Specialization of the motor system in infancy: From broad tuning to selectively specialized purposeful actions

Hana D'Souza¹, Dorothy Cowie², Annette Karmiloff-Smith³, & Andrew J. Bremner¹

¹Sensorimotor Development Research Unit, Department of Psychology, Goldsmiths, University of London, New Cross, London, SE14 6NW, UK.

²Department of Psychology, University of Durham, Durham, DH1 3LE, UK.

³Centre for Brain and Cognitive Development, Department of Psychological Sciences, Birkbeck, University of London, Malet Street, London, WC1E 7HX, UK.

CORRESPONDENCE TO: Dr Andrew J. Bremner, Department of Psychology, Goldsmiths, University of London, London, SE14 6NW, United Kingdom; Tel. +44 (0) 207 078 5142; email: a.bremner@gold.ac.uk; Hana D'Souza, Department of Psychology, Goldsmiths, University of London, London, SE14 6NW, United Kingdom; hana.dsouza@gmail.com

RESEARCH HIGHLIGHTS

- We provide new insights into the developmental process of motor specialization, by which motor abilities, similarly to cognitive and perceptual functions, start out broadly tuned to their goal, becoming progressively more tailored to action goals over the first year of life.
- During purposeful action, 9-month-old infants activated multiple redundant limbs, whereas 12-month-olds were much more likely to restrict their goal-directed movements to a single arm.
- Increased specialization of limb movements was associated with greater selective attention and motor experience.
- Our findings indicate that extraneous movements accompanying purposeful actions have the potential to provide early motor markers of later neurocognitive deficits.

ABSTRACT

In executing purposeful actions, adults select sufficient and necessary limbs. But infants often move goal-irrelevant limbs, suggesting a developmental process of motor specialization. Two experiments with 9- and 12-month-olds revealed gradual decreases in extraneous movements in non-acting limbs during unimanual actions. In Experiment 1, 9-month-olds produced more extraneous movements in the non-acting hand/arm and feet/legs than 12-month-olds. In Experiment 2, analysis of the spatiotemporal dynamics of infants' movements revealed developmental declines in the spatiotemporal coupling of movements between acting and non-acting arms. We also showed that the degree of specialization in infants' unimanual actions is associated with individual differences in motor experience and visual attention, indicating the experience-dependent and broad functional nature of these developmental changes. Our study provides important new insights into motor development: as in cognitive domains, motor behaviours are initially broadly tuned to their goal, becoming progressively specialized during the first year of life.

Keywords: motor development, extraneous movements, motor overflow, specialization, infancy, reaching, action

Towards the end of the first year of life there are marked improvements in the ways in which infants use their limbs to act on the world across a range of behaviours, including manual exploration, reaching, intermanual coordination, and locomotion (e.g., Adolph & Berger, 2011; Fagard, 2000; Kimmerle, Ferre, Kotwica, & Michel, 2010; von Hofsten, 2007). Developmental theorists generally agree that such developments involve the increasing specificity and differentiation of the motor system during infancy (e.g., Gesell, 1954; Gibson & Pick, 2000; Sporns & Edelman, 1993; Thelen, 1985). The development of an ability to select efficient goal-directed movements has often been studied during reaching, focusing particularly on changes in the characteristics of movements within limbs (e.g., Bhat & Galloway, 2007; Konczak & Dichgans, 1997; Thelen et al., 1993; von Hofsten, 1991; for a review see Hadders-Algra, 2013). Yet, less attention has been paid to more broad-scale changes in action across the body, i.e., the transition from a less specialized state of motor selection in which multiple limbs are activated, to one in which only relevant limbs are selectively activated in the service of a goal (Soska, Galeon, & Adolph, 2012). In this report we focus on the development of limb selection during purposeful action; specifically, we examine for the progressive development of an ability to select only a single hand/arm during unimanual actions.

A growing body of evidence suggests that, early in development, the brain is "broadly tuned" to the environment (see Johnson, 2011, for an overview). In other words, it starts out functionally diffuse, with the response properties of neural regions being less selective to particular stimuli (i.e., less specialized). Thus, patterns of activation in the adult brain are more localised than those in the infant brain. Brain activation becomes increasingly specialized over developmental time through interactions between various brain regions and the environment (Edelman, 1987; Elman et al., 1996; Fair et al., 2007; Johnson, 2011; Supekar, Musen, & Menon, 2009). It is likely that motor development is yoked to brain development and follows a similar trajectory, with motor ability being initially "broadly tuned" and becoming specialized over time. This gradual specialization likely happens in interaction with the environment through perception-action cycles (Gibson & Pick, 2000; Sporns & Edelman, 1993), leading to a better fit between action and environment.

Variable and overall movement abundance in early infancy is a hallmark of typical development, and the spontaneous movement repertoire of an infant provides insight into the early development of her/his nervous system (Einspieler & Prechtl, 2005). However, whereas spontaneous movements are typically assessed in the absence of interaction with an object, the goal of the current paper is to focus on how infants come to select particular limbs during purposeful actions towards an object. Previous work has shown that, in infants of 4.5-7.5 months of age (with no observed developmental changes in this period), actions with one hand are often accompanied by goal-irrelevant movements in other limbs, such as clenching, splaying, or wiggling of the fingers and toes (Soska et al., 2012). The production of these extraneous movements in the first months of life contrasts with the skilled purposeful movements of adults and even young children. However, developmental changes in extraneous movements in infancy remain undocumented and poorly understood. Thus we began our investigation with the hypothesis that the large prevalence of extraneous movements observed in young infants (Soska et al., 2012) may reflect a lack of specificity of the motor system, and that developmental decreases in such extraneous movements will signify the increasing specialization of the infant motor system. More specifically, we hypothesised that specialization (and thus reduction of extraneous movements) would occur particularly at the end of the first year of life, as part of the major developments in reaching, intermanual coordination, and locomotion seen at this time (e.g., Adolph & Berger, 2011; Fagard, 2000; Kimmerle et al., 2010; von Hofsten, 2007).

5

In Experiment 1, we began by measuring extraneous movements in non-acting limbs during unimanual reaching, predicting a developmental decrease in such movements between 9 and 12 months of age. Experiment 1 also investigated two factors which we expected to be related to the ability to select appropriate movements in infancy: individual differences in (i) motor experience, and (ii) selective attention. We were expecting decreases in extraneous movements to be specifically linked to the emergence of motor skills in which the use of the limbs is differentiated, such as walking with assistance. Therefore, we asked parents to report on their child's motor experience. We predicted that infants' motor experience would correlate negatively with extraneous movements in task irrelevant limbs.

The ability to visually select stimuli in the environment is known to be closely linked to motor processes in adult humans and non-human animals, with overlapping brain areas involved in attention shifts and movement preparation (Astafiev et al., 2003; Corbetta et al., 1998; Perry & Zeki, 2000; see also Allport, 1989; Rizzolatti & Camarda, 1987). Attention has also been implicated in the modulation of extraneous movements across the lifespan (for a review see Addamo, Farrow, Hoy, Bradshaw, & Georgiou-Karistianis, 2007). For example, children who are more easily distracted also produce more extraneous movements (Waber, Mann, & Merola, 1985). Furthermore, extraneous movements in adults increase when their attention is diverted (Baliz et al., 2005). A relationship between attention and extraneous movements has also been found in infants: looking more at an object during unimanual exploration was associated with fewer extraneous movements (Soska et al., 2012). To explore the relationship between attention and extraneous movements in more depth, we administered a well-established attention task (Gap-Overlap) to assess selection of visual information (Hood & Atkinson, 1993; Johnson, Posner, & Rothbart, 1991), predicting that greater difficulty with visual selection in the Gap-Overlap task would be associated with a higher prevalence of extraneous movements.

