Postural Control in Children with Developmental Coordination Disorder

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ABSTRACT

The development of static balance is a basic characteristic of normal motor development. Most developmental motor tests include a measure of static balance. Children with Developmental Coordination Disorder (DCD) often fail this item. This study reviews the balance problems of children with DCD. The general conclusion is drawn that under normal conditions static balance control is not a problem for children with DCD. Only in difficult, unattended, or novel situations such children seem to suffer from increased postural sway. These findings raise the question of what happens when balance is lost. The present study addresses the strength of correlation between the electromyography (EMG) and force plate signals in one-leg stance over epochs of stable and unstable balance. Four groups of children were involved in the study: two age groups and a group of children with DCD and balance problems and their controls. The results show a clear involvement of tibialis anterior and peroneus muscles in the control of lateral balance in all conditions and groups. The group of children with DCD and balance problems, however, showed a weaker coupling between EMG and corrective force compared with control children, indicating non-optimal balance control. An evaluation of the existing data in terms of evidence of specific structural deficits associated with DCD provided converging evidence that suggests cerebellar involvement.

KEYWORDS
clothesiness, balance, loss of balance, force plate, EMG

INTRODUCTION

Not too long ago, evolution brought us upright stance, which opened new possibilities for functional behavior but also posed new problems for the control of locomotion and balance. Infants are not born with such capacities. In the first place, infants lack the strength for upright posture (Thelen et al., 1984). Additionally they completely lack experience with the pull of gravity and with vision, two sources of sensory information that are intimately involved in control of posture. Then there is the use of sensory feedback, which is essential for postural control. In a fully developed motor system, it is multi-modal and redundant, and intrinsically intertwined with anticipatory action, generating the reactive forces that are needed to counteract the—potential—loss of balance. The main sensory systems that are involved in the control of balance are the visual, kinesthetic, and vestibular systems, and pressure receptors of the somatosensory system. From a developmental point of view, the degree of postural control and
balance acts as a constraint on the development of specific motor skills. Development of postural control has been extensively described in the work of Woollacott and Shumway-Cook (e.g. Shumway-Cook & Woollacott, 1985; 1990). In the present study, I will review problems of balance control in children with Developmental Coordination Disorder (DCD) and present data of an analysis of short epochs of stable and unstable balance selected from trials of one-leg stance.

DEVELOPMENTAL COORDINATION DISORDER

Developmental Coordination Disorder is a classified disorder of problems in motor development (DSM-IV, APA, 1994). DCD can be briefly defined as 'poor motor performance in daily activities that is not consistent with the child's age and intelligence, and is not due to a medical condition'. The DSM-IV classification of DCD is commonly assumed equivalent to the Specific Developmental Disorder of Motor Function (SDDMF) in the ICD-10 classification (WHO, 1992). Swedish researchers (e.g. Gillberg & Gillberg, 1988) use the term DAMP (deficits in perceptual, attentional, and motor function) in addition to other categories of disorders, such as DCD and ADHD (attention deficit hyperactivity disorder). The authors point at the common co-occurrence of ADHD, DCD, and perceptual deficits, the prevalence of purer cases of AD(H)D and DCD being less than that of DAMP (Landgren et al., 1996). In any case, there is no clear evidence of neurological impairment in these children, although the problems may reflect a non-optimal form of brain function, such as in minor neurological dysfunction (MND) (Hadders-Algra, 2003). The prevalence of DCD is 3 percent to 6 percent, with 2 to 3 times as many boys than girls affected.

The DSM classification of DCD explicitly states that the condition significantly interferes with activities of daily living or academic achievement or both. A review of 41 case studies of DCD (aged 4 to 16 years) lists, in order of reported frequency, dressing (e.g. tying shoelaces), drawing and writing, locomotion, constructional play and the use of cutlery and scissors, speech, and ball skills/outdoor play as the major activities in daily living and school work that meet limitations (Geuze, 2005a,b). These limitations obviously are age dependent, as the 4-year-old child will not have yet learned to write. Balance and postural control are assumed fundamental to many of these tasks.

