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From movement kinematics to social cognition: the case of autism

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The way in which we move influences our ability to perceive, interpret and predict the actions of others. Thus movements play an important role in social cognition. This review article will appraise the literature concerning movement kinematics and motor control in individuals with autism, and will argue that movement differences between typical and autistic individuals may contribute to bilateral difficulties in reciprocal social cognition.

1. Introduction

Already in the earliest descriptions of autism a variety of movement atypicalities have been noted including atypical postural control, gait, upper limb movements and fine motor control. However, these neurologically important signs have not been investigated as much as the social impairments in autism. Recent research has significantly advanced our understanding of the contribution of movements to socio-cognitive function. This literature suggests that processes such as action perception, prediction and interpretation are critical to social communication. For instance, these processes may be facilitated between two individuals who move similarly and impeded between individuals who move differently. In this paper, §2 briefly summarizes the literature suggesting that autistic and typical individuals move differently; §3 examines the contribution of one's own movement patterns to the perception, prediction and interpretation of the movements of others and, finally, §4 proposes that movement differences between typical and autistic individuals may contribute to bilateral difficulties in reciprocal social cognition. If so, autistic individuals will have difficulties perceiving, predicting and interpreting the actions of typical individuals and, conversely, typical individuals will have difficulties perceiving, predicting and interpreting the actions of individuals with autism. This interpretation goes some way towards the increasing recognition that the roots of the social difficulties that autistic individuals experience, are deeply embedded in compromised interactions, and are not solely due to processing deficits that are internal to the autistic person.

2. Are movements atypical in autism?

Autism spectrum disorder¹ (henceforth autism) is a developmental disorder characterized by impaired communication and social interaction, and restricted and repetitive interests [1]. Movement atypicalities have been linked with autism as far back as the work of Kanner [6] and Asperger [7], who noted motor abnormalities such as 'sluggish' reflexes, 'clumsy' gait and an absence, from an early age, of anticipatory postures when being picked up.

While most studies have focused primarily on social impairments in autism, there are also a number of reviews that have focused on movement atypicalities and abnormalities in areas of the brain relating to movement such as the cerebellum, striatum and brainstem (e.g. [3,8–10]). Here we give a brief overview of behavioural differences between autistic and typical individuals that have been noted with regard to various different types of movement. As this literature has been reviewed in depth elsewhere [11–16], we briefly summarize the main findings. These illustrate the wide-range of movement atypicalities that have been linked to all forms of autism.

(a) A note on movements and actions

When reviewing the literature concerning movements and actions, there are many possibilities for sub-categorizing the topic. For example, Gowen & Hamilton [12] decompose actions into constituent computational processes including motor planning, feed-forward control and motor execution; in doing so they demonstrate the utility of this approach in starting to isolate particular computational processes that may drive atypical movements in autism. In a review of the action understanding literature, Kilner [17] describes actions at four, non-independent, hierarchically organized levels: (i) the kinematic level: the trajectory and the velocity profile of the action; (ii) the motor level: the processing and pattern of muscle activity required to produce the kinematics; (iii) the goal level: the immediate purpose of the action; and (iv) the intention level: the overall reason for executing the action. This approach is particularly useful in illustrating that, due to the non-independence between different levels of the action hierarchy, an atypicality at one level (e.g. atypical goal identification) can impact upon other levels (e.g. atypical kinematics). Although both approaches are useful to bear in mind throughout this article, this paper will initially adopt a functional perspective in order to demonstrate that atypical movements are not restricted to one functional domain such as handwriting but may impact on many aspects of everyday life for individuals with autism.

(b) Posture and balance

At least 11 studies, to date, have investigated differences between autistic individuals² and non-autistic individuals in terms of postural control [19–29]. In an early study, Kohen-Raz *et al.* [25] measured autistic and typical participants' (aged 6–20 years) weight distribution while standing on stable and unstable surfaces with or without the benefit of vision. Autistic participants were generally less stable in their posture and typically exhibited a tendency to put most of their weight on one heel/toe. Similar patterns have been observed in subsequent studies of postural sway; for instance, autistic children demonstrate abnormalities when standing and looking straight ahead [21,22,30], standing while dual-tasking [31], standing with eyes closed [27,32], standing on unstable surfaces [27] and standing on a sway-referenced platform [26]. With a view to investigating the development of postural control in autism, Minshew *et al.* [26] recruited participants ranging from 5 to 52 years; they concluded that the development of postural control was delayed in autistic participants and differed from typical postural control even in adulthood.