Experiment 1

We measured extraneous movements in non-acting limbs while infants were reaching with one hand (i.e., unimanually) for a ball. Measures of the proportion of unimanual reaches accompanied by extraneous movements were taken: (i) across the entire duration of the reach, and (ii) within +/- 100 ms of the onset of the unimanual reach to the ball, comparing groups of 9- and 12-month-old infants. Extraneous movements were also analysed in relation to infants' motor experience measured by parental report (how long infants had been sitting without support, crawling, standing with assistance, and walking with assistance), and infants' visual attention as measured by the Gap-Overlap task.

Methods

Participants

Two age groups, 9- and 12-month-olds, were tested in this experiment. The final sample size for Experiment 1 is presented in Table 1. Six additional infants were tested but excluded from analysis due to: (i) experimenter error (one 12-month-old), (ii) producing fewer than four unimanual reaches (three 9-month-olds, one 12-month-old), and (iii) not reaching for objects at all (one 9-month-old). The sample size in this study was consistent with sample sizes used in comparable studies (e.g., Adolph, 2000; Bhat & Galloway, 2007). The infants were recruited via a database of parents who expressed an interest in participating in developmental studies. Ethical approval was gained from the institutional research ethics committee. Prior to testing, informed consent was obtained from all parents. Testing only took place if the infant was awake and alert. The participants were given a small gift (e.g., a T-shirt) in return for their participation.

--Table 1 about here--

Procedure and materials

Reaching task. The infant was placed in an infant seat (Bébépod Flex, Prince Lionheart Inc., Santa Maria, CA, U.S.), and secured into place with adjustable straps around the waist so that movement of the trunk was restricted. There were 12 reaching trials in total. On each trial, the infant was presented with a 3.5 cm-diameter ball at the body midline, and at arm's length (calibrated for each infant) so that the infant could just grasp it without leaning forwards. The above measures were taken to prevent any potential compensatory movements in non-acting limbs resulting from changes in posture. The size of the ball was selected to induce unimanual reaching (see, e.g., Fagard, 2000). The ball's colour (white, orange, blue, green) was varied in a fixed random order between trials in order to maintain the infants' interest. Two video cameras operating at 100 Hz were used to record the infants 'movements, each facing the infant either side of the midline. The movements were then coded offline.

For the purpose of the current study, only unimanual reaches were analysed. A coder selected unimanual reaches using the following discrimination criteria from Corbetta and Thelen (1996). Unimanual reaches had to comprise a unilateral extension of one arm (the acting arm) towards the target which was followed by contact with the target. To be counted as a unimanual reach, the other non-reaching arm was required to either remain still or produce non-target-oriented movements which remained at least a fist size away from the ball. On average, each 9-month-old contributed 8.3 unimanual reaches (SD = 2.7). Each 12-month-old contributed on average 9.8 unimanual reaches (SD = 2.1).

For each unimanual reach, the coder identified the timings of: (i) the onset of the reach (i.e., the moment when any part of the acting hand and/or arm from the fingertips to the shoulder started a continuous trajectory which ended in target contact), and (ii) contact (i.e., the moment when the hand touched the target for the first time in the trial). Next, the coder identified for each unimanual reach, whether any extraneous movement occurred. For the

purpose of Experiment 1, an extraneous movement in the hand/arm was any non-target oriented movement of the non-acting hand and/or arm (from the fingertips to the shoulder) which did not come within one fist size of the target ball. Most of these movements included one or more of the following: clenching, lifting, splaying, or wiggling of the fingers, twisting of the wrist, twisting or jerking of the arms. An extraneous movement in the feet/legs was any movement in a foot and/or leg (from the toes to the hips). Most of these movements included one or more of the following: clenching, lifting, splaying, or wiggling of the toes, flexion, extension, or rotation at the ankles, rotation at the knee, jerking of the leg (Soska et al., 2012). The feet/legs never touched the target or came within one fist's size of it. If there was an extraneous movement during the reach, then we coded whether it was tightly linked to the onset of movement in the acting limb (starting +/- 100 ms around the onset of the reach of the acting hand/arm). To compute inter-rater reliability, a second coder independently scored whether the reach was unimanual in 20% of all the reaches. Subsequently, the second coder scored for the presence of movement in the limbs not involved in the unimanual reach and whether they were tightly linked to the onset of the reach in 20% of the data. Inter-rater reliability was over 90%.

We calculated the proportion of unimanual reaches accompanied by extraneous movement for each infant. The measure was computed separately for the non-acting hand/arm and (the average across both) feet/legs. Furthermore, we calculated a proportion of reaches in which extraneous movement onset was tightly linked to the onset of movement in the acting limb (+/- 100 ms around the onset of the reach in the acting hand/arm). The proportion of unimanual reaches accompanied by these reach-onset-locked extraneous movements was computed separately for the non-acting hand/arm and (the average across both) feet/legs. Since these data were proportional, they were arcsine transformed prior to inferential analyses. Raw data are presented in the figures.

Gap-Overlap task. The infant was placed on his or her parent's lap approximately 65 cm from a 20" screen. The experimenter monitored and recorded the infant's looking behaviour from an adjacent room, via a video camera operating at 25 Hz. The eye movements were then manually coded offline. Before each trial, an attractive centrally-located stimulus (an "attention-getter") was displayed on the screen to attract the infant's attention. This was a square of black and white geometrical shapes changing in size (zooming in and out) accompanied by an interesting sound. Once the infant was looking at the attention-getter, the experimenter manually initiated a trial. On each trial, the attention-getter first disappeared and was replaced by a central fixation stimulus. After 800 ms, a peripheral target appeared on the left or right side of the screen and remained displayed for 1200 ms. In the Gap trials, the central fixation stimulus disappeared 200 ms prior to the onset of the peripheral target, thus leaving the screen blank for 200 ms before the appearance of the target. In the Overlap trials, the peripheral target appeared while the central fixation stimulus remained onscreen, leading to an overlap in time between these two stimuli. The central fixation and peripheral target stimuli were selected from a pool of four stimuli (pictures of balls visually matched on colour, attractiveness, and size [5.3 x 5.3 cm]). The pairs of pictures were presented to the infants in a pseudorandom order. Throughout the study, each stimulus was used an equal number of times as a central fixation and a peripheral target. The central fixation and the peripheral target were never the same stimuli within any given trial.

The Gap-Overlap task consisted of three blocks. In each block, eight Gap and eight Overlap trials were presented, thus 16 trials in each block, and 48 trials in total. The order of presentation of the Gap and Overlap trials was randomized within each block. Trials were considered invalid if: (i) the infant did not look at the central stimulus immediately before the presentation of the peripheral target; and/or (ii) the infant did not look at the peripheral target within the duration of the trial. Inter-rater reliability calculated over 20% of the data was 98% for the validity of trials and 92% for saccadic reaction times. We decided *a priori* to exclude reaction times under 150 ms or over 1200 ms (e.g., see Csibra, Tucker, & Johnson, 1998; Wass, Porayska-Pomsta, & Johnson, 2011).

