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1	Targeted Treatment Protocol in Patellofemoral Pain (TIPPs): Does Treatment Designed
2	According to Subgroups Improve Clinical Outcomes in Patients Unresponsive to
3	Multimodal Treatment?
4	
5	Hayri Baran Yosmaoğlu, Emel Sonmezer, Manolya Ozkoslu, Ezgi Sahin, Senay Çerezci, Jim
6	Richards, James Selfe, Jessie Janssen
7	
8	Background: Targeted intervention for subgroups is a promising approach for the management
9	of patellofemoral pain.
10	Hypothesis: Treatment designed according to subgroups improves clinical outcomes in
11	patients unresponsive to multimodal treatment.
12	Study Design: A prospective crossover intervention.
13	Level of Evidence: Level III
14	Methods: PFP patients (n=61, mean age: 27±9 years) were enrolled. PFP patients received
15	standard multimodal treatment three times a week for 6 weeks. Patients not responding to
16	multimodal treatment were then classified into one of 3 subgroups "strong", "weak and tight"
17	and "weak and pronated foot" using six simple clinical tests. They subsequently were
18	administered a further 6 weeks of targeted intervention designed according to subgroup
19	characteristics. Visual Analog Scale (VAS), Perception of Recovery Scale (PRS), EQ-5D-5L,
20	and S-LANSS were used to assess pain, knee function and quality of life before and after the
21	interventions.
22	Results: 36% of the patients (21 patients) demonstrated recovery following multimodal
23	treatment. However, over 70% (29 patients) of these non-responders demonstrated recovery
24	after targeted treatment. The VAS, PRS, S-LANSS, and EQ-5D-5L scores improved
25	significantly after targeted intervention compared to after multimodal treatment (p<0.001). The

VAS score at rest was significantly lower in the weak and pronated foot, and weak and tight subgroups (p=0.011, p=0.008) respectively. Post-treatment pain intensity on activity was significantly lower in the "strong" subgroup (p=0.006).

29 Conclusion: Targeted treatment designed according to subgroup characteristics improves30 clinical outcomes in patients unresponsive to multimodal treatment.

31 **Clinical Relevance:** Targeted intervention could be easily implemented following six simple 32 clinical assessment tests to subgroup patients into one of three subgroups (strong, weak and 33 tight, weak and pronated foot). Targeted interventions applied according to the characteristics 34 of these subgroups have more beneficial treatment effects than a current multimodal treatment 35 program.

36

37 Key words: Rehabilitation, knee injuries, patella, treatment outcome, pain perception

38

39 INTRODUCTION

40 Patellofemoral pain (PFP) is a chronic musculoskeletal problem that causes persistent anterior 41 knee pain.^{2,3,6,8,14,15,20,21,25,26,32,33,49} Despite its widespread use in clinics, it is difficult to suggest 42 that the current multimodal treatment approach leads to successful outcomes in the majority of 43 patients with PFP, only 46% of patients' knees were pain free at discharge.² This indicates that 44 over half of PFP patients do not respond to treatment and may continue their lives with chronic 45 anterior knee pain.

Identification of the factors leading to these low treatment success rates has consistently been a priority of previous International Patellofemoral Pain Research Retreats.^{4,10,12,52} The most important factor affecting the success of treatment that has emerged is that patients have a variety of musculoskeletal and biomechanical differences. The current multimodal treatment, therefore, may not affect the heterogeneous PFP patient population with the same efficiency. 51 Clinically subgrouping PFP patients and delivering targeted treatments has been strongly 52 recommended for future investigations of patellofemoral pain treatment from the International Patellofemoral Pain Research Retreats.^{4,12,52} An overview of previously published PFP 53 subgroups and the methods used to derive subgroups in PFP identified patients with PFP.³⁹ 54 They exhibit different anthropometric and biomechanical characteristics and do not form a 55 homogeneous group. There are 3 subgroups in the PFP population: "strong", "weak and tight" 56 and "weak and pronated foot".³⁸ The purpose of this study was to assess the clinical outcomes 57 58 of targeted treatments designed according to the characteristics of the three subgroups of PFP patients.³⁸ The hypotheses were that the assessment and subgroup classification is clinically 59 60 feasible, and that targeted treatments designed according to the characteristics of the three subgroups of PFP patients would show clinical benefits over and above a multimodal 61 62 intervention.

