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Impaired conditioned pain modulation in young female adults with long-standing patellofemoral pain

a single blinded cross-sectional study

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1	Impaired conditioned pain modulation in young female adults with long-		
2	standing patellofemoral pain – a single blinded cross sectional study		
3			
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25			

ABSTRACT 26

49

Objective: Patellofemoral pain (PFP) is common among young individuals. Female 27 adolescents with PFP present typically with localised mechanical hyperalgesia 28 around the knee but the effect of central pain mechanisms are unknown. This study 29 aimed to compare temporal summation of pain, conditioned pain modulation (CPM), 30 and widespread hyperalgesia in young female adults with PFP and age-matched 31 pain-free controls. 32 Design: Cross-sectional study. 33 Setting and subjects: Twenty young female adults (19 - 21 years) with long-34 standing PFP were compared with 20 pain-free controls from the same population-35 based cohort 36 Methods: Cuff algometry was used to assess the pain detection threshold. Temporal 37 summation of pain was assessed by recording the pain intensity on a visual 38 39 analogue scale during repeated cuff pressure stimulations at pain tolerance intensity on the lower leg. CPM was recorded as the increase in cuff pain detection threshold 40 in response to experimental conditioning pain imposed on the contralateral arm. 41 Handheld pressure algometry was used to assess pressure pain thresholds (PPT) 42 on the knee, shin, and forearm. The examiner was blinded to the type of subject 43 assessed. 44 **Results:** Compared with pain-free controls, young females with PFP did not show a 45 decrease in cuff pain thresholds (P<0.40) or facilitated temporal summation (P<0.15), 46 but had a lower CPM response (P<0.04) and lower PPTs (P<0.005). 47 Conclusions: Young female adults with long-standing PFP demonstrated impaired 48 CPM. This is important as PFP, a peripheral pathology, might have important central

- 50 components which need to be studied in order to understand its extent and
- 51 therapeutic implications.
- 52 Key words: patellofemoral pain; central sensitization; CPM; hyperalgesia;
- 53 experimental pain
- 54

55 INTRODUCTION

Patellofemoral pain (PFP) is a common knee condition among individuals who 56 participate in repetitive knee loading activities (1, 2). The prevalence of PFP is 57 more than twice as high among females compared to males and data from 58 sports medicine clinics suggest that PFP is the most common knee condition 59 and may account for 25% of all consultations regarding knee pain (3, 4). PFP 60 is defined as pain anteriorly around the patella with pain that increases during 61 prolonged sitting, squatting, kneeling, and stair climbing (5). Prospective 62 studies have highlighted a poor long-term prognosis with only 1/3 being pain 63 free 12 months after initiation of treatment (6). 64

Mechanical hyperalgesia was recently demonstrated by reduced pressure pain 65 thresholds (PPTs) assessed around the knee and on the tibialis anterior muscle in 66 female adolescents with PFP compared with pain free controls (7). The distal 67 hyperalgesia observed at the tibialis anterior muscle may reflect segmental 68 spreading of hyperalgesia (8). In some chronic musculoskeletal pain conditions with 69 70 widespread hyperalgesia such as osteoarthritis (OA) and fibromyalgia, temporal summation of the pain perception to repetitive pressure pain stimulations appears to 71 be facilitated compared with pain free controls, which is thought to be the result of 72 facilitated central mechanisms (9, 10). Furthermore, studies have demonstrated 73 reduced sensory perception to thermal stimulation and vibration (11, 12), indicating 74 altered sensory function in patients with PFP. 75

76

Painful stimulation evokes a multisegmental hypoalgesia often referred to as
 conditioned pain modulation (CPM); a manifestation of the descending modulatory

79	effects characterised by attenuated pain response to a painful test stimulus when
80	another painful conditioning stimulus is applied (13). CPM is a proxy of the
81	effectiveness of the endogenous analgesia system. Previous studies have shown
82	impaired CPM in both knee and hip OA (9, 10, 14) as well as other non-arthritic
83	chronic pain conditions (15, 16). Impaired CPM is clinically important as it may be
84	associated with a higher risk of developing chronic post-operative pain (17).
85	Collectively, identification of sensitised central mechanisms and widespread
86	hyperalgesia appears to be clinically important and may be associated with higher
87	risk of long-standing pain (17-21). However, it has never been investigated among
88	young female adults with PFP.
89	
90	The aims of this study were to assess 1) temporal summation of cuff-induced
91	pressure pain, 2) CPM assessed by cuff-algometry, and 3) widespread mechanical
91 92	pressure pain, 2) CPM assessed by cuff-algometry, and 3) widespread mechanical hyperalgesia in young female adults with PFP compared with age matched healthy
92	hyperalgesia in young female adults with PFP compared with age matched healthy
92 93	hyperalgesia in young female adults with PFP compared with age matched healthy pain-free controls. It was hypothesised that young adults with PFP would have
92 93 94	hyperalgesia in young female adults with PFP compared with age matched healthy pain-free controls. It was hypothesised that young adults with PFP would have increased temporal summation of pain and impaired CPM compared with pain-free
92 93 94 95	hyperalgesia in young female adults with PFP compared with age matched healthy pain-free controls. It was hypothesised that young adults with PFP would have increased temporal summation of pain and impaired CPM compared with pain-free controls and that PPTs around the knee and at sites remote from the area of self-
92 93 94 95 96	hyperalgesia in young female adults with PFP compared with age matched healthy pain-free controls. It was hypothesised that young adults with PFP would have increased temporal summation of pain and impaired CPM compared with pain-free controls and that PPTs around the knee and at sites remote from the area of self- reported knee pain would be lower among young female adults with PFP compared