(c) Gait

At least seven separate studies have assessed gait or the 'style of walking' in autistic children and adults, and a number of atypicalities have been observed [33–37]. For example, Nobile *et al.* showed that, compared with typical individuals, autistic children (6–14 years) exhibited trunk postural abnormalities, difficulties in walking in a straight line, a marked loss of smoothness (an increase in the jerkiness of movement) and, in general, a stiffer gait in which the usual fluidity of walking was lost. In a comprehensive review of gait atypicalities in autistic children, Kindregan *et al.* [13] found that the most commonly reported atypicalities concerned step width, step and stride

length, reduced velocity and increased time in the stance phase of gait. On the basis that increased step width provides a wider base of support, and reduced velocity and step and stride lengths help a walker to keep their centre of gravity within this base of support, they argue that together these results suggest a tendency for individuals with autism to augment their stability during walking—and, therefore, that autistic children have a more unstable gait compared with typical children. Extending this research into the adolescent years, Weiss *et al.* [38] found that 16- to 19-year olds with autism differed from typical controls with respect to various spatio-temporal aspects of gait, including step and stride length, foot positioning, cadence, velocity and step time. Hallett *et al.* [35] report mild clumsiness of gait and reduced range of motion of the ankle in autistic adults.

(d) Upper limb movements

Paradigms investigating upper limb movements in autism typically measure arm movement preparation and execution times and kinematic parameters, i.e. parameters referring to joint motions and angles at specific points in a movement and typically reported in terms of the velocity, acceleration (change in velocity) and jerk (change in acceleration) of a particular point on the body. Such studies have revealed differences between autistic and typical individuals [39–45]. To illustrate, Glazebrook *et al.* [41,42] found that adults with autism required more time both during movement initiation and execution for manual aiming movements, while Rinehart and colleagues have reported that autistic children [40] and young adults [46] require more time to prepare point-to-point movements (moving from one point in space to another).

Further work uses the reach-to-grasp task where, upon presentation of a cue, participants move their hand from a start position to grasp a target object. Using such a task, Stoit *et al.* [45] found that autistic children and adolescents exhibited longer movement times from the start of the movement to the grasp of the object. Yang *et al.* [47] found that children with autism showed significantly longer movement times for reach-to-grasp actions and executed their movements with more jerky kinematics. In line with this, Cook *et al.* [48] demonstrated that high-functioning adults with autism make more jerky movements that proceed with greater acceleration and velocity, even when these movements are not goal directed and are thus relatively unconstrained.

(e) Fine motor control

Fine motor control has typically been examined through analysis of handwriting in those with autism. While these studies have generally revealed autistic individuals to have atypical handwriting, the specific details of how handwriting deviates from the norm vary somewhat across studies [49–53]. In a comprehensive review of the literature concerning handwriting produced by children with autism, Kushki *et al.* [15] note consistent atypicalities in the overall legibility of handwriting and letter formation. For example, autistic children have been found to produce more poorly formed letters, though they do not exhibit difficulties in correctly aligning and spacing letters [51]. Macrographia (atypically large handwriting) has also been noted in both children [53] and adults with autism [50]. These features have been related to atypical movement kinematics [53]. Johnson *et al.* [53] demonstrated that handwriting-related movements were considerably

larger, peak velocity was significantly greater and movement trajectory more variable, in autistic children. An analysis of the velocity of movements suggested that autistic children may require higher energy input to achieve the same smoothness of movement as typical controls.

(f) Summary

Compared with typical individuals, children and adults with autism have, on average, been reported to exhibit increased instability during both standing and walking, atypical kinematics with respect to various movements, poor fine motor control as illustrated by atypical handwriting and, when making goal-directed or point-to-point arm movements, increased preparation and execution times. These findings, which are highly reliable and robust over many studies, suggest that, at a low level of cognitive processing, autistic individuals are likely to make movements which deviate from those made by individuals without autism. Adopting a bottom-up view, it is plausible that these 'low level' movement differences might impact on 'higher level' processing. This does not rule out that separate difficulties also exist at a higher level. However, it is possible that a bottom-up account would result in a parsimonious explanation of at least some of the symptoms of autism. In §3, we consider how the established movement atypicalities may influence higher level processes such as the perception, prediction and interpretation of others' actions, and how in turn this may disrupt very high level social interaction.

3. Movements influence socio-cognitive processes

Perceptual and motor systems are tightly linked: action influences perception and perception influences action. Research over the past few decades has demonstrated that this reciprocal relationship between action and perception may play a role in wider socio-cognitive functions, including action prediction, estimation of others' mental states, imitation and the development of positive social attitudes.

