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Pain-Phenotyping in Osteoarthritis: Current Concepts, Evidence, and Considerations towards a Comprehensive Framework for Assessment and Treatment

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1 Pain-Phenotyping in Osteoarthritis: Current Concepts,

2 Evidence, and Considerations towards a Comprehensive

3 Framework for Assessment and Treatment

4 Abstract

5 <u>Objectives:</u> Pain as central symptom of osteoarthritis (OA) needs to be addressed as part

6 of successful treatment. The assessment of pain as feature of disease or outcome in

7 clinical practice and drug development remains a challenge due to its

8 multidimensionality and the plethora of confounders. This article aims at providing

9 insights into our understanding of OA pain-phenotypes and suggests a framework for

10 systematic and comprehensive assessments.

11 <u>Methods:</u> This narrative review is based on a search of current literature for various

12 combinations of the search terms "pain-phenotype" and "knee OA" and summarizes

13 current knowledge on OA pain-phenotypes, putting OA pain and its assessment into

14 perspective of current research efforts.

15 <u>Results:</u> Pain is a complex phenomenon, not necessarily associated with tissue damage.

16 Various pain-phenotypes have been described in knee OA. Among those a phenotype

17 with high pain levels not necessarily matching structural changes and a phenotype with

- 18 low pain levels and impact are relatively consistent. Further subgroups can be
- 19 differentiated based on patient reported outcome measures, assessments of

20 comorbidities, anxiety and depression, sleep, activity and objective measures such as

21 quantitative sensory testing.

22 <u>Conclusions:</u> The complexity of both OA as disease and pain in OA prompt the definition

23 of a set of variables that facilitate assessments comparable across studies to maximize

24 our understanding of pain, as central concern for the patient.

25

- 26 Key words: phenotypes, osteoarthritis, osteoarthritis pain, drug development, patient
- 27 reported outcomes
- 28
- 29

30 Abbreviations

- 31 ADAMTS5 A disintegrin and metalloproteinase with thrombospondin motifs 5
- 32 CPM Conditioned pain modulation
- 33 FDA Food and drug administration
- 34 NGF Nerve growth factor
- 35 NMDA N-methyl-D-aspartate
- 36 OA Osteoarthritis
- 37PROPatient reported outcome
- 38 QST Quantitative sensory testing
- 39 PPT Pressure pain thresholds
- 40 TS Temporal summation
- 41 WOMAC Western Ontario and McMaster Universities Osteoarthritis Index
- 42
- 43

44 Introduction

45 Osteoarthritis (OA) is a complex multifactorial disease and global health care challenge 46 affecting more than 500 million people.¹ Not only is OA a major cause of reduction in 47 quality of life and activities of daily living, with substantial socio-economic impact,^{2, 3} but has also been associated with increased mortality.⁴ Total joint replacement is typically 48 49 the 'last resort' but approximately 20% of patients remain symptomatic after the 50 procedure.⁵ In the absence of treatments that can halt or reverse the OA process, and 51 despite much research over decades, there remains a huge unmet medical need. 52 53 For a "treatment of OA" claim for a medication that targets the underlying 54 pathophysiology, regulatory authorities require benefits on how patients feel, function 55 or (their joints) survive.⁶ While structural changes are objectively quantifiable, validly 56 assessing non-structural outcomes (i.e., pain or function) remains complex. 57 Previous research has established the concept of OA-phenotypes.^{7, 8}, i.e., the existence of 58 observable patient characteristics that systematically differ between groups of patients 59 affected by OA. Phenotyping thereby allows a stratification of a heterogeneous patient 60 population and may be reflective of different underlying pathologic mechanisms 61 defining different endotypes.^{9, 10} The existence of different OA pain-phenotypes¹¹ adds 62 an additional layer of complexity. 63 This narrative review aims at summarizing key concepts of pain-phenotyping, 64 presenting current evidence. Pain is the most important symptom of OA and its 65 treatment central to patients' well-being. The manuscript tries to capture the complexity 66 of OA-pain that underlines the need for personalized and targeted management 67 approaches based on a better understanding of pain-phenotypes and underlying 68 mechanisms. We argue that a better understanding of these aspects is crucial for

69	designing meaningful future trials and measuring treatment success. The ultimate goal
70	is to establish a framework for systematic and comparable pain assessments in OA
71	patients, with the intention of developing and allocating targeted treatments that meet
72	patients' and societies' expectations.
73	
74	
75	Pathophysiology of pain in OA
76	Pain is defined as "an unpleasant sensory and emotional experience associated with, or
77	resembling that associated with, actual or potential tissue damage". ¹² This definition
78	underlines the complex interaction of pain triggers with biological, psychological and
79	social factors (see Supplementary Table 1). ¹³ This definition also makes abundantly
80	clear that the absence of a structural correlate does not disqualify a sensation as pain,
81	and that pain can persist despite the normalization of structure. It remains unclear why
82	or which patients transition from acute to persistent or chronic pain. ¹⁴ In principle,
83	chronicity should be assumed in most OA patients with a typical pain duration of >6
84	months; indeed a "chronic pain" phenotype is consistently reported. ^{15, 16}
85	

86 Pain perception, processing and transition to chronic pain are the result of experience-87 driven neuro-structural changes¹⁷, neuro-immunologic crosstalk^{18, 19} and (epi)genetic 88 mechanisms.^{20, 21} In principle, pain perception occurs in several "morphologic layers". 89 Peripheral joint nociceptors are activated by mechanical, thermal or chemical stimuli 90 such as cytokines or chemokines released as part of inflammatory processes and 91 cartilage degradation in OA. This can also trigger vascularization and ingrowth of 92 additional nociceptors perpetuating the stimulus.²² Continuous or repetitive stimulation 93 of nociceptors can reduce activation thresholds leading to peripheral sensitization with

