 The experimental conditions were in accordance with the Helsinki Declaration, and all participants
109 provided informed written consent to participate in the study. Approval to conduct the study was
110 obtained from the Bioethics Commission at the University of Rzeszow, Poland on April 12, 2018
111 (No 4/12/2018), and all methods were performed in accordance with the relevant guidelines and
112 regulations.

113 **Procedures**

114 The participants were randomly assigned to the study group (n=30) and the control group (n=30).
115 Randomisation was performed using the software MATLAB (MathWorks, Inc. 2018,
116 Massachusetts) with RARtool interface. The person responsible for randomization also managed
117 the list of patients divided into groups and informed the physiotherapist in charge of training which
118 patients were assigned to the experimental group and which to the control group. The individual
119 administering the exercises did not participate in patient examinations or assessments. Initial and
120 final assessments were conducted by a physiotherapist who was blinded to the subjects' group
121 assignments and had no involvement in their training. The list of patients divided into groups was
122 maintained by the researcher overseeing randomization. This list was decoded after the final
123 assessment of the last qualified patient during the final examination. Prior to the start of the
124 program, the research team was instructed not to disclose any information regarding assessments
125 or the course of training. The rehabilitation programme applied in the control group comprised the
126 standard physiotherapy procedure, physical therapy and exercise performed at home. The
127 rehabilitation programme applied in the study group comprised the standard physiotherapy
128 procedure, physical therapy, elements of manual therapy and exercise performed at home. The
129 evaluation was performed twice, before the start and at the end of the rehabilitation, i.e., after three
130 months.

131 **Rehabilitation protocol**

132 Postoperative rehabilitation was initiated on the 15th day after the surgery and was continued for
133 three months, with 40-minute therapy sessions held twice a week. Until then, patients performed
134 exercise at home following instruction received before they were discharged from the
135 Neurosurgery Ward. These included: abdominal and thoracic breathing exercises; active and
136 passive lower limb exercises; upper and lower limb exercises; learning to stand up; learning to
137 stand upright.

138 The rehabilitation programmes were designed to comprise a few components, in the study group -
139 kinesitherapy, physical therapy, home exercise, and elements of manual therapy, and in the control
140 group - kinesitherapy, physical therapy, and home exercise. Kinesitherapy (identical in both
141 groups) included the following: plantar flexion and dorsiflexion of the foot; knee flexion; extension
142 on a chair; walking on all fours; tightening the transverse abdominal muscle - deep stabilisation;
143 hip muscle inhibition; stretching the gluteal muscles and piriformis muscle; pelvic mobilisation
144 through stretching of the multifidus muscle; adductor ball squeeze; weighted breathing; drawing
145 the knees to the chest; eccentric training of the quadriceps muscles; concentric and eccentric
146 training of the gastrocnemius muscle and the tibialis anterior muscles using TheraBand; as well as
147 lumbar flexion and extension using exercise ball. Physical therapy (identical in both groups)
148 included the following: local laser treatment; low-frequency magnetic field; local cryotherapy;
149 electrotherapy: TENS in the operated area.

150 Manual therapy (only in the study group) included the following elements.

151 Myofascial techniques focused on the multifidus and oblique abdominal and small pelvic muscles
152 and aimed to relax tensions and improve spinal stabilization (15% of the duration of a single
153 therapy unit) [6,30].

154 Hip joint mobilization focused on improving mobility and stability of the hip joint in order to
155 increase the lower-body capacity (15% of the duration of a single therapy unit) [16, 19, 39].

156 Post-isometric relaxation (PIR) was applied to improve activation of the transversus abdominis
157 and multifidus muscle particularly in the context of lumbar spine stabilization. The technique was
158 designed to increase range of movement and improve spinal stability. Majority of the patients had
159 an abnormal posture and often maintained a sitting position, which led to muscle tension disorders
160 particularly of the transversus abdominis and in the multifidus muscle [23]. Post-isometric
161 relaxation produces such positive effects as reduced muscle tension and increased range of motion,
162 which is particularly important in the treatment of patients with back pain.

163 The duration of this intervention accounted for 15% of the total time per therapy session [17, 23,
164 29, 37].

165 Trigger point therapy focused on trigger points on the gluteal muscles (small, medium, large),
166 piriformis muscle, muscles in the ischial and tibial group and multifidus muscle, in order to relieve
167 the pain and improve mobility (15% of the duration of a single therapy unit) [30, 31].

168 Fascial techniques were applied in the area of the cervical, thoracic and lumbar spine and in lower
169 extremities, in order to improve tissue flexibility and relieve tension (20% of the duration of a
170 single therapy unit) [5].

171 Diaphragm mobilization aimed to improve respiratory function and core stability (10% of the
172 duration of a single therapy unit) [33].

173 Neuromobilization of the sciatic nerve aimed to improve nerve mobility, which can help reduce
174 pain and improve lower limb muscle function (10% of the duration of a single therapy unit) [18].