The main characteristics of DCD in the motor domain are poor postural control (moderate hypotonia or hypertonia, poor distal control, static and dynamic balance), difficulty in motor learning (learning new skills, planning of movement, adaptation to change, automatization), and poor sensorimotor coordination (coordination within/between limbs, sequencing of movement, use of feedback, timing, anticipation, strategic planning). Consequently, such children are slow, inaccurate, non-fluent, and variable in their movements (Geuze, 2005b). A general taxonomic distinction in the motor domain is that between gross and fine motor skills, with balance control specifically related to gross motor skill. The well-known Movement Assessment Battery for Children (M-ABC test, Henderson & Sugden, 1992) uses these classes of motor skills and has a further distinction between static and dynamic balance skills. These two types of balance are only weakly correlated, the former being more constrained compared with the latter because with a fixed base of support, the possibilities for a correction of loss of balance are much more limited.

It has been argued that perceptual difficulties are the cause of the motor problems in DCD (Laszlo & Bairstow, 1993). A meta-analysis of Wilson and McKenzie (1998) showed that DCD is associated with poor visual perception only in tasks involving visual-spatial processing, whether or not the task involved a motor component. A link
to pure visual perceptual deficits could not be
demonstrated. A second outcome of the meta-
analysis was a somewhat weaker association with
kinesthetic sensitivity. Vision and kinesthesis are
important sources of feedback in the control of
balance.

From these characteristics of DCD, one should
not conclude that children with DCD have, to a
large extent, a similar pattern of deficits. These
children are not a homogeneous group. On the
contrary, the common finding is that only about
half the children are affected in a specific skill or
function (Geuze et al., 2001). Correlations
between specific motor tasks are usually weak,
which implies that the child can fail on certain
motor tasks and succeed on others, suggesting that
there may be subgroups with a more homogeneous
set of symptoms, possibly with a common under-
lying defect. A review of studies that used cluster
analysis of performance on 5 to 6 motor tasks
(including static balance) (Visser, 2004) shows
that typically 4 to 5 subtypes of DCD are found.
Apart from a subtype that is poor on all tasks,
another subtype is found with specific problems in
about half the tasks, including static balance.

From the above it is clear that postural control
and balance is one of the domains of perceptual
motor performance in which many children with
DCD can be impaired. The present study addresses
the question: in what respect is the control of static
balance different between control children and
children with DCD. As this disorder does not
permit a reliable diagnosis before the age of 5
years, I will concentrate on the age range of 5 to
12 years.

A review of static balance control in children with
DCD

The control of static balance has been studied
in children by using force plate or kinematic
recordings, sometimes in combination with
electromyography (EMG) measures. This approach
has been used for both spontaneous sway in a
standard posture and responses to an external
mechanical or visual perturbation. As a background
for the evaluation of the atypical characteristics of
balance control in children with DCD, a brief
outline of development of static balance in typically
developing children is presented, followed by an
exhaustive review of static balance research in
DCD.

Development of static balance in typically
developing children.

Until the age of 3 years, development of static
balance is characterized by early visual predomi-
nance that gradually gives way to a greater
involvement of somatosensory and vestibular input
up to the age of 10 years, when a more adult like
type of control is reached (Foudriat et al., 1993;
Shumway-Cook & Woollacott, 1985). This
development is paralleled, between 4 and 8 years,
by a decreasing level of EMG activity in most
postural muscles measured in different postures:
lower trapezius, pectoralis major, posterior deltoid,
erector spinae, tibialis anterior, gastrocnemius,
peroneus longus and semitendinosus, but not in
upper trapezius, teres major and rectus abdominis
(Williams et al., 1983). In that age range, with
perturbation of posture a transition is found from
variable to structured response synergies
(Shumway-Cook & Woollacott, 1985, 1990) and
adaptations to sensory conditions improve
(Foudriat et al., 1993). Forssberg and Nashner
(1982) evaluated automatic postural adjustments to
anterior and posterior displacements and rotations
of the support surface and visual compensation for
postural sway from recordings of torque, sway,
and EMG responses of gastrocnemius, tibialis
anterior, hamstrings and quadriceps muscles.
Seventeen children aged 1½ to 10 years participated in the study.
Younger children (aged 1½ to 7½ years, n =
14) qualitatively showed responses similar to those
of adults, even when deprived of specific sensory
input. The children, however, showed greater
variability and could not suppress systematically
the influence of inputs derived from the support
surface or from vision when these provided
inappropriate orientation information, due to the
motion of the platform. For standing on two legs,
measures of postural sway show a rapid decline of
postural sway between 3 and 6 years, and a slower
one up to the age of 11 years (Usui et al., 1995). A
longitudinal study of static balance in children
aged 5 to 10 years revealed a transition around the
age of 6 years in the postural sway velocity of the
centre of pressure, a marker of strategy of reactive
forces to loss of balance (Kirshenbaum et al.,
2001).