94 primary hyperalgesia (an abnormally increased sensitivity to pain at the site of tissue 95 damage) or allodynia (pain from otherwise non-noxious stimuli such as light touch). which may be present in OA.¹⁶ Nociceptor activity is transmitted via C-fibers (slow, 96 97 burning pain) or A-delta fibers (fast, sharp pain) to the cell body situated in the dorsal 98 root ganglion of the spinal cord. The activity is further transmitted to higher systems, 99 whereas inhibitory and excitatory influences from the local cellular environment as well 100 as thalamic centers, brainstem and cerebral cortex modulate the pain perception^{17, 23}, 101 explaining the interrelation between pain and affect^{17, 24}, but also the impact from 102 expectation, observed in placebo and nocebo phenomena^{25, 26}. 103 Based on the above mechanisms, primarily three types of pain have been discerned 104 (with some overlap) in OA: 105 I) Nociceptive pain is triggered by tissue damage and often responsive to NSAIDs. ²⁷ Pain 106 in OA was thought to be purely nociceptive ²⁸ with inflammation as potential 107 pathophysiologic trigger and driver of pain. ^{29, 30} The innate immune system, ³¹ and 108 especially macrophages play crucial roles in knee OA-pain through induction of 109 inflammatory mediators, ³² growth factors ³³ and proteinases, ³⁴ and are reciprocally 110 stimulated via nociceptor-secreted neuropeptides.³⁵ They also impact pain processing at 111 the level of dorsal root ganglia and literature supports their role in pain sensitization 112 and neuropathic pain. ^{36, 37} Preclinical animal models evaluating anti-ADAMTS5 (a disintegrin and metalloproteinase with thrombospondin motifs 5), ³⁸ or antibodies 113 targeting Toll-like Receptors ^{39, 40} in knee OA support the idea of neuroinflammatory 114 115 mechanisms in OA-pain. Similarly, the neurotrophin NGF (nerve growth factor) has been 116 implicated in OA-pain and inflammation ^{33, 41}. NGF is increased in OA joints and promising clinical results for pain relief have been reported in humans and animals ⁴²⁻⁴⁴. 117 NGF is released in response to mechanical stress and inflammation⁴⁵, its role in the 118

context of inflammation however is not fully understood yet⁴⁶, which may explain the
safety concerns that finally led to a negative benefit risk evaluation for an anti-NGF
antibody by the FDA (food and drug administration)⁴⁷. In addition, histamine receptors
have been implicated in nociception and chronic pain. Subtypes are expressed in the
peripheral and central nervous system and play a role in the modulation of nociceptive
transmission. ⁴⁸

125

126 II) Nociplastic pain is a result of central dysregulation and sensitization, and refers to 127 "pain that arises from altered nociception despite no clear evidence of actual or 128 threatened tissue damage causing the activation of peripheral nociceptors or evidence 129 for disease or lesion of the somatosensory system causing the pain" (IASP 130 (International Association for the Study of Pain) definition).^{12, 49, 50} Yet, links between 131 disease duration and measures of central sensitization seem weak ⁵¹ and most patients 132 improve markedly after joint replacement, suggesting a peripheral driver of the pain experience. 52 133 134 Nociplastic pain is decoupled from the pathology at the joint level though also associated 135 with neuroimmunologic changes. In view of the impact of central pain modulation, 136 treatments such as patient education, sleep hygiene, and psychological treatment ⁵³ or, 137 centrally acting substances such as NMDA (N-methyl-D-aspartate) antagonists 53, 138 cannabis-based medicines⁵⁴, tricyclic antidepressants, 5-hydroxytryptamine-139 noradrenaline reuptake inhibitors and gabapentinoids⁵³ ⁵⁵ may be beneficial as adjuncts 140 in improving this type of pain. Similarly, sympatholytics may be beneficial in nociplastic 141 and possibly neuropathic pain. 53 142

143 III) Neuropathic pain is typically associated with structural nerve damage⁵⁶, the 144 morphologic correlate of which currently remains elusive in OA and may be related to 145 comorbidities rather than OA (e.g., diabetes, lumbar radiculopathy, etc.). A recent 146 matched pair approach in a cohort of knee OA patients suggested a potential 147 neuropathic pain component in 8.2% (based on PainDETECT). These patients differed 148 from their likely non-neuropathic counterparts (matched for pain intensity) in having a 149 higher degree of functional impairment and more painful joints but generally less 150 pronounced radiographic joint changes. 57

151

152 Methods

This narrative review is based on a non-systematic search of current literature in Ovid
MEDLINE® using the search terms "pain-phenotype" and "knee osteoarthritis" in
various combinations to identify articles covering the area of interest. To evaluate
potential surrogate measures for pain-phenotypes PubMed ® was searched for
biomarkers evaluated in the context of OA. The search was then expanded to crossreferenced biomarkers and interventions.

159

160 Studies examining knee OA pain-phenotypes

161 The relevance of the different mechanisms for pain perception in OA underlines the 162 importance of distinguishing the predominant pain type or mechanism for a successful 163 treatment allocation especially in relation to nociceptive vs non-nociceptive pain. This 164 distinction can be achieved via pain-phenotyping, i.e., the differentiation of patient 165 clusters based on observable traits associated with differences in pain experience. 166

167 Table 1 may be placed here

168

169 Various studies have used phenotyping approaches to characterize pain-phenotypes in 170 OA as summarized in Table 1. Murphy et al.⁵⁸ cross-sectionally evaluated the co-171 occurrence of centrally mediated symptoms in older adults with hip or knee OA and 172 identified three pain-phenotypes. Those with the highest pain levels also showed high 173 levels of depression and fatigue, low sleep quality and a high burden of comorbidities 174 potentially indicating a higher overall impact from central mechanisms of pain 175 perception. Patients in this cluster had the highest disease impact on health-related 176 quality of life. The second cluster had intermediate levels of depression and fatigue, low 177 levels of pain and good sleep, possibly indicative of a mixed peripheral and central pain-178 phenotype. The third cluster had overall low levels of pain, fatigue or depression, but a 179 poor sleep quality. This could be patients with a predominantly nociceptive pain type.⁵⁸ 180 However, because this evaluation was cross-sectional, directionality and mechanisms 181 cannot be discerned. 182 Finan et al.⁵⁹ also evaluated patient reported outcome (PRO) information on 183 anxiety/depression symptoms, sleep and pain catastrophizing but included the 184 congruence between pain and structural changes versus quantitative sensory testing 185 (QST). They dichotomized pain (cut-off 4.22 out of 20 on WOMAC (Western Ontario and 186 McMaster Universities Osteoarthritis Index) pain subscale score) and radiographic 187 grade (Kellgren-Lawrence 1 and 2 vs. 3 and 4) resulting in four combinations. The high-188 pain groups trended towards higher impact in psychosocial function, which was 189 significant for patients with high-pain and low radiographic grade. The most notable 190 finding was that the high-pain and low Kellgren-Lawrence group exhibited 191 hypersensitivity on several QST modalities at unaffected anatomic sites, suggesting a

