Cerebellar-dependent delay eyeblink conditioning in adolescents with Specific Language Impairment

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Abstract Cerebellar impairments have been hypothesized as part of the pathogenesis of Specific Language Impairment (SLI), although direct evidence of cerebellar involvement is sparse. Eyeblink Conditioning (EBC) is a learning task with well documented cerebellar pathways. This is the first study of EBC in affected adolescents and controls. 16 adolescent controls, 15 adolescents with SLI, and 12 adult controls participated in a delay EBC task. Affected children had low general language performance, grammatical deficits but no speech impairments. The affected group did not differ from the control adolescent or control adult group, showing intact cerebellar functioning on the EBC task. This study did not support cerebellar impairment at the level of basic learning pathways as part of the pathogenesis of SLI. Outcomes do not rule out cerebellar influences on speech impairment, or possible other forms of cerebellar functioning as contributing to SLI.

Keywords Specific Language Impairment · Cerebellum · Eyeblink conditioning · Conditioned learning · Adolescence

Specific Language Impairment (SLI) is diagnosed in children with language impairments without an obvious cause (i.e. hearing loss, cognitive or physical impairment). The overall prevalence rate for SLI in epidemiologically ascertained samples of healthy children is 7.4% (8% for boys and 6% for girls) (Tomblin et al. 1997). Grammatical deficits are widely recognized as persistent clinical markers of SLI (Rice et al. 2009a, 1995; Rice and Wexler 1996; Rice 2000; Tager-Flusberg and Cooper 1999). Deficits in motor speech production are relatively independent of SLI, appearing in 5–8% of young affected children although clinically ascertained samples can have a higher percentage (Shriberg et al. 1999). Group comparisons of affected and age-matched control children reveal subtle differences in performance on motor tasks (Bishop 2002; Zubrick et al. 2007; Zelaznik and Goffman 2010), longer latencies on perceptual/cognitive tasks (Leonard et al. 2007), and possible attentional deficits (Lum et al. 2007).

Children with SLI show overlap with other developmental disabilities. Two are of interest here. One is reading impairment. Studies yield an estimate of 50% of young children with SLI who subsequently develop reading impairments (Catts 2004). The language symptoms of SLI are also evident in children with Autism Spectrum Disorders (ASD). Recent studies report similar grammar phenotypes of children with ASD and children with SLI (Roberts et al. 2004; Kjelgaard and Tager-Flusberg 2001). A recent study of children with a history of SLI found the prevalence of ASD in this sample higher than that of the general population (Conti-Ramsden et al. 2006).
The pathogenesis of SLI is unknown. The biological underpinnings are attracting new attention as candidate genes are discovered for SLI, such as KIAA 0319 for several language and reading phenotypes (Rice et al. 2009b) and for ASD, such as CNTNAP2, based on a phenotype of late language emergence during the toddler period (Alarcón et al. 2008). In turn, CNTNAP2 polymorphisms in children with SLI showed quantitative associations with language-related behavioral phenotypes in a sample of children with SLI (Vernes et al. 2008). Further, the region of association coincides with a region reported for language delays in children with autism (Alarcón et al. 2008). Both KIAA 0319 and CNTNAP2 are expressed in the developing human cortex, pointing toward pathways in brain functioning as important elements in the pathogenesis of language impairments. On the other hand, a recent investigation of the first degree relatives of children with SLI and children with ASD did not find support for similar genetic loading for language (Lindgren et al. 2009).

Recent theories posit a role for the cerebellum in the pathogenesis of SLI (Hill 2001), with a particular emphasis on subtle impairments in motor output and attentional processes as concomitant indicators of possible cerebellar involvement. The cerebellum has been thought to be solely a part of motor control, but more recently it has been implicated in cognition (Rapoport et al. 2000), selective attention (Akshoomoff and Courchesne 1992; Yamaguchi et al. 1998), and language skills (Leiner et al. 1991), including morphological deficits in adults with cerebellar injuries (Justus 2004). The complex interconnections of the cerebellum with frontal cortical processes warrants more investigation of the cerebellum’s effects on children’s development (Diamond 2000). A cerebellar deficit hypothesis, based in the underlying neural substrate, has been proposed for developmental dyslexia (Nicolson et al. 2001), in which cerebellar impairment is predicted to affect several pathways of behavioral development. One direct manifestation of cerebellar impairment is predicted to be impaired articulatory skill which in turn leads to reading impairments. Another hypothesized pathway is motor skill impairment and/or balance impairment that leads to writing impairments, and a third pathway predicts that problems with automatising skill and knowledge lead to reading and spelling problems.