(a) Action and perception

Watching another person perform a movement evokes activity (often referred to as 'motor resonance') in one's own motor system. Evidence for this claim comes from a variety of fields: single cell recording studies have found that neurons in the motor system of the macaque (subsequently labelled 'mirror neurons') fire when the monkey passively observes an action [54], and research using a range of neuroimaging methods including functional magnetic resonance imaging (fMRI), transcranial magnetic stimulation, magnetoencephalography (MEG) and electroencephalography (EEG) provides strong evidence for similar responses to action execution and action perception in the human brain. fMRI experiments have identified overlapping activity for action perception and execution in a network of regions (subsequently referred to as the human mirror neuron system (MNS)), including the inferior frontal gyrus (e.g. [55]), inferior parietal lobe [56,57], ventral and dorsal premotor cortex [58,59], anterior intraparietal sulcus [60,61] and the superior temporal sulcus [62]. Furthermore, cross-modal repetition suppression, where a reduced response is seen for observation following execution

or vice versa, has been observed in both frontal [63] and parietal MNS areas [64].

Studies using MEG and EEG have also shown that sensorimotor oscillatory activity in both the 8–12 Hz (μ) and 15–30 Hz (β , *beta*) ranges is attenuated both when observing and executing actions [65–70]. However, electrical activity is not simply suppressed during action execution but is modulated *dynamically* [71,72]. Correspondingly, studies have demonstrated that sensorimotor oscillatory activity is also modulated *dynamically* during action observation according to the kinematics of the observed movement [73–76]. For example, Press *et al.* [76] demonstrated that *beta* power was dynamically modulated according to the acceleration profile of an observed arm movement, mirroring what would be expected during execution of the same action. Such automatic activation of the motor system during action observation can influence behaviour; that is, observing others' actions can interfere with ongoing action selection and execution such that we automatically imitate actions we observe [77–85].

Just as perception influences action, action influences perception. For example, inducing a motor load through performance of a concurrent task has been shown to modulate perceptual judgements about the weight of an object being lifted by an actor [86] or speed of a walker [87]. Similarly, perceptual judgements can be impaired through application of disruptive transcranial magnetic stimulation to motor regions [88]. Furthermore, in clinical populations, deficits in action production resulting from either cortical lesions and/or apraxia are correlated with deficits in action recognition [89–91]. Thus, there is widespread evidence that the motor and visual systems are intrinsically linked and mutually influence each other.³

(b) The importance of being similar: a worked example

Several theoretical accounts of the relationship between the visual and motor systems predict that the more similar two people are in their action execution the more likely they are to engage in motor resonance when observing each other's actions [93–96]. The following worked example describes such a situation in detail and elucidates how such effects might come about.

This example concerns three people, Fred, Jill and George. When Fred performs a reach-to-grasp movement he typically accelerates his hand towards the object until he has covered 50% of the distance, then begins to gradually decelerate. Jill performs this action with the same kinematics as Fred. George is different; George continues to accelerate his hand forward until he has covered 65% of the distance to the object. Fred has made movements like this for most of his life. He has a wealth of experience of observing his kinematic profile and simultaneously activating the motor codes for executing this reach-to-grasp movement (i.e. experience of simultaneously *seeing* and *doing*). This vast amount of experience means that for Fred the visual representation of a reach-to-grasp movement with a 50% acceleration phase has become tightly associated with his motor programmes for executing a reach-to-grasp movement [97]. Consequently, when Fred sees Jill make this movement it automatically activates his motor codes for executing a reach-to-grasp movement. By contrast, Fred has very little experience with seeing reach-to-grasp movements that follow George's (unusual) kinematic profile. Thus for Fred the visual representation of reach-to-grasp movements with George's

kinematics is only weakly associated with his motor code for executing a reach-to-grasp movement and, therefore, George's movement only weakly activates Fred's motor system. It can, therefore, be seen that movements that are more similar to one's own movements are more likely to result in motor resonance (e.g. Fred and Jill's movements) than those that are dissimilar (e.g. Fred and George's movements).

The argument that movement similarity boosts motor resonance is not merely theoretical: various laboratories have tested this hypothesis. For example, Cross *et al.* [98] trained expert dancers to learn complex whole-body dance sequences that were not in their motor repertoire prior to training. They found that motor system activity during passive observation of videoed dance sequences covaried as a function of the observer's ability to execute the dance move; greater activity was seen for movements that the dancer had mastered. Thus, motor resonance increased as participants' own movements became increasingly similar to the videoed movements.

