1	Kinematic signatures of learning that emerge in a real-world motor skill task	
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24 Abstract

25 The neurobehavioral mechanisms of human motor-control and learning evolved in free 26 behaving, real-life settings, yet to date is studied in simplified lab-based settings. We demonstrate 27 the feasibility of real-world neuroscience, using wearables for naturalistic full-body motion-28 tracking and mobile brain-imaging, to study motor-learning in billiards. We highlight the 29 similarities between motor-learning in-the-wild and classic toy-tasks in well-known features, such 30 as multiple learning rates, and the relationship between task-related variability and motor learning. 31 Studying in-the-wild learning enable looking at global observables of motor learning, as well as 32 relating learning to mechanisms deduced from reductionist models. The analysis of the velocity 33 profiles of all joints enabled in depth understanding of the structure of learning across the body. 34 First, while most of the movement was done by the right arm, the entire body learned the task, as evident by the decrease in both inter- and intra- trial variabilities of various joints across the body 35 36 over learning. Second, while over learning all subjects decreased their movement variability and 37 the variability in the outcome (ball direction), subjects who were initially more variable were also more variable after learning, supporting the notion that movement variability is an individual trait. 38 39 Lastly, when exploring the link between variability and learning over joints we found that only the 40 variability in the right elbow supination shows significant correlation to learning. This demonstrates the relation between learning and variability: while learning leads to overall reduction in movement 41 variability, only initial variability in specific task relevant dimensions can facilitate faster learning. 42

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44 Author Summary

45 This study addresses a foundational problem in the neuroscience: studying the mechanisms of motor control and learning in free behaving, real-life tasks, where our brains and bodies operate 46 47 in on a daily basis and which contains the richness of stimuli and responses for what our nervous system evolved. We used the competitive sports of pool billiard to frame an unconstrained real-48 49 world skill learning experiment which is amenable to predictive modelling and understanding. Our data-driven approach unfolds the structure and complexity of movement, variability, and motor-50 learning, highlighting that real-world motor-learning affects the whole body, changing motor-51 control from head to toe. Crucially, we are enabling novel hypothesis driven experimental 52 53 approaches to study behavior where it matters most - in real life settings.

54 Introduction

55 Motor learning is a key feature of our development and daily lives, from a baby learning to roll, to an adult learning a new sport, or a patient undergoing rehabilitation after a stroke. The 56 process of learning a real-world motor skill is usually long and complex, and difficult to quantify. 57 As a result, real-world motor learning is rarely studied, and most of the motor learning literature 58 59 focuses on relatively simple tasks, performed in a lab setup or an MRI scanner, such as force-field 60 adaptations (e.g. Diedrichsen et al., 2005; Howard et al., 2015; Shadmehr and Mussa-Ivaldi, 1994; Smith et al., 2006), visuomotor perturbations (e.g. Haar et al., 2015; Krakauer et al., 2000; Mazzoni 61 62 and Krakauer, 2006; Taylor et al., 2014), and sequence-learning of finger tapping or pinching tasks 63 (e.g. Clerget et al., 2012; Ma et al., 2011; Reis et al., 2009; Yokoi et al., 2018).

64 These reductionistic tasks enable to isolate specific features of the motor learning and 65 tackle them individually. While this plays an important role in our understanding of sensorimotor 66 control and learning, it addresses a very restricted range of behaviors that do not capture the full 67 complexity of real-world motor control and may overlook fundamental principles of motor control and learning in real-life (Faisal et al., 2010; Ingram and Wolpert, 2011; Wolpert et al., 2011). It is 68 only in natural behavioral settings that neuroscientific mechanisms are subject to evolutionary 69 70 selection pressures and it is the operation in these contexts for which the nervous system has been 71 designed (Hecht et al., 2014). Over the past decade there were few important efforts in this direction. One line of research devised more complex tasks for skill learning (e.g. Abe and Sternad, 72 2013; Cohen and Sternad, 2009; Shmuelof et al., 2012), but those were still computer based toy-73 74 tasks which try to emulate real-world tasks. The other line used actual real-world tasks such as 75 juggling (e.g. van Beers et al., 2013; Hecht et al., 2014; Ono et al., 2015; Sampaio-Baptista et al., 76 2014, 2015; Scholz et al., 2009), but these studies analyzed only anatomical and functional MRI 77 changes following learning and did not address behavior or neural activity during the learning 78 process.