The current theoretical models of possible impaired cerebellar functions for SLI or dyslexia are rather sketchy in nature, invoking a relatively wide range of related symptoms and an unspecified locus of cerebellar dysfunction (see a recent paper by Nicolson and Fawcett (Nicolson and Fawcett 2007) for an over-arching conceptual framework and discussion of some of the issues in differentiating the functions of components of cortical and cerebellar functioning). In contrast, there is evidence of a precise dysfunction in cerebellar processing in children with autism (Sears et al. 1994), utilizing classical eyeblink conditioning (EBC) as an index of basic learning pathways linked to a motor response. This experimental task has been widely used to probe cerebellar function in humans and nonhuman mammals.

In the most common form of the EBC procedure, termed single-cue delay EBC, a conditioned stimulus (CS; e.g., 400 ms tone) is paired with a co-terminating unconditioned stimulus (US; e.g., 50 ms corneal airpuff). Repeated CS-US pairings elicit the development of an adaptive conditioned blink response (CR), which occurs in healthy individuals just prior to the US. The delay EBC procedure appears to be the purest assay of the functional integrity of the cerebellum, with the circuitry and synaptic mechanisms being well studied and identified in nonhumans (Kim and Thompson 1997; Steinmetz 2000; Christian and Thompson 2003) compared to other forms of EBC, such as trace conditioning (Christian and Thompson 2003). In addition, the cerebellar networks mediating this form of associative learning appear to be conserved across mammals (rat: (Rogers et al. 2001); human: (Gerwig et al. 2007)). The single-cue delay procedure has been used to examine developmental and clinical conditions associated with cerebellar abnormalities and motor disruptions, such as aging (e.g., (Woodruff-Pak and Thompson 1988; Woodruff-Pak et al. 1999)), autism(Sears et al. 1994), schizophrenia (e.g., (Brown et al. 2005)) and drug abuse (Skosnik et al. 2008). In their study of children with autism ages 7–22 years, Sears et al. (1994) report that affected children differed from controls with faster rates of conditioning at a young age and maintained that rate as age increased whereas control subjects showed increasingly fast rates of conditioning with age. In addition, the autism group showed a more rapid and significant decline in the amplitudes of conditioned responses during an extinction phase. The topography of the conditioned responses of the autism group differed from the control group. Overall, the patterns of motor learning associated with cerebellar functioning differentiated the children with autism from the control group.

Somewhat different methods for evaluating the EBC were used in a study of 13 adolescents and young adults with dyslexia, ages 13–24 years, and 13 same-age controls (Nicolson et al. 2002). The conclusion was that the dyslexic group performed poorly relative to the control group, showing either no conditioning or poor timing of CRs and/or abnormally low orienting responses. The general conclusion of the study is that the outcomes provide evidence of fundamental differences in the way people with dyslexia learn.
In sum, the underlying cerebellar pathways involved in the EBC have been mapped in great detail, with converging evidence from studies of humans and other mammals. Research during the past 25 years strongly supports the role of the cerebellar cortex and deep nuclei in the acquisition and timing of the CR during delay EBC. In relation to learning and behavioral occurrence of the CR, lesion, neural unit recording, and reversible inactivation studies have provided compelling evidence that the memory trace for delay EBC resides in regions of the cerebellar deep nuclei (anterior lateral interpositus nucleus ipsilateral to the trained eye; for review see (Christian and Thompson 2003; Steinmetz 2000)). While it is difficult to assess the effect of interpositus lesions on EBC performance in humans (since in many clinical cases, the cerebellar cortex is also affected; (McGlinchey-Berroth et al. 1995; Topka et al. 1993; Daum et al. 1993; Solomon et al. 1989; Lye et al. 1988), it has been suggested that most of the patient studies demonstrating delay EBC deficits after cerebellar insult include damage to the deep nuclei (Schugens and Daum 1999).