175 During the clinical trials no significant adverse events or complications were identified in the
176 groups investigated. Regular monitoring of the participants' health status and their active
177 cooperation with the treatment team made it possible to detect any possible adverse symptoms
178 early, however none of the patients reported health problems.

179 **Outcome measures**

180 Effectiveness of the rehabilitation programme was evaluated by measuring: the static balance on
181 the force plate from Advanced Medical Technology Inc.(AMTI) [34]; the functional performance
182 using the Oswestry Disability Index (ODI) [7] and Roland-Morris Disability Questionnaire
183 (RMDQ) [4, 10, 12]; lumbar spine range of motion with the original Schober's test [18]; and the
184 intensity of pain on the Visual Analogue Scale (VAS) [34].

185 The measurement of static balance performed using AMTI involved continuous assessment of the
186 Centre of Pressure (COP) of the foot. The analyses took into account the measures of the Average
187 Load Point Y determining the anterior–posterior coordinates Y (Y average, in cm), the Average
188 Load Point X, determining the lateral coordinates X (X average, in cm), Average COP velocity (V
189 average, in cm/s), Path Length (cm) of the COP measured during the trial, and Area Circular, i.e.,
190 the area defined by the COP during the trial (cm²). Stabilography measurements, 30 seconds each,
191 were performed in course of trials with double-leg stance and eyes open/closed, and with single-
192 leg stance (right/left leg) with eyes open/closed. To avoid fatigue, the trials were separated with
193 intervals of 30 seconds. To minimise any interferences or noise, the assessments were performed
194 in a closed room. During the trials, the participants were instructed to stand on the platform and
195 focus their gaze on a red target placed in front on the wall, at a distance of 4 feet (1.2 m). The force
196 plate AMTI employed in the present study has been shown to be a valid instrument of a gold
197 standard quality [34].

198 The participants' functional status was performed using: Oswestry Disability Index (ODI) [7] and
199 Roland-Morris Disability Questionnaire (RMDQ) [4, 10]. ODI contains ten questions, each with
200 six answers scored from 0 to 5. When completing the questionnaire, the patient answers questions
201 related to pain intensity, independence, lifting objects, walking, sitting, standing, sleeping, social
202 life, work, housework and travel. The total score for all ten items can range from 0 to 50, zero
203 points reflecting the poorest and 50 points showing the best functional performance in the daily
204 live [4]. RMDQ is a self-report tool comprising 24 items designed to measure pain-related
205 disability associated with low back pain. The questions address the patient's activity, pain,
206 dependence on others, and emotional status. Items are scored either 0 (if left blank) or 1 (if
207 endorsed), and the total RMDQ score is in the range between 0 (corresponding to 'no disability')
208 and 24 (reflecting 'maximum disability') [4, 10].

209 Lumbar spine range of motion was assessed using Schober's test, which is performed with a tape
210 measure held over the spine between the lumbosacral junction and 10 cm above it [12]. The
211 intensity of pain was assessed with the Visual Analogue Scale (VAS), consisting of a 10cm line,
212 with two end points representing 0 ('no pain') and 10 ('pain as bad as it could possibly be'). The
213 patient is instructed to rate their current level of pain by placing a mark on the line [12, 35].

214 **Sample size**

215 The target sample size of 60 participants was selected based on the value computed using a
216 minimum sample size calculator. It was assumed in the calculations that 30,000-50,000 lumbar
217 discectomy procedures are performed annually in Poland, and 80% power of the test was adopted
218 in calculating the minimum sample size. An assumption of 80% was made, as this is the lowest
219 accepted statistical power which, in the case of a low number of participants, allows for the results
220 to be extrapolated to a wider population. With an assumed 15% drop-out rate during the initial
221 examination and a predicted 20% rate of those who may not complete the programme, it was
222 determined that 60 participants would be enrolled for the intervention. These assumptions are
223 based on experience of other researchers [35].

224 **Data analysis**

225 The acquired data were subjected to statistical analyses computed using the software IBM SPSS
226 23. The distribution of the variables was calculated using the Kolmogorov-Smirnoff test ($n > 100$).
227 Measures of location and dispersion were calculated depending on the agreement between the
228 distribution of the variables and the theoretical distribution. The differences between the groups in

229 the first measurement point were examined using the Mann-Whitney test, whereas the differences
230 between the first and the second measurement were examined using Wilcoxon test. Effectiveness
231 between groups was calculated using Student's t-test.

232

233 **Results**

234 **Flow of included and patients' characteristics**

235 146 patients were examined consecutively upon admission to the Neurosurgery Clinical Hospital.
236 Sixty of these patients met the inclusion criteria. Out of the 86 patients who did not qualify for the
237 program, 140 did not meet the inclusion criteria, and 6 refused to participate (Figure 1). All eligible
238 patients participated in therapy sessions and completed the program. No adverse medical events

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239 occurred during the program, and all participants completed the final examination. Recruitment
240 for the study began in January 2019 and lasted for six months.

