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33 **Abstract**

34 Gait analysis has widely been accepted as an objective measure of function and clinical outcome.  
35 Ambulatory accelerometer-based gait analysis has emerged as a clinically more feasible  
36 alternative to optical motion capture systems but does not provide kinematic characterization to  
37 identify disease dependent mechanisms causing walking disability. This study investigated the  
38 potential of a single inertial sensor to derive frontal plane motion of the pelvis (i.e. pelvic  
39 obliquity) and help identify hip osteoarthritis (OA) related gait alterations. Patients with  
40 advanced unilateral hip OA (n=20) were compared to patients with advanced unilateral knee OA  
41 (n=20) and to a healthy control group (n=20). Kinematic characterization of frontal plane pelvic  
42 motion during gait demonstrated decreased range of motion and increased asymmetry for hip OA  
43 patients specifically.

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46 **Keywords:** ambulatory gait analysis, inertial sensor, osteoarthritis, outcome assessment, frontal  
47 plane pelvic motion, pelvic obliquity, Trendelenburg, performance-based test.

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65 **Introduction**

66 Gait analysis has widely been accepted as an objective measure of function, allowing researchers  
67 and clinicians to better understand biomechanical alterations in the presence of hip osteoarthritis  
68 (OA) and to quantitatively evaluate the functional success of total hip arthroplasty (THA) and  
69 rehabilitation strategies [1-3]. Besides pain relief, functional improvement following surgery has  
70 become more important for the new generation of younger and generally more active hip OA  
71 patients. Therefore, it has been advocated to supplement longitudinal follow-up studies with  
72 objective assessment of function like gait analysis [2, 4]. In clinical gait analysis, a skin marker  
73 based optical motion capture (MOCAP) system provides a non-invasive approach and is  
74 regarded as the gold standard. Unfortunately, a MOCAP system is not feasible for routine use  
75 because it is time consuming, expensive, artificial and limited to a single gait cycle. Advances in  
76 miniaturization and cost of ambulatory motion sensors have emerged accelerometer-based gait  
77 analysis as a potential ambulatory alternative to MOCAP systems [5]. In previous studies, a  
78 single accelerometer positioned at the dorsal side of the pelvis has been advocated for optimal  
79 clinical feasibility to derive spatiotemporal gait parameters (e.g. cadence, step length) based on  
80 heel strike (HS) events in the antero-posterior acceleration signal [6, 7]. These spatiotemporal  
81 gait parameters can discriminate gait between healthy subjects and OA patients [2-4, 8] and have  
82 demonstrated responsiveness to changes postoperatively [2, 3]. However, spatiotemporal gait  
83 parameters lack kinematic characterization to identify the mechanisms causing typical gait  
84 disturbances in hip OA patients such as Trendelenburg's gait. To supplement ambulatory  
85 spatiotemporal gait analysis with kinematic characterization outside the MOCAP laboratory, the  
86 use of a gyroscope in conjunction with an accelerometer (i.e. inertial sensor) has been advocated  
87 [9]. With an inertial sensor, spatiotemporal gait parameters and dynamic orientation angles of  
88 underlying body segments can be determined.

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90 The primary aim of the study was to investigate the potential of a single inertial sensor  
91 positioned at the dorsal side of the pelvis to derive clinically relevant frontal plane gait  
92 kinematics in patients with hip OA, supplementary to spatiotemporal gait parameters. We  
93 hypothesized that motion of the pelvis in the frontal plane (i.e. pelvic obliquity) could accurately  
94 be characterized from a single inertial sensor positioned at the dorsal side of the pelvis [10], and

95 that it would be decreased in patients with hip OA [11]. A second aim was to investigate whether  
96 gait kinematics of pelvic obliquity are influenced by a main effect of osteoarthritis on gait or  
97 related to hip OA specifically. Therefore, hip OA patients were compared to knee OA patients  
98 and it was hypothesized that pelvic obliquity would be decreased in hip OA patients only, as a  
99 result of compensating for abductor muscle weakness and pain of the affected hip joint during  
100 the single limb supportive gait phase [1, 12]. A third aim was to investigate gait kinematics of  
101 pelvic obliquity in a healthy cohort to provide reference data and to investigate the influence of  
102 demographic variability.