Nine-month-olds contributed an average of 14.2 valid Gap trials (SD = 4.7) and 13.4 valid Overlap trials (SD = 4.7). Twelve-month-olds contributed on average 15 valid Gap trials (SD = 4.9) and 14.1 valid Overlap trials (SD = 4.4). The "Gap effect" (the difference in reaction times between Gap and Overlap trials) was computed for each infant as a measure of efficiency of disengaging from a central visual stimulus to orient to a peripheral one. Outliers below and above 2 *SD* were excluded from the data set. The Gap effect was 86 ms (SD = 39 ms) for 9-month-olds and 81 ms (SD = 30 ms) for 12-month-olds.

Motor experience scoring. The amount of experience with motor skills was reported by parents in a custom interview (reporting on sitting without support, crawling, standing with assistance, walking with assistance, standing alone, walking alone; Wijnhoven et al., 2004). The parents were encouraged to use baby books, calendars, pictures, and videos to facilitate their memories (Adolph, 2002). An experimenter also confirmed that the infants could perform the motor skills listed above. A motor experience score was computed based on experience with a range of skills which were present in more than half of the infants tested in each age group. The resultant skills which were included in this measure were: sitting without support, crawling, standing with assistance, walking with assistance. The number of months' experience with each of these skills was summed for each infant to yield a "motor experience score". Outliers below and above 2 *SD* were excluded from the data set. On average, the motor experience score was 7.1 (*SD* = 4.5) for 9-month-olds and 12.1 (*SD* = 4.6) for 12month-olds.

Results

Reaching task

Throughout the reach, we identified significantly more extraneous movements in the non-acting hand/arm in 9-month-olds compared to 12-month-olds, t(36) = 3.27, p = .002, d = 1.09 (Figure 1a). Nine-month-olds also moved their feet/legs during a greater proportion of unimanual reaches than 12-month-olds, t(36) = 2.78, p = .009, d = 0.93 (Figure 1b).

--Figure 1 about here--

Traces of motor activity in extraneous limbs which are closely linked to the onset of purposeful movement have been observed in children and adults (Koerte et al., 2010; for a review see Addamo et al., 2007). Such movements have been characterized as reflecting a motor command which overflows from one limb to others (Addamo et al., 2007). This "motor overflow" in children and adults is typically observed during difficult motor tasks, and very much smaller in amplitude than the extraneous movements in infants documented here and elsewhere (Soska et al., 2012). In order to investigate the presence of motor overflow, we examined the extent to which the onsets of infants' extraneous movements were tightly linked to reach onsets by reporting extraneous movements with an onset within a window of +/- 100 ms around reach onset (henceforth, "tightly linked extraneous movements"). Within this window, 9-month-olds continued to show a higher proportion of unimanual reaches accompanied by the onset of extraneous movements in the non-acting hand/arm than 12-month-olds, t(36) = 5.36, p < .001, d = 1.79 (Figure 1a). A trend in the same direction was also observed with tightly linked extraneous movements in feet/legs, t(36) = 1.81, p = .078, d = 0.61 (Figure 1b).

Extraneous movements, selective attention, and motor experience

A multiple regression analysis was conducted to evaluate whether age, selective attention, and/or motor experience predicted extraneous movements in the non-acting

hand/arm. Using the Enter method, it was found that the linear combination of all three predictors explained a significant amount of the variance in both overall and tightly linked extraneous movements in the non-acting hand/arm (overall: $R^2 = .52$, F(3, 29) = 10.33, p < .52.001; tightly linked: $R^2 = .61$, F(3, 28) = 14.60, p < .001). Age and selective attention made a significant contribution to the prediction equation, while motor experience did not, for overall extraneous movements in the non-acting hand/arm (age: t(29) = -3.81, p < .001; selective attention: t(29) = 3.48, p = .002; motor experience: t(29) = 0.72, p = .479), and also for tightly linked extraneous movements in the non-acting hand/arm (age: t(28) = -5.21, p < .001; selective attention: t(28) = 2.98, p = .006; motor experience: t(28) = 1.09, p = .283). To confirm that selective attention explained a unique proportion of variance, we conducted a hierarchical regression that initially only included age as a predictor. The addition of selective attention as a predictor led to a significant increase in the proportion of variance explained (overall: change in $R^2 = .24$, F(1,30) = 14.47, p < .001, Table 2a; tightly linked: change in R^2 = .16, F(1,29) = 11.14, p = .002, Table 2b). Thus, the greater the Gap effect (i.e., the more difficulty infants had with visual selection), the more extraneous movements (overall, as well as tightly linked to movement onset) they produced in their non-acting hand/arm.

--Table 2 about here--

A multiple regression analysis was conducted to evaluate whether age, selective attention, and/or motor experience predicted extraneous movements in feet/legs. Using the Enter method, it was found that the linear combination of all three predictors explained a significant amount of the variance in overall extraneous movements in the feet/legs, but this time not in extraneous movements which were tightly linked to movement onset (overall: R^2 = .40, F(3, 29) = 6.47, p = .002; tightly linked: $R^2 = .18$, F(3, 29) = 2.14, p = .117). For overall extraneous movements in feet/legs, only motor experience made a significant contribution to the prediction equation, while age and selective attention did not (motor experience: t(29) = -3.00, p = .005; age: t(29) = -1.31, p = .202; selective attention: t(29) = 1.36, p = .185; see Table 3). Thus, the more experience infants have with motor skills, the fewer overall extraneous movements in feet/legs they produced.

--Table 3 about here--

To disentangle which type of motor experience (out of: sitting without support, crawling, standing with assistance, and walking with assistance) predicts overall extraneous movements in feet/legs, we carried out additional multiple regression analyses. Using the Enter method, it was found that the linear combination of all four predictors explained a significant amount of the variance in overall extraneous movements in feet/legs, $R^2 = .50$, F(4, 26) = 6.57, p < .001. Crawling and walking with assistance made a significant contribution to the prediction equation, while sitting without support and standing with assistance did not (crawling: t(26) = -2.24, p = .034; walking with assistance: t(26) = -2.15, p = .041; sitting without support: t(26) = -0.10, p = .923; standing with assistance: t(26) = 1.77, p = .089; see Table 4a). This therefore suggests that locomotor experience is predictive of the decrease of extraneous movements in feet/legs but not, as shown earlier, in the hands/arms. A stepwise regression revealed that crawling and walking with assistance do not contribute a unique proportion of variance, with crawling being a significant predictor in this entry method, $R^2 = .42$, F(1, 31) = 22.62, p < .001, t(31) = -4.76, p < .001, see Table 4b.

--Table 4 about here--

Experiment 2

Experiment 1 revealed a developmental decrease in extraneous movements during reaching between 9 and 12 months of age. We also observed that a large proportion of 9-month-olds' unimanual reaches were accompanied, at their onset (+/- 100 ms), by extraneous movements in the non-acting hand/arm. These onset-locked movements are likely the developmental precursor of the motor overflow seen in older children and adults during demanding actions (Addamo et al., 2007).

In adults, when a unilateral motor command is generated in one hemisphere, the interhemispheric connections of the corpus callosum usually inhibit the corresponding area in the other hemisphere (Grefkes, Eickhoff, Nowak, Dafotakis, & Fink, 2008). Thus, it has been argued that symmetrical control of the two hands is surmounted via inhibitory processes (Dennis, 1976; Duque et al., 2007). However, inhibitory processes are limited in infancy and some have suggested (e.g., Fagard, 1998; Goldfield & Michel, 1986; Goldfield & Wolff, 2004) that symmetrical activation of the hands is a general principle of action in early infancy. Therefore, we should also witness more symmetrical spatiotemporal congruency between acting and non-acting arms in young infants compared to older infants. Experiment 2 tested this hypothesis by examining the spatiotemporal congruency between the acting and nonacting arms. Using motion capture, we measured in fine detail the spatiotemporal coupling between movements in acting and non-acting hands/arms during the action of shaking a rattle with a single hand in 9- and 12-month-olds. We predicted the presence of spatiotemporal coupling (symmetrical about the body midline) in 9-month-olds, which would be significantly reduced in 12-month-olds. A key advantage of the rattling action is that it allows greater confidence that any extraneous movements are an unintended outcome of the action. Many studies of reaching behaviour (Experiment 1 included) have to make assumptions about

15

whether infants intended the reach towards objects to be with one or two hands (see Fagard & Pezé, 1997). With unimanual rattle shaking, intention to act bimanually is extremely unlikely.

