Severe receptive language disorder in childhood—familial aspects and long-term outcomes: results from a Scottish study

Ann Clark, Anne O’Hare, Jocelynne Watson, Wendy Cohen, Hilary Cowie, Rob Elton, Jamal Nasir, Jonathan Seckl

Background and aims: Little is known about the familial characteristics of children with severe receptive specific language impairment (SLI). Affected children are more likely to have long-term problems than those with expressive SLI but to date they have only been described as small cohorts within SLI populations. We therefore aimed to describe the clinical and familial characteristics of severe receptive SLI as defined by a rigorous phenotype and to establish whether non-word repetition showed a relationship with language impairment in these families.

Methods: Cross-sectional study of children who met ICD-10 (F80.2) criteria for receptive SLI at school entry, their siblings and genetic parents with standardised measures of language and non-verbal IQ, phonological auditory memory and speech sound inventory.

Results: At a mean of 6 years after school entry with a severe receptive SLI, the 58 participants had a normal mean and standard deviation non-verbal IQ, but only 3% (two) had attained language measures in the normal range. One third still had severe receptive language impairment. One third of siblings not known to be affected had language levels outside the normal range. Phonological auditory memory was impaired in most family members.

Conclusion: Severe receptive SLI is nearly always associated with an equally severe reduction in expressive language skills. Language impairment in siblings may go undetected and yet they are at high risk. Family members had weak phonological auditory memory skills, suggesting that this could be a marker for language acquisition difficulties. Receptive SLI rarely resolves and trials of therapy are urgently needed.

Specific language impairment (SLI) is a developmental disorder where children fail to acquire language at the normal rate but for whom there is no identifiable medical or neurological aetiology. SLI is common, affecting 6–8% of children at school entry. For many preschool children with SLI the prognosis is good, with 44% showing resolution, particularly if they have expressive language impairment only. However, the prognosis is believed to be much poorer for children with receptive language impairment in which comprehension is also affected. Follow-up studies of children with SLI which have included those with receptive problems have suggested that there are persisting language, literacy and behaviour problems in later childhood and adult life despite speech and language therapy and placement in specialised educational settings such as language units. The level of non-verbal intelligence and language comprehension is considered to have a role in this poorer prognosis.

An active ascertainment study of SLI suggested that only a minority of affected children have an accompanying speech sound or pronunciation difficulty or have been identified as having language delay by their parents. Presumably parents are then unable to bring concerns of poor language acquisition to the attention of primary care professionals during developmental surveillance. The process of identification is further frustrated by the difficulties inherent in identifying abnormalities of language acquisition in surveillance consultations, even when the health visitors have been specially trained to do so.

The acquisition of language is highly heritable and particularly so for those children with the slowest rates of language development. Therefore, it might be reasonable to measure language skills in siblings of families in which there is a proband already identified as having SLI. However, the majority of familial studies of SLI have defined caseness on the basis of a language composite measure which combines both receptive and expressive levels and have therefore included a majority of children with expressive difficulties only. Preliminary conclusions suggest that although there is a high familial incidence of SLI, the rate is much less for children with receptive disorder.

In this report our objective is to describe the receptive and expressive language of a group of children with carefully defined receptive SLI some 6 years after entry to school. We also examined familial characteristics to determine the incidence of language impairment in siblings and whether this was recognised by their parents. We measured phonological auditory memory, through non-word repetition for other family members, which has been shown to be heritable in twin studies. Non-word repetition has also acted as a quantitative phenotype to establish linkage to chromosomes 19 and 16 in a large genetic study of children with SLI and we aim to establish whether non-word repetition showed a relationship with language impairment in these families.

METHODS
Participants
Ethical approval was granted by the three health authorities in South, Central and South Eastern Scotland. Speech and language therapists and paediatricians referred English-speaking monolingual children with SLI including a receptive component to the study. All available biological parents and siblings of the probands also took part.