101 METHODS

102

103 Subjects

The design was a cross-sectional study nested within a population-based cohort. 104 Young female adults diagnosed with PFP were matched to a gender- and age-105 matched comparison group of pain-free controls. Both young adults with PFP and 106 pain-free controls were recruited from the same population-based cohort (the 107 Adolescent Pain in Aalborg 2011) (22). This cohort has been followed since 2011 108 and consisted of 2200 adolescents between 15 and 19 years of age. From these, 109 153 were diagnosed with PFP using previously described inclusion and exclusion 110 criteria (23, 24). In short, the patients with PFP were required to have an insidious 111 onset of anterior knee or retropatellar pain of more than 6 weeks duration and 112 provoked by at least two of the following daily activities: prolonged sitting or kneeling, 113 squatting, running, hopping, or stair walking; tenderness on palpation of the patella, 114 pain when stepping down or double leg squatting; and worst pain during the previous 115 week of more than 3 cm on a 10 cm visual analogue scale (VAS). Exclusion criteria 116 were concomitant injury or pain from the hip, lumbar spine, or other knee structures; 117 previous knee surgery; self-reported patellofemoral instability; knee joint effusion. 118From the 153 adolescents with PFP, 121 were enrolled in a randomised trial (24). 119 Participants who were previously diagnosed with PFP in the original trial (23) were 120 included in a telephone interview to inquire if they still had knee pain and if so they 121 122 were invited to participate in the current study. Pain-free controls were randomly recruited from the same population-based cohort by telephoning a random sample 123 with approximately the same age, gender, and sports participation as the PFP group. 124 The inclusion criteria for pain-free controls were: No current self-reported 125

musculoskeletal pain; no self-reported prior surgery in the lower extremity; no self reported neurological or other medical conditions. The study was conducted in
 accordance with the Helsinki Declaration and was approved by the local ethics
 committee in the North Denmark Region (N-20110020).

130

131 **Protocol**

All parameters were collected by an examiner who was blinded towards group 132 allocation (PFP vs. pain-free controls). Data was collected from the side of the most 133 painful knee among those with PFP and the same side matched on dominance on 134 pain-free controls. Manual pressure algometry, cuff pressure algometry, temporal 135 summation of pain, and CPM were assessed in a sequence on a single day with 136 approximately 3-5 minutes between each test. The reliability of manual PPT 137 measurements performed on pain-free young adults has previously been 138 investigated and found to be acceptable for sites around the hand and head 139 (intraclass correlation coefficients (ICC) ranging from 0.69 to 0.88) (25) In pain-free 140 adults, the reliability of computer controlled cuff-algometry for assessing pressure 141 algometry, temporal summation of pain, and CPM has been found to be good to 142 excellent with ICCs ranging from 0.60-0.89 (26). The primary outcome was temporal 143 summation of pain measured as the change in VAS during repeated cuff stimulations 144 on the lower leg. Secondary outcomes included 1) cuff pressure pain sensitivity 145 recorded on the lower leg, 2) CPM with the outcome being the change in cuff pain 146 sensitivity on the lower leg after cuff-induced arm pain (the conditioning stimulus), 147 and 3) PPTs at the patella, the tibialis anterior muscle, and the lateral epicondyle. 148 149

150 **Pressure algometry**

PPTs were assessed using a hand-held pressure algometer (Somedic Sales AB, 151 Sweden) with a stimulation area of 1 cm^2 placed perpendicular to the skin. Pressure 152 was applied at a rate of 30 kPa/s which was verified using the inbuilt digital indicator 153 on the algometer. The individuals were instructed to indicate when the sensation 154 changed from a sensation of pressure to the first sensation of pain. Measurements 155 were done with the individuals resting in a reclining position and the knee slightly 156 flexed at 15 degrees. PPT was measured at sites close to the knee to reflect 157 localised hyperalgesia and on the contralateral site distant to the knee to investigate 158 widespread hyperalgesia (8-10). The PPTs were measured twice at each site and 159 the average was calculated and used for the analyses. Three assessment sites were 160 located on: 1) The knee at the centre of the patella. 2) The muscle belly of the tibialis 161 anterior muscle 5 cm distal to the tibial tuberosity. 3) The elbow, on the lateral 162 epicondyle of the humerus. 163