For standing on one leg, a fast decline is found
between 6 and 9 years, and a slower one up to the
age of 11 years (Usui et al., 1995). Anticipatory
postural control, preceding voluntary arm
movement while standing, is mature by 4 to 6
years (Shumway-Cook & Woollacott, 1985, 1990).

From this brief overview, it can be concluded
that automatic postural control improves up to the
age of about 10 years, with qualitative changes at
the level of integrated processing of sensory input
around the age of 6 years, and improvement of
dealing with conflicting sensory input up to the
age of 8 years.

POSTURAL SWAY, VISUAL FEEDBACK AND DCD

The problems of balance and postural control
of children with DCD may now be evaluated
against this background of normal development.
Recently a number of studies appeared that
specifically addressed postural control and static
balance in children with DCD (see Table 1).

Forseth and Sigmundsdson (2003) measured the
time to loss of balance in children selected for
hand-eye coordination problems. This group can
be considered a subgroup of DCD. The average
duration of standing on one leg in balance
decreased dramatically when closing the eyes and
when balancing on a beam compared to the floor,
both in the DCD-HECP group (vision vs no-vision
floor 32 vs 9 s, beam 11 vs 3 s) and the control
group (vision vs no-vision floor 50 vs 21 s, beam
24 vs 4 s). The control group was able to maintain
balance considerably longer than the DCD-HECP
group. On the floor with vision, the non-preferred
leg contributed to the group difference only in the
condition stork stand.

Wann et al. (1998) studied postural sway by
recording the head movement of the subjects while
standing with eyes open and with eyes closed. The
investigators did not find an age effect, but
children with DCD (n=6) displayed significantly
more postural sway than matched controls (n=6) in
the condition eyes closed. Comparing vision to no-
vision, the mean peak-peak amplitude in AP
direction increased in 3 control subjects and in 4
children with DCD. Remarkably, the other subjects
(showing a decrease of AP sway without vision)
were those with the largest sway amplitude in the
eyes-open condition.

Geuze (2003) and Przysucha and Taylor
(2004) used force-plate measurements while
standing with eyes open and closed. The authors
specifically selected a subgroup of DCD: children
with DCD and balance problems (DCD-bp). For
quiet standing on two legs, Geuze found only a
slightly increased sway for the DCD-bp group that
was not significant: for lateral and AP directions,
the increase was 4.7% and 6.6% respectively. For
quiet standing on two feet, Przysucha and Taylor
reported slightly increased lateral sway (9%, ns),
and significantly increased AP sway (25%) and
area of COP (42%) in the DCD-bp group. The
Romberg coefficient (sway without vision relative
to sway with vision) was not found to differentiate
between groups in the two studies. Thus, the three
studies indicate equal dependence on vision for the
control of quiet standing with eyes open and eyes
closed.
### TABLE 1

Characteristics of six studies addressing the differences in postural control between children with and without DCD