While evidence clearly suggests that the cerebellar deep nuclei are involved in the acquisition of the CR, several studies have also identified a possible role of the cerebellar cortex in the timing and gain of CRs (Lavond and Steinmetz 1989; McCormick and Thompson 1984; Logan 1991; Perrett et al. 1993). In humans, the involvement of the cerebellar cortex in modulating the CR has been supported using positron emission tomography (Blaxton et al. 1996; Logan and Grafton 1995), functional MRI (Ramnani et al. 2000; Dimitrova et al. 2002) and clinical samples with cerebellar insults (Daum et al. 1993; Woodruff-Pak et al. 1996). More recently, it has been demonstrated that in cerebellar patients, damage to the anterior lobe was significantly correlated with timing deficits during delay EBC as assessed via MRI (Gerwig et al. 2005). One possible conclusion that can be drawn from these studies is that a dynamic interaction between the cerebellar cortex and deep nuclei underlies both the acquisition and the accurate timing of the CR, with the cortex potentially playing a crucial role in modulating the timing of the response.

Studies of children with autism document a pattern of EBC different in the affected versus control children. Although the language abilities of the children in the previous study (Sears et al. 1994) were not described in the research report, the outcomes suggest that the EBC is a candidate for a cerebellar pathway that could be impaired in children with language impairments, to the extent that children with SLI and children with autism share underlying brain pathways for language impairments. Further, previous studies of children and young adults with dyslexia (Nicolson et al. 2002) and dyslexia subsequent to prenatal alcohol exposure (Coffin et al. 2005) report impaired EBC for the affected group. The study reported here addresses the need for a study using the delay eyblink conditioning paradigm to test the functional integrity of the cerebellum in subjects diagnosed with SLI. This paradigm consisted of both an acquisition and extinction period to examine the learning and un-learning of the response, following important precedents in the literature that document the full pattern of motor response learning. The hypotheses were that the children with SLI would exhibit fewer CRs than controls (assuming subtle motor learning impairments) and the timing of the CR should also be impaired with worse timing, as in the previous study of children with autism. The participants in this study did not have speech impairments, providing an opportunity to investigate whether the hypothesis of cerebellar impairment is supported in a sample of children with language impairments without speech impairments.