164

165 **Computer-controlled cuff pressure algometry**

Cuff pressure pain detection thresholds (PDT) and cuff pressure pain tolerance 166 (PTT) were assessed by a computer-controlled cuff pressure algometer (Nocitech, 167 Denmark and Aalborg University, Denmark). Computer controlled cuff algometry 168 have previously been widely used to study central pain mechanisms (9, 10, 27, 28) 169 and has the advantage of being user independent. A 13-cm wide silicone tourniquet 170 cuff (VBM, Germany) with an equal-sized proximal and distal chamber was wrapped 171 around the lower leg on the side with the worst knee pain. The cuff was mounted 172 with a 5 cm distance between its upper rim and the tibial tuberosity. The cuff 173 pressure was increased with a rate of 1 kPa/s simultaneously in both chambers and 174 the maximal pressure limit of the system was 100 kPa which may cause some 175

participants to reach 100 kPa before reaching PTT. The participants used an 176 electronic VAS to rate their pressure-induced pain intensity and a button to release 177 the pressure. The electronic VAS was sampled at 10 Hz. Zero and ten cm extremes 178 on the VAS were defined as "no pain" and "maximal pain", respectively. The 179 participants were instructed to rate the pain intensity continuously on the electronic 180 VAS from the first sensation of pain and to press the pressure release button when 181 the pain was intolerable. The pressure value when the subject rated the sensation of 182 pain as 1 cm on the VAS was defined as the PDT and the pressure recorded when 183 the subject terminated the cuff inflation was defined as the PTT. 184

185

Temporal summation of cuff-induced pressure pain

Temporal summation was assessed by the computer-controlled cuff algometer 187 (Nocitech, Denmark). Ten cuff pressure stimuli (1-s duration and 2-s interstimulus 188 interval) were delivered to the lower leg by simultaneous inflation of both cuff 189 chambers at an intensity equivalent to PTT recorded during the assessment of the 190 cuff pain sensitivity. In the period between stimuli, a constant non-painful pressure of 191 5 kPa was kept, thus ensuring that the cuff did not move. The participants were 192 instructed to rate the pain intensity continuously on the electronic VAS. The mean 193 VAS score during the 1-s interval between stimulations after each of the 10 stimuli 194 was extracted and then normalised by subtraction of the mean VAS scores from the 195 first stimulation. 196

197

198 **Conditioned pain modulation**

Experimental tonic pain was induced in the contralateral arm by cuff-induced pain (conditioning stimulation), and assessment of cuff PDT and PTT was performed on

the lower leg before and during the conditioning stimulus on the arm. A 7.5-cm-wide 201 tourniquet cuff (VBM, Germany) was wrapped around the left arm with the lower rim 202 of the cuff placed 3 cm proximal to the cubital fossa. The computer-controlled cuff 203 algometer maintained a constant pressure at 60 kPa. The CPM effect was 204 expressed as the percentages increase of PDT and PTT, respectively, from baseline 205 to the conditioning assessments. If subjects reached 100 kPa as PTT before 206 conditioning cuff-pain these were excluded from further analysis. A-priori it was 207 expected that some subjects would reach 100 kPa and therefore the CPM effect 208 using PTT was only included as an explorative outcome. 209

210

211 Self-reported outcomes

The following clinical self-reported measures were used: 1) Patellofemoral

213 Osteoarthritis Outcome Score (KOOS)(29), worst pain intensity during the last four

weeks, and current pain measured on a 0-10 numeric rating scale (NSR), 2)

symptom duration (months), 3) most painful knee (right/left), 4) uni- or bilateral pain

(yes/no), and pain localisation measured using the Navigate pain app (30).

217

218 Statistical analysis

The sample-size was based on the primary outcome of detecting a difference in

normalized VAS during temporal summation from stimuli 1 to stimuli 10 of 1.5 cm(10).

221 Common standard deviation was estimated to be 1.5 cm and with a power 0.80 and

alpha at 0.05 this corresponds to a sample-size of minimum 16 in each group.

223

All analyses were defined a-priori. The primary analysis was a comparison between

groups in the change in VAS during temporal summation. Secondary analyses

included comparisons of cuff PDT, cuff PTT, CPM, and PPTs at the centre of the 226 patella, m. tibialis anterior and the lateral epicondyle. Unpaired t-tests were used for 227 all comparisons except for the PPTs where a two-way ANOVA was used with group 228 and site as factors. All calculations were performed using Stata version 11 229 (StataCorp, College Station, Texas, USA). Mean values and 95% confidence 230 intervals (CI) are reported if data were normally distributed and otherwise they are 231 232 presented as median and interquartile range (IQR). P-values less than 0.05 were considered significant. 233

235 **RESULTS**

- The female adults with PFP had a median symptom duration of 6 years and reported
- in general intermittent episodes (Table 1) of peri-patellar pain (Figure 1).
- Table 1: Demographics and patient reported outcomes