192 propensity towards central pain sensitization. In contrast, the other discordant group 193 with low-pain and high-Kellgren-Lawrence were the least pain-sensitive. ⁵⁹ 194 Similarly, Egsgaard et al.⁶⁰ aimed at identifying pain profiles in patients with OA based 195 on psychological measures, QST, Kellgren-Lawrence grade and biomarkers. Compared 196 to controls, the four resultant clusters had higher disease impact on physical 197 functioning, quality of life and pain response. In the order of pain impact (low to high), 198 the cluster of patients with overall low pain sensitivity and higher CPM (conditioned 199 pain modulation) than controls had the lowest pain. The next lowest pain cluster 200 showed increased temporal summation at the arm only (TS) and CPM and pressure pain 201 thresholds (PPT) comparable to controls, potentially indicative of an early stage of 202 chronification. Two clusters showed reduced PPTs, enhanced TS and reduced CPM. In 203 addition, one of those clusters was characterized by greater hyperalgesia, lower general 204 health and pain catastrophizing. While both of these clusters showed alterations in pain 205 thresholds quantifiable with QST, the one additionally affected by lower general health 206 and pain catastrophizing reported the highest values on the three WOMAC subscales, 207 suggesting an additive effect on pain experience.⁶⁰ 208 In addition to psychological measures, radiographic OA grade and patient 209 characteristics, Kittelson et al. included extensor strength in their approach to pain-210 phenotyping of the OAI (osteoarthritis initiative) database ⁶¹, as well as a community 211 sample that comprised participants with symptomatic OA and healthy older adults as 212 controls. ⁶² In both samples they identified four pain-phenotypes, one primarily 213 characterized by a high burden of comorbidities, one by a high level of psychological 214 distress and pain, and one with high extensor strength and a low overall burden of 215 disease. Participants from the community sample in this latter group often had a history of knee trauma or surgery.⁶² A fourth pain-phenotype was identified in both analyses; in 216

the OAI, this fourth phenotype was characterized by a high proportion of joint line and
pes anserine tenderness⁶¹. In the community sample, the fourth phenotype was
differentiated by low target knee PPTs.⁶²

220

221 Reducing heterogeneity due to differences in OA severity, Frey-Law et al.⁶³ analysed 222 pain-phenotypes in patients scheduled for knee arthroplasty and identified five 223 phenotypes based on psychological assessments, patient characteristics, QST, pain 224 characteristics, function and quality of life. One pain-phenotype exhibited low pain 225 sensitivity but high PPTs at the target knee. Another exhibited average pain sensitivity 226 to all tested stimuli. In contrast, three clusters showed high sensitivity to pain. These 227 three clusters differed in their sensitivity to TS, heat and pressure pain, and punctate 228 pain, respectively. There was no relevant impact from the other evaluated 229 characteristics except a predominance of males in the low pain group. Interestingly, in 230 the high pain sensitivity group, high punctate and high heat and pressure pain 231 sensitivity translated into higher clinical pain levels, while TS did not ⁶³. 232 Evaluating thermal measures of QST as potential indicators of central sensitization and 233 neuropathic pain and their correlation with pain levels, pain characteristics and 234 function, Wright et al.⁶⁴ compared a community sample of patients with painful knee OA 235 to pain-free volunteers. Patients with OA displayed lower PPTs than pain-free 236 volunteers at the index knee but not at other sites. In addition, patients with OA showed 237 cold pressure pain on average at higher temperatures than pain-free controls at the 238 index and contralateral knee, as well as a distant site. This cold hyperalgesia was 239 pronounced in a subgroup of 44% of patients. These patients also had a tendency 240 towards reduced thresholds for pressure and thermal pain at sites other than the target 241 knee, higher pain levels, higher functional impact and higher PainDETECT scores.

Despite the differences in QST between the groups, there were no differences in
 psychological impact. ⁶⁴

In the only longitudinal study to date to assess pain susceptibility by Carlesso et al.⁶⁵, 244 245 four distinct phenotypes were identified among people with or at risk of knee OA who 246 were free of persistent knee pain at baseline. Interestingly, the group that was the most 247 sensitized based upon PPT measures had a 2-fold higher risk of developing persistent 248 knee pain compared with the group that had the least sensitization based upon PPT and 249 TS. Further, the group that exhibited TS was not at increased risk for developing 250 persistent knee pain. ⁶⁵ The other factors that were examined (i.e., widespread pain, pain 251 catastrophizing, depressive symptoms, poor sleep) did not differentiate between the 252 groups, and thus did not contribute to risk of developing persistent knee pain. 253 Heat and cold hyperalgesia have recently further been evaluated by Carlesso et al.⁶⁶ in 254 an analysis of pain-phenotypes in patients presenting with knee OA. The analysis was 255 based on the IMMPACT recommendations for pain-phenotyping, i.e., "pain variability, 256 intensity and qualities, somatization, anxio-depressive symptoms, sleep, fatigue, pain 257 catastrophizing, neuropathic pain, and quantitative sensory tests". ⁶⁷ The three pain 258 classes separated based on PRO information (consistent high, intermediate or low 259 disease impact). The results for QST were less clear. Temperature sensitivity and PPTs 260 separated the least affected from the two other classes. Only TS was significantly 261 different for all the classes.⁶⁶ TS has also been demonstrated to separate clusters in 262 other cohorts ^{59, 60, 63, 65}, and to potentially predict acute postoperative pain intensity and 263 chronic postsurgical pain. 68,69

Two studies evaluated clinical pain-phenotyping and included imaging. In a community
sample of older adults, Pan et al. identified three subgroups of patients with knee pain. ⁷⁰
A predominantly female class including patients with high local pain, a high burden of