<table>
<thead>
<tr>
<th>study</th>
<th>N</th>
<th>groups*</th>
<th>age</th>
<th>task /- variables</th>
<th>measures</th>
<th>major sign. problems in DCD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Forseth &amp; Sigmundsson</td>
<td>12</td>
<td>DCD-HECP (estimated &lt;15th perc)</td>
<td>10-11y</td>
<td>one-leg stance</td>
<td>time to loss of balance</td>
<td>control group kept balance longer</td>
</tr>
<tr>
<td></td>
<td>12</td>
<td>non-DCD-HECP (estimated &gt;15th perc)</td>
<td>10-11y</td>
<td>- vision yes/no</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Geuze 2003</td>
<td>24</td>
<td>DCD &lt; 15th perc. + balance problems</td>
<td>6-12 y</td>
<td>stand still</td>
<td>sway lateral &amp; AP</td>
<td>- in 1-leg stance more lateral sway and more</td>
</tr>
<tr>
<td></td>
<td>24</td>
<td>non-DCD &gt; 10th perc</td>
<td>6-12 y</td>
<td>- 1, 2 legs</td>
<td>EMG recovery time</td>
<td>dependence on vision</td>
</tr>
<tr>
<td></td>
<td>16</td>
<td>and 2 age groups: younger group</td>
<td>6-7 y</td>
<td>- vision yes/no</td>
<td></td>
<td>- EMG more co-contraction</td>
</tr>
<tr>
<td></td>
<td>24</td>
<td>older group</td>
<td>10-12 y</td>
<td>- perturbation</td>
<td></td>
<td>- only slight problems in 2-leg stance and</td>
</tr>
<tr>
<td>Johnston et al. 2002</td>
<td>32</td>
<td>DCD &lt; 15th perc. + reference data</td>
<td>8-10 y</td>
<td>stand + arm lift</td>
<td>EMG timing and activation</td>
<td>adaptation to perturbation of stance</td>
</tr>
<tr>
<td></td>
<td>32</td>
<td>non-DCD &gt; 15th perc</td>
<td>8-10 y</td>
<td>- L, R arm</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Przysucha &amp; Taylor 2004</td>
<td>20</td>
<td>DCD &lt; 5th perc. + balance problems</td>
<td>6-11 y</td>
<td>stand still</td>
<td>sway lateral &amp; AP</td>
<td>lack of anticipatory postural muscle activation</td>
</tr>
<tr>
<td></td>
<td>20</td>
<td>non-DCD &gt; 5th perc</td>
<td>6-11 y</td>
<td>- vision yes/no</td>
<td></td>
<td>and poor timing</td>
</tr>
<tr>
<td>Wann et al. 1998</td>
<td>6</td>
<td>DCD &lt; 5th perc. non-DCD nursery adults</td>
<td>10-12 y</td>
<td>stand still</td>
<td>sway AP amplitude</td>
<td>dependence on vision</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td></td>
<td>10-12 y</td>
<td>- vision yes/no</td>
<td>coupling and phase diff.</td>
<td>not different between groups</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td></td>
<td></td>
<td>amplitude of swinging room</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>6</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Williams et al. 1983</td>
<td>6</td>
<td>DCD (mot.awkward) + no motor problems</td>
<td>4,6,8y</td>
<td>static postures</td>
<td>EMG average amplitude</td>
<td>- DCD more postural sway without vision,</td>
</tr>
<tr>
<td></td>
<td>14</td>
<td></td>
<td></td>
<td>- 7 postures</td>
<td></td>
<td>but non-conclusive for entrainment</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>- 12 muscles</td>
<td></td>
<td>- large individual differences</td>
</tr>
</tbody>
</table>

Note: * Percentiles indicate cut-off criterion of the M-ABC test score for group selection. Indicates a group of children selected for hand-eye coordination problems (HECP) selected or a sum score > 10 on 5 items of the M-ABC test.

Wann et al. (1998) addressed the visual dependence more specifically by using a swinging room. When the environment moves slowly (in the frequency range of normal postural corrections, 0.1 to 1 Hz), this movement can entrain postural sway, a phenomenon termed visual-induced sway. Nursery children were found to depend upon vision as a major source of postural information, whereas older children did not. Two DCD children clearly showed postural control problems and visual...
dependence similar to the nursery children. The other four did not differ from the control children.