79 Here we are taking a novel data-driven approach to study behavior where it matters most – in natural real-life settings. The paradigm in which we study real-world motor learning is the game 80 81 of pool table billiards. Billiards is a real-world task ideally suited to neurobehavioral study as 82 motion tracking in terms of movement in space, the natural constraints of game playing, and divisibility into trials captures the style of reductionistic lab-based motor learning tasks. Billiards 83 is also a natural task which is complex and involves many different sub-tasks (precision, alignment, 84 85 ballistic movements, high-level sequential planning) which requires complex skills. To tackle the 86 complexity of the high dimensional task space of this real-world task, we applied naturalistic 87 approaches and developed a neurobehavioral database of real-world motor learning behavior. This 88 includes the full body movement during the entire learning period, as well as the measurements of 89 task performance (balls movement on the table). This enabled us to quantify the trends of changes in each of them separately during the entire learning process, and to look for correlations betweenthe changes in the body movement and the performance in the task.

We structured the results as follows: We ground our results in previous work on reductionistic lab tasks, to show that our unconstrained task and its task goal (directional error of the target ball relative to the pocket it is meant to go in) displays the well-known features of human motor learning. We then characterize full-body movement structure during the task, and how learning changes the kinematics of all joints over trials. Next, we compare across subjects to map their performance, learning rates, and motor variability, and how initial variability and learning rates are linked.

99 **Results**

30 right-handed volunteers, with little to no previous experience playing billiards, performed 300 repeated trials (6 sets of 50 trials each with short breaks in-between) where the cue ball and target ball were placed in the same locations, and subjects were asked to shoot the target ball towards the far-left corner pocket (Figure 1A). During the entire learning process, we recorded the subjects' full body movements with a 'suit' of inertial measurement units (IMUs; Figure 1B), and the balls on the pool table were tracked with a high-speed camera to assess the outcome of each trial (Figure 1C).

107 Movement and Learning in a real-world pool task

108 The ball tracking data showed a double exponential learning curve for the decay in the 109 directional error of the target ball (relative to the direction from its origin to the center of the target 110 pocket) over trials (Figure 1D). The direction of the initial trials error was consistent across subjects 111 as they tended to hit the center of the target ball and shot it forward towards the center of the table. 112 For measuring success rates and intertrial variability we divided the trials into blocks of 25 trials 113 (each experimental set of 50 trials was divided to two blocks to increase the resolution in time). 114 The learning curve over blocks (Figure 1E) emphasized the reduction in the inter-subject variability 115 during learning (decreasing error bars). The success rate over blocks (percentage of successful trials 116 in each block; Figure 1F) showed similar learning to the directional error. The learning was also 117 evident in the intertrial variability in the shooting direction which decayed over learning (Figure 1G). Since learning also occurred within a block (especially during the first block) and the 118 variability might be driven by the learning gradient, we corrected for it by calculating intertrial 119 variability over the residuals from a regression line fitted to the ball direction in each block (while 120 121 the learning curve is exponential, within the small blocks of 25 trials it is almost linear). This corrected intertrial variability showed the same pattern (Figure 1H). Overall, the task performance 122 123 data suggested that subjects reached their peak performance on the fifth experimental set (blocks 124 9-10, trials 200-250) and are doing the same (or even slightly worse) on the last experimental set

- 125 (blocks 11-12, trials 250-300). Thus, we refer to the last two experimental sets (blocks 9-12, trials
- 126 201-300) as the 'learning plateau'.



