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Wireless multichannel vibroarthrographic recordings for the assessment of knee osteoarthritis during three activities of daily living

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Published in: **Clinical Biomechanics**

DOI (link to publication from Publisher): 10.1016/j.clinbiomech.2019.11.015

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Publication date: 2020

Document Version Accepted author manuscript, peer reviewed version

Link to publication from Aalborg University

Citation for published version (APA):

Madeleine, P., Andersen, R. E., Larsen, J. B., Arendt-Nielsen, L., & Samani, A. (2020). Wireless multichannel vibroarthrographic recordings for the assessment of knee osteoarthritis during three activities of daily living. Clinical Biomechanics, 72, 16-23. Advance online publication. https://doi.org/10.1016/j.clinbiomech.2019.11.015

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54 **1. Introduction**

Osteoarthritis (OA) is the most prevalent joint disease (Vos et al., 2012) and a global issue 55 resulting in chronic pain and impaired mobility. Knee OA represents a scientific challenge 56 accounting for 83% of total OA burden (Vos et al., 2012). Further, the biomechanics of the 57 knee joint are of particular interest due to its weight bearing role, high injury rate and 58 degenerative processes leading to OA (Maffulli et al., 2011). Altered hamstring-quadriceps 59 60 muscle balance and kinematics have been reported during gait in knee OA (Hortobagyi et al., 2005; O'Connell et al., 2016). These studies confirm the importance of assessing activity of 61 62 daily living (ADL) in line with the OsteoArthritis Research Society International recommendations for testing physical function in patients with OA (Dobson et al., 2013). 63 Beside biomechanical assessments of ADL in knee OA, vibroarthrography (VAG) of 64 the knee, i.e., measuring the vibrations reflecting knee crepitus during joint motion has also 65 been used as a non-invasive diagnostic tool as a proposed surrogate model for roughness, 66 softness or lubrication of the cartilage surface (Rangayyan and Wu, 2009; Wu et al., 2010). 67 Since the publication of pioneer work of Blodgett (1902) and Walters (1929), the study of the 68 knee joint VAG signal has gained in sensitivity due to improvements in micro-electronics and 69 specificity due to advanced signal processing (Andersen et al., 2018; Krecisz and 70 Baczkowicz, 2018). Similar to the progress made in surface electromyography (Frigo and 71 72 Crenna, 2009) or mechanomyography (Madeleine et al., 2007), technological advances have 73 also enabled to record multi-channels VAG of the knee joint (Andersen et al., 2018; Befrui et al., 2018; Wiens et al., 2016). In these studies, two to eight miniature accelerometers have 74 been attached over the skin of the knee of participants enabling to assess spatial dependencies 75 of the VAG signals by calculating VAG topographical maps. Variations in the internal 76 pressure distribution applied to cartilage and synovial fluid explain non linearity and spatial 77 dependencies of the compound VAG signal (Neu et al., 2008; Wu et al., 2016). We have 78

recently showed non-uniform distribution of VAGs during knee flexion-extension movement 79 (Andersen et al., 2018). More specifically, combining linear and nonlinear parameters has 80 improved our understanding of the VAG signals. As such, the use of multichannel VAG 81 recordings and advanced processing approaches has been suggested to discriminate between 82 knee OA patients and asymptomatic participants and between different types of ADL 83 (Andersen et al., 2018). However, no studies have used multichannel VAG to delineate 84 85 differences among knee OA patients and asymptomatic participants during ADL. Studies assessing the changes in VAG in knee OA patients compared with 86 87 asymptomatic participants have shown high accuracy, sensitivity and specificity (Wu, 2015). Especially, the existing body of VAG literature has revealed increased amplitude, absolute 88 variability and frequency contents in knee OA patients compared with asymptomatic 89 90 participants (Baczkowicz et al., 2017; Baczkowicz and Majorczyk, 2016; Tanaka and 91 Hoshiyama, 2012). Changes in the regularity of the VAG have also been reported confirming that nonlinear analyses provide genuine VAG information (Wu et al., 2016). As previous 92 93 clinical studies using multichannel VAG have only investigated source localisation or classification issues during knee flexion-extension (Rangayyan and Wu, 2009; Wu et al., 94 2010), information concerning the spatial dependencies of linear and nonlinear parameters 95 during ADL is lacking. 96

The purposes of this study were to collect and analyse wireless multichannel VAG topographical maps and characteristics in knee OA patients and asymptomatic participants during ADL. We hypothesised (i) that higher VAG amplitude, variability and frequency contents as well as changed VAG regularity would characterise knee OA patients compared with asymptomatic participants (Baczkowicz et al., 2017; Wu et al., 2016), (ii) that VAG recordings would differentiate between ADL types (Andersen et al., 2018) and (iii) the presence of non-uniform distribution of VAGs (Andersen et al., 2018). If confirmed, the

present technique could be used in clinical practise to objectively assess motor functionduring some typical ADL.