Compared with quiet standing on two legs, standing on one leg resulted in an increase in AP and lateral sway (Geuze, 2003). Not surprisingly, it is the lateral control of balance differentiating the groups. In the eyes open condition, the DCD group had 35% larger sway in this direction than did the control group. In the eyes-closed condition, this difference increased up to 70% (interaction significant) after selecting the successful epochs when the children were in balance. It should also be noted that children with DCD-bp failed the task significantly more often (i.e. could not maintain one-leg balance) than the control children did.

We can conclude that difficulties in balance control show up more clearly with greater task constraints. Developmental Coordination Disorder is associated with larger postural sway and with failing more difficult balance tasks. For the majority of such children, this problem seems not to be due to greater dependence on vision.

MUSCLE CONTROL IN CHILDREN WITH DCD

Williams et al. (1983) recorded EMG activity of upper and lower trapezius, pectoralis major, teres major, posterior deltoidus, rectus abdominis, erector spinae, semitendinosus, tibialis anterior, soleus, gastrocnemius, and peroneus longus muscles in seven different static postures, such as lying down prone on elbows, pivot prone, on all fours, full kneel, half kneel, normal standing, and standing on one leg. The authors found large individual differences, but on average motorically awkward children displayed greater amounts of muscular activity than did control children matched for age, specifically in the lower trapezius, posterior deltoid, rectus abdominis, gastrocnemius, and peroneus longus muscles. Moreover, the gradual decrease of activation level did not decrease in the atypical children between the age of 4 and 8 years. It should be noted that these results are based on a qualitative evaluation of the recorded EMG, i.e. without calibration of the force-EMG relationship.

Johnston et al. (2002) studied anticipatory postural adjustments in response to voluntary arm movement (lifting the straight arm) to a visual target. The muscles investigated included the anterior deltoid, upper and lower trapezius, serratus anterior, latissimus dorsi around the shoulder girdle, and of the trunk the internal oblique muscles bilaterally, external oblique muscle contralaterally, rectus abdominis, and erector spinae. Children with DCD demonstrated altered muscle timing during voluntary goal-directed pointing, when compared with the non-DCD group. The authors demonstrated that, relative to the activation of the anterior deltoid muscle, children with DCD activated only two of five trunk muscles (contralateral internal oblique, erector spinae) in the anticipatory period (~50 to +150 ms around anterior deltoid onset). All four anterior trunk muscles were later in onset. Shoulder muscles (except serratus anterior) and the posterior trunk muscle, erector spinae, showed earlier activation. Children with DCD were also slower to respond to the visual stimulus and to complete the movement to the goal. The authors conclude that these data support the hypothesis that altered postural muscle activity can contribute to poor proximal stability and consequently to poor upper limb coordination of children with DCD.

Geuze (2003) studied differences in EMG activation patterns between children with and without DCD. In the DCD group, when children were well within balance while standing on one leg, co-contraction between the tibialis anterior and the peroneus muscles involved in controlling the ankle joint occurred 2.2 times more often, and slightly more significant peak activations were found in these muscles. In two-leg stance, when balance was unexpectedly perturbed by a ball lightly hitting the back, no difference between the
groups was found for muscular response onset, amplitude of response, and recovery time over repeated trials. Only the first response was different, with longer recovery times for the DCD group. It is concluded that under normal conditions, static balance control is not a problem for children with DCD. Only in novel or difficult situations are such children at risk for losing balance. It is suggested that an analysis of sway and EMG just before losing balance might shed light on why children with DCD tend to lose their balance sooner and more frequently than control children do.

**Losing balance: correlational analysis between EMG and force plate responses**

Epochs of stable and unstable balance of children with and without DCD and balance problems were compared on balance control. Children were selected from three primary schools. Teachers selected one or two children who, in the teacher's opinion, were worst in the class in general motor skills and who might have balance problems. The teachers were asked to check their choice with the physical education (PE) teacher if the school had one. For the control group, one or two children from the same class were selected by random choice (see Geuze 2003 for details). The unaffected children consisted of a younger group (n = 12; m/f 8/4) aged 6.0 to 8.3 years and an older group (N = 14; m/f 7/7) aged 9.1 to 11.6 years. From these groups, a control group was selected (n = 13; m/f 9/4) with an age range of 6 to 11 years (mean age 9.0 years) that was matched for age and nearly for gender with the children with DCD-bp (n = 13; m/f 10/3). None had severe learning difficulties, from which it may be assumed that the IQ was over 80. All children participated in the study of Geuze (2003).