In adults, the interhemispheric inhibition responsible for suppressing bilateral motor activation is down-regulated as the effort required for a motor response increases (Perez & Cohen, 2008; Tinazzi & Zanette, 1998). Thus, we also investigated in the infants the effect of effort (speed of shaking) (Bodwell, Mahurin, Waddle, Price, & Cramer, 2003; Morrison, Hong, & Newell, 2011) on spatiotemporal congruency during rattle shaking. If increased speed of shaking is related to increases in between-arm congruency in infants, this would suggest that the developmental suppression of extraneous movements in infancy is driven at least in part by inhibitory processes.

Methods

Participants

The 9- and 12-month-olds recruited for Experiment 1 were also asked to participate in Experiment 2. The sample size for Experiment 2 is presented in Table 1. In Experiment 2, ten infants (in addition to those reported in Table 1) were tested but excluded from analysis because: (i) they produced fewer than four shaking sequences (six 9-month-olds, two 12-month-olds) and (ii) due to equipment failure (one 9-month-old, one 12-month-old). The sample size in this study was consistent with sample sizes used in comparable studies (e.g., Adolph, 2000; Bhat & Galloway, 2007).

Procedure and materials

In Experiment 2, the infant was placed in the same infant seat as used in Experiment 1, and secured with adjustable straps around the waist so that movement of the trunk was restricted. The rattle (which was 19 cm in length, and 6.7 cm in diameter at its widest point) was presented to the infant. Following extensive piloting with several types of rattle, this particular rattle was selected as being the one which produced the greatest amount of unimanual shaking in the infants. We alternated between giving the infants each of two different versions (one was opaque, the other transparent, but they were otherwise identical) of the same rattle in order to maintain their engagement in the task. At the beginning of each trial, the experimenter demonstrated the shaking of a rattle and placed the rattle in the infant's left or right hand to facilitate unimanual shaking. The side of presentation was presented in a novel pseudorandom order for each participant (with the constraint that the rattle could not be placed in the same hand more than twice consecutively). There were 6 shaking trials in total, each lasting 30 seconds.

Movement of the arms during shaking was recorded using an eight-camera OptiTrack motion capture system operating at 100 Hz (NaturalPoint, Inc., Corvallis, OR, U.S.). Six of the cameras recorded position–time data from both arms while two cameras served as video cameras (100 Hz). This allowed us to obtain motion capture data synchronized with video footage. Cameras were placed surrounding the infant. The reference frame of the system was set relative to the infant, such that the *x*-axis (horizontal) corresponded to the left-right axis of the body (left shoulder to right shoulder). We observed very little trunk rotation, which might otherwise have misaligned the body left-right axis of the body, and we observed very little forward trunk sway which might otherwise have misaligned the body and room vertical axes.

To capture the position of the arms, two custom-made rigid bodies were constructed. Each was made up of an array of four reflective markers (each 15.88 mm in diameter) placed in fixed positions on a small non-reflective plastic board (550 x 550 mm). The rigid bodies were each mounted on a velcro strap which was used to secure them to the infants' forearms (one on each arm).

From the video records, the coder selected unimanual shaking sequences which were at least 2 seconds long, when one hand was shaking the rattle on one side of body while the other hand was not touching the rattle and free to move on the other side of the body. The motion capture data were analysed offline. At each frame the 3D positions of the centre of each of the rigid bodies were calculated using the system's analysis software ("Tracking Tools", NaturalPoint Inc., Corvallis, OR, U.S.). Further analyses were performed with customized Matlab routines (Mathworks Inc., Natick, MA, U.S.). For each shaking sequence, the position of the centre of the rigid body on *x*-, *y*-, and *z*-axis was plotted and then visually inspected to identify shaking sequences which did not contain sufficient data for further analysis (e.g., due to occlusion of motion capture markers). Segments of data with significant artefacts (e.g., spikes) were detected visually and deleted. Next, the data were interpolated using a cubic spline function and filtered using a second-order low pass Butterworth filter operating at 8 Hz.

As a measure of spatiotemporal congruency between the arms, we calculated the correlation at each time point between the positions of the acting and non-acting arms on the horizontal (*x*) and vertical (*y*) axes for every shaking sequence (following Fagard & Pezé, 1997). We focused on the *x*- and *y*-axes, because shaking did not typically involve significant movement in depth (the *z*-axis). On the vertical (*y*) axis, a higher positive correlation indicates greater spatiotemporal congruency (e.g., as one arm moves up, the other also moves up, see Figure 2a). On the horizontal (*x*) axis, negative correlations indicate greater spatiotemporal symmetry about the body midline (e.g., as one arm moves right, the other moves left, see Figure 2b). Outliers above and below 2 *SD* were excluded from the data set. Because the data were bounded between -1 and 1, they were arcsine transformed. The raw data are presented in the figures.

The average speed of the shaking arm was computed for each shaking sequence. In order to examine the role of shaking speed in spatiotemporal coupling between acting and non-acting arm movements, a median split (Mdn = 287 mm/sec) was performed on speed of

the shaking arm to divide shaking sequences into a slower half (M = 187 mm/sec, SD = 64 mm/sec; henceforth "slow") and a faster half (M = 431 mm/sec, SD = 120 mm/sec; henceforth "fast").

Correlations in the vertical and horizontal axes for every sequence were used in the statistical analyses. In total, 242 unimanual shaking sequences were available for the analyses (9-month-olds: 126 shaking sequences; 12-month-olds: 116 shaking sequences). On average, each 9-month-old contributed 9 shaking sequences (SD = 5.9). Each 12-month-old contributed on average 6.1 shaking sequences (SD = 4.9).

Results

Vertical axis

The shaking sequences were entered into a 2 x 2 analysis of variance (ANOVA) examining the effect of Age (9- versus 12-month-olds) and Speed (of the acting arm; slow versus fast) on correlation scores in the vertical axis. The ANOVA revealed a main effect of Age, indicating more positive correlation scores in the shaking sequences of 9-month-olds than those of 12-month-olds, F(1, 238) = 6.90, p = .009, $\eta_p^2 = .03$. Furthermore, there was a main effect of Speed indicating that correlation scores were more positive with speed, F(1, 238) = 20.51, p < .001, $\eta_p^2 = .08$. There was also an interaction between Age and Speed, F(1, 238) = 8.07, p = .005, $\eta_p^2 = .03$. At a slow speed, 9-month-old's shaking sequences showed more positive correlation scores than those of the 12-month-old's triangle and Speed, f(1, 238) = 0.66 (*p*-value Bonferroni corrected) (see Figure 2c). But there was no significant difference between Ages for fast speed. While 9-month-olds did not show a difference in correlation scores between slow and fast speed movements, correlation scores at 12 months were significantly more positive for fast speed compared to slow speed movements, t(114) = 4.56, p < .001, d = 0.87 (*p*-value Bonferroni corrected) (see Figure 2c). One-sample *t*-tests of correlation scores against zero (i.e., no correlation; see Figure 2c) revealed that correlation scores were significantly more positive than zero for both slow and fast speeds in 9-month-olds, t(51) = 3.17, p = .012, d = 0.44; t(73) = 6.52, p < .001, d = 0.76 (*p*-values Bonferroni corrected). This was also true for 12-month-olds, but only for fast speed movements, t(46) = 4.58, p < .001, d = 0.67 (*p*-value Bonferroni corrected).