Proband eligibility was determined by the phenotype of the ICD-10 receptive language disorder (F80.2) (box 1).

Abbreviations: CELF-R, Clinical Evaluation of Language Fundamentals-Revised; EAT, Edinburgh Articulation Test; NVIQ, non-verbal IQ; NWR, non-word repetition test; SD, standard deviation; SU, specific language impairment
Box 1 ICD-10 (F80.2) criteria for receptive specific language impairment (SLI)

- A. Language comprehension, as assessed on standardised tests, is below the 2 standard deviation (SD) limit for the child’s age.
- B. Receptive language skills are at least 1 SD below non-verbal IQ as assessed on standardised tests.
- C. There are no neurological, sensory or physical impairments that directly affect receptive language nor is there a pervasive developmental disorder.
- D. Most commonly used exclusion clause: non-verbal IQ is below 70 on a standardised test.

This eligibility had to be relevant at the time the child was assessed by a multi-disciplinary team as requiring special educational support on entering school, which is usually at the age of 5 years in Scotland. Criterion A was established from a retrospective analysis of the standardised assessment results recorded in the child’s clinical notes. Adherence to criterion B for most children was again based on their having a documented measure of non-verbal IQ (NVIQ) from their clinical notes at the time that they entered school. For education authority, this measure was not available because it did not form part of the assessment screen for language unit entry. Therefore, these children were regarded as provisionally eligible based on the referring clinician’s judgement with acceptance contingent on a child having a measured normal NVIQ during the cross-sectional component of the study. Criterion C was addressed by both retrospective analysis of clinical case notes and current clinical history conducted by the researchers. This established whether the child had had a normal hearing test at the time of diagnosis. As the children were participating in a genetic study, we elected to exclude those who had been born preterm, those who had epilepsy and those born to epileptic mothers because of the reported associations in these conditions with language delay. No clinical investigation such as karyotype or brain imaging was conducted by the research team, but no child required to be excluded on the basis of abnormal results of such investigations that had been conducted by the clinical team or referrer.

Assessments

The parents were interviewed before the assessments to establish whether they considered that they themselves or the probands’ siblings had or had had a language learning impairment, which was defined as a difficulty in speech and language and/or literacy skills. Receptive and expressive language skills for the probands and siblings who were aged 5–16.9 years were assessed using the Clinical Evaluation of Language Fundamentals-Revised (CELF-R test. This assessment consists of six subtests, three of which combine to give a composite receptive language score (RLS) and three a composite expressive language score (ELS). There are relatively few standardised tests for assessing language abilities in school-aged children and the CELF-R is widely used and accepted in the UK for this purpose. However, the test is standardised on American children and information on UK norms is limited, although a study of 20 Scottish children aged 12 years showed that receptive subtest scores were not significantly different. NVIQ was assessed using the age-appropriate version of the Raven’s Matrices.

Auditory memory was measured for all family members aged over 4 years using a research version of a non-word repetition test (NWR). This task is a measure of phonological auditory memory and a poor performance is considered a good clinical indicator of SLI as it can continue to be impaired even when the overt language deficit has resolved. All family members had their speech sound inventory determined by the Edinburgh Articulation Test (EAT). This allowed for a qualitative judgement as to whether any potential errors made in the non-word memory test were typical of a person’s habitual speech sounds to ensure that the speaker was not penalised for making accent-specific productions. The standardisation of the EAT is based on a sample of children aged 3–6 years. All speakers above the age of 6 years who had a raw score of 52 or lower (ie, the standard score equivalent of 85 at 6 years) were considered to have a speech sound disorder in accordance with the authors’ suggestion. The measure of the CELF language scores and the NWR have both been informative in a systematic genome-wide analysis in SLI.