103

## 104 **Materials and Methods**

105 Gait was studied in 20 patients with unilateral end stage hip OA and 20 patients with unilateral  
106 end stage knee OA (table 1). These patients were randomly recruited from the outpatient clinic if  
107 they were listed for a total joint replacement by an orthopaedic surgeon. All patients reported  
108 activity limitation because of OA and scored 3 or 4 on the Kellgren-Lawrence radiographic  
109 osteoarthritis index [13]. Patients with polyarthritis or any other condition affecting gait, except  
110 single joint osteoarthritis, were excluded from this study. Furthermore, gait was studied in 80  
111 healthy participants (age range 19-77yrs; mean 40.0yrs  $\pm$ 16.0; m/f=39/41) who had no joint pain  
112 and no medical history of lower extremity joint surgery. A control group of 20 healthy subjects  
113 was selected from this healthy cohort by age and gender for comparison with the osteoarthritis  
114 patient groups. However, a significantly higher body mass index (BMI) for knee OA patients  
115 was found compared to this control group (table 1).

116

### 117 Data acquisition

118 The study methods were in accordance with a previously published study [4]. Briefly, all  
119 participants were invited to walk 20 meters along a straight flat corridor at their own preferred  
120 speed. A 3D inertial sensor (41x63x24mm; 39g; Microstrain Inertia Link) was used. The sensor  
121 was positioned at the dorsal side of the pelvis, centrally between both posterior superior iliac  
122 spines. At this position, a single inertial sensor allows heel strike detection from the antero-  
123 posterior acceleration signal [6, 7] and kinematic characterization of pelvic motion [10]. Using  
124 automated algorithms in Matlab, spatiotemporal gait parameters were derived: 1) speed (m/s); 2)  
125 cadence (steps/min); 3) step time (s); 4) step length (m); 5) step time irregularity (coefficient of

126 variance:  $100\% * SD/mean$ ) and 6) step time asymmetry ( $100\% * \text{abs}(\text{left-right})/((\text{left+right})/2)$ )  
127 [2, 4, 14]. Dynamic orientation angles of the pelvis were obtained through the inertial sensor's  
128 inbuilt fusion algorithms of acceleration, angular rate and magnetic field vector measurements  
129 and compared to gold standard MOCAP system measures. The waveform of pelvic obliquity  
130 during gait was further characterized to allow assessment of asymmetry. Kinematic gait  
131 parameters of pelvic obliquity included: a) range of motion (ROM, °); b) asymmetry ( $100\% * \text{abs}(\text{left-right})/\text{mean}$ ) and c) pelvic obliquity at heel strike (POHS;  $100\% * (\delta / \text{ROM})$ ) in which  $\delta$   
132 represents the ROM of pelvic obliquity between consecutive heel strikes (figure 1). The pelvic  
133 obliquity at heel strike indicates the orientation of the pelvis in frontal plane for which a value of  
134 50% represents a horizontal pelvic position. Capturing asymmetry of pelvic obliquity and the  
135 pelvic obliquity at heel strike from a single inertial sensor is a novel approach with no previous  
136 results reported in literature.  
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### 139 Statistical analysis

140 Data were analyzed using SPSS version 17.0. To compare mean values of gait parameters  
141 between patients with hip OA, patients with knee OA and the matched control group, a one-way  
142 ANOVA test was used. The interactions between gait parameters and the demographic  
143 covariates gender, age, height, weight and BMI in these three groups were calculated with  
144 MANCOVA which provides the level of statistical significance for the interaction (p-value) and  
145 the proportion of variance accounted for by the interaction (partial  $\text{Eta}^2$ ). Because previous  
146 studies have found that variance of walking speed can significantly influence kinematic gait  
147 parameters, especially in patients with osteoarthritis [15, 16], we performed analysis of  
148 covariance to compare speed-adjusted mean values of the range of motion of pelvic obliquity. In  
149 our cohort of 80 healthy participants, the association between the demographic variables gender,  
150 age, height, weight and BMI with individual gait parameters were measured using multiple linear  
151 regression analysis which provides the level of statistical significance (p-value) and the strength  
152 of the association (beta standardized coefficient).

153

### 154 **Results**

155 Characterization of pelvic obliquity during gait by a single inertial sensor attached at the dorsal

156 side of the pelvis provided waveforms that were qualitatively and quantitatively comparable to  
157 MOCAP measures (figure 2).