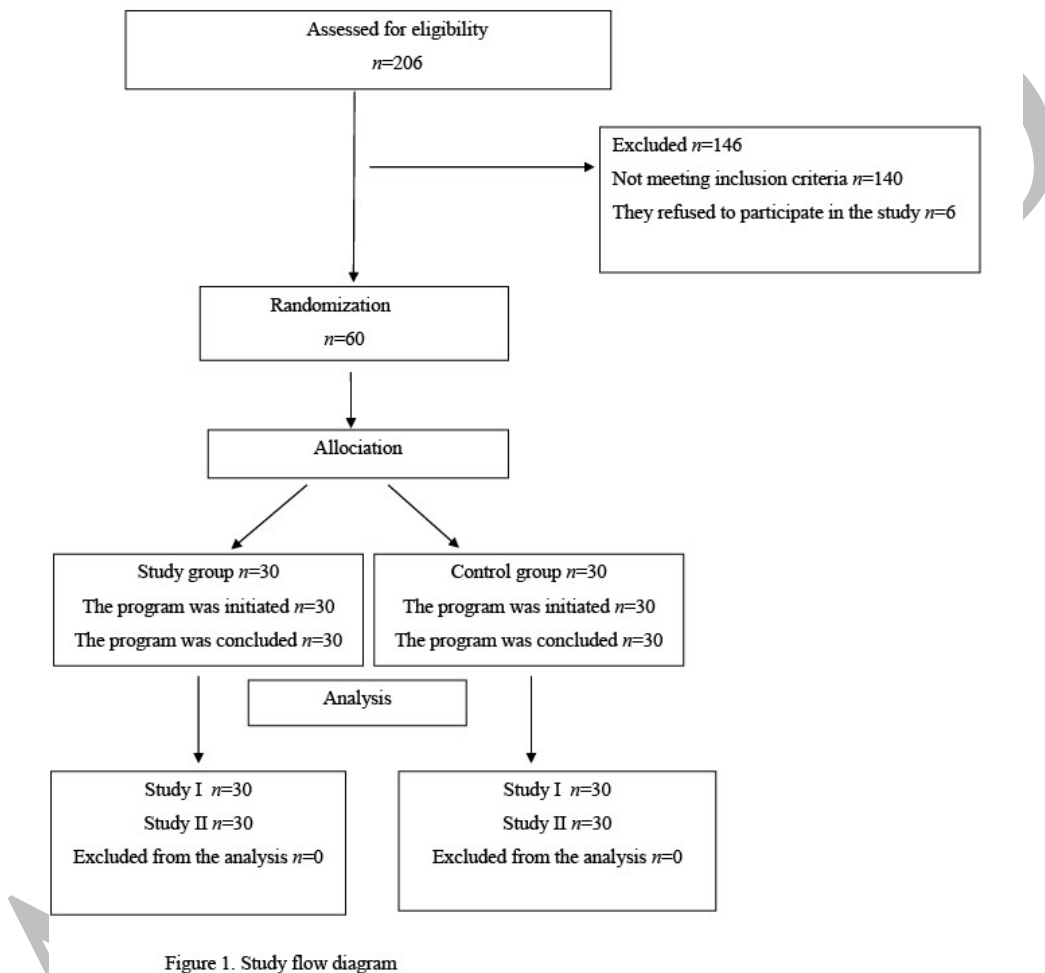


Figure 1. Study flow diagram

241

242

Figure 1. Study flow diagram

243 Patients in the study group (n=30) and the controls (n=30) were matched for the demographic
 244 characteristics. The mean age of patients was 47±15 years in the study group, and 48±14 years in
 245 the control group. Each group comprised 18 female and 12 male participants. In terms of
 246 anthropometrics (age, height, weight, BMI), there were no statistically significant differences
 247 between the groups. The characteristics of the study participants are shown in Table 1

248 Table 1 Characteristics of study participants

Variable	Study group N=30	Control group N=30	Z	P
Age (years) X/SD	47/15	48/14	0.761	0.447
Body height (cm) X/SD	166/12	167/16	0.861	0.541
Body weight (kg) X/SD	62/4	64/3	0.653	0.365
BMI (kg/m ²) X/SD	26.3/13	25.8/9	0.871	0.531

X- mean; SD - standard deviation; Z – Mann-Whitney test; p - statistical significance

249

250

251 **Homogeneity of the patient groups (differences between the study group and the control**
 252 **group before the start of rehabilitation)**

253 No statistically significant differences were found between the groups prior to the rehabilitation
 254 programme, in pain intensity reflected by VAS, functional status assessed with ODI and RMDQ,
 255 spine range of motion measured using Schober's test, as well as most balance parameters
 256 (p>0.05),(Table 2).