**Methods**

**Participants**

There were a total of 43 participants, four groups of children (N=31) and one adult control group (N=12). The children were grouped as follows: 6 young controls (age range=9.1–11.11, mean=10.1, s.d.=1.1), 10 older controls (age range=17.3–19.9; mean age=18.30, s.d.=0;11), 5 young SLI (age range=9.7–12.8; mean age=11;7, s.d.,=1;5), 10 older SLI (age range=14.6–19.9; mean=17;0, s.d.=2.1) Collapsed across ages, there were 11 males in the control groups and 12 males in the SLI group. All children were participants in an ongoing longitudinal study documenting long-term language growth patterns. Children in the SLI group entered the longitudinal study as affected children, initially identified by clinical speech/language pathologists. The children were experimentally screened for speech impairments. The criterion was defined by a passing score on a probe screening for articulation competency with consistent use of final –t, -d, and –z (Rice and Wexler 2001) and only minor mispronunciations, such as distortions of r/,/l/ and blends, on the Goldman-Fristoe Test of Articulation-2 (GFTA-2) (Goldman and Fristoe 1986). The hearing criterion at entry was passage of a screening at 25 dB HL at 1,000, 2,000, and 4,000 Hz. Children with diagnoses of autism or autism spectrum disorders were excluded, as were children with nonverbal intelligence levels below 85 on the Columbia Mental Maturity Scales (CMMS) (Burgemeister et al. 1972) or the Wechsler Intelligence scale for Children (WISC): Nonverbal Scale (Wechsler 2001) and only minor mispronunciations, such as distortions of r/,/l/ and blends, on the Goldman-Fristoe Test of Articulation-2 (GFTA-2) (Goldman and Fristoe 1986). There were a total of 43 participants, four groups of children (N=31) and one adult control group (N=12). The children were grouped as follows: 6 young controls (age range=9.1–11.11, mean=10.1, s.d.=1.1), 10 older controls (age range=17.3–19.9; mean age=18.30, s.d.=0;11), 5 young SLI (age range=9.7–12.8; mean age=11;7, s.d.,=1;5), 10 older SLI (age range=14.6–19.9; mean=17;0, s.d.=2.1) Collapsed across ages, there were 11 males in the control groups and 12 males in the SLI group. All children were participants in an ongoing longitudinal study documenting long-term language growth patterns. Children in the SLI group entered the longitudinal study as affected children, initially identified by clinical speech/language pathologists. The children were experimentally screened for speech impairments. The criterion was defined by a passing score on a probe screening for articulation competency with consistent use of final –t, -d, and –z (Rice and Wexler 2001) and only minor mispronunciations, such as distortions of r/,/l/ and blends, on the Goldman-Fristoe Test of Articulation-2 (GFTA-2) (Goldman and Fristoe 1986). The hearing criterion at entry was passage of a screening at 25 dB HL at 1,000, 2,000, and 4,000 Hz. Children with diagnoses of autism or autism spectrum disorders were excluded, as were children with nonverbal intelligence levels below 85 on the Columbia Mental Maturity Scales (CMMS) (Burgemeister et al. 1972) or the Wechsler Intelligence scale for Children (WISC): Nonverbal Scale (Wechsler 2001) and only minor mispronunciations, such as distortions of r/,/l/ and blends, on the Goldman-Fristoe Test of Articulation-2 (GFTA-2) (Goldman and Fristoe 1986). The hearing criterion at entry was passage of a screening at 25 dB HL at 1,000, 2,000, and 4,000 Hz. Children with diagnoses of autism or autism spectrum disorders were excluded, as were children with nonverbal intelligence levels below 85 on the Columbia Mental Maturity Scales (CMMS) (Burgemeister et al. 1972) or the Wechsler Intelligence scale for Children (WISC): Nonverbal Scale (Wechsler 2001) and only minor mispronunciations, such as distortions of r/,/l/ and blends, on the Goldman-Fristoe Test of Articulation-2 (GFTA-2) (Goldman and Fristoe 1986). The hearing criterion at entry was passage of a screening at 25 dB HL at 1,000, 2,000, and 4,000 Hz. Children with diagnoses of autism or autism spectrum disorders were excluded, as were children with nonverbal intelligence levels below 85 on the Columbia Mental Maturity Scales (CMMS) (Burgemeister et al. 1972) or the Wechsler Intelligence scale for Children (WISC): Nonverbal Scale (Wechsler 2001) and only minor mispronunciations, such as distortions of r/,/l/ and blends, on the Goldman-Fristoe Test of Articulation-2 (GFTA-2) (Goldman and Fristoe 1986).
Woodcock Reading Mastery Tests-Revised (Woodcock 1987), Word Identification subtest. As summarized in Table 1, the archival data files show that at initial testing the group of children with SLI were equivalent to the control children in total number of articulation errors but at significantly lower levels of performance on omnibus language assessment, vocabulary, and word identification levels. The ages of the children at initial testing for each measure are provided in Table 1. They were somewhat older at initial testing for reading, given the ages at which reading instruction started in the schools. Also summarized in Table 1 are assessments concurrent with the eyeblink data collection. As at the initial testing, the two groups do not differ on total number of articulation errors. The concurrent assessments confirm the SLI group continues to perform lower on omnibus language assessment, vocabulary (although the SLI group’s mean vocabulary performance was within normal range), and word identification. Further, persistent grammatical deficits were apparent on an experimental grammaticality judgment task administered at the time of the experiment. Control children were more likely to detect the ungrammaticality of sentences such as *Where is the dog playing?* or *Is the dog playing?* Thus, the SLI group’s language performance was consistently low over time, although the mean level of speech production did not differentiate the groups at outset in the larger parent study when they were much younger, nor at the time of the eyeblink data collection. Also, as expected, the SLI group’s mean performance on a word identification task documents a persistent risk for reading impairment. Because the study was focused on detailed documentation of speech, language, and reading acquisition, no measures of motor performance were included in the longitudinal protocol. In addition, 12 adult controls (mean age=25.67, s.d. 2.05) were recruited via fliers from the local community. Prior to the experimental task, all participants completed an audiology pure tone screening at 500 Hz, 1,000 Hz, 2,000 Hz, and 4,000 Hz; criterion for participation was defined as passing at 25–30 dB in at least one ear for each of the frequencies.