Figure 1. *Experimental setup and task performance*. (**A**) 30 right-handed healthy subjects performed 300 repeated trials of billiards shoots of the target (red) ball towards the far-left corner. (**B**) Full body movement was recorded with a 'suit' of 17 wireless IMUs (Xsens MVN Awinda). (**C**) The pool balls were tracked with a high-speed camera. Dashed lines show the trajectories of the cue (white) and target (red) balls over 50 trials of an example subject. (**D**) The trial-by-trial directional error of the target-ball (relative to the direction from its origin to the centre of the target pocket), averaged across all subjects, with a double-exponential fit (red curve). Grey lines mark the range of successful trials. (**E**) The mean absolute directional error of the target-ball. (**F**) The success rates. (**G**) directional variability. and (**H**) directional variability corrected for learning (see text). (**E**-**H**) presented over blocks of 25 trials, averaged across all subjects, error bars represent SEM.

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The full body movements were analyzed over the velocity profiles of all joints, and not the 128 129 joint angles profiles, as those are less sensitive to potential drifts in the IMUs and have proven to 130 be more robust and reproducible across subjects in natural behavior (Thomik, 2016). In the current 131 data we can also see this robustness across trials (Figure 2A). The covariance matrix over the velocity profiles of the different joints, averaged across the initial block trials of all subjects, showed 132 133 that most of the variance in the movement is in the right arm, and specifically in the right shoulder 134 (Figure 2B). This is a signature for the naivety of the subjects, as pool billiards guide books 135 emphasize that the shooting movement should be from the elbow down while the shoulder should 136 be kept still. The covariance of the velocity profiles averaged across the initial block of the learning 137 plateau (trials 201-225) showed similar structure with an overall decrease relative to the initial trials but an increase in the variance of right elbow rotation (Figure 2C). 138



Figure 2. *Velocity profiles and covariance*. (**A**) Velocity profiles in 3 degrees of freedom (DoF) for each joint (blue: flexion/extension, red: abduction/adduction; green: internal/external rotation) averaged across subjects and trials over the first block of trials (1-25) in the inner circle (grey background) and the first block after learning plateau (201-225) in the outer circle (white background). The joints of the right arm, which do most of movement in the task, are highlighted in orange box. (**B**,**C**) The variance covariance matrix of the velocity profiles of all joints averaged across subjects and trials (**B**) over the initial block (1-25) and (**C**) first block after learning plateau (201-225). The order of the DoF for each joint is: flexion/extension, abduction/adduction, internal/external rotation.

140 On the group level, the velocity profiles of all joints (including the joints of the right arm 141 that carry most of the movement variance) showed only minor changes following learning. For 142 example, the flexion/extension of the right elbow showed a decrease in velocity from the initial 143 trials to the trials of the learning plateau (Figure 2A).

144 The generalized variance (GV; the determinant of the covariance matrix (Wilks, 1932)) over the velocity profiles of all joints increased fast over the first ~30 trials and later decreased 145 146 slowly (Figure 3A), suggesting active control of the exploration-exploitation trade-off. The covariance over the initial block, the block over the peak GV, and first block after learning plateau 147 (Figure 3B), shows that the changes in the GV were driven by an increase in the variance of all 148 149 DoFs of the right shoulder, and the negative covariance between the abduction/adduction and 150 internal/external rotation of the right shoulder to the flexion/extension of the right shoulder and wrist. The internal/external rotation of the right elbow showed a continuous increase in its variance, 151 which did not follow the trend of the GV. Principal component analysis (PCA) across joints for the 152 velocity profiles per trial for each subject, showed that while in all trials ~90% of the variance can 153 be explained by the first PC, there is a slow but consistent rise in the number of PCs that explain 154 more than 1% of the variance in the joint velocity profiles (Figure 3C). The manipulative 155 156 complexity, suggested by Belić and Faisal (2015) as way to quantify complexity for a given number of PCs on a fixed scale (C = 1 implies that all PCs contribute equally, and C = 0 if one PC explains 157 all data variability), showed cleaner trajectory with the same trend (Figure 3D). This suggests that 158 over trials subjects use more degrees of freedom in their movement. 159