(c) The importance of being similar: repercussions for socio-cognitive processes

As discussed in §3b, movements that are similar to one's own movement patterns are more likely to result in motor resonance. A number of studies suggest that a by-product of this motor resonance is the facilitation of various socio-cognitive functions, including action perception, prediction, interpretation and imitation. This point is illustrated here with various examples from the literature.

(i) Movement similarity and action perception

Casile & Giese [99] used motor training to ascertain the contribution of movement similarity to perception. Participants learned a novel upper-body movement while blindfolded, meaning that they received verbal and haptic, but not visual, feedback. Before and after training point-light stimuli were used to test the visual recognition of the learned movement. Despite the absence of visual stimulation during training, participants demonstrated an enhanced ability to visually recognize the trained movement. Furthermore, visual recognition performance after training correlated strongly with the accuracy of the execution of the learned movement. Thus, the more similar a participant's executed movements were to the observed movement, the better their visual recognition performance.

(ii) Movement similarity and action prediction

Aglioti *et al.* [100] demonstrated that professional basketball players could predict the success of free shots at a basket earlier and more accurately than individuals with comparable visual experience (coaches or sports journalists) but reduced motor experience. Moreover, Aglioti and colleagues found that only basketball players showed time-specific motor activation during observation of erroneous shots. They suggest that individuals who can move more similarly to the observed stimuli (i.e. basketball players) are more successful in their predictions, and that such results are a function of enhanced motor resonance.

(iii) Movement similarity and the mental state of confidence

Theoretical accounts predict that motor similarity should promote mental state inference [94]. Patel *et al.* [101] tested this hypothesis with respect to a particular mental state: confidence.

In an initial execution condition, participants performed a visual discrimination task wherein they successively viewed two images, one a target and one a foil. Participants indicated whether the first or second image contained the target by picking up a marble and placing it in the appropriately labelled slot, and subsequently rated their confidence in their decision. In this phase of the experiment, increasing confidence was associated with faster movements. In an ensuing observation task, participants watched a series of video clips showing the hands of anonymized actors performing the execution task and judged how confident they considered the actor to be. Patel and colleagues found that participants' judgements depended upon their own movement speed in the execution condition—if a participant watched an actor who moved faster than themselves then they were more likely to rate this actor as being confident, whereas movements performed slower than a participant's own movements were more likely to be rated as low in confidence. Participants were therefore more likely to accurately estimate confidence for movements that were similar in speed to their own movements.

(iv) Movement similarity and behavioural imitation

Kilner *et al.* [82] demonstrated that behavioural imitation of observed movements is greater for movements that are similar to one's own. Kilner *et al.* tracked participants' arm movements while they executed vertical sinusoidal arm 'waving' movements. Simultaneously, participants watched a video of an actor making incongruent horizontal movements. The video was experimentally manipulated such that the arm moved either with typical human kinematics (in a smooth, fluid manner) or at constant velocity (i.e. like a traditional robot). They found that observing videos of a person moving with human kinematics interfered with participants' on-going actions such that they subtly imitated the observed movement. By contrast, there were no subtle signs of imitation for the constant velocity movements. Thus, imitation was enhanced for movements that were similar to the participants' own movements relative to movements that were dissimilar.

(v) Movement similarity and positive affect

Movement similarity has been associated with positive affect. For example, Kirsch *et al.* [102] found that participants reported greater enjoyment and interest when observing dance movements from within their own motor repertoire, and an associated body of literature suggests that behavioural correlates of motor resonance such as movement synchronicity and automatic imitation may be intrinsically rewarding. For instance, Hove & Risen [103] demonstrated that participants who tapped synchronously with an experimenter liked the experimenter more than participants who tapped asynchronously. They argued that synchronicity of movements between interactants can promote the development of positive attitudes. Similarly, numerous studies have demonstrated that being imitated increases positive evaluations of interactions [104–107], and after being imitated people are more helpful, increase the amount they donate to charity [108], and feel closer to others [109]. Thus, a number of studies support the notion that movement similarity and behavioural correlates of motor resonance, such as movement synchronicity and automatic imitation, promote positive affect.

(vi) How important is motor resonance?

The literature described in §3c(i–v) shows that, compared with people who move in dissimilar ways, people who move in similar ways will probably experience more fluid action perception and prediction, be better at estimating each others' mental states, be more likely to imitate each other and be more inclined to develop positive affective ties to each other. It is possible that these diverse benefits of motor similarity are all due to enhanced motor resonance.