	Pain free (n=20)	Patellofemoral pain (n=20)	P-values		
Age [years]*	20.5 (20.0-21.0)	20.0 (19.0-21.0)	0.26		
Weight [kg]	61.7 (7.4)	63.8 (8.3)	0.40		
Height [cm]	169 (5)	170 (5)	0.53		
Sports participation (% yes)	75%	80%	0.61		
Duration of symptoms (years)*	N/A	6 (4.5-7)	-		
Worst pain last four weeks [NRS]	0 (0-0)	7 (5.5-8.0)	<0.0001		
KOOS symptoms	96 (5)	79 (11)	<0.0001		
KOOS pain	99 (2)	74 (11)	<0.0001		
KOOS activity	100 (1)	84 (10)	<0.0001		
KOOS Sport	98 (3)	59 (23)	<0.0001		
KOOS QoL	97 (7)	55 (18)	<0.0001		
PainDetect*	0 (0-0)	7.5 (4.5-11.0)	<0.0001		
Self-reported description of pain from					
PainDetect					
Persistent pain with slight fluctuations (n)		4			
Persistent pain with pain attacks (n)		3			
Pain attacks without pain between them (n)		12			
Pain attacks with pain between them (n)		1			
* Median and interquartile range. 0 to 100, best to worst scale. PFOOS: Patellofemoral Osteoarthritis					

* Median and interquartile range. 0 to 100, best to worst scale. PFOOS: Patellofemoral Osteoarthrit Outcome Score

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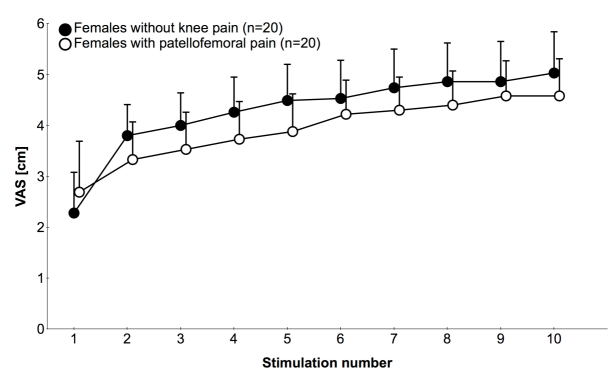
- Figure 1: The 20 small figures show the participants usual self-reported pain, while the large picture
- show the average pain location of the 20 young adults with patellofemoral pain.



The VAS scores following the ten repeated cuff stimulations showed a progressive increase in both groups illustrating the temporal summation of pain. The analysis showed no signification difference between groups in the increase in VAS from stimulus 1 to 10 (0.9 cm (95%CI: -0.5; 2.3 cm, t(38)=1.48, P=0.15) (Figure 2). Figure 2: Mean (+1.96*SE, N=20) of the visual analogue scale (VAS) scores after 10 cuff pressure pain stimulations at the pain tolerance intensity in females with patellofemoral pain (open symbols)

and pain free controls (solid symbols).

Temporal summation of pain



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244

255 Cuff pain sensitivity

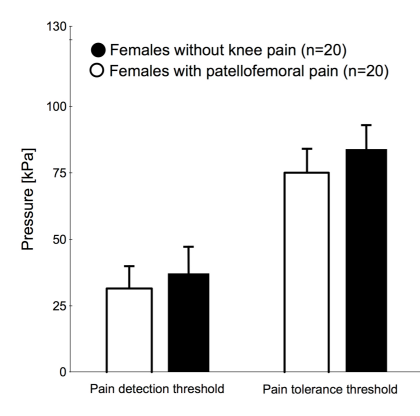
There were no significant differences in PDT (-5 kPa (95%CI: -18; 7 kPa, t(38)=0.85,

257 P=0.40)) or PTT (-8 kPa (95%CI: -21; 6, t(38)=1.11, P=0.27) between young female

adults with PFP and pain-free controls (Figure 3).

259

- Figure 3: Mean ((+ 1.96*SE, N=20) cuff pain detection threshold (PDT) and pain tolerance (PTT) 261
- 262 threshold in females with patellofemoral pain (open symbols) and pain free controls (solid symbols).
- here. 263



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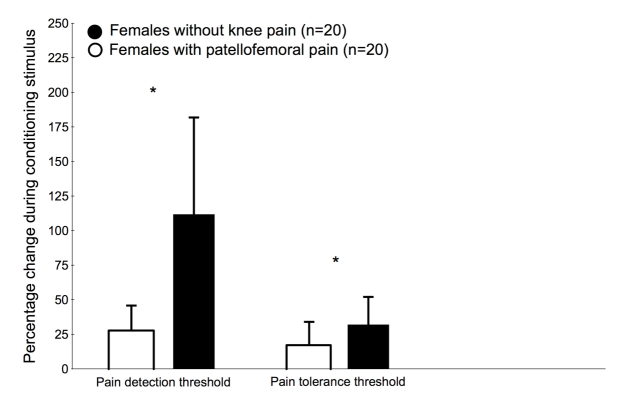
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Conditioning pain modulation 266

Young female adults with PFP had a 78% (95%CI: 4; 151%, t(38)=2.15, P<0.04) 267 lower CPM response in their PDT (Figure 4). The explorative CPM assessments 268 based on PTT measurements excluded 11 pain-free controls who reached 100 kPa 269 before experimental cuff-pain tolerance was reached but showed a 20% lower PTT 270 response among young female adults with PFP compared to pain-free controls 271 (95%CI: 1; 39%, t(27)=2.24, P<0.04) (Figure 4). 272 273

274

- Figure 4: Percentage increase (+1.96*SE, N=20) in pain detection threshold (PDT) and pain tolerance
- 277 (PTT) from before, to during (CPM) the experimental tonic pain was induced in the contralateral arm
- 278 in female with patellofemoral pain (open symbols) and pain-free controls (solid symbols). The PDT
- 279 includes 18 individuals in each group as the measurement system malfunctioned during collection of
- 280 data. * denotes significant differences (P<0.04).