63 METHOD

64 Design

65 A prospective crossover intervention study design was used (Figure 1).

66 **Participants**

67 Patients aged between 18 and 40 attending a physiotherapy outpatient clinic at a University 68 Hospital with a clinical diagnosis of patellofemoral pain were approached for eligibility in this 69 study. Eligibility criteria were based on previously defined PFP criteria.^{7,38,47} Subjects were 70 excluded if they had any of the following: previous knee surgery, clinical evidence of ligamentous instability and/or internal derangement, a history of patellar subluxation or 71 72 dislocation, joint effusion, true knee joint locking and/or giving way, bursitis, patellar or iliotibial tract tendinopathy, Osgood Schlatter's disease, Sinding-Larsen Johansson Syndrome, 73 74 muscle tears or symptomatic knee plicae, serious co-morbidity which would preclude or affect 75 compliance with the assessment, or were pregnant.

76

77 Subgroup Classification Method

Quadriceps and Hip Abductor muscle strength ³¹, Patellar glide test^{44,54}, Quadriceps length⁵³, Gastrocnemius length⁵³, and Foot posture index³⁶ assessments were performed to classify all consenting patients into one of three subgroups (strong, weak and tight, weak and pronated foot) using the algorithm derived from the work by Selfe et al.³⁸

82

83 Intervention

84 Multimodal Treatment

The multimodal treatment program was designed based on the usual exercise and modalities used in local clinics.^{20,21,32,49} All patients received standard, supervised, 60 min multimodal treatment three times a week for 6 weeks. Table 1 shows the details of the multimodal rehabilitation program.

89 Targeted Treatment

90 Patients who did not respond to multimodal treatment were assigned to one of the treatment 91 groups "strong", "weak and tight", and "weak and pronated foot". They then followed a further 6 weeks, 45 min targeted intervention program administered three times a week. The targeted 92 93 treatment program was designed according to the key deficits identified in each patient by the 94 subgrouping clinical assessment tests. The patients in the "strong" subgroup had no muscle 95 strength deficit therefore, the intervention program for this subgroup was targeted at improving neuromuscular control and coordination ability using proprioceptive exercises such as 96 progressive balance exercises, and knee braces^{46,47} which have been shown to offer 97 improvements in movement control in patients with PFP,⁴¹ reductions in patellofemoral 98 reaction forces⁴⁴ and have been shown to reduce pain at 6 and 12 months during a PFP 99 rehabilitation program.⁴⁸ In the "weak and tight" subgroup, the exercise program consisted of 100

101 Closed Kinetic Chain (CKC) muscle strengthening and stretching, and weight management 102 advice, as a larger body mass index was identified as a potentially relevant clinical feature in 103 this subgroup.³⁸ In the "weak and pronated foot" subgroup, muscle weakness and abnormal foot 104 alignment were identified as the key factors. Therefore, the intervention program included CKC 105 strengthening exercises and foot orthoses.^{5,24} Table 2 shows the details of each of the specific 106 targeted intervention programs.

107 **Outcome measures**

Pain during activity measured using the Visual Analog Scale (VAS) was the primary outcome
 measure of this study ¹⁹. Activity was specified by patients.