267 emotional problems and limited structural changes was identified, while another class 268 was dominated by males with low disease impact but definite structural changes. The 269 third class was healthy overall with limited signs of structural OA and low levels of knee 270 pain, assumed by the authors to comprise participants with early OA. Pain levels 271 between the high and low pain groups consistently differed over 10.7 years and were 272 not necessarily correlated with the presence of radiographic signs of OA. 273 In another cohort study of community dwelling adults, Burston et al.⁷¹ evaluated the 274 impact of anxiety and depression on incident knee pain. They report an odds ratio (OR) 275 of 1.71 for incident knee pain at twelve months in individuals with baseline anxiety 276 (adjusted for depression), and a 1.66 OR in patients with baseline depression (adjusted 277 for anxiety). These insights complement a preclinical OA model that demonstrated 278 astrocyte activation as potential correlate of altered pain perception in animals with 279 elevated baseline anxiety-like behaviour reversible after introducing a centrally acting 280 anxiolytic. 71

281

In summary, the above-described studies clearly demonstrate the existence of several
OA pain-phenotypes, which seem differentiable based on objective measures and PRO
information. Many approaches suggest a low pain-phenotype as well as a phenotype
with high pain perception and impact. Interestingly, few articles on OA and OA painphenotypes specifically report pes anserine tenderness^{61, 72, 73}, which may confound OApain perception and OA pain-phenotyping.

Furthermore, the observed differences and similarities in previous OA-phenotyping
analyses underline the importance of the choice of input variables for the allocation of
clusters in phenotyping.⁷² The observed differences in pain perception and painphenotypes do not necessarily correlate with the extent of radiographic changes. There

seems to be a certain overlap between structural OA and OA pain-phenotypes if imaging
is included as an input variable. ^{61, 62, 70, 72} Whether imaging information dominates
differences between phenotypes, or if pain-phenotypes are associated with structural
changes assessed on imaging merits further investigation.
To further differentiate pain-phenotypes, the degree of altered neurobiological

signalling appears to be particularly relevant; specific questionnaires and QST measures,

especially TS and PPTs or thermal sensitivity appear to be important.

299

300 Limitations of existing tools to identify OA pain-phenotypes

301

302 Pain measurement in OA studies primarily focuses on questionnaires that inquire about 303 the intensity, pain on movement and a limited range of pain characteristics to capture 304 the pain experience (Supplementary Table 1). However, most of these questionnaires do 305 not differentiate the underlying pain mechanism(s) at play in any given individual. 306 Further highlighting the complexity of OA, numerous biomarkers (as potential 307 indicators of pathophysiologic mechanisms in OA) and interventions have been 308 evaluated in the context of structural and symptom (pain) OA outcomes (Supplemental 309 Table 2). Patients with different pain-pheno- and endotypes may report similar pain 310 intensity and dimensions. These pain measures therefore may not be suitable to 311 categorize patients but should be used as outcome measures to explore treatment 312 effects. To identify different pain-pheno- and -endotypes, assessments should include 313 clinical/biological information as well as medical history (e.g., burden of comorbidities, 314 signs of dysfunctional pain experience or pain quality, sleep, anxiety and depression, 315 physical activity and assessment of somatosensory function by QST, see Supplemental 316 Table 3). Given the above-described convergence of structural OA and OA pain317 phenotypes if imaging or performance measures are added to the clustering, the318 selection of input variables has to be carefully considered.

Comorbidity impacts pain ⁷⁴ and various measures are used to estimate the burden of 319 320 comorbidity (comprehensively summarized by Stirland et al. 75). It is however vital to 321 consider a score's "original purpose and the outcomes for which it is validated".⁷⁵ 322 Scores developed to predict mortality (e.g., Charlson Comorbidity Index) may be 323 unsuitable to reflect the burden of comorbidity and its impact on physical functioning. 324 Affective states such as anxiety, depression or pain catastrophizing influence pain 325 modulation and perception of pain. While there are diagnostic criteria and tools to 326 identify and grade anxiety and depression, a consensus regarding how to measure 327 catastrophizing has not yet been reached. ⁷⁶ Measures of emotional dysregulation or 328 positive and negative affect can also be useful. ^{77, 78} Kinesiophobia has been reported as 329 predictor of disability impacting quality of life in various pain conditions; it has been 330 associated with chronic pain and thus may also present a useful addition.^{79,80} 331 Exercise can positively influence pain⁸¹; pain and activity may have a reciprocal 332 relationship in some individuals; it may therefore be misleading to assess one without 333 the other. ^{82, 83} This results in methodologic challenges. Objective performance tests are 334 subject to day-to-day variability and reflect what patients are able to do under 335 observation rather than what they habitually do in their free-living environment. The 336 domain of activity, in the future, may best be captured using digital devices that allow 337 the measurement of indicators in the free-living environment like step count, activities 338 at a certain heart rate or radius of mobility. Similarly, objective assessment of sleep 339 structure may be obtained using wearable technology.^{84, 85} Measuring elements of sleep 340 is increasingly recognized as an important aspect to understanding the pain experience 341 since sleep and pain are also closely inter-related; pain may disrupt sleep, and sleep

disturbance negatively impacts descending pain inhibitory pathways, heightens pain
sensitivity and attenuates opioid analgesia. ^{84, 86-88} These examples underline the
importance of systematically assessing pain and potential confounders in an integrative
approach.

346

347 Considerations for a broader collection of pain measures

348 This summary highlights the complexity of the pain experience as multidimensional 349 physical and psychological phenomenon, as well as of the plethora of assessment tools. 350 It also suggests the existence of different patterns of observable traits, OA pain-351 phenotypes, which likely reflect different underlying mechanisms contributing to the 352 overall pain experience. Striving for the development of a personalized and targeted 353 management of OA, pain is a critical factor, and central to patients' well-being. OA-pain 354 is associated with multiple pathophysiological mechanisms reflected in distinct pheno-355 and endotypes. This implies the need to systematically define those pain-pheno- and 356 -endotypes independent of the underlying OA pheno- and -endotype.