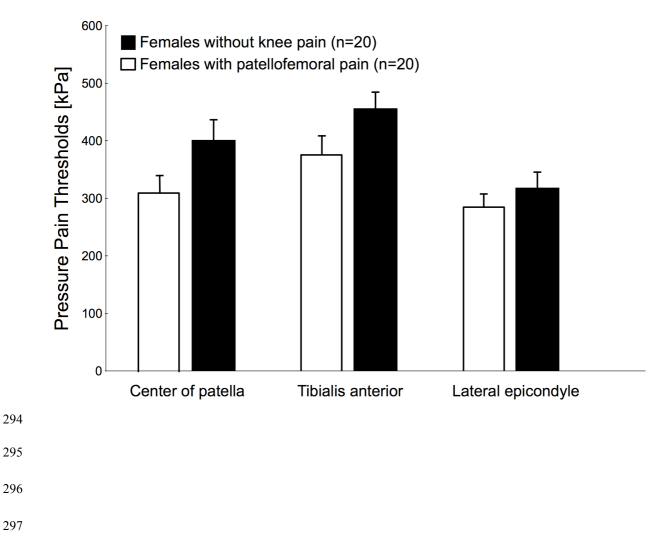
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282

- 283 **Pressure pain sensitivity**
- 284 There was a significant effect of group (PFP vs. pain-free controls) on PPTs (F_{1, 114} =
- 285 8.2, P < 0.005) (-68 kPa, (95%CI: -115; -21 kPa)) and PPT site (F_{2, 114} = 7.1, P <
- 286 0.001), (Figure 5).

- 288
- 289
- 290
- 291

- 292 Figure 5: Mean (+1.96*SE, N=20) handheld pressure pain threshold (PPT) on the center of the
- 293 patella, tibialis anterior and the lateral epicondyle.



298 **Discussion**

Young female adults with long-standing PFP were characterised by impaired CPM
assessed by PDT and spreading hyperalgesia, but contrary to our main hypothesis,
they showed no signs of facilitated temporal pain summation. This is the first study to
provide evidence for an altered pain processing among young adults with PFP.

303

Temporal summation of pain

305 Contrary to the a priori hypothesis, young female adults with long-lasting chronic PFP did not have facilitated temporal summation. Previous studies have shown a 306 facilitated temporal summation of pain in patients with knee OA and in other 307 musculoskeletal pain disorders such as fibromyalgia, chronic low back pain and 308 whiplash (31-34). The difference in results may potentially be explained by the 309 difference in study populations. The current study population reported knee pain for 310 an average of six years, which is similar to previous studies on knee OA (10), but 311 they are indeed much younger (≈45 years younger). The young adults with PFP 312 developed knee pain while they were in their early teens while patients with knee OA 313 developed knee pain in their mid-50s. Likewise the young adults with PFP presented 314 slightly lower peak pain intensities compared to sensitised adult patients with knee 315 OA typically reporting peak pain during the last 24 hours of 8 on a NRS (10). This is 316 important because higher peak pain is associated with a more facilitated temporal 317 summation (10). The pain reported by young adults with PFP is normally associated 318 with patellofemoral joint loading (e.g. stair walking or squatting) and rarely they 319 report pain at rest (35). Patients with knee OA often report pain at rest and also 320 during walking. Collectively the lack of facilitating temporal summation may suggest 321

that long pain duration is not the only factor, which is required to cause changes in
 temporal summation.

324

325 Conditioned pain modulation

Young female adults with PFP had a less efficient CPM similar to what have been 326 observed in older adults with knee OA (31). Less-efficient CPM mechanisms have 327 previously been reported in patients with musculoskeletal pain conditions, such as 328 myofascial temporomandibular disorders (36), chronic low back pain (37), and 329 fibromyalgia (38) but this study is the first to report impaired CPM in a younger 330 patient population. A reduced potency of the descending control makes the entire 331 neuroaxis more vulnerable to pain (39). However, an important finding is that the 332 CPM response was highly variable among the young female adults with PFP. Some 333 had no change in PDT during the test stimulus while others had responses similar to 334 pain-free controls. Earlier studies have linked a less efficient CPM response to 335 poorer long-term outcome after thoracotomy (17). Although pure speculation, this 336 may also be the case for young female adults with PFP who are known for having a 337 high degree of chronicity with only 1/3 being pain-free one year after treatment (40, 338 41). Eleven pain-free controls reached maximum in their pain tolerance threshold 339 assessment before the conditioning stimulus was applied which made it impossible 340 to compare the effect of the conditioning stimulus on their pain tolerance threshold. 341

342

343 CPM and temporal summation of pain are both considered part of central pain 344 processing but reflect two different mechanisms. Conditioned pain modulation 345 originates from the activation of brainstem inhibitory projections that, in turn, act to 346 postsynaptically inhibit spinal and trigeminal wide dynamic-range neurons (42). The

inability of the noxious conditioning stimulus to increase pain thresholds indicates a
potential deficiency in the body's endogenous pain modulatory ability. Temporal
summation is thought to be a facilitating mechanism that mimics the initial phase of
the windup process in dorsal horn neurons seen in animals (43). Therefore, the data
from this study suggests that mainly the inhibitory mechanism is affected in young
female adults with PFP.