**EBC stimuli and procedure**

Participants completed a 133-trial delay EBC paradigm similar to (Sears et al. 1994) and (Brown et al. 2005). Initially, eight US alone trials were presented with an intertrial interval (ITI) of 15 s. Without interruption, the

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**Table 1** Speech, language, and reading mean scores and standard deviations per group

|                      | SLI Mean | SLI SD | Control Mean | Control SD | p value | df | Cohen’s d effect size |
|----------------------|----------|--------|--------------|------------|---------|----|-----------------------|
| **Initial testing**  |          |        |              |            |         |    |                       |
| GFTA total errors    | 10.20    | 7.61   | 6.44         | 6.86       | 0.158   | 29 | 0.55                  |
| GFTA age             | 6, 11    | 1, 6   | 4, 7         | 1, 7       |         |    |                       |
| Omnibus languagea    | 73.53    | 7.76   | 119.69       | 17.27      | 0.000   | 29 | 3.81                  |
| Omnibus age          | 6, 9     | 1, 7   | 4, 7         | 1, 7       |         |    |                       |
| Vocabularyb          | 83.87    | 15.47  | 106.44       | 12.22      | 0.000   | 29 | 1.85                  |
| Vocabulary age       | 6, 9     | 1, 7   | 4, 7         | 1, 7       |         |    |                       |
| Word ID              | 83.93    | 11.41  | 104.94       | 12.30      | 0.000   | 29 | 1.71                  |
| Word ID age          | 7, 7     | 1, 0   | 5, 11        | 1, 0       |         |    |                       |
| **Concurrent testing** |         |        |              |            |         |    |                       |
| GFTA total errors    | 2.07     | 1.71   | 2.06         | 2.65       | 0.996   | 29 | 0.00                  |
| Omnibus languagec    | 83.33    | 15.38  | 107.50       | 10.33      | 0.000   | 29 | 2.34                  |
| Grammar judgements   | 0.82     | 0.16   | 0.97         | 0.04       | 0.001   | 29 | 3.75                  |
| Vocabularyd          | 97.67    | 9.48   | 111.69       | 8.28       | 0.000   | 29 | 1.69                  |
| Word ID              | 81.47    | 13.63  | 103.81       | 11.79      | 0.000   | 29 | 1.89                  |

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a Omnibus language score is the summative standard score for Test of Early Language Development (TELD-2 or TELD-3), Test of Language Development-Primary 2nd Edition (TOLD P:2), or Clinical Evaluation of Language Fundamentals 3rd edition (CELF-3)

b Vocabulary score is the summative standard score for the Peabody Picture Vocabulary Test-R (Revised) or Peabody Picture Vocabulary Test-3

c Omnibus language score is the summative standard score for Clinical Evaluation of Language Fundamentals 3rd Edition (CELF-3)

d Vocabulary score is the summative standard score for the Peabody Picture Vocabulary Test-3
acquisition phase immediately followed and consisted of 10 trial blocks (mean ITI=15 s, range=10–20 s), each containing 10 CS-US paired trials. Paired CS-US trials consisted of a 400 ms, 1,000 Hz tone (80 dB SPL) with a co-terminating 50 ms air puff. The procedure concluded with an extinction phase consisting of 25 CS alone trials presented across blocks (mean ITI=15 s; range=10–20 s). To maintain attention throughout the procedure, participants watched the movie Milo and Otis with no sound. Subjects were observed through a closed circuit monitor in order to observe alertness. The experiment was briefly suspended if signs of fatigue were observed so that the examiner could interact with the participant.