357

358 We therefore suggest the systematic collection of additional pain-related data, such as 359 pain quality, including potential signs of sensitization and other altered neurobiological 360 mechanisms, burden of comorbidity, presence of anxio-depressive psychopathology, 361 sleep quality and physical activity as a minimal set of assessments. Other aspects such as 362 pain catastrophizing, kinesiophobia, dysregulation of affect, etc. may play an important 363 role. At the moment there is however less consensus about their independent relevance 364 and optimal tools for the assessment of these concepts. Similarly, the potential 365 application of this additional pain-related data necessitates further evaluation. The 366 individual use of the PRO information could lead to unnecessary fragmentation of the

367 patient population. The use of patient response or patient characteristics patterns in 368 form of phenotypes for subgroup analyses or treatment allocation though could support 369 drug development. Pain-phenotyping could be specifically valuable to discriminate 370 treatments without any effect on pain, from those that target specific pain processes. 371 QST allows valuable additional insights into pain processing. Necessary expertise, 372 equipment and time for valid assessments may be challenging, thereby impacting the 373 implementability of comprehensive QST protocols in large multicenter trials. 374 Nevertheless, future research may guide the construction of targeted somatosensory 375 assessment-batteries based on their discriminative value e.g., in combination with PROs, 376 which would allow a broad implementation and add relevant scientific value to OA 377 trials.

378

379 One challenge has been the comparability of various PROs that focus on slightly different 380 clinical domains. Georgopoulous et al. have recently demonstrated, that harmonized results of the 4 most widely used PROs for pain assessment produce similar patient 381 382 acceptable symptom states and are thus comparable.⁸⁹ To increase our knowledge 383 about pain-phenotypes from published and future studies, a similar concept to generally 384 interpret and compare PRO results could be applied, leveraging established cut off 385 values ⁷¹. Alternatively cut off values such as tertiles or quartiles of the original score 386 range could be used. ⁶⁵ The latter approach is based on the assumption, that for a score 387 e.g., ranging from 0-100 with 100 denoting high impact from a given pathology, people 388 who score between 0-25 or 0-33 are less likely to be impacted, compared to those 389 scoring between 66-100 or 75-100. While on a granular level, the different scores may 390 convey different nuances of patient experience (and thus allow focus in a specific 391 project), a separation in tertiles or quartiles in principle allows the clear identification of highly vs marginally affected individuals for comparison with other studies. This could
also facilitate the implementation of systematic PRO-based assessments in clinical
practice to allow individualized treatment approaches.

395

396 The legacy of numerous failed trials, the increasing cost pressure on healthcare systems, 397 and the public and individual health burden of OA are concerning. Given the increase in 398 mechanistic understanding, the field is under a certain pressure to develop medicines 399 that address patients' symptoms and halt or reverse OA. One prerequisite for the 400 development of worthwhile treatments is the establishment of clinical endpoints that 401 provide a meaningful reflection of disease modification and long-term patient benefit. 402 This can only be accomplished if we better understand and measure pain in OA which 403 could also give further insights in the pain structure relationship. However, to achieve 404 real progress, data need to be comparable. Systematic generation of data that allow OA 405 pain-phenotyping may be one piece of the puzzle towards a "treatment of OA".

406

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412

413 Author contributions

414 AH and FS have collected the information for the tables and performed literature

415 research. All authors have been involved in the analysis and interpretation of the data

416 and contributed to the final manuscript.

417

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425 **Competing interests**

- 426 Franziska Saxer is employee and shareholder of Novartis, she is affiliated to the University
- 427 Basel and member of the European Union Medical Devices Expert Panel section
- 428 Orthopaedics, traumatology, rehabilitation, rheumatology
- 429 Alexa Hollinger is a medical fellow at Novartis, she is affiliated with the University Hospital
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442

443 **REFERENCES**

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Table 1: Summary of key OA pain-phenotyping studies

Pain in knee OA

Version 1, 15.04.2023

1/18

Reference	Population	Sample Size (for comparative studies n of OA patients : n of controls)	Phenomenon	Comparator	Approach	Evaluation	Population Characteristics	Phenotypes/Groups
Murphy et al. ⁵⁸ 2011	older adults (≥65y) with hip or knee OA and signs of primary fatigue	129 (69% knee OA)	relationship among pain, fatigue, and physical activity	na	Hierarchical agglomerative cluster analysis Cross sectional Community sample	Patient Characteristics	61% female Age: 72.2 (+/-9.8), range 65 to 90 y BMI: 30.5 +/-5.9 kg/m ² , range 21.5 to 49.9 Self-reported duration of pain (months) 132.1 (146.5) range 0 to 708	no significant differences in patient characteristics
						Brief Fatigue Inventory	BFI total 4.5 (2.0) range 0.25 to 8.75	Cluster I: 36% highest scores on all measures - high stiffness, high disability, TUG 13.5 +/- 8.9 Cluster II: 30% subclinical depression, moderate fatigue, moderate fatigue, moderate illness burden, overall low pain, low sleep disturbance - stiffness moderate.
						WOMAC	WOMAC pain 7.9 (3.4) range 2 to 20 WOMAC stiffness 3.3 (1.7) range 0 to 8 WOMAC disability 20.9 (10.3) range 3 to 42	
						 5 times daily NRS pain assessment Illness burden (41 somatic symptoms) Timed up-and-go test Activity measured via Actiwatch Pittsburgh Sleep Quality Index (PSQI) Center for Epidemiologic Studies Depression Scale (CES-D) 		disability low, TUG 10.5 +/-2.1 s <u>Cluster III:</u> 34% relevant sleep disturbance, mild pain, low fatigue and depression scores, low illness burden - low stiffness, moderate disability, TUG 10.2 +/- 2.3s

Reference	Population	Sample Size (for comparative studies n of OA patients : n of controls)	Phenomenon	Comparator	Approach	Evaluation	Population Characteristics	Phenotypes/Groups
Finan et al. ⁵⁹ 2013	Baseline of study to evaluate psychological treatments in OA patients with/without insomnia		Association between self- reported levels of pain with measures of central sensitization in the absence of moderate-to- severe radiographic evidence of pathologic changes of knee OA	na.	cross-sectional multivariate general linear modeling	 Patient Characteristics STAI CES-D PCS PSQI Radiographic disease severity (Kellgren/Lawrence) QST PPT CPT Mechanical phasic pain Thermal phasic pain Sensitivity to tonic pain CPM 	66.7% female Age: 61.05 +/- 8.93 y BMI: 30.94 +/- 5.85 kg/m ²	Low pain/low knee OA grade (21.2%): overall lowest BMI High pain/high knee OA grade (28.3%): reduced distant (and local) PPT vs low pain groups, high rate of depression, anxiety, sleep disturbance and pain catastrophizing, overall highest BMI Low pain/high knee OA grade (23.90%): overall oldest group High pain/low knee OA grade (26.5%): significantly increased pain response to distant mechanical phasic stimuli and thermal phasic pain compared to high knee OA groups, reduced distant PPT vs low pain groups, high rate of depression, anxiety, sleep disturbance and pain catastrophizing, overall youngest group no differences in CPM or QST measures locally, education and income as significant covariates