353

354 **Pressure and cuff pain sensitivity**

Young adults with PFP had lower PPTs but showed no difference in either PDT or 355 PTT measured with the cuff algometer. The reason might be that cuff algometry 356 primarily captures deep tissue hyperalgesia while mechanical PPTs measure 357 hyperalgesia of superficial structures and muscles (8, 44). Reduced efficiency of the 358 CPM system may explain the widespread hyperalgesia. However, the present 359 population may have a lower degree of central sensitization compared to patients 360 with knee OA who are often characterised by facilitated temporal summation, 361 widespread hyperalgesia, and an inefficient CPM system (10). An important aspect 362 when interpreting these results is that this population is much younger than previous 363 studies on older adults with chronic pain. Although not heavily researched it appears 364 that changes in pain processing is dependent on the age of the individual (45). 365 Emerging evidence suggests that there might be some critical periods during 366 adolescence and childhood where pain experiences might induce long-lasting and 367 specific effects not observed among adults (45). However, it does appear that PPTs 368 may change in response to recovery. A recent study demonstrated that adolescents 369 with PFP deeming themselves as recovered after 3 months of exercise therapy had 370

- a 68-76 kPa larger improvement in PPTs around the knee and tibialis anterior
- compared to adolescents with not recovered after treatment (46).
- 373

374 Strengths and limitations

A strength of the study is that all the participants were recruited from a large, well-

defined, population-based cohort that have been followed for three years.

377 Recruitment of a population-based sample suggests that our data may be

378 generalizable to young female adults with long-standing PFP. An examiner blinded

to group allocation was used to minimise the risk of detection bias which is a

380 significant strength.

381 The present findings may not apply to the male population of young adults with PFP, as only females were included. The results may only apply to female adults with PFP 382 who developed knee pain during their early teens and not those who develop knee 383 pain during adulthood. Hormonal status of the participants was not assessed which 384 may introduce an unsystematic bias and reduce the difference in pain sensitivity 385 between groups. No reliability studies have been performed among this population 386 and no data exist for the minimally clinically important change. This makes it difficult 387 to interpret the relative difference between groups. 388

389

390 Clinical implications

Based on the large variation in CPM response among the young female adults with PFP it seems likely that altered central processing of pain is only present within a subgroup. It is known from a previous randomised trial among adolescents with PFP that there is a subgroup of adolescents who does not respond favourably to the

current best available evidence-based treatment, exercise therapy (23). It may be
that this subgroup is characterised by facilitated central mechanisms and treatment
among this subgroup should move away from the mechanical paradigm focusing
purely on improving strength and restoring lower extremity alignment. Instead
normalisation of the hyperexcitability of the nervous system should be targeted.
Interestingly, exercise-induced hypoalgesia may be affecting the facilitated central
mechanisms in the subgroup with efficient exercise therapy (47).

This study demonstrated that young female adults with long-standing patellofemoral pain were characterized by impaired conditioned pain modulation. This is the first study to provide evidence of an altered pain processing among young female adults with patellofemoral pain which is important as patellofemoral pain might have an important pain processing component which needs to be studied in order to understand its extent and therapeutic implications.