Figure 3. *Variance and Complexity*. (**A**) The trial-by-trial generalized variance (GV), with a double-exponential fit (red curve). (**B**) The variance covariance matrix of the right arm joints velocity profiles averaged across subjects and trials over the initial block (trials 1-25), the second block (trials 26-50), in which the GV peaks, and first block after learning plateau (block 9, trials 201-225). The order of the DoF for each joint and the colorbar are the same as in Figure 2. (**C**) The number of principal components (PCs) that explain more than 1% of the variance in the velocity profiles of all joints in a single trial, with an exponential fit (red curve). (**D**) The manipulative complexity (Belić and Faisal, 2015), with an exponential fit (red curve). (**A**,**C**,**D**) Averaged across all subjects over all trials.

161 As a measure of task 162 performance in body space, 163 we defined a measure of Velocity Profile Error (VPE) 164 165 for each joint in each trial 166 (see methods). For all joints, VPE showed a clear pattern 167 168 of decay over trials in an 169 exponential learning curve (Figure 4A). A proximal-to-170 distal gradient in the time 171 172 constant of these learning 173 curves was observed across the right arm, from the 174 shoulder to the elbow and 175 176 the wrist rotation which 177 showed very slow learning 178 (the other wrist angles had very low VPE from the start, 179 thus did not learn much). 180 181 Intertrial variability in joint 182 movement was measured 183 over the VPEs in each block. Learning was also evident in 184 the decay over learning of 185 186 the VPE intertrial variability 187 over most joints across the 188 body (Figure 4B).

189 **Inter-subject differences** 190 in variability and learning 191 We found 192 substantial differences 193 between subjects in their 194 initial errors, final errors, 195 intertrial variability, and 196 learning, which are



Figure 4. *Learning over Joints.* (**A**) Velocity Profile Error (VPE) reduction across all joints. The trial-by-trial VPE for all 3 DoF of all joints, averaged across all subjects, with an exponential fit. The time constants of the fits are reported under the title. The color code of the DoF is the same as in figure 2 (blue: flexion/extension; red: abduction/adduction; green: internal/external rotation). (**B**) Velocity Profile Error (VPE) intertrial variability over blocks of 25 trials, averaged across all subjects.

197 overlooked in the group average results. One subject, who had low initial errors, showed no

198 learning, i.e. did not reduce her error over trials from the first block (trials 1-25) to the learning

199 plateau (trials 201-300). For all other subjects the final errors were smaller than the initial errors

200 (Figure 5A). There was a significant correlation between the initial and the final errors, meaning

subjects with higher initial errors tended to have higher final errors as well.



Figure 5. Variability and learning across subjects. (A) Correlation between subjects' mean absolute directional error over the first block (trials 1-25) and the learning plateau (trials 201-300). (B) Correlation between subjects' directional variability over first block (corrected for learning trend, see text) and over the learning plateau (C) Correlation between subjects' mean absolute directional error over the learning plateau and their learning (D) Correlation between subjects' VPE variability over the first block (corrected for learning trend, see text) and their learning (E) Correlation between subjects' VPE variability (in logarithmic scale) over the first block and the learning plateau for the right arm joints. (F) Correlation between subjects' VPE variability (in logarithmic scale) over the first block and their learning for the right arm joints. (A-F) Correlation values are Spearman rank correlation, regression lines are linear fits with 95% confidence intervals.

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While over learning most subjects decreased their intertrial variability in the outcome (ball direction; Figure 1H & 5B) there was some tendency (though non-significant) for subjects who were initially more variable to be also more variable after learning (Figure 5B). The intertrial variability of the joint velocity profiles, which also decreased over learning (Figure 4B), showed a clearer and stronger correlation between the initial and the final intertrial variability (Figure 5E & S1 Fig). While this phenomenon was observed in various joints across the body, and dominant in the abduction across the spine joints, it was most dominant in the right shoulder abduction and 210 rotation, the two joint angles that do most of the movement and carry most of its variance (Figure 2).