Version 1, 15.04.2023

3/18

Reference	Population	Sample Size (for comparative studies n of OA patients : n of controls)	Phenomenon	Comparator	Approach	Evaluation	Population Characteristics	Phenotypes/Groups
Egsgaard ⁶⁰ et al. 2015	full spectrum from no clinical OA to clinical OA, randomly selected from pre-existing database 40-80y controls with no OA and little or no pain	280 (216:64)	identification of knee pain profiles identification of marker patterns correlating to pain profiles	non-OA knees largely independent of pain	Principal Components Analysis (PCA) with clustering using Ward's method with squared Euclidean distance	Patient Characteristics	64% female Age: 61.7 +/- 10.0 y BMI: 33.9 +/- 7.0 kg/m ²	Principal components: PC1: physical health questionnaires PC2: peripheral, central, and spreading sensitization, PC3: biochemical markers, PC4: pain catastrophizing, PC5: temporal summation.
						 OA grade Comorbidities Number of painful joints Pain duration Pain localization WOMAC Lequesne functional index EQ-5D Pain catastrophizing QST PPT TS CPM Biomarkers VICM CIM CRP CRPM CIIIM 		Profile A (12.5%): moderate impact on WOMAC/ Lequesne, low to moderate catastrophizing, near normal TS, high CPM and PPT as potential sign of resilience, still reduced QoL Profile B (27.3%): moderate impact on WOMAC/ Lequesne, low to moderate catastrophizing, near normal TS, moderate CPM but reduced PPT, reduced QoL Profile C (39.4%): moderate impact on WOMAC/ Lequesne, low to moderate catastrophizing, increased TS, reduced CPM and

				PPT, reduced QoL, CRP near normal Profile D (18.9%): higher impact on WOMAC and especially Lequesne, increased catastrophizing, increased TS, reduced CPM and PPT, reduced QoL
				Profile E (1.9%): outlier cluster, not reported in detail
				controls low impact on WOMAC/Lequesne, moderate CPM and PPT, low TS

Reference	Population	Sample Size (for comparative studies n of OA patients : n of controls)	Phenomenon	Comparator	Approach	Evaluation	Population Characteristics	Phenotypes/Groups
Kittelson et al. ⁶¹ 2016	OAI from the incident and progression cohort	3494	Knee OA pain- phenotypes based on 1) knee OA pathology 2) psychological distress 3) altered pain neurophysiology 4) relation to patient characteristics	na	Latent Class Analysis cross sectional cluster analysis (4-year follow- up visit) with some longitudinal information	Patient Characteristics	OA 59.2% female Age: 64.9 +/-9.0 y BMI: 28.9 +/- 5.0 kg/m ²	<u>Class 1:</u> on average older than all other classes, higher proportion of females, slowest walking speed, high level of comorbidities <u>Class 2:</u> on average older than class 3/4, high levels of knee joint tenderness, weak extensor strength
						 Numeric Pain Rating Scale (NPRS) WOMAC Radiographic severity of knee OA MVIC Tenderness of the knee joint Modified Charlson Comorbidity Index Number of pain sites (as surrogate for central sensitization) CES-D Modified version of the coping strategies questionnaire- catastrophizing subscale 20-meter timed walking test at self-selected walking speed Health seeking behavior (unstructured question) 	similar symptom duration and health seeking behavior	weak extensor strength and high proportion of pes anserine tenderness <u>Class 3:</u> highest pain level, psychological distress, highest number of painful sites and more severe radiographic OA <u>Class 4:</u> mild radiographic OA, low levels of pain and comorbidity, highest average extensor strength
Kittelson et al. ⁶² 2021	Recruitment from community (healthy elderly) and orthopaedic clinics (OA patients)	183 (152:31)	Knee OA pain- phenotypes based on 1) multimorbidity 2) psychological distress 3) pain sensitivity	healthy community dwelling elders	Latent Profile Analysis Cross sectional Community sample	Patient Characteristics	OA 64.5% female, control 64.5% female Age: OA 65.2 +/- 8.5 y, control 64.9 +/- 9.0 y	Group 1 (9% of pt with knee pain): characterized by high FCI scores (upper gastrointestinal, osteoporosis, heart disease, asthma), slower walking speed than group 2/4 ("weakness and

50-8	85y	4) knee impairment or pathology			BMI: OA 30.2 +/- 6.0 kg/m ² , control 26.7 +/- 4.6 kg/m ²	heightened pain sensitivity with multimorbidity") Group 2 (63% of pt with knee pain): low PCS and FCI (vs group 1 and 3), higher target knee PPT
				 Visual analog scale (VAS) WOMAC pain ICOAP Normalized knee extensor strength at maximum voluntary isometric contraction (MVIC) Functional Comorbidity Index (FCI) Pain Catastrophizing Scale (PCS) Walking speed Health seeking behaviour (unstructured question) Symptom duration QST PPT (target knee) 	similar symptom duration and health seeking behavior	and lower extensor strength than healthy elderly or group 4 ("weakness and heightened pain sensitivity") <u>Group 3 (11% of pt with knee pain)</u> : characterized by pain catastrophizing, higher pain ratings than group 2/4 ("weakness and heightened pain sensitivity with pain associated distress") <u>Group 4 (17% of pt with knee pain)</u> : characterized by high PPT vs all other groups, otherwise similar to healthy elderly, highest proportion of pt with previous knee surgery or
				Tampa Scale for Kinesiophob Comorbidity Index, PPT (regin Radiographic Severity of Kne evaluated but excluded based with pain intensity (Spearman	onal/distant), e Osteoarthritis, CES-D I on weaker correlation	trauma ("normal strength, low pain sensitivity")