158 In patients with hip OA and knee OA, the accelerometer based spatiotemporal gait parameters  
159 demonstrated significant differences compared to the control group (table 2). Patients with hip  
160 OA demonstrated significantly decreased step length and walking speed compared to the control  
161 group. Step time irregularity and step time asymmetry were not significantly different between  
162 patients with hip OA and the control group. Patients with knee OA demonstrated significantly  
163 decreased walking speed, decreased cadence, increased step time irregularity and increased step  
164 time asymmetry. Comparing spatiotemporal gait parameters between patients with hip and knee  
165 OA demonstrated only a significantly higher step time asymmetry for knee OA patients.

166 Kinematic gait parameters of pelvic obliquity demonstrated lower range of motion (ROM) of  
167 pelvic obliquity in both hip OA and knee OA patients compared to the control group (table 2).  
168 After statistical correction for the variance of walking speed between groups, the range of motion  
169 of pelvic obliquity at a walking speed of 1.13m/s demonstrated significantly lower outcomes for  
170 hip OA patients compared to controls (ROM pelvic obliquity:  $5.6^{\circ} \pm 2.1$  vs.  $8.0^{\circ} \pm 2.4$ ;  $p < 0.01$ ;  
171 respectively) and compared to knee OA patients (ROM pelvic obliquity:  $5.6^{\circ} \pm 2.1$  vs.  $7.3^{\circ} \pm 2.3$ ;  
172  $p < 0.01$ ; respectively) whereas no significant difference was observed between knee OA patients  
173 and controls anymore. Furthermore, significantly higher asymmetry of pelvic obliquity was  
174 found for hip OA patients compared to controls and compared to knee OA patients ( $32.2\% \pm 25.6$   
175 vs.  $15.9\% \pm 13.1$  and vs.  $16.1\% \pm 12.4$ ;  $p < 0.05$  respectively). Knee OA patients demonstrated  
176 significantly higher pelvic obliquity at heel strike (POHS) compared to controls ( $73.6\% \pm 22.4$  vs.  
177  $50.2\% \pm 15.4$  respectively) however no significant difference was observed compared to hip OA  
178 patients. Analysis of demographic variability (MANCOVA) in the groups of hip OA patients,  
179 knee OA patients and healthy controls demonstrated only a significant interaction between BMI  
180 and POHS ( $\text{Eta}^2 = 0.08$ ;  $p < 0.05$ ).

181 Results of gait parameters in all healthy subjects and results of multiple linear regression analysis  
182 between gait parameters and demographic variables are shown in table 3. Multiple analysis of  
183 covariance for the demographic variables age, gender, height, weight, BMI with gait parameters  
184 demonstrated a significant negative effect of age on the range of motion of pelvic obliquity (beta  
185 standardized coefficient= -0.33).

186 **Discussion**

187 This study investigated the potential of a single inertial sensor positioned at the dorsal side of the  
188 pelvis to characterize frontal plane pelvic motion (i.e. pelvic obliquity) during gait,  
189 supplementary to spatiotemporal gait parameters, and describes its clinical relevance for patients  
190 with hip OA. First, measures of pelvic obliquity assessed with an inertial sensor were compared  
191 to a MOCAP system. In a previous study [17], assessment of pelvic kinematics during gait by a  
192 MOCAP system with reflective markers attached onto a rigid plate at the dorsal side of the pelvis  
193 demonstrated good accuracy compared to single markers placed over the anterior and posterior  
194 superior iliac spines (ASIS and PSIS). A more recent study by Borhani et al. [18] demonstrated  
195 that reflective markers on a rigid plate provide more accurate results with less skin artefacts,  
196 especially in overweight and obese patients. In this study, a single inertial sensor was placed at  
197 the dorsal side of the pelvis and qualitatively and quantitatively comparable waveforms for  
198 pelvic obliquity during gait were found between inertial sensor based measures and MOCAP  
199 system based measures with reflective markers placed over the anterior and posterior superior  
200 iliac spines. These findings are in accordance to results of previous validation studies  
201 demonstrating good accuracy and reliability for the assessment of trunk motion measured by  
202 inertial sensors and MOCAP systems [9, 10, 19-21].