106

107 **2. Methods**

108 2.1. Design

The present investigation was a cross-sectional study involving patients suffering from knee OA and asymptomatic participants. The study was conducted according to the ethical guidelines of the Helsinki Declaration and was approved by the North Denmark Region Committee on Health Research Ethics (VN-20160081). All participants provided written informed consent.

114 2.2. Participants

115 Twenty knee OA patients (11 males and 9 females) were recruited from a database at the

116 Centre for Clinical and Basic Research (CCBR, Aalborg, Denmark) and 20 asymptomatic

117 participants (10 males and 10 females) were recruited from the dwelling community (Table

118 1). Knee OA patients were diagnosed in accordance with American College of Rheumatology

119 classification (Kellgren and LAWRENCE, 1957). Participants were screened for inclusion by

120 a medical doctor at CCBR. Inclusion criteria for knee OA patients included age 18-80,

121 clinically diagnosed knee OA with Kellgren-Lawrence grade ≥ 2 , self-reported pain during

122 walking and BMI <35, no use of painkillers in the 24 hours prior to experimentation.

123 Inclusion criteria for asymptomatic participants were age 18-80, no diagnosed knee OA, no

self-reported pain during walking and BMI <35, no use of painkillers in the 24 hours prior to

experimentation. Exclusion criteria were pregnancy, drug addiction, lack of ability to

126 cooperate and, participation in other pain trials throughout the study period.

127 2.3. Experimental protocol

All participants participated in one session and they all completed the entire session. The 128 same experimenter (R.E.A.) conducted all tests. The participants performed three different 129 types of ADL in a counterbalanced order: (i) 5 repetitions of sit to stand movement (ii) Stairs 130 descent (10 stairs). (iii) Stairs ascent (10 stairs) in line with the recommendations for testing 131 physical function in patients with OA (Dobson et al., 2013). The sit to stand exercise were 132 carried out at a slow pace (60-s were allowed for the five repetitions). Arms were maintained 133 134 along the body side through the sit to stand exercise (Malling and Jensen, 2016). Hands were not used during raising movement from the chair. Stairs descent and ascent were carried out 135 136 without using the hand railing at the slowest speed that the participants were comfortable with while maintaining balance. Pain intensity was assessed using a visual analogue scale 137 ("0": no pain and "10": worst pain imaginable) after sit to stand and stairs descent-ascent. 138

139 2.4. Vibroarthrographic recording

VAG recording was carried out using a custom-made device based on a Trentadue wireless 140 multichannel recorder (OT Bioeletronica, Torino, Italy), a custom 16 channel accelerometers 141 adaptor and micro machined accelerometers LIS344ALH (ST microelectronics, Geneva, 142 Switzerland). The setup has a sensitivity of 600 mV/g and 0-1800 Hz linear transmission. The 143 recording probe is composed of an accelerometer chip supporting board set up to only record 144 acceleration in the orthogonal direction. The probe weight is approx. 0.75 g with wire and has 145 an 8.5×7 mm size. The VAG device contains a 10-500 Hz band-pass filter. Gain was set to 3 146 147 and the VAGs were sampled at 2000 Hz. The VAGs were recorded using a custom script (IOIVibcorder, Aalborg University, Aalborg, Denmark) implemented in Matlab 2016a (The 148 MathWorks, Inc, Natick, Massachusetts, United States). 149

During ADL tests, the recording device was placed in a belt bag around the waist of the

151 participant with wires attached to the thigh allowing natural movement. Eight accelerometers

152 were placed on the most painful knee of the knee OA patients (right knee for all patients but

one due to knee surgery). The accelerometers were placed accordingly (right knee for all but 153 one) for the asymptomatic participants. Accelerometers were attached to the skin with double 154 side tape. Four accelerometers separated by 1-2 cm were placed on the participant's patella in 155 a square configuration. One accelerometer was placed on the tibial tuberosity below the 156 patella, two were placed respectively on the lateral side of the knee 1-2 cm from the lateral 157 epicondyle and on the medial side of the knee 1-2 cm from the medial epicondyle of femur 158 159 towards the patella. The last accelerometer was placed above the knee over the quadriceps tendon in line with our previous study (Andersen et al., 2018), see Fig. 1. Special attention 160 161 was given to ensure that motion did not loosen the accelerometers attachment.