However, such effects may also be mediated by a visual experience route. To illustrate, imagine you have had a well-spent afternoon mastering the art of balancing a teaspoon on the tip of your finger. In doing so, you have learned that success is associated with a particular pattern of muscle contractions. Now imagine your friend attempts this complex balancing act. After watching only their initial bodily positioning, you successfully predict that the teaspoon will fall. According to the motor resonance account, observing your friend's initial positioning activates the corresponding motor codes within your system, generating a forward model (a prediction of the sensory consequences of the pattern of muscle contractions) from which you can predict the probability of success. However, while mastering the art of teaspoon balancing, in addition to motor experience, you also received visual experience. For example, you may have learned that the sight of your finger being at a particular angle relative to the ground and a certain distance from your body is highly predictive of success. If your motor system were temporarily lesioned you would still be able to use this visual experience to estimate your friend's chances of success.

Thus, both motor and visual experiences are important in our processing of others' actions. For many of the studies discussed above it has been empirically demonstrated that motor resonance adds predictive power over and above that contributed by the visual system alone [100,110]. However, when thinking about the repercussions of movement atypicalities in clinical populations, it is important to remember that if an individual tends to move differently compared with typical individuals they will have both different motor *and* visual experience of actions.

(d) Summary

Whether due to the natural development of their movements throughout their lifetime, or intense targeted training (e.g. dance classes), people who move similarly to each other will have comparable motor and visual experiences. Conversely, motor and visual experience is less comparable for individuals who move differently. Further, similar motor and visual experiences appear to facilitate socio-cognitive processes, including action perception, prediction, estimation of mental states, imitation and the development of positive affective ties. Thus, these processes are probably enhanced for people that move similarly and (relatively) impaired for those who do not.

4. Atypical movements and socio-cognitive function in autism

In 1996, Leary & Hill [111] published a controversial comment on the autism literature. They suggested that autism

research had virtually ignored movement atypicalities, instead focusing on social and communicative problems. They argued that social descriptions of behaviours such as 'a failure to cuddle', 'socially inappropriate gestures' and 'an indifference to affection' could be recast in terms of neurological motor symptoms such as 'abnormal posture and tone', 'dyskinesia' and 'marked underactivity'. Critically, they asserted that the application of a social context to motor behaviours diverts attention from the possible neurological explanations and thus hinders appropriate treatment interventions. Although Leary & Hill's [111] focus concerned social *interaction*—actions and reactions that occur between people—they also commented on social *cognition*—internal processes relating to the perception, prediction and interpretation of others:

Many individuals who experience movement disturbance report differences in internal mental processes, such as perception, changes in attention, consciousness, motivation, and emotion [112–115]. [111, p. 40]

Sections 2 and 3 summarized the literature demonstrating that autistic individuals move differently from typical individuals, and argued that socio-cognitive tasks such as perceiving, predicting and interpreting others may be made more difficult between people who move differently compared with those who move similarly. This section elaborates on Leary & Hill's comment by making the case that—at least in part due to movement differences—autistic individuals may have difficulties in perceiving, predicting and interpreting the actions of typical individuals, and, conversely, typical individuals may have difficulties perceiving, predicting and interpreting the actions of autistic individuals. I conclude by highlighting in §4d–g outstanding questions to be addressed by research in this area.

(a) Movement similarity and action perception in autism

Using motion-tracking technology, Cook *et al.* [48] examined the relationship between movement kinematics and action perception in autism. Adults with autism and typical individuals matched in terms of age, gender and intelligence performed simple sinusoidal arm 'waving' movements while the kinematics (velocity, acceleration and jerk) of their movements were recorded. Autistic individuals produced arm movements that were more jerky, and which proceeded with greater acceleration and velocity (figure 1), than those produced by typical individuals. The magnitude of these kinematic atypicalities was significantly positively correlated with autism symptom severity as measured by the Autism Diagnostic Observation Schedule semi-structured questionnaire [116]. Such results are consistent with reports from other laboratories of atypically jerky arm [47] and whole-body [36] movements in autism.