Reference	Population	Sample Size (for comparative studies n of OA patients : n of controls)	Phenomenon	Comparator	Approach	Evaluation	Population Characteristics	Phenotypes/Groups
Frey-Law et al. ⁶³ 2017	Baseline of TANK (TENS After New Knee) study NCT01364870 ≥30y scheduled for primary total knee joint replacement	218	QST pain sensitivity profiles in advanced knee OA	na	Principal Components Analysis (PCA) and Principal Axis Factoring (PAF) with clustering using Ward's method with squared Euclidean distance	Patient Characteristics	54.6% female (50% in control group) Age: not reported BMI: not reported	Low Pain Sensitivity Profile (18.3%): low QST based standardized pain sensitivity before and after adjustment for age and sex Average Pain Sensitivity Profile (38.5%): average QST based standardized pain sensitivity, after adjustment for age and
	scheduled for primary total knee joint				 Pain intensity (rest and movement) via 21-point NRS Pain duration Analgesic medication State-Trait Anxiety Inventory (STAI) anxiety subscale Geriatric Depression Scale (GDS), 5-item version PCS KOOS SF-36 QST PPT HPT and HPToI Punctate Pain Intensity via VAS TS via tonic heat stimulus 		Profile (38.5%): average QST based standardized	

Commented [MB1]: Especially?

Pain in knee OA

		pain levels also in KOOS, at rest, gait and range of movement
		no relevant impact from other assessments apart from sex. Men were allocated predominant in low pain sensitivity cluster. After adjustment higher pain sensitivity for non- white and/or hispanic individuals

Commented [MB2]: Maybe rephrase to avoid misunderstanding. As I understand it, men are predominant in the low pain sens. cluster.

Pain in knee OA

Reference	Population	Sample Size (for comparative studies n of OA patients : n of controls)	Phenomenon	Comparator	Approach	Evaluation	Population Characteristics	Phenotypes/Groups
Wright et al. ⁶⁴ 2017	adults with painful knee OA pain-free volunteers (≥50y)	120 (80:40)	widespread cold, pressure, and heat hyperalgesia in OA patients differences in QST measures, levels of pain, pain characteristics, and perceived function in patients with wide-spread cold hyperalgesia	pain free control OA patients with and without wide-spread cold hyperalgesia	Standard statistics Cross sectional Community sample	Patient Characteristics	OA 55% female, control 60% female Age: OA 64, range 50 to 86 y; control 64, range 51 to 86 y OA 38% obese, control 10% obese	no significant differences in patient characteristics
						Brief Fatigue Inventory	BFI total 4.5 (2.0) range 0.25 to 8.75	<u>OA vs pain free:</u> sign. higher index knee PPT in OA (pressure
						WOMAC	OA WOMAC pain, 18.5/50 OA WOMAC function, 53.4/250	hyperalgesia: 22.50% index knee, 16.25% contralat. knee, 3.75% distant site) sign. higher CDT at index
						Short-Form Health Survey (SF-36)	43.75% (n=35) cold hyperalgesic based on 12.25°C cut off	and contralat. knee (cold hypoesthesia 11.25% index knee, 17.50% contralat. knee, 17.50% distant site; cold hyperalgesia 47.50% index knee, 37.50% contralat. knee, 43.75% distant site) sign. higher overall WDT in OA, no differences in HPT (heat hypoesthesia 11.25% index knee, 17.50% contralat. knee, 17.50% distant site; heat
						 PainDETECT Pain quality assessment scale (PQAS) QST PPT CDT CPT WDT HPT 		

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				hyperalgesia 47.50% index knee, 37.50% contralat. knee, 43.75% distant site)
				Cold hyperalgesic vs non-hyperalgesic OA patients:
				sign. lower cold detection and cold pain threshold at all sites cold-hyperalgesic vs non-cold hyperalgesic OA patients, no difference between non-hyperalgesic OA patients vs pain-free controls
				sign. lower warmth detection threshold at index knee and distant site (cold hyperalgesic patients vs all others), sign. lower warmth detection threshold at contralateral knee (cold hyperalgesic patients vs pain free controls, but not vs other OA patients), lower heat pain threshold at all sites (cold hyperalgesic patients vs other OA patients), but no difference between cold hyperalgesic patients and controls.
				sign. higher index knee and contralat knee PPT, no sign. difference at distant site
				no differences in SF36 based on cold hyperalgesia in OA patients, higher WOMAC pain and disability in patients with cold

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				hyperalgesia, correlation between cold hyperalgesia and PainDETECT scores and surface and paradoxical subscores in pain quality assessment scale

Reference	Population	Sample Size (for comparative studies n of OA patients : n of controls)	Phenomenon	Comparator	Approach	Evaluation	Population Characteristics	Phenotypes/Groups
Pan et al. ⁷⁰ 2019	Recruitment from community (healthy elderly) and orthopedic clinics (OA patients) 50-85y	Tasmanian Older Adult Cohort Study	963	knee pain- phenotypes in an older population	Latent Class Analysis Cross sectional Community sample	Patient Characteristics	50% female (sampling strategy) Age: 62.8 +/- 7.4 y BMI: 27.7 +/- 4.6 kg/m ²	Class 1 (25%): highest proportion of females, on average more emotional problems, higher burden of comorbidity, more severe knee pain and more painful sites, lower knee structural damage, lower education
	<u>əu-əəy</u>					 WOMAC pain Number of painful sites MRI characteristics (cartilage defects, bone marrow lesions, effusion-synovitis) Radiographic presence of knee OA Education level Single mental health item from the short form-8 4-item comorbidity questionnaire (heart attack, diabetes, hypertension, rheumatoid arthritis) 		Class 2 (20%): more males, higher level of education, fewer painful sites or structural knee abnormalities, lower levels of pain Class 3 (50%): overall lowest prevalence of knee pain, comorbidities, radiographic OA, structural damage and low BMI consistently WOMAC and painful sites Class 1 > Class 2 > Class 3 over average 10.7 y