110 The Perception of Recovery Scale was measured using a 7-point Likert scale ranging from 111 "completely recovered" to "worse than ever". Patients were classified as "recovered" if they 112 rated themselves as "completely recovered" or "strongly recovered". Patients rating themselves 113 in one of the other five categories from "slightly recovered" to "worse than ever" were 114 categorised as "not recovered".³⁵

The EQ-5D-5L was used as a self-reported generic measure of health and quality of life.
Patients rated their overall health on the day of the interview on a 0–100 hash-marked, vertical
visual analogue scale (EQ-5D-5L-VAS). A higher EQ-5D-5L-VAS score indicating better
health status.²²

Neuropathic Pain was measured using The Self-Administered Leeds Assessment of Neuropathic Symptoms and Signs (S-LANSS) questionnaire. The S-LANSS comprises a 5item questionnaire regarding pain symptoms and two items for clinical signs involving selfadministered sensory tests for the presence of allodynia and decreased sensation to pinprick. This was used to discriminate the small number of patients who may have neuropathic knee pain from those with nociceptive pain.⁴² The possible scores range from 0 to 24, with a score of 12 or greater considered to be suggestive of neuropathic pain.²⁸ Finally, a single leg hop test 126 was used to determine functional performance.¹ Distance was measured from toe to heel and
127 the mean score of three repetitions was recorded.

128 Data analysis

129 A sample size calculation was performed based on the minimal detectable change on the pain 130 VAS. Data from a previous study indicates that the VAS scores in patients with PFP was $4.3 \pm$ 1 cm,⁹ with 30% of the maximum score of the VAS-pain considered to be the detectable change, 131 132 the sample size for each treatment subgroup was determined to be 8 patients to achieve a 90% 133 power at the 0.05 level of significance. Data were not normally distributed when analysed with 134 the Kolmogorov-Smirnov test Consequently, non-parametric tests were indicated. Therefore 135 the "Wilcoxon signed rank test" was used to compare pre and post treatment outcomes with an 136 alpha value of 0.05. In addition, the mean of rank scores, standard errors and Z scores were 137 reported, along with descriptive statistics to describe the general features of the subjects. All 138 statistical analysis was conducted using SPSS 21.0.

139

140 **RESULTS**

Of the 128 patients who were screened, 95 were included in the present study. Of these 61 patients completed the multimodal treatment (Figure 1) (Table 3). Twenty-one patients (36%) demonstrated recovery following multimodal treatment (Phase I) and were discharged. 40 Patients (64%) not responding to multimodal treatment were administered a further 6 weeks of targeted intervention designed according to subgroup characteristics (phase 2). Twenty-nine (72.5%) patients demonstrated recovery following targeted intervention (phase II) and 11 (27.5%) patients did not respond to either of the treatment approaches (Table 4).

Pain intensity (VAS) at rest and during activity, and Perceived Recovery Scale (PRS), were
significantly improved after targeted intervention (p<0.001) (Table 5). S-LANSS, EQ-5D-5L
and EQ5D-5L-VAS scores were significantly improved following targeted intervention

compared to pre-targeted treatment scores (p = 0.001, p<0.001, p = 0.02), respectively (Table
5).

153 Within the three subgroups, the findings showed that PRS score was significantly improved 154 after targeted treatment compared to pre-targeted treatment levels in the "strong", "weak and 155 tight", and "weak and pronated foot" subgroups (p=0.005, p=0.001, p=0.004) respectively. 156 VAS pain intensity at rest was also significantly lower after targeted intervention in the "weak 157 and pronated foot" and "weak and tight" subgroups (p=0.011, p= 0.008) respectively, however 158 within the "strong" subgroup, no change was seen between pre-treatment and post treatment (p 159 = 0.245) (Table 6). However, pain intensity during activity was significantly lower after 160 treatment in the "strong" (p=0.006), the "weak and pronated foot" and "weak and tight" 161 subgroups; although these reductions were not statistically significant (p=0.059, p=0.06) 162 respectively (Table 6).

Other measures including quadriceps length test, S-LANSS, EQ5D-5L, and EQ5D-VAS were significantly improved in the "weak and tight" subgroup. S-LANSS, EQ5D-5L, and patellar mobility were significantly improved in the "weak and pronated foot" subgroup. In the "strong" group only gastrocnemius length was significantly different between pre- and post-targeted treatment (p=0.03). Results for outcome measures are shown in Table 7.