162 *2.5. Data analysis*

163 Data preprocessing and VAG parameter extraction were carried out using Matlab.

Preprocessing consisted of conversion of VAG signals into SI units (ms⁻¹) and digital filtering 164 using a bandpass FIR filter using a Kaiser windowed, 10-500 Hz (1453-points, beta: 5.6533). 165 Epochs containing the beginning and end of the recorded ADL were extracted and the 166 outcome parameters were processed across time. A recent literature review conducted by the 167 authors [1] has shown that six parameters are likely to thoroughly depict the characteristics of 168 the VAG signal. Thus, we computed the following parameters (Table 2) over the extracted 169 epochs: (i) averaged rectified values (ARV); (ii) mean power frequency (MPF), (iii) variance 170 171 of means squared (VoMS), (iv) form factor (FF), (v-vi) the % of determinism and recurrence 172 (%DET and %REC). Recurrence quantification analysis (RQA) was applied using the zscored data (Nalband et al., 2016). The %REC parameter is the percentage of recurring points 173 in the recurrence matrix below the tolerance threshold (see below). The %DET parameter is 174 the percentage of recurrence points forming diagonal lines in the recurrence plot of at least 175 length 2. %REC and % DET increases as the signal becomes more regular (Liu et al., 2004). 176 The embedding dimension, delay and tolerance values were as defined in (Andersen et al., 177

2018) to allow for easier comparison. Using the nearest neighbour approach (Kennel et al., 178 1992) the embedding dimension was set to 5. Using an approach based on the drop of auto 179 correlation function below 0.2, the delay parameter was set to 19 ms. Using a %REC 180 minimization optimization method tolerance was set to 0.2839. 181 2.6. Statistical analysis 182 Statistical analysis carried out using SPSS version 23 (IBM Corp., Armonk, NY, USA). A 183 184 linear mixed model with group (knee OA patients and asymptomatic participants), ADL (sit to stand, stairs descent and stairs ascent) and *location* (1-8) as within subject factors for each 185 186 of the parameters. All interactions between factors were included in the model. To allow for residuals with unequal variance, a repeated factor associated with patient type, ADL type and 187 location was added to the model. When a significant effect was observed, a Bonferroni 188 adjustment was performed for a pairwise comparison. Data are presented in the results 189 190 section as mean (SE). P values < 0.05 were considered significant.

191

192 **3. Results**

193 *3.1. Participant characteristics*

194 The demographic data showed that the asymptomatic participants were older than the knee

195 OA patients but similar in terms of gender distribution, body height and body mass (Table 1).

196 Table 3 shows the overall results of the statistical analysis.

197 *3.2. Differences between knee osteoarthritis patients and asymptomatic participants*

198 *Group* played a significant role for ARV and %REC. Higher ARV (0.535 (0.033) mm*s⁻² vs.

199 $0.399 (0.033) \text{ mm}^{*}\text{s}^{-2}$, P = 0.006) and lower %REC (0.120 (0.048) % vs. 0.345 (0.049) %, P

= 0.001) were found for knee OA patients compared with asymptomatic participants (Fig. 2i and 2v).

202 *3.3. Differences among activities of daily living*

203 ADL played a significant role for all parameters except %REC (Fig. 2i-iv and 2vi). ARV

- were lowest during sit to stand $(0.187 (0.027) \text{ mm}^{*}\text{s}^{-2})$, intermediate during stairs ascent
- 205 $(0.502 (0.024) \text{ mm}^{*}\text{s}^{-2})$ and highest during stairs descent $(0.703 (0.026) \text{ mm}^{*}\text{s}^{-2})$, P < 0.001).
- VoMS were smaller during sit to stand $(0.270 (0.035) \text{ mm}^{4*}\text{s}^{-8})$ than both stairs ascent (0.907)
- 207 (0.052) mm^{4*}s⁻⁸) and stairs descent (1.070 (0.050) mm^{4*}s⁻⁸, P < 0.001). MPF were higher
- during sit to stand (123.7 (3.7) Hz) than both stairs ascent (65.4 (3.1) Hz) and stairs descent