257 Table 2. Results of the measurements of patients' capacities in the initial examination before the
 258 start of the rehabilitation programme in the study and the control group

Variable	Control group N=30	Study group N=30	Z	P
VAS [points] X/SD	6/3	6/3	0.398	0.691
ODI [points] X/SD	26/10	27/4	0.637	0.608
RMDQ [points] X/SD	12.9/1.78	14.01/2.1	0.1451	0.781
Schober's Test [cm] X/SD	11/2	12/1	2.116	0.1521
Area circular EO [cm ²] X/SD	2.89/1.9	2.91/2.18	0.33	0.76
Path length EO [cm] X/SD	120.6/21.8	129/22.9	1.67	0.07

Area circular EO [cm ²] X/SD	2.19/1.44	2.18/1.53	-0.02	0.982
Path length EO [cm] X/SD	33.41/9.03	33.36/8.97	-0.03	0.976
Area circular EC [cm ²] X/SD	3.90/2.11	2.67/1.67	-2.24	0.025
Path length EC [cm] X/SD	59.00/36.82	38.34/16.79	-2.01	0.044
Area circular EOSR [cm ²] X/SD	7.39/1.82	9.44/8.17	-0.56	0.574
Path length EOSR [cm] X/SD	99.98/33.40	99.23/23.99	0.98	0.326
Area circular EOSL [cm ²] X/SD	7.68/2.21	8.73/8.27	-0.09	0.929
Path length EOSL [cm] X/SD	96.22/28.86	88.27/27.43	-0.16	0.871
Area circular ECSR [cm ²] X/SD	7.59/3.02	7.47/1.18	0.98	0.329
Path length ECSR [cm] X/SD	111.03/27.82	111.92/32.14	-0.38	0.701
Area circular ECSL [cm ²] X/SD	7.10/1.58	7.15/1.87	0.23	0.819
Path length ECSL [cm] X/SD	89.61/26.21	89.61/15.09	1.06	0.290
X- mean; SD – standard deviation; VAS - Visual Analogue Scale; ODI - Oswestry Disability Index; RMDQ - Roland-Morris Disability Questionnaire; EO – Eyes Open; EC – Eyes Closed; EOSR - Eyes Open Right Leg; EOLL - Eyes Open Left Leg; ECRL - Eyes Closed Right Leg; ECLL - Eyes Closed Left Leg; Z – Mann-Whitney test; p - statistical significance				

259

260 Results of the study group before and after rehabilitation

261 The statistical analysis showed significant decrease in the intensity of pain on VAS ($p < 0.01$),
 262 significant improvement in functional performance assessed with ODI ($p < 0.05$) and RMDQ
 263 ($p < 0.01$), and significantly increased spine range of motion in Schober's test ($p < 0.05$), in the study
 264 group after rehabilitation (Table 3).

265 Table 3. Results of the study group before and after rehabilitation

Variable	Study group before rehabilitation N=30	Study group after rehabilitation N=30	Z	P
VAS [points] X/SD	6/3	3/2	4.718	0.0041
ODI [points] X/SD	27/4	25/12	3.819	0.021
RMDQ [points] X/SD	14.01/2.1	11.4/1.8	4.781	0.00417
Schober's Test [cm] X/SD	12/1	14.4/2.1	4.991	0.01993

X- mean; SD – standard deviation; VAS - Visual Analogue Scale; ODI - Oswestry Disability Index; RMDQ - Roland-Morris Disability Questionnaire; Z- Wilcoxon test statistic; p - statistical significance

266

267 In the study group, statistically significant differences were observed between pre- and post-
 268 therapy measurements in most of the balance parameters evaluated ($p < 0.005$). The least significant
 269 differences in the results acquired before and after the therapy were found in the trials with Eyes
 270 closed, left leg; in this case statistically significant differences ($p < 0.005$) were only identified in
 271 the values Y average and Path length, (Table 4).