Eyeblinks were recorded using pairs of electromyo- graphic (EMG) electrodes (8 mm Ag/AgCl; Model TD-23; MedAssociated, St Albans, VT). Bipolar recording electrodes, with conductive gel applied, were placed on the orbicularis palpebrarum muscle below the left eye. A ground electrode was placed on the forehead. All electrode impedances were maintained below 10 kΩ. The US consisted of a 10 p.s.i. (50 ms duration) puff of medical-grade air presented to the left eye with tubing affixed to eye-glass rims and positioned 1 cm away from the inner canthus of the eye. Foam ear inserts were used for presentation of the CS tones (E-A-RLINK, Aearo Company Auditory Systems, Indianapolis, IN). EMG data were continuously recorded at 1.0 kHz with a Sensorium EPA-6 bioamplifier (highpass filter=1 Hz, 12 dB/octave; lowpass filter=300 Hz, eighth order elliptic; gain=5,000) and acquired using Brainvision software (v. x.x, Richardson, Texas).

EBC data processing

Individual trials were epoched from the continuous EMG data file (using Brainvision Edit software), and filtered (10 Hz high pass filter; 6 dB/octave) before being rectified and smoothed using a 41-point Gaussian weighted moving average. Data were then entered into the Data-Munch software program for further analysis (King and Datamunch 1999). For each subject, responses were recorded as blinks if the amplitude exceeded five standard deviations above the baseline (baseline window for each trial=125 ms before CS presentation). CRs were recorded if the blink occurred between 100 and 350 ms after CS onset (corresponding to a period beginning 250 ms before US onset). The onset latency was calculated as the point in time where the conditioned response exceeded 5.0 standard deviations from the baseline. The peak latency is the time point for the maximal value for that conditioned response. Trials in which spontaneous blinks occur within a window from 75 ms before CS presentation to 25 ms following CS onset were labeled bad trials and excluded from further analysis.

Statistical analysis

The primary dependent measures for the eyeblink procedure were percentage of CRs, CR onset latency, CR peak latency, CR amplitude, and UR amplitude. For each of these variables, the majority of the analyses were conducted with a mixed model ANOVA with two levels of a group factor (i.e. control, SLI) and 10 levels of a within-subjects block factor. The group factor was a between-subjects factor when within-session effects were examined whereas it was a within-subjects factor for analyses across session. An analogous analytical model was used to examine extinction data, except that the block factor had only 5 levels. For the primary dependent variables, effect sizes are reported by partial η², where small effect sizes are less than 0.06, moderate effect sizes range from 0.06 to 0.14, and large effect sizes are greater than 0.14 (Cohen 1973). All statistical tests used an alpha level of p<0.05 to determine significance (two-tailed). If Mauchly’s Test of Sphericity was significant, Greenhouse-Geisser was used to determine significance. All tests were performed using the software package SPSS 14.0.

Results

Age effects

Young controls (age 9–12), older controls (Catts 2004; Roberts et al. 2004; Kjelgaard and Tager-Flusberg 2001; Conti-Ramsden et al. 2006; Rice et al. 2009b; Alarcón et al. 2008; Vernes et al. 2008) and adult controls (Akshoomoff and Courchesne 1992; Yamaguchi et al. 1998; Leiner et al. 1991; Justus 2004; Diamond 2000; Nicolson et al. 2001; Nicolson and Fawcett 2007; Sears et al. 1994) were compared in order to examine possible learning effects of age. A repeated measures ANOVA for percentage of CRs during the acquisition phase revealed a main effect of block, F(9, 17)=3.12 p=0.02; partial η²=.62, but no significant main effect of age (partial η²=.02) and no block×age interaction (partial η²=.40). The extinction phase revealed a main effect of block, F(4, 22)=8.81 p<0.001; partial η²=.62, but no significant main effect of age (partial η²=.04) and no block×age interaction (partial η²=.16). For CR peak latencies there was a main effect of block, F(9, 17)=4.42 p=0.001; partial η²=.15, but no main effect of age (partial η²=.11) or block×age interaction (partial η²=.06). No statistical effects were revealed with regards to CR onset, CR amplitude, UR amplitude, or during the extinction phase. In summary, the conditioned responses were learned at a similar rate and did not differ between the age groups; timing and amplitudes also did not differ between the groups. For further analysis, the young
controls and the older controls were combined to form one control group.