Horizontal axis

The shaking sequences were entered into a 2 x 2 ANOVA examining the effect of Age (9- versus 12-month-olds) and Speed (of the acting arm; slow versus fast) on correlation scores in the horizontal axis. The ANOVA revealed a main effect of Age, indicating more negative correlation scores in the shaking sequences of 9-month-olds than those of 12-month-olds, F(1, 238) = 9.01, p = .003, $\eta_p^2 = .04$. There was no main effect of speed or interaction between Age and Speed.

One-sample *t*-tests of correlation scores against zero (i.e., no correlation; see Figure 2d) revealed that correlation scores were significantly negative with respect to zero for fast speed movements in 9-month-olds, t(73) = -2.72, p = .032, d = 0.32 (*p*-value Bonferroni corrected). But slow movements in 9-month-olds and both slow and fast movements in 12-month-olds were not significantly different from zero.

--Figure 2 about here--

General discussion

We have documented for the first time a substantial decrease in extraneous movements accompanying unimanual object-directed reaching between 9- to 12-months of age (Experiment 1). This developmental trend is reflected in a simultaneous decrease in the spatiotemporal congruency of movements between the arms during unimanual shaking (Experiment 2). We argue that both findings reflect a wider developmental process of gradual motor specialization over the first year of life, in which infants' motor responses to action goals become increasingly tailored to their purpose, resembling the developmental processes of specialization seen in other domains including language and face processing (Gervain & Mehler, 2010; Lewkowicz & Ghazanfar, 2009; Maurer & Werker, 2014; Pascalis et al., 2005; Scott & Monesson, 2010; Werker & Tees, 1984; for a review see Johnson, 2011). The "broad tuning" of the motor system early in development is likely shaped through interactions with the environment (Gibson & Pick, 2000; Sporns & Edelman, 1993). For example, in Experiment 1, 9-month-olds activated multiple limbs even though only one of the hands successfully retrieved the object. It is likely that feedback about which limb was successful at retrieval over many repetitions gives rise to the ability to retrieve an object unimanually without activating any other limb. This would lead to a decrease in extraneous movements by 12 months of age. However, the current paper focused specifically on arm movements in reaching and shaking. Future research should explore the emergence of specialization in different motor sub-domains, which may differ in developmental timing.

What developmental processes underlie this motor specialization? The increases in inter-limb congruency with speed observed in Experiment 2 point to a role for inhibition in the modulation of extraneous movements (Addamo et al., 2007; Hoy, Fitzgerald, Bradshaw, Armatas, & Georgiou-Karistianis, 2004; Perez & Cohen, 2008). It may be that brain changes underlying the development of inhibitory connections between two hemispheres can explain the changes in extraneous movements observed in Experiments 1 and 2. The main brain structure responsible for interhemispheric inhibition is the corpus callosum, and this develops across early life reaching its adult size and myelination in later adolescence when extraneous movements typically decline to an adult level (for a review see Addamo et al., 2007; Giedd et al., 1999). It has also been implicated in the development of a range of motor behaviours such

as laterality (e.g., Sacco, Moutard, & Fagard, 2006). Moreover, the development of the corpus callosum and developmental decreases in extraneous movements might both be related to the development of intermanual coordination abilities (Muetzel et al., 2008). However, it is very likely that other neurodevelopmental changes (e.g., pruning in ipsilateral corticospinal projections; Eyre, Taylor, Villagra, Smith, & Miller, 2001; Martin, 2005) contribute to explaining the wider range of changes in motor specialization.

In adults, the brain areas involved in movement preparation overlap substantially with those implicated in selective attention (Astafiev et al., 2003; Corbetta et al., 1998; Perry & Zeki, 2000; see also Allport, 1989; Rizzolatti & Camarda, 1987). Here, we found that infants with a greater ability to disengage from a familiar visual stimulus and shift attention to a new event were better able to produce movements more specifically tailored to their action goals (i.e., fewer extraneous movements), indicating an overlap in early life between processes of selective attention and movement (e.g., Bacher & Robertson, 2001; Robertson & Johnson, 2009). The emerging ability to shift attention between sensory stimuli which occurs during the first months of life (Colombo, 2001; Richards & Casey, 1992) likely provides the crucial foundation for the selective processes required in the motor skills, which continue to be perfected well beyond infancy (Addamo et al., 2007; Koerte et al., 2010).

Finally, our finding of a link between greater motor experience and fewer extraneous movements in the feet/legs suggests that motor specialization (as with specialization in other domains; Johnson, 2011) is an experience-dependent process. It is interesting to note that the locomotor skills were the most related to a reduction in extraneous movements in feet/legs. Given that the acquisition of locomotor skills places a particular burden on learning to move the feet/legs independently, the particular coupling between motor skills and extraneous movements of the feet/legs reinforces the view that motor learning is specific to the mode of action (Adolph, 2000). Further research will be needed to determine how motor experience

interacts with the neural mechanisms described above and which, we suggest, underlie the development of motor specialization. A better understanding of the developmental processes underlying motor specialization has great clinical significance since aggravated extraneous movements have been described in various clinical populations including children with attention-deficit/hyperactivity disorder (e.g., D'Agati, Casarelli, Pitzianti, & Pasini, 2010; MacNeil et al., 2011) and autism (e.g., Jansiewicz et al., 2006).

Why do infants produce extraneous movements during purposeful action? In fact we consider it highly likely that, far from being extraneous, such movements serve adaptive purposes in early life. It may be that such movements are a vestige of our evolutionary past. Movements in the feet/legs during reaching and manipulation (Soska et al., 2012) could be driven by a phylogenetically older, quadrupedal system of movement (see Dietz, 2002). It is possible that the dramatic decline in such extraneous feet/legs movements observed here may signify the progression to a more recently evolved mode in which the manual system operates independently of quadrupedal movement (Dietz, 2002). Similarly, but without the proposal of an independent manual system, it may be that infants become gradually more competent at controlling redundant degrees of freedom and developing efficient movement synergies (see Bernstein, 1967; Sporns & Edelman, 1993). Irrespective of this question, however, it is likely that extraneous movements play an adaptive role in ontogenetic development. It may be that the broad motor tuning reflected by extraneous movements facilitates specialization by enabling the selection of the most efficient movements for a given action as sensorimotor experience progresses. A further possibility is that extraneous movements provide motor activity and reafferent sensory feedback which is crucial to activity dependent processes of development in the nervous system (see Blumberg, 2015).

The current paper provides important new insights into the processes whereby infants become able to select appropriate limb movements in the service of purposeful action. In the

SPECIALIZATION OF THE MOTOR SYSTEM IN INFANCY

early stages of learning to act on the world, the infant motor system appears to be "broadly tuned", but over the first year of life, motor responses become progressively more specialized. We argue that this broad motor tuning in early infancy likely fulfils an adaptive function by providing young infants with a wide repertoire of responses to their environment from which they can select the most effective over the coming months of life. The process of motor specialization documented here has wide-reaching implications for the development of a range of motor abilities such as object exploration, locomotion, intermanual coordination, the emergence of lateralised action, and tool use.