272 Table 4. Analytical results of balance measurement in the study group before and after
 273 rehabilitation

Study group		Before			After			Z	P
		Mean	Median	SD	Mean	Median	SD		
Eyes open, both legs	X average	-0.27	-0.45	1.20	-0.09	-0.05	1.05	1.35	0.176
	Y average	-3.41	-3.08	3.47	-2.36	-2.18	3.75	2.36	0.018
	Area circular	2.18	1.84	1.53	1.99	1.83	1.21	2.39	0.016
	Path length	33.36	33.80	8.97	31.98	31.12	8.65	3.35	0.001
	V average	1.21	1.19	0.34	1.18	1.18	0.24	2.27	0.023
Eyes closed, both legs	X average	0.58	-0.29	3.03	0.37	-0.29	2.95	1.36	0.173
	Y average	-2.69	-2.88	1.96	-2.05	-2.06	2.12	2.93	0.003
	Area circular	2.67	2.39	1.67	2.33	2.15	1.28	2.89	0.003
	Path length	38.34	40.31	16.79	36.92	38.33	16.41	4.19	<0.001
	V average	1.50	1.37	0.77	1.46	1.34	0.51	1.24	0.213
Eyes open, right leg	X average	5.85	7.13	3.15	5.30	6.56	2.85	4.01	<0.001
	Y average	-1.89	-1.22	4.07	-1.72	-1.21	3.89	1.98	0.046
	Area circular	7.39	6.90	1.82	7.06	6.78	1.79	2.93	0.003
	Path length	99.98	100.27	33.40	95.08	98.54	29.40	2.93	0.003
	V average	3.54	3.53	1.28	3.35	3.47	1.05	1.78	0.075
Eyes open, left leg	X average	-6.47	-7.39	3.23	-5.87	-6.17	2.83	3.34	0.001
	Y average	-2.16	-1.41	4.06	-1.88	-1.41	3.18	1.42	0.154
	Area circular	8.73	7.13	8.27	8.43	7.27	8.19	1.86	0.061
	Path length	88.27	94.77	27.43	82.51	89.39	23.46	3.82	<0.001
	V average	3.50	3.34	1.32	3.29	3.29	1.08	2.39	0.017
Eyes closed, right leg	X average	7.08	7.66	1.89	6.85	7.04	1.80	2.52	0.012
	Y average	-0.63	-0.62	1.32	-0.51	-0.62	1.06	1.99	0.046
	Area circular	7.47	6.88	1.18	7.26	6.74	1.05	2.36	0.017
	Path length	111.92	104.00	32.14	109.17	99.29	32.49	2.98	0.003
	V average	3.73	3.47	1.07	3.68	3.36	1.10	1.82	0.067
Eyes closed, left leg	X average	-6.16	-4.81	2.57	-5.84	-4.81	2.23	1.85	0.062
	Y average	-0.36	-2.32	2.91	-0.27	-2.06	2.83	1.99	0.046
	Area circular	7.15	8.13	1.87	7.03	7.77	1.64	1.82	0.068
	Path length	89.61	82.12	15.09	87.73	82.12	13.93	2.54	0.011

	V average	2.99	2.74	0.50	2.88	2.74	0.61	1.75	0.079
Z- Wilcoxon test statistic; SD - standard deviation; X average - average load point X which determined lateral coordinates X (cm); Y average - average load point Y which determined the anterior–posterior coordinates Y (cm); V average - average COP velocity (cm/s); p significance level; p < 0.05 reflects statistically significant relationship; p < 0.01 reflects highly significant relationship; p < 0.001 reflects very highly significant relationship									

274

275 **Results of the control group before and after rehabilitation**

276 The statistical analysis showed significant decrease in the intensity of pain on VAS (p<0.05),
 277 significant improvement in functional performance assessed with ODI (p<0.05) and RMDQ
 278 (p<0.05), and significantly increased spine range of motion in Schober’s test (p<0.05) in the
 279 control group after rehabilitation (Table 5).

280 Table 5. Results of the control group before and after rehabilitation

Variable	Control group before rehabilitation N=30	Control group after rehabilitation N=30	Z	P
VAS [points] X/SD	6/3	4.7/1.2	3.814	0.0371
ODI [points] X/SD	26/10	23/12	1.861	0.041
RMDQ [points] X/SD	12.9/1.78	10.8/1.51	7.991	0.031
Schober’s Test [cm] X/SD	11/2	13.6/1.8	6.113	0.0213
X- mean; SD – standard deviation; VAS - Visual Analogue Scale; ODI - Oswestry Disability Index; RMDQ - Roland-Morris Disability Questionnaire; Z- Wilcoxon test statistic; p - statistical significance				

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282

283 In the control group statistically significant differences were observed between pre- and post-
 284 therapy measurements only in a few balance parameters (p<0.005). Statistically significant
 285 differences between the results of measurements before and after the therapy were found in the
 286 trial with eyes closed and double-leg stance, where the differences were identified only in the
 287 values of Y average and Path length (p<0.005). In the trials with eyes open and right leg stance
 288 there was a difference in the value of Area circular, whereas in the trials with eyes open and left
 289 leg stance differences were found in the values of Y average, Path length and V average (p<0.005).
 290 Corresponding results were observed in trials with eyes closed and left leg stance, in the parameters
 291 Path length and V average (p<0.005), (Table 6).