In order to examine possible age effects on learning, younger SLI participants (age 9–12) were compared to older SLI participants (Catts 2004; Roberts et al. 2004; Kjelgaard and Tager-Flusberg 2001; Conti-Ramsden et al. 2006; Rice et al. 2009b; Alarcón et al. 2008; Vernes et al. 2008). A repeated measures ANOVA for percentage of CRs during acquisition revealed a main effect of block, $F(5, 9)=3.12$, $p=0.05$; partial $\eta^2=.62$, but no significant main effect of age (partial $\eta^2=.01$) and no block×age interaction (partial $\eta^2=.15$). The extinction phase revealed a main effect of block, $F(4, 10)=3.09$, $p=0.049$; partial $\eta^2=.61$, but no significant main effect of age (partial $\eta^2=.10$) and no block×age interaction (partial $\eta^2=.20$). For CR peak latencies there was a main effect of block, $F(5, 9)=14.35$, $p=0.005$; partial $\eta^2=.96$, but no main effect of age (partial $\eta^2=.80$) or block×age interaction (partial $\eta^2=.56$). No statistical effects were revealed with regards to CR onset, CR amplitude, UR amplitude, or during the extinction phase. In summary, the conditioned responses were learned at a similar rate and did not differ between the age groups; timing and amplitudes also did not differ between the groups. For further analysis, the younger SLI participants and the older SLI participants were combined to form one SLI group.

SLI vs controls

Percent CRs, CR onset latencies, CR peak latencies, CR amplitudes, and UR amplitudes during each 10-trial block (plus extinction) were calculated for each group. For % CRs (Fig. 1), a repeated measures ANOVA revealed a main effect of block, $F(9, 21)=5.70$, $p<0.001$; partial $\eta^2=.71$. However, there was no main effect of group (partial $\eta^2=.03$) and no block×group interaction (partial $\eta^2=.53$). For CR peak latency, there was also a main effect of block, $F(9, 21)=5.16$, $p<0.001$; partial $\eta^2=.15$, but no main effect of group (partial $\eta^2=.01$) and no block×group interaction (partial $\eta^2=.02$). There were no significant effects with regards to CR onset, CR amplitude, and UR amplitude. The extinction phase revealed a main effect of block, $F(4, 26)=7.97$, $p<0.001$; partial $\eta^2=.22$, but no significant main effect of group (partial $\eta^2=.01$) or block×group interaction (partial $\eta^2=.03$). Thus, the two groups learned the response at similar rates and the timings of the responses did not differ. No differences regarding PPVT scores were observed when the SLI participants were compared to the control participants ($p=.187$). There were no significant correlations found when the PPVT was correlated to mean CR% (Fig. 2), CR peak latencies, CR onset latencies, CR amplitudes, and UR amplitudes. There were also no significant values when each of the conditioning variables were split to first half (blocks 1–5) and second half (blocks 6–10). In order to examine variance between the groups, a one way ANOVA was used to compare the variance in percent CRs between the SLI and control groups. The ANOVA revealed no significant difference between the groups $F(1, 29)=0.96$, $p=0.34$; partial $\eta^2=.23$. This finding suggests that the two groups’ variance was not significantly different.

Discussion

Overall, this is the first study to employ delay eyeblink conditioning, a cerebellar-dependent task, with children with SLI, in order to evaluate hypotheses generated from models of cerebellar impairment contributing to speech,
language, or reading impairment. The results suggest that the cerebellar circuitry involved in delay eyeblink conditioning, using the fundamental single-cue delay EBS procedure, is not affected in this group of SLI participants without speech impairments but with persistent language and reading impairments. The advantage of the EBC procedure used here is that the underlying cerebellar neural pathways are well documented in particular detail. The outcomes do not support cerebellar impairment at the level of basic motor learning pathways as part of the pathogenesis of SLI, at least in the classic SLI profile of the children in this sample. In agreement with past studies (Sears et al. 1994) it was reported there were no age effects in the control groups on measures of conditioning—children ages 9–11 are as proficient as young adults. The findings here for children with SLI do not replicate the earlier report (Sears et al. 1994) of EBC differences in a group of children with autism versus control children. The methods of the two studies are very similar so presumably the lack of replication may be attributable to group differences. The findings also do not replicate the earlier report of EBC differences in a group of adolescents and young adults with dyslexia (Nicolson et al. 2002). The lack of replication in this case may be attributable to methodological differences or sampling differences. The methods of Nicolson et al. involved a CS of 800 ms, longer than usually used in EBC studies, which could influence responses; the number of learning trials was smaller than usual, with 42 paired tone-airpuff trials instead of the 100 trials used in this study; and the percentage of CR differences between groups were not reported. The sample of persons with dyslexia is also different from this study. The participants with dyslexia in Nicolson et al. scored in the normal or above normal levels on verbal tasks. It is quite possible that persons with reading impairments without language impairments are different from persons with reading and language impairments.