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212 Learning was defined as the difference between the initial error (over the first block: trials 1-25) and the final error (over the learning plateau: trials 201-300) normalized by the initial error. 213 214 There was no significant correlation between the learning and the final error (as subjects who started 215 worse could have learn more but still not perform better after learning), but there was a trend that 216 more learning leads to smaller final errors (Figure 5C). We tested if higher levels of initial task-217 relevant motor variability (variability in the directional error of the target ball) in this complex real-218 world task could predict faster learning across individual, as found in simple lab experiments (Wu 219 et al., 2014). We indeed found that individuals with higher intertrial variability in the directional 220 error of the target ball over the first block showed more learning (Spearman rank correlation r=0.64, 221 p < 0.001; Figure 5D). Importantly, this is the corrected intertrial variability (as in Figure 1I) which 222 is calculated over the residuals from a regression line fitted to the ball direction to correct for the 223 learning that is happening within the block. As a control we also tested for correlation with the 224 initial variability in the target ball velocity, task-irrelevant motor variability, and found no correlation (Spearman rank correlation r=0.06, p=0.77). Next, we tested the link between learning 225 226 and initial variability over the joint velocity profiles of the right arm (Figure 5F). We found that the 227 only joint angle where the intertrial variability showed significant correlation to learning was the right elbow rotation (Spearman rank correlation r=0.47, p=0.0086), which is the arm supination. 228 229 We further tested the link over the full body kinematics (S2 Fig) and found no other joint that 230 showed this correlation. Thus, while learning leads to overall reduction in movement variability, 231 only initial variability in specific, task-relevant, dimensions can facilitate/predict learning.

Discussion 232

233 In this paper we introduce a new paradigm for studying naturalistic motor learning during 234 whole-body movement in a complex real-world motor skill task. Our results present new insights 235 into motor learning in the real-world. While the learning curves in this in-the-wild paradigm are 236 within the same range of those reported in reductionistic motor adaptation tasks (e.g. McDougle et al., 2015; Smith et al., 2006) we find that this learning is taking place not only in the task relevant 237 238 joints but across the entire body. Also, we found that task relevant initial variability in the ball 239 direction (movement outcome) can predict learning, like in toy tasks (Wu et al., 2014), and so can 240 the initial variability in the right arm supination which is the task relevant joint angle variability.

241 While pushing towards real-world neuroscience, we started here with a relatively 242 constrained version of the real-world task, asking subjects to perform repeated trials of the same 243 pool shot. This was to enable analysis using well developed methods of laboratory studies in toy-244 tasks. Nonetheless, it is a major step in the direction of a naturalistic study. First, we allow fullbody unconstrained movement. Second, we do not use any artificial go cue and allow self-paced
movement and as many preparatory movements as the subject needs for each shoot. Third, subjects
receive natural somatosensory feedback. And last, we do not perturb the feedback to induce
learning.

249 Fundamentals of real-world motor learning

Across all subjects, we found that motor learning is a holistic process - the body is affected 250 251 as a whole from learning the task. This was evident in the decrease in the VPE and the intertrial 252 variability over learning (Figure 4). This result should not come as a surprise considering decades 253 of research in sport science showing this relationship. For example, baseball pitcher's torso, pelvis, 254 and leg movements are directly associated with ball velocity (Kageyama et al., 2014; Oliver and 255 Keeley, 2010; Stodden et al., 2006). Recently it was also demonstrated with full-body motion 256 capture in a ball throwing task (Maselli et al., 2017). And yet, unlike baseball pitches, basketball 257 throws, or any unconstrained overarm throw, where the whole body is moving, in a pool shot the 258 shooting arm is doing most of the movement and there is very little body movement. Thus, the 259 whole-body learning is not trivial and suggestive that even in arm movement toy-tasks there is a whole-body learning aspect which is overlooked. 260

We also found a proximal-to-distal gradient in the learning rates over the right arm joints (Figure 4A). This is especially interesting in light of the well-known phenomenon of proximal-todistal sequence in limb movements in sports science (Herring and Chapman, 1992) and in rehabilitation (Twitchell, 1951). While there are records of proximal-to-distal sequence at multiple time scales (Serrien and Baeyens, 2017), our results are the first to suggest that this gradient also occur over repetitions as part of the learning process.