Statistical analysis

Quantitative measurements were approximately normally distributed, so parametric methods were used throughout. Associations between different measurements within each category for the study participants were conducted by two-sample t tests or Pearson correlation as appropriate. Multiple linear regression was used to test for the effects of different factors on NVIQ after adjusting for age. Paired t tests with the family as the unit, were used to compare probands with other family members using the mean scores for all of those tested in the case of siblings.

RESULTS

A total of 355 children were referred to the study. Of these, 266 were excluded, 141 because, although they had a receptive language impairment, it was less marked than 2 SD below the mean and therefore did not meet the ICD-10 (F80.2) criteria. A further 78 children were excluded because their NVIQ was below the normal range and 47 were excluded under criterion C. Of the remaining 89 families invited onto the study, 22 declined and nine subsequently withdrew, leaving 58 probands and their families to participate. There were no children from ethnic minorities. There was no significant difference in age and gender of the participants who declined or withdrew. The participant group characteristics are shown in table 1.

The proband adherence to ICD-10 (F80.2) criteria had been established during the late preschool year at the approach of school entry with the exception of one child who had a late diagnosis of SLI at the age of 8 years. All probands were receiving or had recently had specialist educational support. Parents reported language and literacy difficulties for 35 of the siblings and four of the mothers gave this as a self-report and

<table>
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<th>Table 1 Participant group characteristics</th>
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<tr>
<td><strong>Proband</strong> <em>(n = 58)</em></td>
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<tr>
<td><strong>Age (study entry), years</strong></td>
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<tr>
<td><strong>Multi-disciplinary assessment of special educational needs to study entry, mean (SD)</strong></td>
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10 of the fathers. Table 2 shows the results of the assessment battery for the probands.

Proband phenotype
At a mean of 6 years from the initial multi-disciplinary assessment and diagnosis of receptive SLI, cross-sectional study demonstrated that the proband’s language levels remained significantly impaired. A total of 35% of the probands had receptive language measures that were below 2 SD from the mean and thus continued to meet ICD-10 (F80.2) criteria for receptive SLI. An additional 28 (52%) of probands had a receptive deficit that lay between 1 and 2 SD below the mean. Expressive language was even more severely affected and 35 (60%) had expressive language below 2 SD from the mean, with a further 17 (29%) between 1 and 2 SD below. The expressive language was positively correlated with receptive language (p = 0.001, r = 0.60) (fig 1) and only two (3%) of the children had attained both receptive and expressive ability within normal limits at the cross-sectional point of the study. Fourteen of 51 probands had a speech sound disorder. There was no significant correlation between the school entry scores for the Reynell Language Scales\(^\text{4}\) and the CELF-R scores at the point of the study. This remained true after the groups were split around the median follow-up time, so that there was no evidence that those with shorter time intervals of follow-up had more association between the two scores.

The language measures were not related to age (receptive language vs age non-significant, p = 0.829, r = -0.0301; expressive language vs age non-significant, p = 0.303, r = 0.1427). NVIQ was normally distributed and after adjusting for age in a multiple regression there was no significant relationship with language measures.

Family members
The sibling assessment profiles are shown in table 3. A total of 62 of 98 siblings were the correct age to complete the assessment of the CELF-R (20 siblings were too young and 12 were too old for the assessment battery and four were non-compliant). Figure 2 shows that although the siblings had significantly higher language scores than their probands (p > 0.001), the distribution of their standard scores was below that expected in the normal population. Six of the 62 had both a receptive and expressive language standard score below 70 and thus would have met ICD-10 (F80.2) criteria for receptive SLI. Four of these children had been reported to have difficulties in their language acquisition or literacy development by their parents. In addition, one other child had a receptive language standard score below 70, thus fulfilling ICD-10 criteria for receptive SLI but their expressive language was better developed and their parents had not reported any difficulties that they recognised in the acquisition of language or literacy. Therefore, even though four children met criteria for SLI they had not been recognised by their parents to have a language learning difficulty. Three of these children had a measured speech sound disorder on the EAT.