168

169 **DISCUSSION**

The results of our study suggest that the TIPPs subgroups and the algorithm used to classify PFP patients as "strong", "weak and tight", "weak and pronated foot" ³⁸ is valid and clinically implementable. The findings from this study were in agreement with previous work¹³ that reported differential response patterns in outcomes at 12 months in their subgroups. This suggests that targeted interventions based on subgroups, provides an important development in the treatment strategy for patients with PFP.^{4,52}

The "strong" subgroup demonstrated a poor response to multimodal treatment but a a 176 177 significant improvement after targeted treatment was observed. This finding is consistent with Greuel et al.¹⁸ and Gallina et al.¹⁷ who both reported results confirming that motor control of 178 179 the quadriceps is problematic in some PFP patients. One explanation for this is improved 180 neuromuscular control in patients classified as "strong". Since these patients already 181 demonstrated relatively high quadriceps muscle torque, targeted intervention was delivered 182 focusing on progressive development of motor control on unstable surfaces instead of 183 conventional muscle strength exercises. Given that quadriceps strength did not change as a 184 result of the targeted intervention, these progressive balance exercises and patellar bracing has improved motor control and stability.⁴¹ In addition, bracing may reduce patellofemoral forces 185 during activities of daily living and sporting tasks⁴⁴ and improvements within rehabilitation 186 protocols.⁴⁸ This was reflected in the improvement in the other pain related parameters, 187 188 However, since the average pre-treatment VAS pain level at rest in this subgroup was already 189 low a decrease from 1.8 to 0.7 has minimal clinical relevance.

190 Clinically the "weak and tight" subgroup appeared to be the most responsive group to treatment 191 overall with a relatively even split of 52% responding to multimodal treatment and all of the 192 remaining patients responding to targeted intervention. This finding was not surprising as 193 multimodal treatment routinely includes strengthening and stretching exercises. However, 194 closer analysis of the outcomes in the "weak and tight" subgroup suggest that although patients' 195 perception of recovery improved, the VAS activity pain intensity was not significantly 196 decreased after targeted treatment in this subgroup. Considering muscle weakness is the main 197 issue in this subgroup, the probable cause of this unexpected finding is persistent inability to 198 compensate patellofemoral loads especially during relatively high level activities of daily life 199 such as ascending/descending stairs even after the targeted treatment. Targeted intervention 200 consisting of functional strengthening may still be insufficient for high level activities of daily living which demand considerable muscular activity, although it caused approximately a 30%
increase in muscle torque and a significant improvement in perception of recovery in this
subgroup.

204 Findings from the "weak and pronated foot" subgroup suggest that targeted treatment including, 205 foot orthoses and pain free strengthening exercises was also successful in terms of perception 206 of recovery and VAS pain on rest. Although the same improvement was not observed in VAS 207 pain during activity. One explanation for this could be the indirect effect of the foot orthoses 208 on the knee as the patients showed no improvement in strength after targeted treatment. 209 Moreover, optimum correction is very difficult to determine during the intervention of foot 210 orthoses. Special single physiotherapy interventions or combining interventions for patellar 211 taping, mobilisation or manual therapy may have beneficial effects on pain related functional symptoms in PFP.^{11,30,34} However, the therapeutic effects of these applications remain limited 212 213 because PFP patients exhibit a wide variety of structural features and biopsychosocial 214 differences. The biomechanical and anthropometric characteristics of patients were not similar. 215 Foot pronation, for example, was noticeably high in some patients, while some had neutral foot 216 alignment. Similarly, quadriceps muscle strength, which is a predisposing factor or a most common symptom in previous studies^{8,54} has been high in some patients with the remainder 217 218 having considerable muscle weakness. Therefore, specific applications such as foot orthoses, 219 knee braces, tape, and even exercises may not be required by every patient.