292 Table 6. Analytical results of balance measurement in the control group before and after
 293 rehabilitation

Control group		Before			After			Z	P
		Mean	Median	SD	Mean	Median	SD		
Eyes open, both legs	X average	-0.17	-0.16	1.15	-0.22	-0.45	1.33	0.53	0.592
	Y average	-3.51	-3.19	3.42	-3.41	-3.08	3.47	1.1	0.273
	Area circular	2.19	1.84	1.44	2.18	1.84	1.53	0.1	0.916
	Path length	33.41	32.95	9.03	33.36	33.80	8.97	0.53	0.592
	V average	1.26	1.26	0.37	1.21	1.19	0.34	1.46	0.144
Eyes closed, both legs	X average	0.40	0.57	1.49	0.94	0.34	2.68	1.49	0.136
	Y average	-4.69	-4.32	5.25	-2.43	-2.72	2.95	2.06	0.039
	Area circular	3.90	3.45	2.11	3.57	3.03	2.08	1.34	0.179
	Path length	59.00	44.27	36.82	66.17	49.17	39.17	1.6	0.108
	V average	2.21	1.64	1.31	2.21	1.64	1.31	0	1
Eyes open, right leg	X average	3.99	2.75	19.96	4.05	2.75	19.91	0	1
	Y average	-1.22	-1.26	2.71	-0.95	-1.26	2.11	0	1
	Area circular	9.44	7.05	8.17	8.45	7.05	3.60	0	1
	Path length	99.23	97.58	23.99	116.39	108.28	35.30	2.59	0.009
	V average	3.62	3.55	0.94	3.86	3.68	1.20	1.6	0.108
Eyes open, left leg	X average	-6.52	-6.31	3.59	-4.95	-5.81	8.19	0	1
	Y average	-2.16	-1.67	4.07	-0.78	-0.67	3.21	2.1	0.036
	Area circular	7.68	7.05	2.21	12.61	7.45	18.70	1.82	0.067
	Path length	96.30	92.22	28.86	115.54	109.62	36.38	2.52	0.012
	V average	3.47	3.27	0.96	3.92	3.66	1.24	2.02	0.043
Eyes closed, right leg	X average	7.06	7.38	2.32	6.60	7.29	2.82	1.6	0.108
	Y average	-0.65	-1.17	2.43	-1.30	-0.62	3.31	0	1
	Area circular	7.59	6.78	3.02	7.69	6.78	2.90	0	1
	Path length	111.03	106.81	27.82	114.69	110.69	29.19	1.34	0.179
	V average	3.82	3.69	0.97	3.82	3.69	0.97	0	1
Eyes closed, left leg	X average	-6.80	-7.39	3.07	-6.80	-7.39	3.07	0	1
	Y average	-0.54	-0.46	2.49	-1.20	-0.95	3.41	1.34	0.179
	Area circular	7.10	6.93	1.58	8.98	7.58	8.00	1.6	0.109
	Path length	89.61	82.12	26.21	109.70	98.46	37.66	2.52	0.012
	V average	2.93	2.74	1.01	3.66	3.28	1.25	2.8	0.005

Z- Wilcoxon test statistic; SD - standard deviation; X average - average load point X which determined lateral coordinates X (cm); Y average - average load point Y which determined the anterior–posterior coordinates Y (cm); V average - average COP velocity (cm/s); p significance level; p < 0.05 reflects statistically significant relationship; p < 0.01 reflects highly significant relationship; p < 0.001 reflects very highly significant relationship

294

295 **Differences between the study group and the controls after the conclusion of the**
 296 **rehabilitation**

297 Statistical analysis showed significant differences in pain reduction on VAS in favour of the study
 298 group. Decrease in pain intensity was significantly more visible in the study group than in the

299 controls (p=0.0001). Likewise, improvement in functional performance reflected by ODI was
 300 significantly greater in the study group (p=0.0001). On the other hand, no statistically significant
 301 differences were observed in the participants' functional status measured with RMDQ (p=0.0866)
 302 or in spine range of motion in Schober's test (p=0.5878) between the groups (Table 7).

303 Table 7. Differences in the results between the study group and the controls after completion of
 304 the rehabilitation programme

Variable	Effect of rehabilitation Control group N=30	Effect of rehabilitation Study group N=30	T	P
VAS [points] X/SD	1.3/1.8	3/1	4.522	0.0001
ODI [points] X/SD	1/1	4/3	5.196	0.0001
RMDQ [points] X/SD	2.1/1.51	2.59/0.3	1.743	0.0866
Schober's Test [cm] X/SD	2.6/0.2	2.4/2	-0.545	0.5878

X- mean; SD – standard deviation; VAS - Visual Analogue Scale; ODI - Oswestry Disability Index; RMDQ - Roland-Morris Disability Questionnaire; T-test – Student's t-test; p - statistical significance

305
 306 Similarly, significantly greater improvements in balance parameters were found in the study group
 307 (p<0.05). Statistically significant differences between the study group and the controls after the
 308 therapy were identified in the trial with eyes closed, double-leg stance, in the parameters of Area
 309 circular, Path length and mean velocity (V average); in trials with eyes open, left leg stance, in
 310 Path length; and in trials with eyes closed, left leg stance, in Path length and mean velocity, with
 311 significance level reflected by p<0.005, (Table 8).

312 Table 8. Differences in the balance parameters between the study group and the controls after
 313 completion of the rehabilitation programme

After		Study group			Control group			Z	P
		Mean	Median	SD	Mean	Median	SD		
Eyes open, both legs	X average	-0.09	-0.05	1.05	-0.22	-0.45	1.33	0.50	0.615
	Y average	-2.36	-2.18	3.75	-3.41	-3.08	3.47	1.58	0.114
	Area circular	1.99	1.83	1.21	2.18	1.84	1.53	-0.37	0.712
	Path length	31.98	31.12	8.65	33.36	33.80	8.97	-0.87	0.383
	V average	1.18	1.18	0.24	1.21	1.19	0.34	-0.81	0.416
Eyes closed, both legs	X average	0.37	-0.29	2.95	0.94	0.34	2.68	-1.08	0.280
	Y average	-2.05	-2.06	2.12	-2.43	-2.72	2.95	0.72	0.473
	Area circular	2.33	2.15	1.28	3.57	3.03	2.08	-2.36	0.018
	Path length	36.92	38.33	16.41	66.17	49.17	39.17	-3.19	0.001