209 (61.8 (3.2) Hz, P < 0.001). VoMS was also lower during stairs ascent than during stairs

descent (P < 0.032). FF were lowest during sit to stand (2.114 (0.066) a.u.) than both stairs

- 211 ascent (3.367 (0.081) a.u.) and stairs descent (3.500 (0.084) a.u., P < 0.001). %DET were
- lowest during sit to stand (24.750 (2.271) %), intermediate during stairs ascent (42.787
- 213 (2.053) %) and highest during stairs descent (48.804 (2.026) %, P < 0.001).
- 214 *3.4. Differences among location*
- 215 Location played a significant role for all parameters; see Table 4 for the result of the pair
- 216 wise comparisons (Fig. 2i-vi). Lower ARV were recorded on the patella and the tibial
- tuberosity (P \leq 0.05). VoMS and FF were lower on the patella and higher on the medial
- 218 condyle (P < 0.05). Higher MPF were found on the patella and on the tibial tuberosity (P <
- 219 0.05). Higher %REC were recorded on the patella and the medial condyle (P < 0.05). Finally,
- lower %DET were found on the patella (P < 0.05).
- 3.5. Interactions between group, ADL and location
- 222 There were significant $Group \times ADL$ interactions for %DET. The %DET was lowest during
- sit to stand than during stairs ascent and stairs descent as well as lower during stairs ascent
- than stairs descent for and asymptomatic participants (P < 0.001). The %DET was lower
- during sit to stand than during stairs descent and lower during stairs ascent than stairs descent
- for knee OA patients (P < 0.001). There were also significant $ADL \times Location$ interactions
- 227 for ARV, VoMS, FF and %REC. The pair wise comparisons showed that ARV were lower

during sit to stand than both stairs ascent and descent and lower during stairs ascent than stairs descent for *location* 1-8 (P < 0.001). The pairwise comparisons showed that VoMS and FF were lower during sit to stand than both stairs ascent and descent for *location* 1-8 (P < 0.014 and P < 0.001, respectively). The pair wise comparisons showed that %REC were lower during sit to stand than both stairs ascent and descent for *location* 2 (P < 0.001) and during sit to stand than stairs descent for *location* 3 (P < 0.05).

234

235 **4. Discussion**

Spatial dependencies depicted by multichannel VAG recordings from knee OA patients and
asymptomatic participants were investigated for the first time during ADL. Partly in line with
our first hypothesis, higher VAG amplitude and lower VAG regularity characterised knee OA
patients compared with asymptomatic participants. The present study also confirmed as
hypothesised that wireless multichannel VAG recordings can differentiate between ADL
types and depict non-uniform spatial distribution of knee joint VAG.

4.1. Differences in vibroarthrography between knee osteoarthritis patients and asymptomatic
participants

The recordings of VAG provide clinically relevant information related to biomechanical and friction features reflecting the condition of the joint (Shieh et al., 2016; Stoltze et al., 2017). In this study, we computed a series of parameters representing signal amplitude, frequency contents, absolute and relative variability as well as VAG regularity. The ARV of the VAGs

248 were higher in knee OA patients compared with asymptomatic participants in line with

- 249 previous studies (Baczkowicz et al., 2017; Baczkowicz et al., 2019; Baczkowicz and
- 250 Majorczyk, 2016; Tanaka and Hoshiyama, 2012). Contrary to these studies and to our
- 251 hypothesis, the MPF, VoMS and FF of the VAGs did not differ in this population of knee OA
- 252 patients compared with asymptomatic participants. Differences in the studied populations and

the VAG processing mostly explain these differences. The %REC of the VAGs were lower in 253 knee OA patients compared with asymptomatic participants underlining that the VAGs were 254 less regular (Liu et al., 2004). Such increases in amplitude and decreases in regularity mostly 255 underlined differences in the internal pressure distribution on the cartilage and in synovial 256 fluid in knee OA (Neu et al., 2008) as well as altered muscle activation (Hortobagyi et al., 257 2005). This is also corroborated by previous VAG studies reporting articular surface with 258 259 chondral lesions and higher friction in knee OA (Baczkowicz et al., 2019; Baczkowicz and Majorczyk, 2016; Stoltze et al., 2017; Wu et al., 2016). Increased roughness of cartilage has 260 261 been shown to alter arthrokinematic motion (Lorenz et al., 2013). The parameters assessing the amplitude and regularity of the VAG signals characterize the biomechanical aspects of 262 movement pattern, e.g., joint loading. The current study also suggest that these parameters are 263 likely to be associated with joint degenerations in OA patients, confirming the importance of 264 using linear and nonlinear analytic methods in VAG studies (Andersen et al., 2018). 265