In a separate perception task, participants watched a series of visual stimuli comprising an image of a human hand that made vertical sinusoidal movements (down and then up) across the computer screen. The velocity profile of the hand was generated by motion-morphing between human-like minimum jerk motion and robot-like constant velocity. Participants also completed a non-biological control condition which featured a falling tennis ball, the velocity profile of which was a motion-morph between gravitational motion and constant velocity. Participants were required to label the movement of the stimulus as 'natural' or 'unnatural'. Results showed that the degree to which kinematic profiles

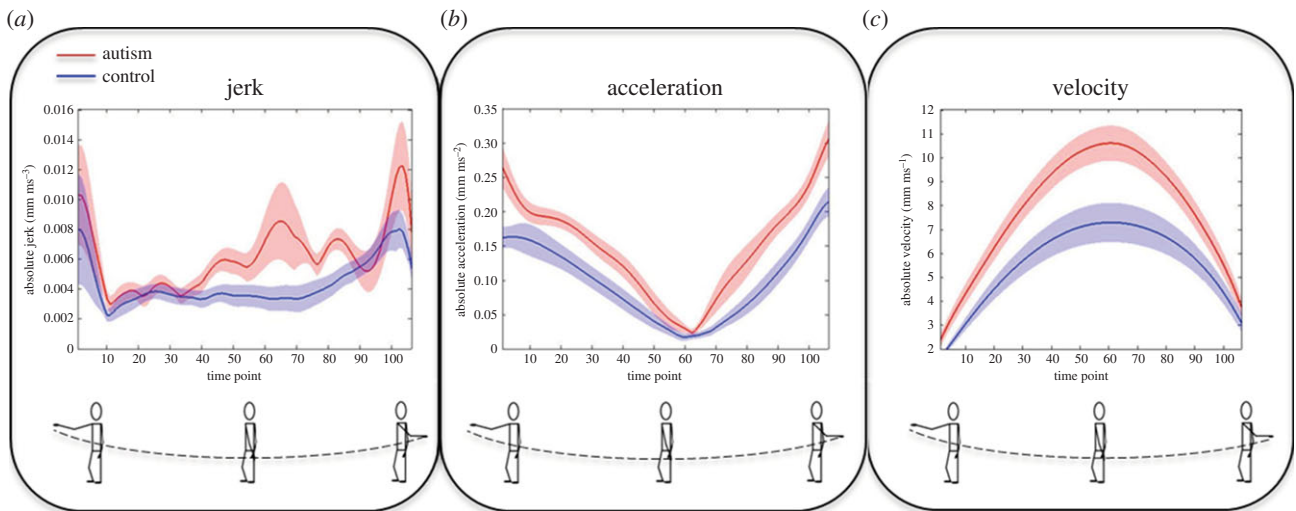


Figure 1. Kinematics of arm movements for autistic and typical individuals. When executing simple sinusoidal arm movements individuals with autism made more jerky movements (a) and travelled with faster absolute acceleration (b) and velocity (c). Mean movement vectors are plotted in red for the autism group and blue for the typical control group. Shaded regions indicate the standard error of the mean. Image adapted from Cook *et al.* [48] (fig. 3; CC BY).

were atypical when *executing* arm movements was significantly correlated with biased responding when *observing* motion of a human hand but not a tennis ball. In other words, the more atypical an autistic participant's kinematics (relative to kinematics exhibited by typical individuals), the less likely they were to classify movements that follow typical kinematics as 'natural'. Such results are consistent with the conclusions of Patel *et al.* [101] drawn from their studies of typical individuals; in the same way that a typical observer's perception of a confident movement was modelled on their own confident movements, autistic individuals' perception of natural movements is likely to be modelled on their own movements.

(b) Movement similarity and imitation in autism

Cattaneo *et al.* [117] investigated the link between action execution and automatic imitation of others' actions in children with autism and a matched group of typically developing children. In an action execution condition, participants were required to pick up a piece of paper and place it in a container, or pick up a piece of food and eat it. During both actions, the activity of the mouth-opening mylohyoid (MH) muscle was recorded using electromyography. In a separate 'observation condition' participants passively observed a typical child pick up a piece of (i) food and place it in their mouth or (ii) paper and place it in a container while activity from the MH muscle was recorded. Cattaneo *et al.* found that during the *execution* condition, MH muscle activity from typical children started to increase several hundreds of milliseconds before their hand grasped the food. It continued to increase during actual grasping, and reached its peak when the child started to open its mouth. MH muscle activity for autistic children was strikingly different: no activity increase was found during the entire reaching and grasping phases; the muscle only became active as the food was brought to the mouth.

These group differences during action execution translated into group differences during action observation: for typical children MH activity was observed when they passively observed another child reach and grasp a piece of food. By contrast, the autistic children did not show MH activation during the observation of either reaching or grasping

phases. Thus, atypical action execution in autistic children (i.e. a lack of anticipatory activation of the MH muscles when bringing food to their own mouth) was associated with atypical imitative responses.