The motor performance of the children was tested at school with the Dutch version of the Movement-ABC test (Henderson & Sugden, 1993; Smits-Engelsman, 1998). The selection criteria for the children with DCD and balance problems (DCD-bp) were (i) M-ABC score ≤ 15th centile; (ii) M-ABC balance subscore > 2; (iii) M-ABC static balance score > 1. The average M-ABC score for the control group was 4.5 (range 1.5–8) and for the DCD-bp group 15.6 (range 8.5–30.5). A typical example of the recordings and displacement of the center of pressure is shown in Fig. 1.

Control of balance when standing still on one leg for 10–30 s was analyzed by cross-correlation of the lateral and AP force plate signals (Fx and Fy) with the muscle activation of the tibialis anterior, peroneus, rectus femoris and semitendinosus muscles. The main function of the peroneus muscle in one-leg stance is to control AP sway, but the muscle also has a lateral component. The tibialis anterior muscle is mainly involved in the control of lateral sway, but has also a small AP component. rectus femoris flexes the hip and semitendinosus flexes the shank and retracts the thigh. The EMG's were filtered (60-1000 Hz) and converted into a true RMS signal (0-35 Hz). Force plate signals were filtered low-pass at 10.5 Hz. These signals were sampled at 100 Hz.

From each recording, one epoch of 10 s was selected, usually from 5 to 15 s after the beginning, and 3 further epochs of 1.5 s. These were epochs when the child was (1) in balance; (2) unstable in balance; (3) close to losing balance; as evident from the path of the Center of Pressure (CoP) in this epoch. The cross-correlation between EMG and force plate signals was calculated over these 4 epochs.

Cross-correlation between EMG and force signals should incorporate delays from the electromechanical coupling in the muscle (typically 50 ms) and transfer of force from the muscle through soft tissues to the force plate. These two were estimated together from adult reactive responses to voluntary loss of balance in AP or in lateral direction to be 50 to 60 ms. With children being slower (70 to 80 ms, Raynor, 1999) the delay time for the analysis was set at 80 ms. Eventual perceptual-motor delay of the
Fig. 1A: Two epochs of static balance of a 9-year-old boy with DCD and balance problems. Original recordings (left) show EMG’s of Tibialis anterior and Peroneus longus muscles and the reactive forces exerted on the forceplate in anterior-posterior (x) and in lateral (y) directions. On the right the displacement of the Center of Pressure (COP) as a measure of postural sway. Top: stability of balance; bottom: balance is lost at the arrow. Vertical scale in arbitrary units.
Fig. 1B: Same for matched control child. Balance is nearly lost at the arrow.
TABLE 2

Cross-correlation (p< .01 for values > .20 and p< .001 for values > .26) between EMG and force plate signals for 1.5s epochs (150 samples) of stable balance (top) and near loss of balance (bottom) for the lower leg muscles and upper leg muscles. Bold figures indicate major group differences.

<table>
<thead>
<tr>
<th>stable</th>
<th>stable</th>
<th>Peroneus</th>
<th>Rectus Femoris</th>
<th>Semitendinosus</th>
</tr>
</thead>
<tbody>
<tr>
<td>groups</td>
<td>N</td>
<td>Fy</td>
<td>Fx</td>
<td>Fy</td>
</tr>
<tr>
<td>6-8 years</td>
<td>12</td>
<td>0.25</td>
<td>0.48*</td>
<td>0.21</td>
</tr>
<tr>
<td>9-11 years</td>
<td>14</td>
<td>0.28</td>
<td>0.57*</td>
<td>0.22</td>
</tr>
<tr>
<td>DCD-bp</td>
<td>13</td>
<td>0.37</td>
<td>0.41**</td>
<td>0.25</td>
</tr>
<tr>
<td>Controls</td>
<td>13</td>
<td>0.33</td>
<td>0.57**</td>
<td>0.22</td>
</tr>
</tbody>
</table>