The functional hop test is often used in clinics to measure functional capability.⁵¹ Considering that there was no increase in quadriceps muscle strength in the "weak and pronated foot", and "strong" subgroups, an improvement in the hop test scores was not expected.

223 Due to the methodological design of this study, patients received 6 weeks of multimodal 224 treatment before 6 weeks of targeted treatment with no intervening washout period. This is a 225 study limitation since the cumulative effects of the previous treatment (multimodal) were

226	ignored. Therefore, the observed difference in some parameters could be the result of regression
227	to the mean.
228	CONCLUSION
229	Both the TIPPs assessment and subgroup classification algorithm are clinically feasible that
230	those with PFP are not a homogeneous group, and have biomechanical and structural
231	differences.
232	
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383 Table 1. Multimodal Treatment Program

MODALITY	APPLICATION TYPE				
Thermotherapy	Cold packs /20 min				
Transcutaneous Electrical Neural Stimulation (TENS)	Conventional mode-20 min				
	50-100Hz, 20-60 pulse/sec				
Therapeutic Ultrasound (US)	1 Watt/cm ² - 5 min/ around knee joint				
Hamstring/tensor fascia lata/ iliotibial band stretching	30sn/5 rep				
Isometric quadriceps strengthening	10 rep x 3 set				
Isometric hip adductor strengthening	10 rep x 3 set				
OKC knee extension exercise	3 sets of patients' 8-10 RM, in painless ROM				
OKC Hip adductor exercise	side lying/ 3 sets of patients' 8-10 RM				
Home based exercise program*					

*RM: Rep *Home b*

RM: Repetition Maximum, rep: repetition, ROM: Range of motion, OKC: Open kinetic chain

*Home based exercise program included the same applications except TENS, NMES, US

390 Table 2. Targeted treatment program

STRONG SUBGROUP					
Progressive balance/proprioception exercises	Standing on one leg on wobble board				
	3 sets of 1 min exercise each leg				
	1-3 sets per session depending on pain				
	Progression*: Eyes closed, bouncing ball against wall, bouncing				
	ball against wall on an unstable surface				
Patellar bracing**	Patient was asked to put on knee brace during ADL				
Activity modification	Activity reduction to fit within envelope of function locally				
	determined and negotiated with individual patient				
WEAK AND TIGHT SUBGROUP					
CKC strengthening exercises	Plie/lunge/single limb squat				
	Pain free ROM				
	10 reps per set/ 1-3 sets depending on pain				
Gastrocnemius and Quadriceps Stretching exercises	30 seconds static stretch x 3 reps x 1 per day				
Weight management strategies	Locally determined and negotiated with individual patient				
WEAK AND PRONATED FOOT SUBGROUP					
CKC strengthening exercises	Plie/lunge/single limb squat				
	Pain free ROM				
	10 reps per set/ 1-3 sets depending on pain				
Foot orthoses	Custom made insole supporting medial longitudinal arch of				
	foot***				
Activity modification	Improve activity levels locally determined and negotiated with				
	individual patient				

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 393 ADL: Activity of Daily Life CKC: Closed Kinetic Chain
 *Progression timing in balance exercise was decided by

*Progression timing in balance exercise was decided by clinician based on patient pain free achievement

** Off the shelf knee support with patellar pad was used (Orthocare© material: 5mm neoprene /SBR /nylon jersey/pk). Brace size was selected by clinician according to patient comfort and patellar coherence (S/M/L/XL sizes were used)

*** Custom Made Insoles are tailored individually based on static and dynamic examination of load distribution on foot. using CAT-CAM free step V.1.3.30

Table 3 Demographic data of patients who participated in the study

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PATIENTS (N=61)	MEAN	SD
AGE (YEAR)	27	9
HEIGHT (CM)	170	8
WEIGHT (KG)	65	13
TIME SINCE SYMPTOMS STARTED	24	28
(MO)		
BMI (KG/M2)	22.5	3
	-	