(c) Is the mirror neuron system broken in autism?

Much of the past decade's literature concerning autism has debated the integrity of the MNS in this population (e.g. [118–123]). Thus, it is important to be clear about the claims made in this paper. Although the difference between the current stance and the broken mirror stance may appear subtle, it is important. Mirror neurons are active both when a person executes a movement and when they observe a movement. Hence, they can be considered a 'link' between the visual and motor system. Indeed, the broken mirror account of autism focuses on the *link between* action observation and execution: its key tenet is that what is broken is the *link between seeing and doing*. The current focus is different: here I focus on action execution, that is, not on any *link between seeing and doing*, but on *the doing itself*. This assumption is neutral as to whether mirror system activity measured independently, is atypical in autism. Several accounts have rivalled the broken mirror theory of autism [118,119,122]. This does not affect the current claim. Even if one assumes that the *link between* action observation and action execution is intact in individuals with autism they may still exhibit atypical imitation, and other socio-cognitive functions, due to atypical movement execution and their subsequently atypical visual and motor experience.

(i) Further questions

Section 4b highlights that reduced similarity between autistic and non-autistic movement kinematics and anticipatory muscle activation may impact on socio-cognitive functions (i.e. biological motion categorization and imitation). Clearly, much further work is required to elucidate the link between various movement differences between autistic and typical individuals (e.g. postural control, gait, fine motor control) and such socio-cognitive functions as intentional inferences, reading emotions from actions, estimating mental states, etc. In addition to widening the scope of this literature, there are

a number of important questions that also need to be addressed by this growing research field.

(d) What is atypical in the social interactions between autistic and non-autistic people?

The thesis outlined above supposes that impaired perception, interpretation and prediction of a typical person's movements can arise because an autistic individual has had a lifetime of visual and motor experience with their own movements—which differ from those of typical individuals. The same applies to the typical individual who encounters an autistic person. That is, most typical individuals have little visual and no motor experience with autistic movement patterns; thus they will probably have poor representations of autistic movements and thus potential deficits in the perception, prediction and interpretation of autistic behaviour. This is an important insight. It suggests that social interaction difficulties lie not with the autistic individual themselves but, rather, with both interaction partners: the autistic person has difficulties perceiving, predicting and interpreting the actions of the non-autistic person and vice versa. This shift in focus away from autistic individuals, towards the *interaction between* autistic and non-autistic people, is consistent with recent calls to develop a 'second person neuropsychiatry' with an increased focus on social interaction [124,125].

The question arises whether social interactions between partners who are both autistic are more fluid and whether such individuals show enhanced motor resonance due to greater movement similarity. A plausible alternative is that each atypical movement pattern is atypical in its own way and therefore dissimilar to every other individual. Preliminary support for the former comes from anecdotal evidence that high-functioning individuals with autism describe social interactions with other autistic individuals to be less effortful and more efficient compared with interactions with non-autistic people [124]. This argument also applies to the comparison of different conditions with neurological movement disorder. Further research is therefore necessary to investigate social interaction and its relationship to movement execution for autistic–autistic dyads and dyads comprising an autistic individual and an individual with a different movement disorder.

(e) Are atypical movements unique to autism?

The answer to this question is assuredly no. There are many conditions in which individuals exhibit movements that are different from those exhibited by typical controls including attention deficit hyperactivity disorder (ADHD), specific language impairment (SLI), Huntington's disease, Parkinson's disease and developmental coordination disorder. Indeed, finding a 'movement signature' that can differentiate individuals with autism from those with other conditions has become an important aim for the field due to its potential to expedite early detection. Initial studies show promise in differentiating autistic and *typical* children on the basis of movement patterns [39]. However, an important goal is to be able to differentiate autistic children from *those with other developmental conditions*, such as ADHD and SLI.

Differentiating autistic movements from those exhibited by children with ADHD is perhaps the most promising avenue in this literature so far [126–133]. MacNeil &

Mostofsky [129] have argued that whereas both children with ADHD and autism show impairments in basic motor control, difficulties with the formation of perceptual-motor action models are specific to autism. In line with this, Ament *et al.* [126] suggest that impairments in motor skills requiring the coupling of visual and temporal feedback to guide and adjust movement can differentiate ADHD, autism and developmental delay. McPhillips *et al.* [134] have begun to extend this line of research to other developmental conditions by comparing children with autism and SLI. However, much further work is required before a 'movement signature' differentiating autism from other conditions can be identified.