267 Variability & learning

268 Intertrial variability is a fundamental characteristic of human movements and its underling neural activity (for review see Faisal et al., 2008). It was recently reported that individuals exhibit 269 270 distinct magnitudes of movement variability, which are consistent across movements and effectors, 271 suggesting an individual traits in movement variability (Haar et al., 2017). Our results show that 272 subjects who were initially more variable tended to be also more variable after learning in many 273 joints across the body (Figure 5E & S1 Fig) and specifically in those of right shoulder that carry 274 most of the variance in the movement. This supports the notion that there is an individual trait in 275 movement variability.

Intertrial kinematic variability is also thought to be critical for motor learning (e.g. Braun et al., 2009; Dhawale et al., 2017; Herzfeld and Shadmehr, 2014; Teo et al., 2011; Wilson et al., 2008). It was suggested that individuals with higher levels of task-relevant movement variability exhibit faster motor learning in both skill learning and motor adaptation error-based paradigms (Wu et al., 2014). The failures to reproduce this result in visuomotor adaptation studies (He et al., 2016; 281 Singh et al., 2016), led to the idea that experiments with task-relevant feedback (which is common 282 in visuomotor studies) emphasize execution noise over planning noise, whereas measurements 283 made without feedback (as in Wu et al., 2014) may primarily reflect planning noise (Dhawale et 284 al., 2017). This is in-line with a recent modeling work in a visuomotor adaptation study (with task-285 relevant feedback) in which subjects with higher planning noise showed faster learning, but the 286 overall movement variability was dominated by execution noise that was negatively correlated with 287 learning (van der Vliet et al., 2018). In our task there were no manipulations or perturbations, thus, 288 task-relevant feedback was fully available to the participants. On the other hand, in real-world there is no baseline variability, and the variability was measured during early learning and therefore is 289 290 probably dominated by planning noise, as subjects explore, regardless of the visual feedback. 291 Indeed, subjects with higher variability in the target ball direction over the first block showed higher 292 learning rates (Figure 5D). Our results straighten the link between variability and learning and are the first to show that it applies to real-world tasks. Moreover, the only joint angle that showed 293 294 significant correlation between initial variability and learning was the right elbow rotation (Figure 295 5F & S2 Fig). Following the idea that task-relevant variability predicts learning, it would suggest 296 that the right elbow rotation is the task-relevant joint angle to adjust during initial learning of a 297 simple pool shoot. Indeed, guide books for pool and billiards emphasize that while shooting one 298 should keep one's body still and move only the back (right) arm from the elbow down. While the 299 elbow flexion movement gives the power to the shoot, the arm supination (also known as 'screwing' 300 in billiards and measured by the elbow rotation in our IMUs setup) maintains the direction of the 301 cue.

302 Conclusions

In this study we demonstrate the feasibility and importance of studying human neuroscience in-the-wild, and specifically in naturalistic real-world skill tasks. While finding similarities in learning structure between our real-world paradigm and lab-based motor learning studies, we highlight crucial differences, namely, real-world motor learning is a holistic full-body process. Looking at the motor behavior over learning across the entire body enabled us to explore the relationship between variability and learning and define task relevant variability that can facilitate learning.

310 Methods

311 *Ethics statement.* All experimental procedures were approved by Imperial College 312 Research Ethics Committee and performed in accordance with the declaration of Helsinki. All 313 subjects gave informed consent prior participating to the study.