<table>
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<tr>
<th>near loss</th>
<th>near loss</th>
<th>Tibialis</th>
<th>Peroneus</th>
<th>Rectus Femoris</th>
<th>Semitendinosus</th>
</tr>
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<tbody>
<tr>
<td>groups</td>
<td>N</td>
<td>Fx</td>
<td>Fy</td>
<td>Fx</td>
<td>Fy</td>
</tr>
<tr>
<td>6-8 years</td>
<td>12</td>
<td>0.25</td>
<td>0.43</td>
<td>0.23</td>
<td>0.36**</td>
</tr>
<tr>
<td>9-11 years</td>
<td>14</td>
<td>0.27</td>
<td>0.45</td>
<td>0.20</td>
<td>0.48**</td>
</tr>
<tr>
<td>DCD-BP</td>
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<td>0.24**</td>
<td>0.40*</td>
<td>0.27</td>
<td>0.36**</td>
</tr>
<tr>
<td>Controls</td>
<td>13</td>
<td>0.35**</td>
<td>0.48*</td>
<td>0.21</td>
<td>0.50**</td>
</tr>
</tbody>
</table>

^ these groups share 11 subjects. Levels of significant differences between younger and older group, or between DCD-BP and control group: * p<.1, ** p<.05, *** p<.01

TABLE 3

Percentage of subjects with significant (p < 0.01) correlations for the 1.5s epochs of stable balance (left) and close to loss of balance (right). Bold indicates group difference > 20%.

<table>
<thead>
<tr>
<th>stable</th>
<th>stable</th>
<th>Peroneus</th>
<th>near loss</th>
<th>Peroneus</th>
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</thead>
<tbody>
<tr>
<td>groups</td>
<td>N</td>
<td>Fx</td>
<td>Fy</td>
<td>Fx</td>
</tr>
<tr>
<td>6-7 years</td>
<td>12</td>
<td>45</td>
<td>36</td>
<td>64</td>
</tr>
<tr>
<td>9-11 years</td>
<td>14</td>
<td>71</td>
<td>43</td>
<td>71</td>
</tr>
<tr>
<td>DCD-bp</td>
<td>13</td>
<td>69</td>
<td>54</td>
<td>69</td>
</tr>
<tr>
<td>Controls</td>
<td>13</td>
<td>77</td>
<td>46</td>
<td>77</td>
</tr>
</tbody>
</table>

^ these groups share 11 subjects
feedback loops, that may range from 30 ms for the M1 reflex to a few hundreds of ms for higher level processing, was not considered in this analysis. Table 2 summarizes the outcome of the analysis.

Strong correlations are found in each group for lateral control. Older subjects compared with young ones show on average a stronger lateral control for the tibialis, peroneus, and semitendinosus muscles. The children with DCD show a weaker lateral control for the tibialis, peroneus, and upper leg muscles, indicating less efficient control. In contrast, older subjects did show an appropriate reduction of tibialis involvement in the control of Fx when balance is at risk. In the more unstable situation, their correlation pattern resembles that of younger children.

Over a 10s period, all groups showed a significant correlation (p < .0001, r > .326) between tibialis and peroneus activation and Fy. In the DCD-bp group, this correlation was significant for 81% of the subjects, whereas for the other groups this was over 90%. The overall percentage of subjects having a significant correlation (p < 0.1) with Fx was 25%—31% for the DCD-bp and younger groups and 14% for the other groups. Table 3 lists the number of subjects with significant (p <. 01) correlations for the 1.5s epochs.

In the DCD-bp group, the lower correlation strength between EMG-activation and reactive forces in balance control could be due to an inconsistency in the timing of muscular activation in balance control.

STRUCUTURAL DEFICITS ASSOCIATED WITH DCD?

The cerebellum is important for movement control and plays a particularly crucial role in balance and locomotion (Morton & Bastian, 2004). This role might point at cerebellar involvement in the motor problems of DCD. The motor problems of children with DCD are varied, however, and without further evidence, such a conclusion is premature.