Overall, the results of this study suggest that children with SLI and controls both learned the response and at a similar rate. The groups did not differ in terms of other learning factors such as peak onset of the CR response or the CR and UR amplitudes. Overall, the findings suggest that the circuitry involving delay eyeblink conditioning, as measured in this procedure, including the cerebellum, is intact in children with SLI.

The cerebellum has also been implicated to be necessary for the learning of vocabulary. PET studies have shown an activation of the cerebellum during verbal short term memory tasks (Paulesu et al. 1993). This has also been verified with a case study of an individual with a cerebellar lesion (Siliveri et al. 1998). Since eyeblink/cerebellar learning and vocabulary knowledge is sufficiently intact for children with SLI, this could explain why it does not predict individual differences in vocabulary outcomes (i.e. PPVT). This is consistent with other studies which have found no correlations with a variety of intelligence performance variables correlated with eyeblink conditioning (Cromwell et al. 1961; Ohlrich and Ross 1968). This study also replicated findings that young adults do not differ from adolescents on the level of eyeblink conditioning (Sears et al. 1994).

Importantly, this sample of children with SLI is without a history of speech impairments. Although reviews sometimes conflate SLI and impairments of speech production (Hill 2001), an epidemiologically ascertained sample of children yielded an estimated 2% of overlap of speech and language impairments; for the children with SLI, speech disorders were evident in approximately 5–8% of the children (Shriberg et al. 1999). The findings here are consistent with the possibility that cerebellar impairment could be implicated in motor speech components instead of the grammar components of children with SLI. At the same time, the study also suggests that learning at the level of the EBC, under cerebellar control, is not likely to be a source of the persistent grammatical deficits characteristic of SLI, given that the participants had grammatical deficits and unaffected EBC. Under this logic, the outcomes suggest that EBC-related cerebellar circuitry is not implicated in a possible cerebellar role in reading impairments, as hypothesized by Nicolson, et al. (Nicolson et al. 2001), given that this sample of children with SLI showed persistent low levels of performance on word identification tasks although there were no group differences on the EBC learning measures. The alternative possibility is that EBC impairments are present in some but not all groups diagnosed with dyslexia.

There are several limitations to the current study which must be considered. First, the delay eyeblink conditioning paradigm circuitry involves the entire cerebellum. It is possible that deficits could be present with specific regions of the cerebellum such as the anterior cortex. In this case, other variations of eyeblink conditioning, such as a long ISI, could be employed to probe anterior cerebellar cortex function. Second, young infants can demonstrate acquisition that is indistinguishable from adults in terms of asymptote of learning and timing of responding in a delay conditioning task (Herbert et al. 2003). Thus, a task involving increased demands on the cerebellum could provide more insight into the possible roles of the cerebellum in SLI. Finally, although the sample size might be considered overly small to detect group differences, it is unlikely to have been insufficient given the small between-group effect size of the primary variable of interest, acquisition phase % CRs (i.e., main effect of group partial $\eta^2$ was .01).
In conclusion, the results of the study indicate participants with SLI acquire a cerebellar-dependent task at a similar rate to controls. SLI participants also show no differences in timing and amplitude of the response. This would suggest that the cerebellum of participants with SLI without speech impairments (but with reading impairments) is not grossly impaired and thus the deficits witnessed with participants with SLI are likely to be localized to other areas on the brain.

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