203  
204 In patients with unilateral end stage hip OA, measures of pelvic obliquity during gait  
205 demonstrated less ROM and higher asymmetry compared to healthy controls. To investigate  
206 whether these gait alterations are due to a main effect of osteoarthritis on gait, or related to hip  
207 OA specifically, gait was also compared to patients with unilateral end stage knee OA. In order  
208 to allow a meaningful comparison of gait parameters between hip OA patients, knee OA patients  
209 and healthy controls, standardization of walking speed was aimed for as a significant influence  
210 of walking speed on kinematic gait parameters has been reported in literature [1, 15]. To avoid  
211 artificial measures of gait with a treadmill, we instructed subjects to walk freely at preferred  
212 speed in a hospital's corridor and a statistical correction for the variation of walking speed  
213 between groups was applied with ANCOVA [16]. After this statistical correction, hip OA  
214 patients demonstrated even lower ROM of pelvic obliquity and twice the amount of pelvic  
215 obliquity asymmetry compared to healthy controls and to patients with knee OA. These findings  
216 could suggest that alterations in pelvic obliquity during gait are not due to a main effect of



217 osteoarthritis on gait, but related to hip OA specifically. Moreover, knee OA patients  
218 demonstrated no significant difference for ROM of pelvic obliquity compared to healthy controls  
219 after correcting for the differences in walking speed. However, significantly higher pelvic  
220 obliquity at heel strike (POHS) was found for knee OA patients compared to healthy controls  
221 whereas no significant difference was observed for POHS between knee OA and hip OA  
222 patients. Interpretation of these findings are made with caution as they can be confounded by the  
223 significantly higher BMI in knee OA patients compared to healthy controls and BMI  
224 demonstrating a significant interaction with POHS.

225 Alterations of frontal plane pelvic motion during gait have been related to hip OA causing pain,  
226 limited range of motion of the hip joint and decreased muscle strength of the hip abductor  
227 muscles [22, 23], often resulting in a limp or gait asymmetry by compensatory mechanisms of  
228 the trunk [24]. During single-limb support in the stance phase of gait, pelvic equilibrium in the  
229 frontal plane is ensured by the hip abductor muscles which help maintaining balance of the trunk  
230 [25]. In patients with hip OA, weakness of the hip abductor muscles can result in two distinct  
231 walking patterns. In “Trendelenburg gait” [26], a pelvic drop on the non-supportive swing limb  
232 with increased hip adduction on the stance limb is found (i.e. Trendelenburg’s sign) [12, 27],  
233 moving the compressive force laterally to the acetabulum [28]. This pelvic drop is frequently  
234 compensated by increased lateral trunk lean, shifting the body’s center of mass towards the  
235 stance limb and shortening the moment arm of the hip abductor muscles, resulting in a typical  
236 “Duchenne gait” [29] or “abductor lurch” [28] with the pelvis level or elevated on the non-  
237 supportive swing limb. When pain arises in the hip joint during walking, there is also  
238 compensatory trunk lean towards the supporting side with significant hip joint load reduction  
239 achieved by a combined sideways shift of the pelvis [30]. These patterns of hip unloading  
240 mechanisms have also been observed in other hip conditions such as Legg Calvé Perthes Disease  
241 (LCPD) [31], congenital hip dislocation [32], and the relationship between severity of hip  
242 abductor muscle weakness and the amount of pelvic drop and compensatory lateral trunk lean  
243 has been demonstrated in patients with cerebral palsy (CP) [33]. A limitation of this study is that  
244 we only measured frontal plane angles at the level of the pelvis and did not obtain the  
245 contribution of compensatory lateral trunk lean from the upper trunk. The aim of the study was  
246 to obtain frontal plane gait kinematics from a single inertial sensor to supplement spatiotemporal  
247 gait parameters derived at the dorsal side of the pelvis for optimal clinical feasibility and

248 reliability. Measuring lateral trunk lean would necessitate the use of a second sensor and may be  
249 less feasible for routine clinical gait analysis.