(f) Are movement atypicalities in other conditions, such as ADHD, associated with socio-cognitive function?

If moving atypically (i.e. different from typical controls) is associated with atypical perception, prediction and interpretation of controls' movements, and, if atypical movements occur in various conditions—from ADHD to Parkinson's disease—the current theory implies that individuals with these conditions might exhibit socio-cognitive atypicalities.

Socio-cognitive function in conditions including ADHD [135], Parkinson's disease [136] and Huntington's disease [137] is an active area of research, and it may be the case that further work in this field uncovers atypicalities in socio-cognitive function that cut across traditional diagnostic labels. However, it should be noted that for many conditions there may be additional factors, such as attentional control and executive function deficits, which feed into both motor control and social cognition impairments (this also applies to autism, see below). It is therefore important that future research attempts to ascertain the relative contribution of these various factors and/or uses tasks with minimal executive function, attention and memory requirements.

In studies where clinical groups are compared, the onset and duration of atypicalities matter. To give an example, if an individual has a sudden insult resulting in atypical movements (e.g. a torn ligament) this is unlikely to impact on socio-cognitive function; for that individual, their lifetime's visual and motor experience with typical movement patterns will probably outweigh the acute episode of atypical movements. This reasoning should apply to Parkinson's disease and other movement disorders acquired in late adulthood. It is likely that the impact of atypical movements on social cognition is a function of the length of time one has experienced atypical movements. At present, further research is required to ascertain the influence of the duration of movement atypicalities, and whether an individual's developmental stage at the time of onset is important.

(g) Are movement atypicalities the root cause of autistic cognition?

Using a bottom-up explanatory framework, can atypical movements in autism be considered the *root cause* of autistic cognition? Such an account is likely to be too simplistic. Rather, I argue that, though movement atypicalities may not explain all features of autistic behaviour, the role of movements in autistic socio-cognitive function should not be overlooked.

Contemporary accounts of autism suggest atypical computations that may pervade many cognitive functions from visual perception to decision-making. Recent examples are the notions of atypical priors [138] and aberrant precision of sensory information [139]. The latter, for instance, proposes that the precision of (i.e. reliability or confidence attributed to) incoming sensory information is too high relative to the precision of prior beliefs. This account provides a compelling explanation for visual perceptual atypicalities in autism: for instance, suggesting that autistic individuals' immunity to many visual illusions [140] may be due to abnormally high precision attributed to incoming sensory information relative to prior beliefs [139]. In addition, it has been argued that this account may help to explain difficulties with social interaction due to the heavy reliance of social interactions on prior beliefs [124]. Although the aberrant precision account has also been extended to repetitive and stereotyped behaviours [139], further work would be required to apply this account to the wide-ranging movement atypicalities documented in §2 of this paper. However, it is not impossible to imagine such an account. With respect to the atypically jerky gait characteristic of autism [36], the ability to walk in a smooth fluid manner is learned and refined during early development [141]. This process can be recast within a predictive coding framework whereby prior beliefs about how to optimally move are refined according to incoming sensory information. Atypically jerky gait in autism could therefore conceivably be due to an imbalance in the precision of incoming sensory information relative to prior beliefs.

(h) A final note

This paper has argued that visual and motor experience with own—atypical—movements in autism can result in the development of atypical (visual and/or motor) representations of movements, which is likely to impact on the perception,

prediction and interpretation of others' movements. Perhaps the most interesting implication of this claim is that the same argument should be true for typical individuals. That is, due to reduced experience with autistic movements, typical individuals may exhibit deficits in the perception, prediction and interpretation of autistic behaviour. Support for this hypothesis comes from a recent study showing poor recognition of autistic emotional facial expressions by typical control observers [142]. The real-world implications of this proposition should not be overlooked: it may be the case that many typical individuals who provide services for individuals with autism are poor at understanding the actions of their autistic service users. Thus, a final suggestion for further research is a comprehensive test of the hypothesis that typical controls exhibit poor perception, prediction and interpretation of autistic movements and an investigation of suitable training programmes.

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Endnotes

¹This review focuses on studies of autism spectrum disorder (referred to as autism for brevity) as defined in the DSM V [1]. Studies focusing exclusively on participants with Asperger's disorder have been excluded given the on-going debate concerning differences in motor function between autism spectrum disorder and Asperger's disorder [2–5].

²'Disability-first' terminology is used throughout in line with the majority preference expressed in a recent survey of the autistic community [18].

³Though note that recent accounts argue that motor system activity has widespread effects on perception that are not restricted to the action domain [92].

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