	V average	1.46	1.34	0.51	2.21	1.64	1.31	-2.43	0.015
Eyes open, right leg	X average	5.30	6.56	2.85	4.05	2.75	19.91	1.54	0.124
	Y average	-1.72	-1.21	3.89	-0.95	-1.26	2.11	0.01	0.994
	Area circular	7.06	6.78	1.79	8.45	7.05	3.60	-1.25	0.212
	Path length	95.08	98.54	29.40	116.39	108.28	35.30	-1.84	0.066
Eyes open, left leg	V average	3.35	3.47	1.05	3.86	3.68	1.20	-1.12	0.261
	X average	-5.87	-6.17	2.83	-4.95	-5.81	8.19	0.22	0.824
	Y average	-1.88	-1.41	3.18	-0.78	-0.67	3.21	-1.30	0.193
	Area circular	8.43	7.27	8.19	12.61	7.45	18.70	-1.18	0.237
	Path length	82.51	89.39	23.46	115.54	109.62	36.38	-3.39	0.001
Eyes closed, right leg	V average	3.29	3.29	1.08	3.92	3.66	1.24	-1.69	0.090
	X average	6.85	7.04	1.80	6.60	7.29	2.82	-0.18	0.853
	Y average	-0.51	-0.62	1.06	-1.30	-0.62	3.31	0.49	0.626
	Area circular	7.26	6.74	1.05	7.69	6.78	2.90	0.47	0.636
	Path length	109.17	99.29	32.49	114.69	110.69	29.19	-1.06	0.290
Eyes closed, left leg	V average	3.68	3.36	1.10	3.82	3.69	0.97	-0.98	0.326
	X average	-5.84	-4.81	2.23	-6.80	-7.39	3.07	1.47	0.141
	Y average	-0.27	-2.06	2.83	-1.20	-0.95	3.41	0.78	0.433
	Area circular	7.03	7.77	1.64	8.98	7.58	8.00	-0.93	0.352
	Path length	87.73	82.12	13.93	109.70	98.46	37.66	-2.67	0.008
	V average	2.88	2.74	0.61	3.66	3.28	1.25	-2.92	0.004
Z – Mann-Whitney test; SD- standard deviation; X average - average load point X which determined lateral coordinates X (cm); Y average - average load point Y which determined the anterior–posterior coordinates Y (cm); V average - average COP velocity (cm/s); p significance level; p < 0.05 reflects statistically significant relationship; p < 0.01 reflects highly significant relationship; p < 0.001 reflects very highly significant relationship									

314

315

316 Discussion

317 The current study was a response to the need for a clearly defined and evidence-based
318 postoperative rehabilitation programme, and it was designed to assess the change in
319 psychophysical performance of patients after lumbar discectomy in relation to the postoperative
320 therapy. In the literature related to these issues there are a number of studies investigating a variety
321 of rehabilitation programmes after lumbar discectomy. These programmes vary in terms of the
322 timing of postoperative interventions introducing kinesitherapy, the type of exercises applied, as
323 well as their intensity and frequency [1, 2, 6, 9, 11, 14, 32, 26, 27]. However, it is difficult to
324 compare the effectiveness such widely varied programmes. Consequently, it is difficult to choose
325 the best one for use in clinical practice while, as it is well known, there is a need to enhance the
326 outcome of discectomy with postoperative rehabilitation [8,16,36]. Nevertheless, many patients
327 receive little or no formal rehabilitation after surgery [36].

328 The study confirmed the effectiveness of the investigated rehabilitation protocol in both groups in
329 relation to all parameters analysed, with the exception of static balance, where the gains were
330 significantly greater in the study group. In this group, significant improvements were observed in
331 almost all balance parameters. By comparison, in the control group the findings showed significant
332 improvements only in the mean values of Y average and Path length in trials with eyes closed and
333 double-leg stance, in the parameter of Area circular in trials with eyes open and right leg stance,
334 and in Y average, Path length and V average in trials with eyes open and left leg stance.
335 Corresponding results were observed in trials Eyes closed and left leg stance, in the parameters
336 Path length and V average. Similar results were reported by Ozkara et al, who showed that
337 rehabilitation of patients after lumbar discectomy reduces pain, improves functional ability and
338 lumbar spine range of motion [11]. There is also contradictory evidence regarding effectiveness
339 of postoperative rehabilitation, for instance contributed by Oosterhuis et al. [9] who conducted a
340 randomised multicentre trial assessing effects of early rehabilitation (exercise therapy) after
341 lumbar disc surgery, and reported no effects of such intervention. The control group in that study
342 comprised patients after the surgery who were not referred to postoperative rehabilitation. The
343 parameters investigated in the study included ODI, back pain and leg pain measured on VAS, as
344 well as recovery level rated on a Likert scale and general physical and mental health evaluated
345 with SF12, all of these were assessed 3, 6, 9,12 and 24 months after the surgery [9]. In 2014,
346 Oosterhuis et al. conducted a systematic review of the literature related to the effectiveness of
347 clinical rehabilitation after lumbar discectomy [38]. The authors identified 22 clinical trials,
348 involving a total of 2503 patients. The authors posed the following research questions: “Is active
349 rehabilitation after the surgery more effective than no treatment?”, as well as “Which type of active
350 rehabilitation is most effective?”. They also assessed effectiveness of interventions relative to
351 when postoperative rehabilitation is initiated. In their overall conclusion, the authors stated that
352 none of the studies demonstrated high or even moderate strength of scientific evidence. All the
353 interventions in their design varied considerably in terms of their content, duration and intensity
354 [20]. In fact, these conclusions provided a motivation for the current study where the patients were
355 randomly divided into two groups, the control group participating in the basic version of the
356 rehabilitation programme and the study group, in addition to the same programme, receiving
357 manual therapy.