250 Spatiotemporal gait parameters demonstrated significant differences for hip OA and knee OA  
251 patients compared to healthy controls. These findings are in accordance to previous studies  
252 comparing gait between healthy subjects and hip OA or knee OA patients prior to arthroplasty  
253 [2, 3, 8]. In patients with unilateral hip OA, step length and walking speed were significantly  
254 decreased compared to healthy controls. Patients with hip OA tend to walk with smaller steps,  
255 and because the step frequency (i.e. cadence) was not significantly different, it results in  
256 decreased walking speed. The disability to walk with larger steps may demonstrate a general  
257 effect of osteoarthritis on gait, as step length and walking speed are also reduced in knee OA  
258 patients, and these spatiotemporal gait parameters do not identify underlying mechanism related  
259 to hip OA specifically. Kinematic gait parameters on the other hand demonstrated significant  
260 lower ROM of pelvic obliquity and almost twice the amount of pelvic obliquity asymmetry  
261 compared to healthy controls after correcting for variance in walking speed, whereas knee OA  
262 patients demonstrated no significant difference for ROM of pelvic obliquity or pelvic obliquity  
263 asymmetry compared to healthy controls. These findings could suggest that alterations in pelvic  
264 obliquity during gait are not due to a main effect of osteoarthritis on gait, but related to hip OA  
265 specifically. Hence, additional assessment of pelvic obliquity during gait could be a clinically  
266 relevant measure of functional outcome following THA. For instance, the abductor-sparing  
267 anterior approach for THA has demonstrated a closer-to-normal ROM of pelvic obliquity during  
268 gait with significantly reduced pelvic obliquity ( $2^\circ$ ) at ipsilateral foot-off compared to patients  
269 with a lateral approach [34]. Furthermore, restoring offset during THA to match that of the  
270 normal contralateral side has been shown to improve abductor strength and to reduce the  
271 incidence of Trendelenburg's gait [35].

272 A third aim was to investigate gait kinematics of pelvic obliquity in a healthy cohort, to provide  
273 reference data and to investigate the influence of demographic variability. Spatiotemporal gait  
274 parameters of eighty healthy participants demonstrated similar results compared to previous  
275 reports [6, 14, 36]. The asymmetry of pelvic obliquity during gait and the pelvic obliquity  
276 measured at heel strike have not been reported previously in literature. According to our results,  
277 asymmetry in pelvic obliquity up to 15% can be regarded as normal and healthy participants

278 demonstrated a perfect horizontal orientation of the pelvis in the frontal plane at heel strike  
279 (POHS = 50.6%). In our healthy cohort, the range of motion of pelvic obliquity was significantly  
280 decreased by ageing, however the effect size was rather small (beta standardized coefficient -  
281 0.33). Still, these findings could hypothetically suggest that measurements of pelvic obliquity  
282 during gait capture decreased physiological functions caused by ageing such as muscle atrophy  
283 resulting in hip abductor weakness. In contrast to previous reports [36, 37], we did not find a  
284 significant correlation between walking speed and range of motion of pelvic obliquity in healthy  
285 subjects. Gard et al. [36] compared the range of motion of pelvic obliquity during gait, measured  
286 by a MOCAP system, in three healthy subjects (age 22-29) walking at eight different walking  
287 speeds between 1.0-2.4m/s at increments of 0.2m/s. Over the range of walking speeds, the range  
288 of motion of pelvic obliquity ranged from 5-20° with a linear increase with the walking speed for  
289 each individual. Furthermore, a study by Michaud et al. [37] investigated the range of motion of  
290 pelvic obliquity during gait in nine persons with transtibial or transfemoral amputation, and  
291 compared their results with results from the study cohort of Gard et al. A linear relationship was  
292 found for range of motion of pelvic obliquity with speed, demonstrating correlation coefficients  
293 all exceeding 0.70. We measured range of motion of pelvic obliquity in eighty healthy subjects  
294 while they walked at preferred speed only. We found inter-subject variability in the range of  
295 motion of pelvic obliquity during gait within a limited range of walking speeds. Because we did  
296 not measure different walking speeds, we cannot truly compare our results with the previous  
297 findings from Gard et al. and Michaud et al. Individual differences in the range of motion of  
298 pelvic obliquity may be multifactorial, but may change with a similar magnitude between  
299 subjects by increasing walking speed.