- 406 407 Table 4. Perception of recovery after treatments

	PHASE 1 MULTIMODAL TREATMENT (N=61)				PHASE 2 TARGETED TREATMENT (N=40)			
PRS	Overall % (n)	Weak and Tight % (n)	Weak and Pronated % (n)	Strong % (n)	Overall % (n)	Weak and Tight % (n)	Weak and Pronated % (n)	Strong % (n)
FULLY IMPROVED	11 (7)	16 (4)	-	9 (2)	7.5 (3)	8 (1)	-	11(2)
GREAT IMPROVEMENT	23 (14)	36 (9)	29 (4)	9 (2)	65 (26)	92 (11)	80 (8)	39 (7)
SOME IMPROVEMENT	48 (29)	36 (9)	57 (8)	55(12)	17.5 (7)	-	20 (2)	28 (5)
NO CHANGE	16 (10)	12 (3)	14 (2)	18 (4)	10 (4)	-	-	22 (4)
A LITTLE WORSE	4 (3)	-	-	9 (2)	0 (0)	-	-	-
408 409 410 411 412 413 414 415 416 417 418 419 420 421 422 423 424 425 424 425 426 427 428 429 430 431 432								

433 Table 5. Outcome measures differences in targeted treatment

	Before Targeted Treatment		After Targeted Treatment			
Outcome Measures (n=40)	Median	Min-Max	Median	Min-Max	Z	р
Perception of recovery	3	3 - 5	2	1 - 4	-5,034	<0.001*
VAS activity (cm)	4.4	0.1 - 8.8	1.8	0 - 7.5	-4.075	<0.001*
VAS rest (cm)	1.7	0 - 7.4	0.5	0 - 7.0	-3.599	<0.001*
S-LANSS	5	0 - 16	0	0 - 24	-3.449	0.001*
EQ5D-5L	7	5 - 10	6	5 - 11	-3.704	<0.001*
EQ5D-VAS	80	30 - 95	85	50 - 100	-2.322	0.020*
Quadriceps muscle strength (Nm/kg)	1,1	0,5-2,1	1,2	0,6-2,3	-3.644	<0.001*
Hip abductor muscle strength (Nm/kg)	1,3	0.7 – 2,6	1,3	0,6 – 1,9	-1.456	0.145
Patellar mobility test (mm)	12	7 - 25	11	2 - 18	-2.062	0.039*
Foot posture index	6	0 - 11	6	0 - 12	-0.372	0.710
Quadriceps length (⁰)	142.7	115 - 156	145.2	128 - 155	-2.150	0.032
Gastrocnemius length (⁰)	19.6	8 - 40	20.5	12.3 - 40	-1.358	0.174
Jump (cm)	90.2	30 - 180	91	38 - 179	-1.472	0.141

435 *p<0.05, VAS: Visual Analog Scale, S-LANSS: The Leeds Assessment of Neuropathic Symptoms and Signs, EQ5DL:
436 European Quality 5 Dimension, °: degree

Table 6. Differences in subgroups before and after targeted treatment (n=40)

BEFORE TREATMENT

 Immediate
 AFTER TREATMENT

Р

Z

		Median	Min-Max	Median	Min-Max		
VAS IN ACTIVITY	Weak and Pronated (n=10)	5.3	0.5 - 8.8	2.7	0.2 - 6.6	-1.886	0.059
	Weak and Tight Group (n=12)	3.7	0.4 - 7.7	3	0-6.5	-1.883	0.060
	Strong Group (n=18)	5.0	0.1-8.2	2.0	0-7.5	-2.741	0.006*
VAS AT REST	Weak and Pronated (n=10)	3.9	0-7.1	0.8	0-3.4	-2.547	0.011*
	Weak and Tight Group (n=12)	1.0	0- 3.5	0.68	0-1.6	-2.667	0.008*
	Strong Group (n=18)	1.8	0-7.4	0.7	0-7	-1.161	0.245
PRS	Weak and Pronated (n=10)	3	3-4	2	2-3	-2.887	0.004*
	Weak and Tight Group	3	3-4	2	1-2	-3.213	0.001*
	(n=12) Strong Group (n=18)	3	3-5	2.5	1-4	-2.830	0.005*