Phonological auditory memory for probands, siblings and parents
Both parents participated in the study for 50 families, only the mother took part for seven, and only the father for one. Forty nine mothers and 41 fathers were available to complete the EAT and none had a speech disorder on this measure. Three mothers had reported language learning difficulties: one an SLI and two dyslexia. Ten of the fathers described language learning difficulties with SLI in one, dyslexia in five, phonological disorder in one and a mixed language learning impairment in three. The probands’ non-word repetition scores as a measure of phonological auditory memory were not related to receptive language ability (NS, p = 0.082, r = 0.25) or age (NS, p = 0.056, r = -0.27) and were significantly related to expressive language (p = 0.005, r = 0.39) and NVIQ (p = 0.013, r = 0.35). The non-word repetition standard scores for the siblings also correlated with expressive language measures (p > 0.001) but did not correlate with the siblings’ receptive language scores. Looking at the family members as a whole, the non-word repetition standard scores were substantially below the normal range for probands, siblings and parents (fig 3). There were significant correlations between the proband’s standard scores and that of their family members (the average of their siblings (p = 0.001, r = 0.50), their mothers (p = 0.002, r = 0.42) and their fathers (p = 0.011, r = 0.37)). There was no significant correlation between the maternal and the paternal scores (p = 0.07, r = 0.28). The parents’ reporting of their own affected status was not reflected in their non-word repetition score (mothers, p = 0.37, t = 0.91; fathers, p = 0.55, t = 0.60). The siblings’ non-word repetition standard scores, however, did correlate with the reports from their parents that they were affected with a language learning difficulty (p = 0.004). Overall, 40 (69%) of the probands had a first-degree relative with either a language learning impairment described from the history or a directly measured impaired language standard score.

**DISCUSSION**

Receptive SLI, in which a child experiences difficulties in acquiring language comprehension despite normal non-verbal intelligence, is an important condition to identify because it has long-term adverse sequelae on language, learning and behaviour.\(^\text{10, 12}\) The definition of the condition is debated both in terms of where one places the exact cut-off points for the exclusionary criteria for language impairment severity and NVIQ and what constitutes a predisposing neurological impairment.\(^\text{11}\) However, in this paper we are reporting on a group of children with a severe form of the disorder which conforms to strict research ICD-10 criteria. This is the largest group of children with severe receptive SLI reported to date in the literature and the 65% participation in this family genetic study was very high compared with that described in the literature for other conditions such as asthma.\(^\text{14}\) This rigour of phenotyping identified a group in whom a language learning difficulty occurred in 69% of first-degree relatives, which is much higher than previously thought to be the case in receptive SLI.\(^\text{17}\)
The ratio of boys to girls in our study group was just over 2 to 1. There is no other directly similar group with which to compare this finding. As the children in this report were identified as being in need of a language unit type level of educational support, the closest comparison might be the cohort of children reported in a number of publications who were a random sample of 50% of all year 2 children attending language units in the UK. The ratio of boys to girls was over 3 to 1 in these studies, but the cohort included children with expressive disorder only, as well as those with moderate learning difficulties and autism spectrum disorders. The other literature that explores gender ratios for SLI includes epidemiological studies, but again the definition of SLI is usually based on a composite of receptive and expressive language skills and is less severe. With this approach the prevalence in boys and girls is very similar at 0.08 and 0.06, respectively.

Although the children were identified by their receptive language impairment, they all had equally severe expressive language impairments. The same pattern was observed in affected siblings. There are a number of studies reviewed by Bishop which suggest that receptive language impairment may be conceptualised as being at the severe end of SLI in contrast to isolated expressive disorder which characterises the milder form. This relationship between receptive and expressive language is in keeping with the apparent genetic similarities of linkage in the different forms of receptive and expressive SLI, although the work in this area to date is not powered per se to compare the genetics between the two.