### 300 **Conclusion**

301 This study demonstrates that ambulatory gait analysis with a single inertial sensor positioned at  
302 the dorsal side of the pelvis allows both spatiotemporal and kinematic characterization of gait.  
303 Focusing on pelvic motion in the frontal plane (i.e. pelvic obliquity), patients with hip OA  
304 demonstrated significantly less range of motion and higher asymmetry compared to healthy  
305 controls and compared to patients with knee OA. Therefore, kinematic characterization of pelvic  
306 obliquity during gait seems to capture hip OA related disability. Pelvic obliquity seems a  
307 valuable biomechanical measure of gait that is independent of time, and could be used to

308 objectively assess functional disability in patients with hip OA and to monitor functional  
309 improvement after total hip arthroplasty.

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316

### 317 **References**

- 318 1. Zeni, J.A., Jr. and J.S. Higginson, *Differences in gait parameters between healthy subjects and*  
319 *persons with moderate and severe knee osteoarthritis: a result of altered walking speed?* Clin  
320 Biomech (Bristol, Avon), 2009. **24**(4): p. 372-8.
- 321 2. Senden, R., et al., *The importance to including objective functional outcomes in the clinical follow*  
322 *up of total knee arthroplasty patients.* Knee, 2011. **18**(5): p. 306-11.
- 323 3. Ornetti, P., et al., *Gait analysis as a quantifiable outcome measure in hip or knee osteoarthritis: a*  
324 *systematic review.* Joint Bone Spine, 2010. **77**(5): p. 421-5.
- 325 4. Bolink, S.A., et al., *Inertial sensor motion analysis of gait, sit-stand transfers and step-up*  
326 *transfers: differentiating knee patients from healthy controls.* Physiol Meas, 2012. **33**(11): p.  
327 1947-58.
- 328 5. Zijlstra, W. and A.L. Hof, *Assessment of spatio-temporal gait parameters from trunk*  
329 *accelerations during human walking.* Gait Posture, 2003. **18**(2): p. 1-10.
- 330 6. Senden, R., et al., *Acceleration-based gait test for healthy subjects: reliability and reference data.*  
331 *Gait Posture, 2009. 30(2): p. 192-6.*
- 332 7. Gonzalez, R.C., et al., *Real-time gait event detection for normal subjects from lower trunk*  
333 *accelerations.* Gait Posture, 2010. **31**(3): p. 322-5.
- 334 8. Constantinou, M., et al., *Spatial-temporal gait characteristics in individuals with hip*  
335 *osteoarthritis: a systematic literature review and meta-analysis.* J Orthop Sports Phys Ther,  
336 2014. **44**(4): p. 291-87.
- 337 9. Ishigaki, N., et al., *Analysis of pelvic movement in the elderly during walking using a posture*  
338 *monitoring system equipped with a triaxial accelerometer and a gyroscope.* J Biomech, 2011.  
339 **44**(9): p. 1788-92.
- 340 10. Zijlstra, A., et al., *A body-fixed-sensor based analysis of compensatory trunk movements during*  
341 *unconstrained walking.* Gait Posture, 2008. **27**(1): p. 164-7.
- 342 11. Lenaerts, G., et al., *Aberrant pelvis and hip kinematics impair hip loading before and after total*  
343 *hip replacement.* Gait Posture, 2009. **30**(3): p. 296-302.
- 344 12. Watelain, E., et al., *Pelvic and lower limb compensatory actions of subjects in an early stage of*  
345 *hip osteoarthritis.* Arch Phys Med Rehabil, 2001. **82**(12): p. 1705-11.
- 346 13. Kellgren, J.H. and J.S. Lawrence, *Radiological assessment of osteo-arthrosis.* Ann Rheum Dis,  
347 1957. **16**(4): p. 494-502.