4.2. Activities of daily living and vibroarthrography

We chose to study sit to stand, stairs descent and ascent, which are considered normal ADL, 267 as well as functions recommended to examine patients with knee OA (Dobson et al., 2013). 268 Many VAG studies have studied open kinetic chain movements most likely to avoid artefacts 269 during to e.g., heel strike (Andersen et al., 2018). On the other hand, these movements do not 270 reflect the biomechanical load applied to the patellofemoral joint occurring during closed 271 kinetic chain movements characterising ADL (Baczkowicz et al., 2019). A cadaveric model 272 study has shown that the contact stress applied to the patellofemoral joint can be up to 16 273 times higher during squat compared with open chain kinetic movement (Cohen et al., 2001). 274 A few studies have investigated sit to stand (Baczkowicz et al., 2019; Shark et al., 2011; 275 Tanaka and Hoshiyama, 2012; Wiens et al., 2016). Baczkowicz et al. (2019) have suggested 276 that the high contact stress would occur along increased kinetic friction and result in higher 277

amplitude, variability and frequency contents of the VAG signal. A biomechanical study has 278 reported correlations between the amplitude of the VAG signal and the estimated relative 279 total knee compressive force (Stoltze et al., 2017). To the best of our knowledge, no studies 280 have investigated VAG during stairs descent or ascent. Stairs descent is usually studied in 281 relation to patellofemoral pain due to increased compressive force applied to the joint 282 (Rathleff et al., 2013). All the computed parameters beside %REC differentiate between the 283 284 three types of ADL. The ARV, VoMS, FF and %DET were lowest during sit to stand compared with stairs ascent and descent. Further, ARV and %DET were higher during stairs 285 286 descent compared with stairs ascent mostly due to increased compressive forces during ADL. The results related to amplitude and variability of the VAG signal during closed chain kinetic 287 movement were in line with Baczkowicz et al. (2019) but differed for frequency contents. 288 Here too, differences in signal processing (epoch length, frequency computation) and 289 290 movement artefacts mostly explain this discrepancy. Overall, the current findings confirmed that wireless VAG recordings can be used to study ADL offering important perspectives for 291 future clinical studies targeting knee OA in ecological environment. 292

293 4.3. Spatial dependencies in vibroarthrography

A novel aspect of the current study relates to its ability in revealing non-uniformity of the 294 VAG spatial distribution during the three studied ADL as well as differences in VAG spatial 295 distribution among knee OA patients and asymptomatic participants. The accelerometer 296 297 location influenced the computed parameters, all showing different patterns of uneven acceleration dampening in agreement with Andersen et al. (2018). When comparing 298 locations, lower VAG amplitudes were found on the patella and the tibial tuberosity. In 299 parallel, lower and higher absolute and relative variability were seen on the patella and the 300 medial condyle, respectively. Finally, higher frequency contents were found on the patella 301 and on the tibial tuberosity while more (%REC) and less (%DET) regular VAG signals were 302

recorded on the patella underlining that underlying knee structures affect the VAG signals. 303 Differences in the VAG maps between lateral and medial side of the knee are likely to be 304 related to the distribution of internal forces during ADL (Stoltze et al., 2017). Confirming 305 recent findings, the computed parameters revealed unique features of the VAG signals 306 underlining the importance of reporting these distinct parameters. Multi-channel VAG 307 recordings open new possibilities enabling to identify the unique signature of a pathological 308 309 knee as well as to assess the effect of interventions based on, e.g., strength training or knee braces. 310