409

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413 References

Boling MC, Padua DA, Marshall SW, Guskiewicz K, Pyne S, Beutler A. A prospective 414 1. 415 investigation of biomechanical risk factors for patellofemoral pain syndrome: the Joint Undertaking to Monitor 416 and Prevent ACL Injury (JUMP-ACL) cohort. Am J Sports Med. 2009;37(11):2108-16. 417 Thijs Y, De Clercq D, Roosen P, Witvrouw E. Gait-related intrinsic risk factors for 2. patellofemoral pain in novice recreational runners. Br J Sports Med. 2008;42(6):466-71. 418 419 Taunton JE, Ryan MB, Clement DB, McKenzie DC, Lloyd-Smith DR, Zumbo BD. A 3. 420 retrospective case-control analysis of 2002 running injuries. Br J Sports Med. 2002;36(2):95-101. 421 Boling M, Padua D, Marshall S, Guskiewicz K, Pyne S, Beutler A. Gender differences in the 4. 422 incidence and prevalence of patellofemoral pain syndrome. Scand J Med Sci Spor. 2010;20(5):725-30. 423 Witvrouw E, Werner S, Mikkelsen C, Van Tiggelen D, Vanden Berghe L, Cerulli G. Clinical 5. 424 classification of patellofemoral pain syndrome: guidelines for non-operative treatment. Knee Surg Sports 425 Traumatol Arthrosc. 2005;13(2):122-30. 426 6. Collins NJ, Crossley KM, Darnell R, Vicenzino B. Predictors of short and long term outcome in 427 patellofemoral pain syndrome: a prospective longitudinal study. BMC Musculoskelet Disord. 2010;11:11. 428 Rathleff MS, Roos EM, Olesen JL, Rasmussen S, Arendt-Nielsen L. Lower mechanical 7. 429 pressure pain thresholds in female adolescents with patellofemoral pain syndrome. J Orthop Sports Phys Ther. 430 2013;43(6):414-21. 431 Graven-Nielsen T, Arendt-Nielsen L. Assessment of mechanisms in localized and widespread 8. 432 musculoskeletal pain. Nat Rev Rheumatol. 2010;6(10):599-606. Graven-Nielsen T, Wodehouse T, Langford RM, Arendt-Nielsen L, Kidd BL. Normalization of 433 9 434 widespread hyperesthesia and facilitated spatial summation of deep-tissue pain in knee osteoarthritis patients after knee replacement. Arthritis Rheum. 2012;64(9):2907-16. 435 436 10 Arendt-Nielsen L, Nie HL, Laursen MB, Laursen BS, Madeleine P, Simonsen OH, et al. 437 Sensitization in patients with painful knee osteoarthritis. Pain. 2010;149(3):573-81. 438 11. Jensen R, Kvale A, Baerheim A. Is pain in patellofemoral pain syndrome neuropathic? The 439 Clinical journal of pain. 2008;24(5):384-94. 440 Jensen R, Hystad T, Kvale A, Baerheim A. Quantitative sensory testing of patients with long 12. 441 lasting Patellofemoral pain syndrome. Eur J Pain. 2007;11(6):665-76. 442 Yarnitsky D, Arendt-Nielsen L, Bouhassira D, Edwards RR, Fillingim RB, Granot M, et al. 13. 443 Recommendations on terminology and practice of psychophysical DNIC testing. Eur J Pain. 2010;14(4):339. 444 Kosek E, Ordeberg G. Lack of pressure pain modulation by heterotopic noxious conditioning 14. 445 stimulation in patients with painful osteoarthritis before, but not following, surgical pain relief. Pain. 446 2000;88(1):69-78. 447 Lewis GN, Rice DA, McNair PJ. Conditioned pain modulation in populations with chronic 15. 448 pain: a systematic review and meta-analysis. The journal of pain : official journal of the American Pain Society. 449 2012;13(10):936-44. 450 Granovsky Y. Conditioned pain modulation: a predictor for development and treatment of 16. 451 neuropathic pain. Curr Pain Headache Rep. 2013;17(9):361. Yarnitsky D, Crispel Y, Eisenberg E, Granovsky Y, Ben-Nun A, Sprecher E, et al. Prediction of 452 17. 453 chronic post-operative pain: pre-operative DNIC testing identifies patients at risk. Pain. 2008;138(1):22-8. 454 Sterling M, Jull G, Vicenzino B, Kenardy J. Sensory hypersensitivity occurs soon after 18. 455 whiplash injury and is associated with poor recovery. Pain. 2003;104(3):509-17. 456 19. Kasch H, Qerama E, Bach FW, Jensen TS. Reduced cold pressor pain tolerance in non-457 recovered whiplash patients: a 1-year prospective study. Eur J Pain. 2005;9(5):561-9. 458 Lundblad H, Kreicbergs A, Jansson KA. Prediction of persistent pain after total knee 20. 459 replacement for osteoarthritis. J Bone Joint Surg Br. 2008;90(2):166-71. 460 Nijs J, Van Oosterwijck J, De Hertogh W. Rehabilitation of chronic whiplash: treatment of 21. cervical dysfunctions or chronic pain syndrome? Clin Rheumatol. 2009;28(3):243-51. 461 Rathleff MS, Roos EM, Olesen JL, Rasmussen S. High prevalence of daily and multi-site pain--462 22. a cross-sectional population-based study among 3000 Danish adolescents. BMC pediatrics. 2013;13:191. 463 464 23. Rathleff MS, Roos EM, Olesen JL, Rasmussen S. Exercise during school hours when added to 465 patient education improves outcome for 2 years in adolescent patellofemoral pain: a cluster randomised trial. Br 466 J Sports Med. 2015;49(6):406-12. 467 24. Rathleff MS, Roos EM, Olesen JL, Rasmussen S. Early intervention for adolescents with 468 Patellofemoral Pain Syndrome - a pragmatic cluster randomised controlled trial. Bmc Musculoskel Dis. 469 2012;13(1):9. 470 25. Cathcart S, Pritchard D. Reliability of pain threshold measurement in young adults. J Headache 471 Pain. 2006;7(1):21-6.