ACKNOWLEDGMENTS

This research was supported by the Department of Psychology, Goldsmiths University of London (bursary awarded to H. D.), and European Research Council Grant No. 241242 (European Commission Framework Programme 7), awarded to A. J. B. The authors thank the infants and parents who took part, Maurice Douglas, and researchers at The Sensorimotor Development Research Unit, Goldsmiths University of London, and the Centre for Brain and Cognitive Development, Birkbeck University of London.

REFERENCES

- Addamo, P. K., Farrow, M., Hoy, K. E., Bradshaw, J. L., & Georgiou-Karistianis, N. (2007).
 The effects of age and attention on motor overflow production—a review. *Brain Research Reviews*, 54(1), 189-204. doi:10.1016/j.brainresrev.2007.01.004
- Adolph, K. E. (2000). Specificity of learning: why infants fall over a veritable cliff. *Psychological Science*, *11*(4), 290-295. doi:10.1111/1467-9280.00258
- Adolph, K. E. (2002). Learning to keep balance. In R. Kail (Ed.), Advances in child development & behavior (Vol. 30, pp. 1-40). Amsterdam: Elsevier Science.
- Adolph, K. E., & Berger, S. E. (2011). Physical and motor development. In M. H. Bornstein
 & M. E. Lamb (Eds.), *Developmental science: An advanced textbook* (6 ed., pp. 241-302). Hillsdale, NJ: Lawrence Erlbaum Associates.
- Allport, A. (1989). Visual attention. In M. I. Posner (Ed.), Foundations of cognitive science (pp. 631-681). Cambridge, MA: MIT Press.
- Astafiev, S. V., Shulman, G. L., Stanley, C. M., Snyder, A. Z., Van Essen, D. C., & Corbetta, M. (2003). Functional organization of human intraparietal and frontal cortex for attending, looking, and pointing. *The Journal of Neuroscience*, 23(11), 4689-4699.
- Bacher, L. F., & Robertson, S. S. (2001). Stability of coupled fluctuations in movement and visual attention in infants. *Developmental Psychobiology*, *39*(2), 99-106.
 doi:10.1002/dev.1034
- Baliz, Y., Armatas, C., Farrow, M., Hoy, K. E., Fitzgerald, P. B., Bradshaw, J. L., & Georgiou-Karistianis, N. (2005). The influence of attention and age on the occurrence of mirror movements. *Journal of the International Neuropsychological Society*, *11*(7), 855-862. doi:10.1017/S1355617705051003

- Bernstein, N. A. (1967). *The co-ordination and regulation of movements*. New York: Pergamon.
- Bhat, A. N., & Galloway, J. C. (2007). Toy-oriented changes in early arm movements III: constraints on joint kinematics. *Infant Behavior and Development*, 30(3), 515-522. doi:10.1016/j.infbeh.2006.12.007
- Blumberg, M. S. (2015). Developing sensorimotor systems in our sleep. *Current Directions in Psychological Science*, 24(1), 32-37. doi:10.1177/0963721414551362
- Bodwell, J. A., Mahurin, R. K., Waddle, S., Price, R., & Cramer, S. C. (2003). Age and features of movement influence motor overflow. *Journal of the American Geriatrics Society*, *51*(12), 1735-1739. doi:10.1046/j.1532-5415.2003.51557.x
- Colombo, J. (2001). The development of visual attention in infancy. *Annual Review of Psychology*, *52*(1), 337-367. doi:10.1146/annurev.psych.52.1.337
- Corbetta, D., & Thelen, E. (1996). The developmental origins of bimanual coordination: a dynamic perspective. *Journal of Experimental Psychology: Human Perception and Performance*, 22(2), 502-522. doi:10.1037/0096-1523.22.2.502
- Corbetta, M., Akbudak, E., Conturo, T. E., Snyder, A. Z., Ollinger, J. M., Drury, H. A., ... Shulman, G. L. (1998). A common network of functional areas for attention and eye movements. *Neuron*, 21(4), 761-773. doi:10.1016/s0896-6273(00)80593-0
- Csibra, G., Tucker, L. A., & Johnson, M. H. (1998). Neural correlates of saccade planning in infants: A high-density ERP study. *International Journal of Psychophysiology*, 29(2), 201-215. doi:10.1016/S0167-8760(98)00016-6
- D'Agati, E., Casarelli, L., Pitzianti, M. B., & Pasini, A. (2010). Overflow movements and white matter abnormalities in ADHD. *Progress in Neuro-Psychopharmacology and Biological Psychiatry*, *34*(3), 441-445. doi:10.1016/j.pnpbp.2010.01.013

- Dennis, M. (1976). Impaired sensory and motor differentiation with corpus callosum agenesis: a lack of callosal inhibition during ontogeny?. *Neuropsychologia*, 14(4), 455-469. doi:10.1016/0028-3932(76)90074-9
- Dietz, V. (2002). Do human bipeds use quadrupedal coordination? *Trends in Neurosciences*, 25(9), 462-467. doi:10.1016/s0166-2236(02)02229-4
- Duque, J., Murase, N., Celnik, P., Hummel, F., Harris-Love, M., Mazzocchio, R., ... Cohen,
 L. (2007). Intermanual differences in movement-related interhemispheric
 inhibition. *Journal of Cognitive Neuroscience*, *19*(2), 204-213.
 doi:10.1162/jocn.2007.19.2.204
- Edelman G. M. (1987). *Neural Darwinism: The theory of neuronal group selection*. New York, NY: Basic Books, Inc.
- Einspieler, C., & Prechtl, H. F. R. (2005). Prechtl's assessment of general movements: a diagnostic tool for the functional assessment of the young nervous system. *Mental Retardation and Developmental Disabilities Research Reviews*, *11*(1), 61-67. doi:10.1002/mrdd.20051
- Elman, J. L., Bates, E. A., Johnson, M. H., Karmiloff-Smith, A., Parisi, D., & Plunkett, K.
 (1996). *Rethinking innateness: A connectionist perspective on development*. Cambridge, MA: MIT Press.
- Eyre, J. A., Taylor, J. P., Villagra, F., Smith, M., & Miller, S. (2001). Evidence of activitydependent withdrawal of corticospinal projections during human development. *Neurology*, 57(9), 1543-1554. doi:10.1212/wnl.57.9.1543
- Fagard, J. (1998). Changes in grasping skills and the emergence of bimanual coordination during the first year of life. In K. J. Connolly (Ed.), *The Psychobiology of the hand* (Vol. Clinics in Developmental Medicine, pp. 123-143). Londres: Mac Keith Press.