In our study we examined the capacity of phonological working memory through an NWR for all family members who were old enough to complete it. This measures short-term storage and verbal processing, has been reported as deficient in SLI and demonstrates a consistent linkage with the SLI1 region of chromosome 16q. It has not been specifically described in receptive language impairment, but our study demonstrates that not only is it reduced in affected probands, but it is generally depressed in siblings and parents. These findings are in keeping with the evidence that although performance on the test by young children can be constrained by the immature phonological and articulatory systems, the highly consistent pattern of associations between non-word repetition and language learning across different ages and groups suggests that speech-motor outputs cannot explain the individual performance. Although many of the adults in our study performed poorly on this measure of non-word repetition, none of them had a speech output problem as formally measured by a standardised articulation test. Although performance on an NWR is highly heritable and associates with poorer language acquisition, it is possible to have difficulties with phonological memory and yet not develop SLI, and so other factors appear involved. However, although our

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**Table 3** Sibling assessment profile expressed in standard scores

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<th>Mean (SD)</th>
<th>Range</th>
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<tr>
<td>Non-verbal IQ (Raven’s)</td>
<td>72</td>
<td>96 (14.72)</td>
<td>75-125</td>
</tr>
<tr>
<td>Language measures (CELF-R)</td>
<td>n</td>
<td>62</td>
<td></td>
</tr>
<tr>
<td>Receptive language composite</td>
<td></td>
<td>89.9 (16.6)</td>
<td>50-128</td>
</tr>
<tr>
<td>Expressive language composite</td>
<td></td>
<td>81.0 (15.3)</td>
<td>50-124</td>
</tr>
<tr>
<td>Auditory memory (NWR)</td>
<td>n</td>
<td>82</td>
<td></td>
</tr>
<tr>
<td>Mean (SD)</td>
<td></td>
<td>81.7 (17.6)</td>
<td>55-125</td>
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CELF-R, Clinical Evaluation of Language Fundamentals-Revised; NWR, non-word repetition test; SD, standard deviation.
understanding of the genetics of speech and language disorders is advancing rapidly since the first reporting of the Fox P2 gene on 7q31 in a family with a severe speech disorder,\textsuperscript{38,39} with subsequently linkage sites reported for SLI,\textsuperscript{19,28,40} our present state of knowledge does not allow for genetic confirmation of SLI. Therefore, our findings suggest that siblings of children affected by receptive SLI should have a formal assessment of their language skills by a speech and language therapist. As our findings confirm that children in families affected by receptive SLI also have expressive difficulties, the identification of affected siblings has implications for their treatment. A recent Cochrane report emphasised that although more research was needed to identify the best way forward for treating receptive language difficulties,\textsuperscript{41} there were significant benefits demonstrable for expressive language difficulties from speech and language therapy. The importance of our findings is that the incidence of language impairment is so high in siblings that they might be best served by undergoing formal language assessment given the difficulties of case recognition and diagnosis.\textsuperscript{11,12,33,42}

In conclusion, we acknowledge that this present study is not prospective and so there is the possibility of a bias towards more severely affected children participating. Nevertheless, the very high rate of continuing severe receptive and expressive difficulties at a mean of 6 years after identification of the SLI at school entry is of concern, particularly if extrapolated into adult life.\textsuperscript{10} We are also unable to comment on outcome and familial and clinical features for children who have either a less severe receptive language impairment or one that might be arising from a wide range of underlying aetiologies, as these children were excluded under the rigorous subject eligibility that we had
to adhere to in order to arrive at an informative phenotype for genetic study. What we have demonstrated is that for families with a child recognised to have a severe receptive language impairment at school entry, there is a high rate of SLI in siblings which may not be evident clinically or apparent to their parents, and that there may be merit in identifying affected siblings through formal language testing. Also poor phonological memory is a characteristic for these family members. Finally, the prognosis for recovery of receptive and expressive language appears poor and affected children are likely to require long-term support.

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