472 26. Graven-Nielsen T, Vaegter HB, Finocchietti S, Handberg G, Arendt-Nielsen L. Assessment of 473 musculoskeletal pain sensitivity and temporal summation by cuff pressure algometry: A reliability study. Pain. 474 2015. Skou ST, Graven-Nielsen T, Rasmussen S, Simonsen OH, Laursen MB, Arendt-Nielsen L. 475 27. 476 Widespread sensitization in patients with chronic pain after revision total knee arthroplasty. Pain. 477 2013;154(9):1588-94. 478 Polianskis R, Graven-Nielsen T, Arendt-Nielsen L. Computer-controlled pneumatic pressure 28 479 algometry--a new technique for quantitative sensory testing. Eur J Pain. 2001;5(3):267-77. 480 Roos EM, Roos HP, Lohmander LS, Ekdahl C, Beynnon BD. Knee Injury and Osteoarthritis 29 481 Outcome Score (KOOS)--development of a self-administered outcome measure. J Orthop Sports Phys Ther. 482 1998;28(2):88-96. 483 Boudreau S, Spence R, Vasov G, Egsgaard L. Feature Extraction APP for Pain Profiles. In: 30. 484 Jensen W, Andersen OK, Akay M, editors, Replace, Repair, Restore, Relieve - Bridging Clinical and 485 Engineering Solutions in Neurorehabilitation. Biosystems & Biorobotics. 7: Springer International Publishing; 486 2014. p. 853-4. 487 Lluch E, Torres R, Nijs J, Van Oosterwijck J. Evidence for central sensitization in patients with 31. 488 osteoarthritis pain: A systematic literature review. Eur J Pain. 2014;18(10):1367-75. 489 Nijs J, Van Houdenhove B, Oostendorp RA. Recognition of central sensitization in patients 32. 490 with musculoskeletal pain: Application of pain neurophysiology in manual therapy practice. Man Ther. 491 2010;15(2):135-41. 492 33. Lemming D, Graven-Nielsen T, Sorensen J, Arendt-Nielsen L, Gerdle B. Widespread pain 493 hypersensitivity and facilitated temporal summation of deep tissue pain in whiplash associated disorder: an 494 explorative study of women. J Rehabil Med. 2012;44(8):648-57. 495 Neziri AY, Curatolo M, Limacher A, Nuesch E, Radanov B, Andersen OK, et al. Ranking of 34. 496 parameters of pain hypersensitivity according to their discriminative ability in chronic low back pain. Pain. 497 2012;153(10):2083-91. 498 Zhang W, Doherty M, Peat G, Bierma-Zeinstra MA, Arden NK, Bresnihan B, et al. EULAR 35. 499 evidence-based recommendations for the diagnosis of knee osteoarthritis. Ann Rheum Dis. 2010;69(3):483-9. 500 36. Bragdon EE, Light KC, Costello NL, Sigurdsson A, Bunting S, Bhalang K, et al. Group 501 differences in pain modulation: pain-free women compared to pain-free men and to women with TMD. Pain. 502 2002:96(3):227-37. 503 37. Peters ML, Schmidt AJ, Van den Hout MA, Koopmans R, Sluijter ME. Chronic back pain, 504 acute postoperative pain and the activation of diffuse noxious inhibitory controls (DNIC). Pain. 1992;50(2):177-505 87. 506 Kosek E, Hansson P. Modulatory influence on somatosensory perception from vibration and 38. 507 heterotopic noxious conditioning stimulation (HNCS) in fibromyalgia patients and healthy subjects. Pain. 508 1997;70(1):41-51. 509 39. Arendt-Nielsen L, Yarnitsky D. Experimental and clinical applications of quantitative sensory 510 testing applied to skin, muscles and viscera. J Pain. 2009;10(6):556-72. Collins NJ, Bierma-Zeinstra SM, Crossley KM, van Linschoten RL, Vicenzino B, van 511 40. Middelkoop M. Prognostic factors for patellofemoral pain: a multicentre observational analysis. Br J Sports 512 513 Med. 2013;47(4):227-33. 514 41. Rathleff MS, Rasmussen S, Olesen JL. [Unsatisfactory long-term prognosis of conservative 515 treatment of patellofemoral pain syndrome.]. Ugeskr Laeger. 2012;174(15):1008-13. Le Bars D, Menetrey D, Conseiller C, Besson JM. Depressive effects of morphine upon lamina 516 42. 517 V cells activities in the dorsal horn of the spinal cat. Brain Res. 1975;98(2):261-77. 518 Arendt-Neilsen L, Graven-Nielsen T. Translational Aspects of Musculoskeletal Pain: From 43. 519 Animals to Patients in Fundamentals of Musculoskeletal Pain . Graven-Nielsen T, Arendt-Neilsen L, Mense S, editors. Seattle: International Association for the Study of 520 Pain: 2008. 521 522 Polianskis R, Graven-Nielsen T, Arendt-Nielsen L. Pressure-pain function in desensitized and 44. 523 hypersensitized muscle and skin assessed by cuff algometry. J Pain. 2002;3(1):28-37. 524 de Lalouviere LLH, Ioannou Y, Fitzgerald M. Neural mechanisms underlying the pain of 45 525 juvenile idiopathic arthritis. Nat Rev Rheumatol. 2014;10(4):205-11. Rathleff MS, Roos EM, Olesen JL, Rasmussen S, Arendt-Nielsen L. Self-reported Recovery is 526 46. 527 Associated with Improvement in Localised Hyperalgesia Among Adolescent Females with Patellofemoral Pain -528 Results from a Cluster Randomised Trial. Clin J Pain. 2015. 529 47. Vaegter HB, Handberg G, Graven-Nielsen T. Isometric exercises reduce temporal summation 530 of pressure pain in humans. Eur J Pain. 2014.