- Fagard, J. (2000). Linked proximal and distal changes in the reaching behavior of 5- to 12month-old human infants grasping objects of different sizes. *Infant Behavior and Development*, 23(3-4), 317-329. doi:10.1016/S0163-6383(01)00047-9
- Fagard, J., & Pezé, A. (1997). Age changes in interlimb coupling and the development of bimanual coordination. *Journal of Motor Behavior*, 29(3), 199-208.
 doi:10.1080/00222899709600835
- Fair, D. A., Dosenbach, N. U. F., Church, J. A., Cohen, A. L., Brahmbhatt, S., Miezin, F. M.,
 ... Schlaggar, B. L. (2007). Development of distinct control networks through
 segregation and integration. *Proceedings of the National Academy of Sciences*, 104(33),
 13507-13512. doi:10.1073/pnas.0705843104
- Gervain, J., & Mehler, J. (2010). Speech perception and language acquisition in the first year of life. *Annual Review of Psychology*, *61*(1), 191-218.
 doi:10.1146/annurev.psych.093008.100408
- Gesell, A. (1954). The ontogenesis of infant behavior. In L. Carmichael (Ed.), Manual of child psychology (2 ed., pp. 335-373). New York: Wiley.
- Gibson, E. J., & Pick, A. D. (2000). An ecological approach to perceptual learning and development. New York: Oxford University Press.
- Giedd, J. N., Blumenthal, J., Jeffries, N. O., Rajapakse, J. C., Vaituzis, A. C., Liu, H., ...
 Castellanos, F. X. (1999). Development of the human corpus callosum during childhood and adolescence: a longitudinal MRI study. *Progress in Neuro-Psychopharmacology and Biological Psychiatry*, 23(4), 571-588. doi:10.1016/S0278-5846(99)00017-2
- Goldfield, E. C., & Michel, G. F. (1986). Spatiotemporal linkage in infant interlimb coordination. *Developmental Psychobiology*, *19*(3), 259-264.
 doi:10.1002/dev.420190311

- Goldfield E. C., & Wolff, P. H. (2004). A dynamical systems perspective on infant action and its development. In A. Slater & J. G. Bremner (Eds.), *Theories of infant development* (pp. 3-29). Malden, MA: Blackwell.
- Grefkes, C., Eickhoff, S. B., Nowak, D. A., Dafotakis, M., & Fink, G. R. (2008). Dynamic intra- and interhemispheric interactions during unilateral and bilateral hand movements assessed with fMRI and DCM. *Neuroimage*, *41*(4), 1382-1394.
 doi:10.1016/j.neuroimage.2008.03.048
- Hadders-Algra, M. (2013). Typical and atypical development of reaching and postural control in infancy. *Developmental Medicine & Child Neurology*, 55(s4), 5-8. doi:10.1111/dmcn.12298
- Hood, B. M., & Atkinson, J. (1993). Disengaging visual attention in the infant and adult. Infant Behavior and Development, 16(4), 405-422. doi:10.1016/0163-6383(93)80001-0
- Hoy, K. E., Fitzgerald, P. B., Bradshaw, J. L., Armatas, C. A., & Georgiou-Karistianis, N. (2004). Investigating the cortical origins of motor overflow. *Brain Research Reviews*, 46(3), 315-327. doi:10.1016/j.brainresrev.2004.07.013
- Jansiewicz, E. M., Goldberg, M. C., Newschaffer, C. J., Denckla, M. B., Landa, R., & Mostofsky, S. H. (2006). Motor signs distinguish children with high functioning autism and Asperger's syndrome from controls. *Journal of Autism and Developmental Disorders*, 36(5), 613-621. doi:10.1007/s10803-006-0109-y
- Johnson, M. H. (2011). Interactive specialization: a domain-general framework for human functional brain development? *Developmental Cognitive Neuroscience*, 1(1), 7-21. doi:10.1016/j.dcn.2010.07.003
- Johnson, M. H., Posner, M. I., & Rothbart, M. K. (1991). Components of visual orienting in early infancy: Contingency learning, anticipatory looking, and disengaging. *Journal of Cognitive Neuroscience*, 3(4), 335-344. doi:10.1162/jocn.1991.3.4.335

- Kimmerle, M., Ferre, C. L., Kotwica, K. A., & Michel, G. F. (2010). Development of role-differentiated bimanual manipulation during the infant's first year. *Developmental Psychobiology*, 52(2), 168-180. doi:10.1002/dev.20428
- Koerte, I., Eftimov, L., Laubender, R. P., Esslinger, O., Schroeder, A. S., Ertl-Wagner, B., ...
 & Danek, A. (2010). Mirror movements in healthy humans across the lifespan: effects of development and ageing. *Developmental Medicine & Child Neurology*, *52*(12), 1106-1112. doi:10.1111/j.1469-8749.2010.03766.x
- Konczak, J., & Dichgans, J. (1997). The development toward stereotypic arm kinematics during reaching in the first 3 years of life. *Experimental Brain Research*, 117(2), 346-354. doi:10.1007/s002210050228
- Lewkowicz, D. J., & Ghazanfar, A. A. (2009). The emergence of multisensory systems through perceptual narrowing. *Trends in Cognitive Sciences*, 13(11), 470-478. doi:10.1016/j.tics.2009.08.004
- MacNeil, L. K., Xavier, P., Garvey, M. A., Gilbert, D. L., Ranta, M. E., Denckla, M. B., & Mostofsky, S. H. (2011). Quantifying excessive mirror overflow in children with attention-deficit/hyperactivity disorder. *Neurology*, *76*(76), 622-628. doi:10.1212/WNL.0b013e31820c3052
- Martin, J. H. (2005). The corticospinal system: from development to motor control. *Neuroscientist*, *11*(2), 161-173. doi:10.1177/1073858404270843
- Maurer, D., & Werker, J. F. (2014). Perceptual narrowing during infancy: A comparison of language and faces. *Developmental Psychobiology*, 56(2), 154-178.
 doi:10.1002/dev.21177
- Morrison, S., Hong, S. L., & Newell, K. M. (2011). Similarity in the dynamics of contralateral motor overflow through increasing frequency of movement in a single limb.
 Experimental Brain Research, 213(4), 403-414. doi:10.1007/s00221-011-2790-0

- Muetzel, R. L., Collins, P. F., Mueller, B. A., Schissel, A. M., Lim, K. O., & Luciana, M. (2008). The development of corpus callosum microstructure and associations with bimanual task performance in healthy adolescents. *Neuroimage*, *39*(4), 1918-1925. doi:10.1016/j.neuroimage.2007.10.018
- Pascalis, O., Scott, L. S., Kelly, D. J., Shannon, R. W., Nicholson, E., Coleman, M., & Nelson, C. A. (2005). Plasticity of face processing in infancy. *Proceedings of the National Academy of Sciences of the United States of America*, 102(14), 5297-5300. doi:10.1073/pnas.0406627102
- Perez, M. A., & Cohen, L. G. (2008). Mechanisms underlying functional changes in the primary motor cortex ipsilateral to an active hand. *Journal of Neuroscience*, 28(22), 5631-5640. doi:10.1523/jneurosci.0093-08.2008
- Perry, R. J., & Zeki, S. (2000). The neurology of saccades and covert shifts in spatial attention: An event-related fMRI study. *Brain*, 123(11), 2273-2288. doi:10.1093/brain/123.11.2273
- Richards, J. E., & Casey, B. J. (1992). Development of sustained visual attention in the human infant. In B. A. Campbell, H. Hayne, & R. Richardson (Eds.), *Attention and information processing in infants and adults: Perspectives from human and animal research* (pp. 30-60). Hillsdale, NJ: Erlbaum.
- Rizzolatti, G., & Camarda, R. (1987). Neural circuits for spatial attention and unilateral neglect. *Advances in Psychology*, *45*, 289-313. doi:10.1016/s0166-4115(08)61718-5
- Robertson, S. S., & Johnson, S. L. (2009). Embodied infant attention. *Developmental Science*, *12*(2), 297-304. doi:10.1111/j.1467-7687.2008.00766.x
- Sacco, S., Moutard, M. L., & Fagard, J. (2006). Agenesis of the corpus callosum and the establishment of handedness. *Developmental Psychobiology*, 48(6), 472-481. doi:10.1002/dev.20162