311 4.4. Limitations

A main limitation of the present study is its cross-sectional design which confined us to study 312 group of 20 participants' differences. This type of design does not allow inferring whether the 313 reported changes in VAGs are a source or a cause of knee OA. However, changes in VAGs 314 appear with ageing, supporting the influence of degenerative processes (Baczkowicz et al., 315 2015). The studied ADL were conducted at self-chosen paces during sit to stand and stairs 316 descent or ascent (Malling and Jensen, 2016; Rathleff et al., 2013). The VAGs were processed 317 across time due to the lack of temporal information beside the onset and offset of movement. 318 Future VAG studies applying segmentation to movement phases and studying the effects of 319 movement artefacts on VAGs are therefore warranted. 320

321

322 **5. Conclusions**

This study revealed spatial dependencies of VAG topographical features in knee OA patients and asymptomatic participants during ADL. Multichannel VAG recordings enabled to differentiate between knee OA patients and asymptomatic participants in terms of VAG amplitude and regularity. The present study also demonstrated the feasibility of wireless

327 multichannel VAG recordings for assessing different ADL types offering new perspectives

328 for ecological biomechanical assessments of the knee joint.

329

330 Acknowledgments

- 331 This study was part of the individualized osteoarthritis interventions (IOI) project and has
- received funding from Innovation Fund Denmark (grant nr. 40-2014-3). The authors are
- 333 grateful to CCBR (Aalborg, Denmark) for the recruitment of participants.

334

335 Declaration of competing interest

- 336 All authors declare no conflict of interest.
- 337

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	Knee Osteoarthritis	Asymptomatic	
Variables	Patients	Participants	
v ar fables	(n=20)	(n=20)	
Age (years)	66.2 (5.2)	70.3 (5.9)*	
Sex (female/male)	9/11	10/10	
Body height (cm)	169.8 (9.0)	169.0 (10.8)	
Body mass (kg)	77.7 (9.9)	81.5 (13.0)	
Body mass index (kg/cm ²)	28.1 (2.7)	27.2 (3.2)	
Kellgren Lawrence score (left/right)	2.6 (0.9)/2.6 (0.5)	NA/NA	
Pain intensity after sit to stand (VAS)	4.8 (2.7)	0 (0)‡	
Pain intensity after stairs descent and ascent	5 5 (2 2)	0 (0)‡	
(VAS)	5.5 (2.3)		

471 **Table 1**. Baseline demographic and clinical characteristics of participants

472 Values are presented as mean (SD). NA: Not available. VAS: Visual analogue scale

473 * P<0.05. ‡ P<0.001

Table 2. List of the extracted vibroarthrographic variables

Variables	Interpretation		
Averaged rectified values (ARV)	Signal amplitude		
Mean power frequency (MPF)	Frequency contents of the signal		
Variance of means squared (VoMS)	Absolute reliability		
Form factor (FF)	Relative reliability		
% of determinism and recurrence (%DET	Changes in periodicity of the time		
and %REC)	series		

Table 3. Results of the statistical analysis on averaged rectified value (ARV), variance of means squared (VoMS), form factor (FF), mean power
frequency (MPF), % of Recurrence (%REC) and % of Determinism (%DET) of the vibroarthrographic signals with group (knee osteoarthritis
patients-asymptomatic participants), activity of daily living (sit to-stand, stairs descent and ascent) and accelerometer location (1-8) as within
factors of the linear mixed model.

	ARV	VoMS	FF	MPF	%REC	%DET
Group	F _{1,40.828} =8.502,	$F_{1,105.292}=1.065,$	F _{1,50.502} =0.309,	F _{1,39.970} =0.862,	F _{1,181.925} =10.632,	F _{1,38.732} =0.075,
	P=0.006	P=0.304	P=0.581	P=0.359	P=0.001	P=0.786
ADL	F _{2,260.230} =313.495,	$F_{2,460.450}$ =160.790,	$F_{2,453.152}=272.893,$	F _{2,412.177} =232.835,	$F_{2,126.192}=0.200,$	F _{2,365.519} =135.774,
	P<0.001	P<0.001	P<0.001	P<0.001	P=0.819	P<0.001
Location	F _{7,127.592} =17.175,	F _{7,141.649} =11.975,	F _{7,147.421} =45.795,	F _{7,178.344} =38.945,	F _{7,156.097} =2.920,	F _{7,145.089} =11.571,
	P<0.001	P<0.001	P<0.001	P<0.001	P=0.007	P<0.001
Group ×	F _{2,260.230} =1.012,	$F_{2,460.450}=2.080,$	$F_{2,453.152}=0.250,$	F _{2, 412.177} =2.697,	$F_{2,126.192}=0.030,$	F _{2,365.519} =5.085,
ADL	P=0.365	P=0.126	P=0.779	P=0.069	P=0.970	P=0.007
Group ×	F _{7,127.592} =1.531,	F _{7,141.649} =0.500,	F _{7,147.421} =1.692,	F _{7,178.344} =1.914,	F _{7,156.097} =2.514,	F _{7,145.089} =0.781,
Location	P=0.162	P=0.833	P=0.115	P=0.070	P=0.018	P=0.604
ADL ×	F _{14,96.408} =2.754,	F _{14,139.014} =2.201,	F _{14,148.128} =3.528,	F _{14,107.429} =1.405,	F _{14,111.443} =2.591,	F _{14,108.071} =0.445,
Location	P=0.002	P=0.010	P<0.001	P=0.163	P=0.003	P=0.956
Group ×	F _{14,96.408} =0.221,	F _{14,139.014} =0.420,	F _{14, 148.128} =0.742,	F _{14,107.429} =0.557,	F _{14,111.443} =0.567,	F _{14,108.071} =0.219,
$ADL \times$	P=0.999	P=0.966	P=0.729	P=0.892	P=0.886	P=0.999
Location						