358 The present study shows that the rehabilitation protocol additionally including manual therapy was
359 more effective than the basic programme without manual therapy. Its superiority was reflected by
360 more pronounced decrease in pain intensity, more visibly improved functional performance as well
361 as improved static balance. It is possible that one of the mechanisms responsible for improvements
362 in the aforementioned functions is associated with the fact that the techniques used in manual
363 therapy, such as fascial manipulation, beneficially affect the structure of soft tissues. Muscle and
364 joint contractures that develop in connection to discogenic pain, especially the accompanying
365 radiculopathy, can lead to restricted mobility of the lumbar spine and to impaired static balance
366 [3,15]. Radicular pain can also be associated with the epidural and muscle scarring which develops
367 after any spinal surgery, including discectomy. Therefore, personalised work with each patient,
368 supplemented with well-matched manual therapy and patient education regarding physical activity
369 after the surgery, speeds up the process of recovery [25].
370 Comparison of the present findings to other results reported in the literature, and generally attempts
371 to compare studies assessing different postoperative rehabilitation protocols, encounter difficulties
372 due to the diverse methodological approaches applied in the scientific research and because of the
373 diversity of the rehabilitation protocols investigated [25]. There are many undefined factors to be
374 considered in the planning of postoperative rehabilitation, such as the type of the intervention or
375 timing (the time point to introduce post-surgery rehabilitation or its duration), whereas the current
376 findings demonstrate the effectiveness of the rehabilitation protocol applied in this group of
377 patients, which is the practical value of our research. The present study provides evidence
378 confirming beneficial effects of the clearly formulated postoperative rehabilitation programme on
379 the psychophysical performance of patients after lumbar discectomy and shows that the
380 rehabilitation protocol can justifiably be implemented in the daily practice.

381 **Practical implications**

382 Techniques of manual therapy applied in addition to the traditional model of postoperative
383 treatment after discectomy, have been shown to be more effective in reducing pain, increasing
384 balance and reducing disability.

385 **Limitations**

386 The present study has some limitations. Firstly, in our study, patients did not receive any pre-
387 rehabilitation. This is due to the fact that in Poland few, if any, patients have any rehabilitation
388 plan prior to discectomy. This may be an important factor, given that research has shown positive

389 effects of pre-rehabilitation e.g., after hip or knee replacement surgeries in the patients' functional
390 performance at 6 months after the surgery [21]. Therefore, it seems that further research on the
391 effectiveness of different types of therapy should also address rehabilitation applied before the
392 surgery. The study also did not include a follow-up evaluation to assess the lasting effects of the
393 postoperative rehabilitation. Therefore, it seems the results of the study generated new questions
394 about the long-term effects of postoperative rehabilitation and thus suggest further directions and
395 subject matter for further research focusing on rehabilitation after disc surgery.

396 **Conclusions**

397 The study showed that the postoperative rehabilitation programmes produced a positive and
398 statistically significant effect in both groups of participants, reflected by improvements in most of
399 the psychophysical parameters investigated, except for balance in the control group. It was shown
400 that the protocol additionally including manual therapy was more effective than the basic
401 programme. Its superiority was reflected by greater pain reduction, more visibly improved
402 functional performance as well as improved static balance. The results of the study are of practical
403 importance, as the protocols tested can be implemented in daily clinical practice in postoperative
404 rehabilitation offered to patients after lumbar discectomy.

405 **Abbreviations:**

406 **ODI**- functional status using Oswestry Disability Index,
407 **RMDQ**- Roland-Moris Disability Questionnaire,
408 **VAS**- Visual Analogue Scale,
409 **HNP** - herniated nucleus pulposus,
410 **AMTI** - the force plate from Advanced Medical Technology,
411 **COP** - the Centre of Pressure of the foot

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416 **Conflicts of Interests:**

417 The authors declare that they have no competing interests.

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