- 348 14. Bautmans, I., et al., *Reliability and clinical correlates of 3D-accelerometry based gait analysis*  
349 *outcomes according to age and fall-risk*. Gait Posture, 2011. **33**(3): p. 366-72.
- 350 15. Bejek, Z., et al., *The influence of walking speed on gait parameters in healthy people and in*  
351 *patients with osteoarthritis*. Knee Surg Sports Traumatol Arthrosc, 2006. **14**(7): p. 612-22.
- 352 16. Astephen Wilson, J.L., *Challenges in dealing with walking speed in knee osteoarthritis gait*  
353 *analyses*. Clin Biomech (Bristol, Avon), 2012. **27**(3): p. 210-2.
- 354 17. Vogt, L., et al., *Cross-validation of marker configurations to measure pelvic kinematics in gait*.  
355 Gait Posture, 2003. **18**(3): p. 178-84.
- 356 18. Borhani, M., A.H. McGregor, and A.M. Bull, *An alternative technical marker set for the pelvis is*  
357 *more repeatable than the standard pelvic marker set*. Gait Posture, 2013. **38**(4): p. 1032-7.
- 358 19. Faber, G.S., et al., *Optimal inertial sensor location for ambulatory measurement of trunk*  
359 *inclination*. J Biomech, 2009. **42**(14): p. 2406-9.
- 360 20. Wong, W.Y. and M.S. Wong, *Trunk posture monitoring with inertial sensors*. Eur Spine J, 2008.  
361 **17**(5): p. 743-53.
- 362 21. Boonstra, M.C., et al., *The accuracy of measuring the kinematics of rising from a chair with*  
363 *accelerometers and gyroscopes*. J Biomech, 2006. **39**(2): p. 354-8.
- 364 22. Rasch, A., N. Dalen, and H.E. Berg, *Muscle strength, gait, and balance in 20 patients with hip*  
365 *osteoarthritis followed for 2 years after THA*. Acta Orthop, 2010. **81**(2): p. 183-8.
- 366 23. Arokoski, M.H., et al., *Hip muscle strength and muscle cross sectional area in men with and*  
367 *without hip osteoarthritis*. J Rheumatol, 2002. **29**(10): p. 2185-95.
- 368 24. Nankaku, M., et al., *Gait analysis of patients in early stages after total hip arthroplasty: effect of*  
369 *lateral trunk displacement on walking efficiency*. J Orthop Sci, 2007. **12**(6): p. 550-4.
- 370 25. Chamnongkitch, S., et al., *Difference in hip prosthesis femoral offset affects hip abductor strength*  
371 *and gait characteristics during obstacle crossing*. Orthop Clin North Am, 2012. **43**(5): p. e48-58.
- 372 26. Trendelenburg F. Dtsch Med Wochenschr 1895;21:21
- 373 27. Reininga, I.H., et al., *Compensatory trunk movements in patients with hip osteoarthritis:*  
374 *accuracy and reproducibility of a body-fixed sensor-based assessment*. Am J Phys Med Rehabil,  
375 2011. **90**(8): p. 681-7.
- 376 28. Amaro, A., et al., *Gluteus medius muscle atrophy is related to contralateral and ipsilateral hip*  
377 *joint osteoarthritis*. Int J Sports Med, 2007. **28**(12): p. 1035-9.
- 378 29. Duchenne, G.B. Physiologie der Bewegungen. Theodor Fischer, Cassel, Berlin; 1885.
- 379 30. Schroter, J., et al., *The 'Entlastungsgang'. A hip unloading gait as a new conservative therapy for*  
380 *hip pain in the adult*. Gait Posture, 1999. **9**(3): p. 151-7.
- 381 31. Westhoff, B., et al., *Computerized gait analysis in Legg Calve Perthes disease--analysis of the*  
382 *frontal plane*. Gait Posture, 2006. **24**(2): p. 196-202.
- 383 32. Stief, F., et al., *Effect of compensatory trunk movements on knee and hip joint loading during*  
384 *gait in children with different orthopedic pathologies*. Gait Posture, 2014. **39**(3): p. 859-64.
- 385 33. Krautwurst, B.K., et al., *The influence of hip abductor weakness on frontal plane motion of the*  
386 *trunk and pelvis in patients with cerebral palsy*. Res Dev Disabil, 2013. **34**(4): p. 1198-203.
- 387 34. Varin, D., M. Lamontagne, and P.E. Beaulé, *Does the anterior approach for THA provide closer-*  
388 *to-normal lower-limb motion?* J Arthroplasty, 2013. **28**(8): p. 1401-7.
- 389 35. Cassidy, K.A., et al., *Effect of femoral offset on pain and function after total hip arthroplasty*. J  
390 Arthroplasty, 2012. **27**(10): p. 1863-9.
- 391 36. Gard, S.A. and D.S. Childress, *The effect of pelvic list on the vertical displacement of the trunk*  
392 *during normal walking*. Gait & Posture, 1997. **5**(3): p. 233-238.
- 393 37. Michaud, S.B., S.A. Gard, and D.S. Childress, *A preliminary investigation of pelvic obliquity*  
394 *patterns during gait in persons with transtibial and transfemoral amputation*. J Rehabil Res Dev,  
395 2000. **37**(1): p. 1-10.