Table 4. Results of the pairwise comparison for average rectified values (ARV), variance of means squared (VoMS), form factor (FF), mean power frequency (MPF), and % of Determinism (%DET) for locations ($P \le 0.05$). In each cell, the mentioned parameters corresponding to the location indicated along the rows was compared with the remaining locations.

	Location 1	Location 2	Location 3	Location 4	Location 5	Location 6	Location 7	Location 8
T		ADV				ADV		
Location 1		ARV<	ARV<	ARV<	ARV<	ARV<	ARV<	ARV< VoMS<
		VoMS< FF<	VoMS< FF<	VoMS< FF<	FF<	VoMS< FF<	VoMS< FF<	FF<
	-	MPF>	MPF>	MPF>	MPF>	MPF>	MPF>	MPF>
		%DET<	%DET<	%DET<	IVII I >	%DET<	%DET<	IVII I >
Location 2		70221	,0221	,0221	ARV>	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	,0221	ARV>
			VoMS<			VoMS<		
	-	-	FF<		FF>	FF<	NS	FF>
			MPF>	MPF>		MPF>		
Location 3					ARV>			
					VoMS>			VoMS>
	-	-	-	NS	FF>	NS	NS	FF>
					MPF<			MPF<
					%DET>			%DET>
Location 4					ARV>			ARV>
					VoMS>			
	-	-	-	-	FF>	NS	NS	FF>
					MPF<			MPF<
Location 5						ARV<	ARV<	
						VoMS<	VoMS<	
	-	-	-	-	-	FF<	FF<	NS
						MPF>	MPF>	
						%DET<	%DET<	
Location 6								ARV>
								VoMS>
	-	-	-	-	-	-	MDE	FF>
							MPF<	MPF< %DET>
Location 7								ARV>
Location /								
	-	-	-	-	-	-	-	FF>
								MPF<
								%DET>
Location 8								
	-	-	-	-	-	-	-	-
· · · · · · · · · · · · · · · · · · ·								

NS: Non significant.

Figure legends:

Fig. 1: Accelerometer locations. Accelerometers were placed over the quadriceps tendon (1), the lateral side of the knee approx. 1-2 cm medial from the epicondyle of femur towards the patella (2), lateral proximal on the patella (3), medial proximal on the patella (4), lateral distal on the patella (5), medial distal on patella (6), the medial side of the knee approx. 1-2 cm medial from the epicondyle of femur towards the patella (7), and the tibial tuberosity (8). Example of the micro machined accelerometer mounted on a printed circuit board used to record the vibroarthrographic signals.

Fig. 2: Average maps of the (i) averaged rectified values (ARV, mm*s⁻²), (ii) mean power frequency (MPF, Hz), (iii) variance of means squared (VoMS, mm⁻⁴*s⁻⁸), (iv) form factor (FF, a.u.), (v) % of recurrence (%REC), and (vi) % of determinism (%DET) of the vibroathrographic signals recorded using eight accelerometers (black dots) during activities of daily living (sit to stand, stairs descent and stairs ascent) among patients with knee osteoarthritis (n=20) and asymptomatic participants (n=20). See Fig. 1 for accelerometer nomenclature.

Figure 1