453 *p<0.05, VAS: Visual Analog Scale, PRS: Perception of Recovery Scale

	Weak and Tight subgroup (n=12)				Weak and Pronated subgroup (n=10)				Strong subgroup (n=18)			
	Before Median (Min- Max)	After Median (Min- Max)	Z	р	Before Median (Min- Max)	After Median (Min- Max)	Z	р	Before Median (Min- Max)	After Median (Min-Max)	Z	р
S-LANSS	5 (0- 11)	0 (0 - 6)	-2.716	0.007*	6 (0-11)	0 (0 – 10)	-2.410	0.016*	5 (0- 169)	1.5 (0 – 24)	-0.947	0.344
EQ5D-5L	7.5 (5-10)	6 (5–9)	-2.556	0.011*	9 (6- 9)	6 (5–11)	-2.203	0.028*	6 (5-10)	6 (5–10)	-1.613	0.107
EQ5D-VAS	80 (50- 90)	90 (50-95)	-2.034	0.042*	80 (50-90)	80 (50-100)	-1.027	0.305	82.5 (30-95)	82.5 (55-100)	-1.444	0.149
Quadriceps muscle strength (Nm/kg)	0.84 (0.51.3)	1.05 (0.6 – 1.4)	-3.061	0.002*	1.06 (0,6-2.1)	1.3 (0.7 – 1.6)	-1.887	0.059	1.2 (0.9 – 1.6)	1.2 (0.9 – 2.2)	-0,893	0.372
Hip abductor muscle strength (Nm/kg)	0.9 (0.7 – 1.4)	1.1 (0.6–1.6)	-1,844	0.065	1.1 (0.7–1.6)	1.2 (0.9–1.6)	-0.593	0.553	1.4 (0.9–2.6)	1.5 (1 –1.9)	-0.259	0.796
Patellar mobility test (mm)	10 (7- 15)	10 (8- 15)	-0.103	0,918	15 (11- 22)	12 (2- 18)	-2.325	0.020*	12 (8- 25)	11 (7- 17)	-0.803	0,422
Foot posture index	5 (0-9)	5.5 (2-10)	-1.725	0.084	7.5 (4-11)	7.5 (2-12)	-0.679	0.497	5 (0-11)	6 (0-12)	-0.178	0.859
Quadriceps length (⁰)	137 (115 – 149)	140 (128 -152)	-2.134	0.033*	140 (118 – 152)	146 (130 -155)	-1.481	0.139	147 (117 – 155)	148 (128 -155)	-0.071	0.943
Gastrocnemius length (⁰)	18.2 (10-26)	17.4 (12.6-27)	-1.295	0.195	21.3 (10-40)	17.3 (12.6-34)	-1.244	0.214	19.6 (8-27)	21.5 (12.3-40)	-2.120	0.034*
Jump test (cm)	79.1 (30-115)	81 (38-115)	-1.718	0.286	85.4 (40-149)	84.2 (65-154)	-1.718	0.086	104.5 (49.3-180.6)	107.2 (57.3-179.3)	-0.305	0.760
*n <0.05 VAS. Viewal	Analog Soula IAN	ICC. The Loads Assa	accurate of	Nounonal	is Comptone and C	Signa FOSDI - F.	non an A.	ality 5 Dim	angion of deaners			

Table 7. Outcome measures in subgroups before and after targeted treatment

*p<0.05, VAS: Visual Analog Scale, LANSS: The Leeds Assessment of Neuropathic Symptoms and Signs, EQ5DL: European Quality 5 Dimension, °: degree