What evidence is there for a cerebellar deficit that contributes to DCD? Three possible sources of information can help to clear a question like this.
1. Direct evidence from fMRI or electrophysiological studies. These are not available yet.
2. Indirect evidence from studies that associate DCD with functional measures assumed to be related to cerebellar function, such as the balance studies reported here.

Among the studies that associate DCD with functional measures assumed related to cerebellar function are those using developmental neurological screening tests. Lundy-Ekman et al. (1991) selected two subgroups of children with DCD: a subgroup of children with cerebellar soft neurological signs and a subgroup with soft basal ganglia signs of dysfunction. The children and their controls participated in tasks of timing and force control. The first subgroup performed poorly specifically on timing tasks—timing is assumed a cerebellar function, whereas the second subgroup performed poorly on the force control task, assumed related to basal ganglia function. Volman and Geuze (1998) applied the same neurodevelopmental test (Touwen's test) to divide their group of 24 children with DCD into cerebellar and basal ganglia subgroups. We did not succeed, however, because the distribution of soft neurological signs did not allow classification of the children with DCD into reliable subgroups of cerebellar or basal ganglia dysfunction. Recently O'Hare and Khalid (2004) investigated the association of abnormal cerebellar function in children with DCD and reading difficulties. In a group of 23 children attending pediatric occupational therapy and diagnosed as DCD all showed soft signs of cerebellar dysfunction. However, according to their Table 4 the mean score of 14 out of 15
components of the Quick Neurological Screening Test is over the upper limit of 'suspicious' score, with only two components that clearly tap cerebellar function. It seems to me that, although cerebellar soft signs may be present in all children, this is by no means a specific finding, as the many other soft neurological signs indicate the involvement of varied neurological structures.

It should be kept in mind that the perceptual-motor system operates as a distributed network in which the cerebellum plays a functional role. Evidence that the cerebellum is involved in timing processes is mainly derived from studies on finger tapping. Ivy and Keele (1989) reported timing deficits in patients with cerebellar lesions. Rivkin et al. (2003) employed fMRI to study specific activation patterns in the brain during alternating finger tapping in a group of 14 children aged 7.9 to 11.3 years. This timing task activated a neural network involving the primary motor cortex, the supplementary and presupplementary motor area, and parts of the cerebellum. The fMRI data of children with DCD are not available.

Visser (2003, 2005) put forward indirect evidence from disorders comorbid to DCD, such as dyslexia (Fawcett & Nicolson, 1995; Nicolson & Fawcett, 2001). For example, co-morbidity was reported in a study of children attending pediatric occupation therapy and diagnosed as DCD, parents reporting reading problems in 70% of cases (O'Hare & Khalid, 2002). Visser argues that problems of automatization are common in comorbid disorders of dyslexia and DCD, and that this indicates cerebellar involvement in these disorders. Motor problems and abnormalities in muscle tone are common symptoms in the majority of subjects with dyslexia (Nicolson et al., 2001). A major finding reported is an automatization deficit of balance control (Fawcett & Nicolson, 1992; Yap & Van der Leij, 1994). Such a deficit will become apparent during the performance of a motor task concurrent with a second non-motor task. According to this dual-task paradigm, a decrease in performance compared with the performance of the single-task condition indicates a lack of automatization of the primary (balance) task. The findings led Fawcett and colleagues to conclude that dyslexia is caused by a general deficit in the ability to automatize fully the skills that affect both reading proficiency and automatized motor skill. Nicolson and Fawcett relate the lack of automatization to cerebellar dysfunction.

CONCLUSION

Many children with DCD show poor postural and balance control, especially in extremely difficult situations. The characteristics of this poor control are likely to be task dependent—especially task difficulty and the availability of sensory information will influence the quality of postural and balance control. The major characteristics of poor control in DCD are an inconsistent timing of muscle activation sequences, co-contraction, a lack of automatization, and slowness of response. Converging evidence indicates that cerebellar dysfunction contributes to the motor problems of children with DCD.

REFERENCES


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