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397 **Legends**

398

399 **Figure 1:** Characterization of pelvic obliquity (PO) during gait demonstrating primary peaks,  
400 secondary components which occur at heel strike (HS), range of motion (ROM) and  $\delta$ .

401 **Figure 2:** waveforms of pelvic obliquity during gait in a healthy subject. Left figure shows a  
402 MOCAP system based measurement of one gait cycle. Right figure shows an inertial sensor  
403 based measurement of one gait cycle.

404 **Table I:** Group demographics. \*p<0.05 Knee OA vs. Control group.

405 **Table II:** Gait parameters of hip OA patients, knee OA patients and the control group. P-values  
406 correspond with level of significance compared to healthy controls.

407 **Table III:** Reference data for gait parameters in healthy subjects demonstrating mean values and  
408 standard deviations (SD), and beta standardized coefficients from multiple linear regression  
409 analysis between gait parameters and demographic variables. \*p<0.05

410 **Tables:**

	<b>Control group n=20</b> male:female = 9:11		<b>Hip OA n=20</b> male:female = 10:10		<b>Knee OA n=20</b> male:female = 9:11	
	Mean	SD	Mean	SD	Mean	SD
Age (years)	61.0	6.1	63.4	8.5	65.4	9.3
Height (cm)	173	8.4	172	9.7	167	9.1
Weight (kg)	77.2	12.7	81.1	17.8	84.2	18.6
BMI	25.8	3.0	27.2	4.9	30.2*	7.3

411 Table I

	<b>Control group</b> <b>n=20</b>		<b>Hip OA</b> <b>n=20</b>			<b>Knee OA</b> <b>n=20</b>			<b>Hip OA</b> <b>vs</b> <b>Knee OA</b>
	Mean	SD	Mean	SD	p-value	Mean	SD	p-value	p-value
<b>Gait parameters</b>									
Speed	1.30	0.15	1.10	0.19	<0.01	0.98	0.19	<0.001	n.s.
Cadence	114.8	8.0	109.7	8.4	n.s.	105.9	11.3	<0.05	n.s.
Step time	0.53	0.04	0.55	0.04	n.s.	0.57	0.06	<0.05	n.s.

Step length	0.68	0.07	0.61	0.09	<0.01	0.55	0.07	<0.001	n.s.
Step time irregularity (%)	0.04	0.03	0.04	0.03	n.s.	0.06	0.03	<0.05	n.s.
Step time asymmetry (%)	2.50	1.84	2.31	1.61	n.s.	5.05	2.30	<0.001	<0.001
RoM pelvic obliquity (°)	8.6	2.8	5.5	1.7	<0.001	6.7	1.8	<0.05	<0.05
RoM pelvic obliquity (°) corrected for speed	8.0	2.4	5.6	2.1	<0.01	7.3	2.3	n.s.	<0.01
PO asymmetry (%)	15.9	13.1	32.2	25.6	<0.05	16.1	12.4	n.s.	<0.05
POHS (%)	50.2	15.4	66.4	24.9	n.s.	73.6	22.4	<0.01	n.s.

412 Table II

	Healthy subjects n=80		Demographic variables				
<b>Gait parameters</b>	Mean	SD	Age	Gender	Length	Weight	BMI
Speed (m/s)	1.29	0.15	0.09	0.28	-0.61	0.74	-0.78
Cadence (steps/min)	113.65	8.34	0.09	-0.03	-0.68	0.23	0.28
Step time (s)	0.53	0.04	-0.09	0.03	0.67	-0.22	0.30
Step length (m)	0.68	0.06	0.07	0.27	0.30	0.90	0.87
Step time irregularity (cv)	4.62	2.90	-0.05	-0.14	0.85	-1.49	1.28
Step time asymmetry (%)	3.13	2.32	0.04	-0.13	0.56	-0.68	0.43
RoM pelvic obliquity (°)	10.1	3.2	-0.33*	0.15	0.47	-0.62	0.50
PO asymmetry (%)	14.9	12.6	0.00	-0.22	0.08	0.02	-0.08
POHS (%)	50.6	14.8	-0.05	0.17	-0.34	0.00	0.25

413 Table III

414