- Scott, L. S., & Monesson, A. (2010). Experience-dependent neural specialization during infancy. *Neuropsychologia*, 48(6), 1857-1861.
 doi:10.1016/j.neuropsychologia.2010.02.008
- Soska, K. C., Galeon, M. A., & Adolph, K. E. (2012). On the other hand: Overflow movements of infants' hands and legs during unimanual object exploration. *Developmental Psychobiology*, 54(4), 372-382. doi:10.1002/dev.20595
- Sporns, O., & Edelman, G. M. (1993). Solving Bernstein's problem: A proposal for the development of coordinated movement by selection. *Child Development*, 64(4), 960-981. doi:10.1111/j.1467-8624.1993.tb04182.x
- Supekar, K., Musen, M., & Menon, V. (2009). Development of large-scale functional brain networks in children. *PLoS Biology*, 7(7), 1521. doi:10.1371/journal.pbio.1000157
- Thelen, E. (1985). Developmental origins of motor coordination: Leg movements in human infants. *Developmental Psychobiology*, *18*(1), 1-22. doi:10.1002/dev.420180102
- Thelen, E., Corbetta, D., Kamm, K., Spencer, J. P., Schneider, K., & Zernicke, R. F. (1993).
 The transition to reaching: Mapping intention and intrinsic dynamics. *Child Development*, *64*(4), 1058-1098. doi:10.1111/j.1467-8624.1993.tb04188.x
- Tinazzi, M., & Zanette, G. (1998). Modulation of ipsilateral motor cortex in man during unimanual finger movements of different complexities. *Neuroscience Letters*, 244(3), 121-124. doi:10.1016/S0304-3940(98)00150-5

von Hofsten, C. (1991). Structuring of early reaching movements: a longitudinal study. *Journal of Motor Behavior*, 23(4), 280-292. doi:10.1080/00222895.1991.9942039

von Hofsten, C. (2007). Action in development. *Developmental Science*, *10*(1), 54-60. doi:10.1111/j.1467-7687.2007.00564.x

- Waber, D. P., Mann, M. B., & Merola, J. (1985). Motor overflow and attentional processes in normal school-age children. *Developmental Medicine & Child Neurology*, 27(4), 491-497. doi:10.1111/j.1469-8749.1985.tb04573.x
- Wass, S., Porayska-Pomsta, K., & Johnson, M. H. (2011). Training attentional control in infancy. *Current Biology*, 21(18), 1543-1547. doi:10.1016/j.cub.2011.08.004
- Werker, J. F., & Tees, R. C. (1984). Cross-language speech perception: Evidence for perceptual reorganization during the first year of life. *Infant Behavior & Development*, 7(1), 49-63. doi:10.1016/S0163-6383(84)80022-3
- Wijnhoven, T. M., de Onis, M., Onyango, A. W., Wang, T., Bjoerneboe, G.-E. A., Bhandari, N., ... Al Rashidi, B. (2004). Assessment of gross motor development in the WHO
 Multicentre Growth Reference Study. *Food and Nutrition Bulletin*, 25(1 Suppl), S37-45.



<u>Figure 1:</u> Experiment 1. Proportion of unimanual reaches accompanied by movements in non-reaching limbs, for 9- and 12-month-olds. (a) Overall movements observed in the non-acting hand/arm throughout the reach, and movements in the non-acting hand/arm with an onset that is tightly linked to the onset of the reach (+/- 100 ms). (b) Overall movements in the feet/legs throughout the reach, and movements in the feet/legs with an onset that is tightly linked to the onset of the reach (+/- 100 ms). Error bars show +/- 1 SE; *p < .05, **p < .01, ***p < .001.



Figure 2: Experiment 2. Correlation scores for unimanual rattle-shaking movements. (a) An example of a rattle-shaking sequence where acting and non-acting arms show large positive correlation on the vertical (y) axis indicating that they were moving up and down in synchrony (the correlation between positions of the arms is .92). (b) An example of a rattle-shaking sequence where acting and non-acting arms show moderate negative correlation on the horizontal (x) axis indicating that they were moving in symmetry about the midline (the correlation between positions of the arms is -.39). (c) Correlation scores on the vertical axis during shaking in 9- and 12-month-olds for slow speed and fast speed of shaking. Error bars show +/- 1 SE; *p < .05, **p < .01, ***p < .001.

| | Age group | n | Sex | Mean age (months) |
|--------------|---------------|-----------------------|---------|---------------------------|
| Exporimont 1 | 9-month-olds | 18 | 9f, 9m | 8.88 (<i>SD</i> = 0.30) |
| Experiment | 12-month-olds | 20 | 8f, 12m | 12.14 (<i>SD</i> = 0.32) |
| Experiment 2 | 9-month-olds | 14 | 8f, 6m | 8.90 (<i>SD</i> = 0.31) |
| Experiment 2 | 12-month-olds | month-olds 19 7f, 12m | 7f, 12m | 12.19 (<i>SD</i> = 0.28) |

Table 1: Participant characteristics in Experiments 1 and 2

Table 2: Hierarchical regression analysis (Enter method) for variables predicting (a) overall movements observed in the non-acting hand/arm during the reach, and (b) movements in the non-acting hand/arm with an onset that is tightly linked to the onset of the reach (+/- 100 ms).

| 1 | `` |
|----|----------|
| 10 | ונ |
| ۱c | 1 |
| 1 | |

| | В | SE B | β | |
|---------------------|--------|-------|---------|--|
| Model 1 | | | | |
| Constant | 2.419 | 0.433 | | |
| Age | -0.138 | 0.041 | 520** | |
| Model 2 | | | | |
| Constant | 1.869 | 0.390 | | |
| Age | -0.136 | 0.034 | 511*** | |
| Selective attention | 0.006 | 0.002 | .487*** | |

Note: R^2 = .27 for Model 1 (p < .01); ΔR^2 = .24 for Model 2 (p < .001). **p < .01, ***p < .001.

(b)

| | В | SE B | β |
|---------------------|--------|-------|--------|
| Model 1 | | | |
| Constant | 1.727 | 0.272 | |
| Age | -0.123 | 0.025 | 661*** |
| Model 2 | | | |
| Constant | 1.431 | 0.251 | |
| Age | -0.122 | 0.022 | 658*** |
| Selective attention | 0.003 | 0.001 | .395** |

Note: R^2 = .44 for Model 1 (*p* < .001); ΔR^2 = .16 for Model 2 (*p* < .01). ***p* < .01, ****p* < .001.

| | В | SE B | β |
|---------------------|--------|-------|-------|
| Constant | 1.697 | 0.473 | |
| Age | -0.059 | 0.046 | 214 |
| Selective attention | 0.003 | 0.002 | .201 |
| Motor experience | -0.045 | 0.015 | 504** |

Table 3: Multiple regression analysis (Enter method) for variables predicting overall movements in the feet/legs during the reach.

Note: $R^2 = .40 (p < .01)$. **p < .01.

Table 4: Multiple regression analysis for motor experience variables predicting overall movements in the feet/legs during the reach: (a) Enter method and (b) Stepwise method.

| | В | SE B | β |
|--------------------------|--------|-------|------|
| Constant | 1.060 | 0.182 | |
| Sitting without support | -0.004 | 0.042 | 019 |
| Crawling | -0.099 | 0.044 | 439* |
| Standing with assistance | 0.069 | 0.039 | .275 |
| Walking with assistance | -0.182 | 0.084 | 378* |
| | | | |

Note: $R^2 = .50 (p < .001)$. *p < .05.

(b)

| | В | SE B | β |
|----------|--------|-------|--------|
| Constant | 1.215 | 0.099 | |
| Crawling | -0.149 | 0.031 | 650*** |

Note: $R^2 = .42 (p < .001)$. ***p < .001.