Reference	Population	Sample Size (for comparative studies n of OA patients : n of controls)	Phenomenon	Comparator	Approach	Evaluation	Population Characteristics	Phenotypes/Groups
Burston et al. ⁷¹ 2019	participants from a community- based cohort study ≥40y	230 (130:100) 3274 for impact of anxiety (351 anxiety at baseline) on incident knee pain at 12 months 3767 for impact of knee pain (1020 with baseline knee pain) on incident anxiety at 12 months	associations between knee pain, pain spread, anxiety, and depression	Non-OA patients	Spearman correlation and linear regression	Patient Characteristics	OA 61.9% female, control 58.2% female Age: OA 60.27 +/- 9.61 y; control 63.06, +/- 8.88 y BMI: OA 27.1 +/- 4.56 kg/m ² , control 30.09 +/- 6.62 kg/m ²	Impact of anxiety (25% of population) anxiety sign. associated with all pain measures and PPTs after adj. for depression odds ratio (OR) for incident knee pain at 12 months in patients with anxiety 1.71 (adj. for depression) OR for incident anxiety at 12 months in patients with knee pain 1.18 (after adj.
						 HADS Intermittent and Constant Osteoarthritis Pain scale (ICOAP) Numeric Rating Scale (NRS) OA severity (Kellgren- Lawrence) QST PPT 		for depression) OR for incident anxiety at 12 months in patients with depression 3.20 Impact of depression (10% of population) OR for incident knee pain at 12 months in patients with depression 1.66 (adj. for anxiety)

Reference	Population	Sample Size (for comparative studies n of OA patients : n of controls)	Phenomenon	Comparator	Approach	Evaluation	Population Characteristics	Phenotypes/Groups
Carlesso et al. ⁶⁵ 2019	MOST population 50-79y having/at risk of developing knee OA without persistent knee pain	852	pain susceptibility phenotype (PSP) based on development of persistent pain at 2 years	na	Latent Class Analysis observational longitudinal	 Patient Characteristics Widespread pain index (WPI) QST PPT TS Coping Strategies Questionnaire (single item for pain catastrophizing) - CES-D 	55% female Age: 67y BMI: 29.5 kg/m ²	Pain susceptibility phenotypes (PSP) PSP 1 (34%): pressure pain sensitivity (~16– 26%), facilitated TS (33– 35%) PSP 2 (31%): pressure pain sensitivity (0–6%), facilitated TS (2–10%), 22% non-caucasian PSP 3 (23%): pressure pain sensitivity (75–89%), facilitated TS (53–58%), 74% female, higher risk of developing incident knee pain PSP 4 (12%): pressure pain sensitivity (0–4%), facilitated TS (82–90%), 26% female, 23% non- caucasian, mean age 70% no relevant differences in other aspects analyzed

Reference	Population	Sample Size (for comparative studies n of OA patients : n of controls)	Phenomenon	Comparator	Approach	Evaluation	Population Characteristics	Phenotypes/Groups
Carlesso et al. ⁶⁶ 2022	orthopaedic specialist confirmed diagnosis of OA ≥40y	343	Pain-phenotype identification based on IMMPACT criteria	na	Latent Class Analysis observational longitudinal	Patient Characteristics	63% female Age: 64y BMI: 32kg/m²	Class 1 (49%): overall low scores in all assessed measures (i.e. low severity) or marginal signs of central sensitization according to QST Class 2 (40%): overall moderate scores in assessed measures, but high pain variability, mixed QST values
						 Modified Pain Detect Questionnaire Hospital Anxiety and Depression Scale (HADS) Patient Health Questionnaire-15 (self- administered version of the Primary Care Evaluation of Mental Disorders (PRIME-MD) diagnostic instrument) Pain Catastrophizing Scale Multidimensional Fatigue Inventory Pain variability (INRS 3 times via text for a week) Average pain intensity (NRS, recall 1 week) Pittsburgh Sleep Quality Index (PSQI) Short form McGill Pain Questionnaire 2 QST 		Class 3 (11%): overall highest scores in assessed measures (except pain variability), QST values for PPT patella, TS, cold pain and CPM heat pain as indicator of relevant central sensitization decreasing function from class 1 to class 3 considering walk fast and climb stairs, no significant difference for sit stand increasing health care utilization of 44% and 240% for class 2 and 3 respectively compared to class 1

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	PPT
	• <i>T</i> S
	• CPT
	• HPT
	CPM (Conditioned
	pain modulation)
	Self-report Charlson
	comorbidity index
	Life Orientation Test-
	Revised scale
	(dispositional
	optimism)
	Chronic Pain Self
	Efficacy Scale
	Kellgren-Lawrence
	grade
	Knee Injury and
	Osteoarthritis
	Outcomes Score
	(KOOS) activities of
	daily living subscale
	Core measures of functional parformance
	functional performance
	(1) transition from sit to
	stand, 2) walk fast and
	3) climb stairs
	Healthcare Utilization
	(via provincial
	insurance system in
	one vicinity)

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Abbreviations:

pt: patients

BMI: Body Mass Index; BFI: Brief Fatigue Inventory; CES-D: Center for Epidemiologic Studies Depression Scale; FCI: Functional Comorbidity Index; GDS: Geriatric Depression Scale HADS: Hospital Anxiety and Depression Scale; ICOAP: Intermittent and Constant O steoarthritis Pain scale KOOS: Knee Injury and Osteoarthritis Outcomes Score; MVIC: Normalized knee extensor strength at maximum voluntary isometric contraction; NRS: Numeric Rating Scale; NPRS: Numeric Pain R ating Scale PCS: Pain Catastrophizing Scale; PRIME-MD: Primary Care Evaluation of Mental Disorders; PQAS: Pain quality assessment scale; PSQI: Pittsburgh Sleep Quality Index; S F-36: Short-Form Health Survey; STAI: State-Trait Anxiety Inventory; WOMAC: Western Ontario and McMaster Universities Osteoarthritis Index;

QST: Quantitative Sensory Testing; CDT: Cold Detection Threshold; CPT: Cold Pain Threshold; CPM: Conditioned pain modulation; HPT: Heat Pain Threshold; HPTol: Heat Pain Tolerance; PPT: Pressure Pain Threshold; TS: Temporal Summation; WDT: Warmth Detection Threshold

CIM: Collagen I Metabolite; CIIIM: Sollagen III Metabolite; CRP: C-Reactive Protein; CRPM: C-Reactive Protein Metabolite; VICM: Citrullinated Vimentin Fragment;

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