314 Experimental Setup and Design. 30 right-handed healthy human volunteers with normal or corrected-to-normal visual acuity (12 women and 18 men, aged 24±3) participated in the study. 315 316 The volunteers, who had little to no previous experience with playing billiards, performed 300 317 repeated trials where the cue ball (white) and the target ball (red) were placed in the same locations and the subject was asked to shoot the target ball towards the pocket of the far-left corner (Figure 318 319 1A). The trials were split into 6 sets of 50 trials with a short break in-between. For the data analysis 320 we further split each set into two blocks of 25 trials each, resulting in 12 blocks. During the entire 321 learning process, we recorded the subjects' full body movements with a motion tracking 'suit' of 322 17 wireless inertial measurement units (IMUs; Figure 1B). Brain activity was recorded with 323 wireless EEG, neural findings reported elsewhere (Haar and Faisal, 2020). The balls on the pool 324 table were tracked with a high-speed camera (Dalsa Genie Nano) to assess the subjects' success in 325 the game and to analyze the changes throughout learning, not only in the body movement and brain 326 activity but also in its outcome – the ball movement (Figure 1C).

327 Full-Body Motion Tracking. Kinematic data were recorded at 60 Hz using a wearable 328 motion tracking 'suit' of 17 wireless IMUs (Xsens MVN Awinda, Xsens Technologies BV, 329 Enschede, The Netherlands). Data acquisition was done via a graphical interface (MVN Analyze, 330 Xsens technologies BV, Ensched, The Netherlands). The Xsens joint angles and position data were 331 exported as XML files and analyzed using a custom software written in MATLAB (R2017a, The 332 MathWorks, Inc., MA, USA). The Xsens full body kinematics were extracted in joint angles in 3 degrees of freedom for each joint that followed the International Society of Biomechanics (ISB) 333 334 recommendations for Euler angle extractions of Z (flexion/extension), X (abduction/adduction) Y 335 (internal/external rotation).

336 *Movement Velocity Profile Analysis.* From the joint angles we extracted the velocity 337 profiles of all joints in all trials. We defined the peak of the trial as the peak of the average absolute 338 velocity across the DoFs of the right shoulder and the right elbow. We aligned all trials around the 339 peak of the trial and cropped a window of 1 sec around the peak for the analysis of joint angles and 340 velocity profiles.

341 *Statistical Analysis.* Trial by trial learning curves were fitted with a single or a double342 exponential learning curve using matlab fit function.

As a measure of task performance in body space, correlation distances (one minus Pearson correlation coefficient) were calculated between the velocity profile of each joint in each trial to the velocity profiles of that joint in all successful trials. The mean over these correlation distances produced a single measure of Velocity Profile Error (VPE) for each joint in each trial.

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$$VPE_{i} = \frac{\sum_{s}^{N_{scs}} corrDist(velProf_{i}, velProf_{s})}{N_{scs}}$$

Thus, VPE in trial *i* was the sum of the correlation distances between the velocity profile in trial *i* (*velProf_i*) and the velocity profiles in successful trials *s* (*velProf_s*), divided by the number of successful trials (N_{scs}).

For measuring success rates and intertrial variability we divided the trials into blocks of 25 351 trials by dividing each experimental set of 50 trials to two blocks. This was done to increase the 352 353 resolution in time from calculating those on the full sets. To improve robustness and account for outliers, we fitted the errors in each block with a t-distribution and used the location and scale 354 parameters (μ and σ) as the blocks' center and variability measures. Similarly, all correlations 355 356 between error, variability, and learning are Spearman's rank correlation coefficients. Regression 357 lines are based on linear regression fits (in logarithmic scale for VPE variability) and are presented 358 with 95% confidence intervals.

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458 Supporting Information



S1 Fig. Correlation between subjects' VPE variability over the first block and over the learning plateau. Presented for all joints in 3 degrees of freedom (DoF) for each joint (blue: flexion/extension, red: abduction/adduction; green: internal/external rotation). Subjects' VPE variability is in logarithmic scale. Correlation values are Spearman rank correlation, regression lines are linear.



S2 Fig. *Correlation between subjects' VPE variability over first block and their learning*. Presented for all joints in 3 degrees of freedom (DoF) for each joint (blue: flexion/extension, red: abduction/adduction; green: internal/external rotation). Subjects' VPE variability is in logarithmic scale. Correlation values are Spearman rank correlation, regression lines are linear.