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Abstract

Objective—Younger siblings of children with autism spectrum disorder (ASD) are at high risk (HR) for developing ASD as well as features of the broader autism phenotype. While this complicates early diagnostic considerations in this cohort, it also provides an opportunity to examine patterns of behavior associated specifically with ASD compared to other developmental outcomes.

Method—We applied Classification and Regression Trees (CART) analysis to individual items of the Autism Diagnostic Observation Schedule (ADOS) in 719 HR siblings to identify behavioral features at 18 months predictive of diagnostic outcomes (ASD, atypical development, and typical development) at 36 months.

Results—Three distinct combinations of features at 18 months were predictive of ASD outcome: 1) poor eye contact combined with lack of communicative gestures and giving; 2) poor eye contact combined with a lack of imaginative play; and 3) lack of giving and presence of repetitive behaviors, but with intact eye contact. These 18-month behavioral profiles predicted ASD versus non-ASD status at 36 months with 82.7% accuracy in an initial test sample and 77.3% accuracy in a validation sample. Clinical features at age 3 among children with ASD varied as a function of their 18-month symptom profiles. Children with ASD who were misclassified at 18 months were higher functioning, and their autism symptoms increased between 18 and 36 months.

Conclusion—These findings suggest the presence of different developmental pathways to ASD in HR siblings. Understanding such pathways will provide clearer targets for neural and genetic research and identification of developmentally specific treatments for ASD.
Keywords
ASD; high-risk siblings; infants; broader autism phenotype; predictors of outcomes

Introduction
Autism spectrum disorder (ASD), characterized by impairments in communication and social reciprocity and the presence of repetitive behaviors and sensory interests, is one of the most prevalent and heritable neurodevelopmental disorders. Compared to the general population, younger siblings of children with ASD are at increased risk for developing ASD (18.7%) or social-communicative and other developmental vulnerabilities. Prospective studies of such high-risk (HR) cohorts provide unique opportunities to investigate the developmental dynamics underlying the emergence of autistic psychopathology as well as resilience. Better understanding of these dynamics could lead to identification of early predictors of ASD in HR siblings, which in turn would enhance screening and diagnostic practices in this cohort and may inform identification of novel treatment targets. Considering the importance of early intervention in shaping brain development and behavioral outcomes in young children with ASD, identification of the affected siblings as soon as their symptoms begin to emerge offers the potential for altering their long-term outcomes in a clinically meaningful manner.

Early identification of HR siblings who are most likely to develop ASD is impeded by the complexity of clinical presentation and developmental trajectories unique to this cohort. A large minority of HR siblings without ASD exhibit broader autism phenotype features, e.g., elevated scores on diagnostic instruments of autism severity and lower verbal skills, limited functional play, imitation, and repetitive behaviors, and therefore it may be difficult to differentiate them in the second year of life from those who eventually develop ASD. Moreover, it is not clear when and how the behavioral symptoms begin to emerge. Symptoms may emerge after a period of relatively typical development either as a result of a loss of skills or a failure to acquire new skills. The conceptualization of onset patterns is further complicated by the finding that some symptoms may show a gradual decrease in frequency (e.g., eye contact), whereas others fail to increase at the rate observed in typical controls (e.g., social vocalizations), suggesting that the departure from typical trajectories in specific domains may follow different patterns and take place during different developmental periods. The complexity of broader autism phenotype expression amongst HR siblings and variable symptom onset patterns may contribute to difficulties identifying features early in development that are consistently associated with ASD among HR siblings.

In a recent study aimed at identifying predictive signs of ASD in HR siblings based on clinical presentation as captured by individual items of the Autism Diagnostic Observation Schedule –Toddler Module (ADOS-T), Macari and colleagues employed a nonparametric decision-tree learning algorithm (Classification and Regression Trees, CART). A combination of several features—including the limited ability to engage in play, paucity of communicative gestures, limited imitation skills, and atypical intonation—
differentiated siblings with ASD from the remaining sample at around the first birthday with a high degree of sensitivity and specificity. While promising with regard to a methodological approach to identifying early prognostic features of ASD, the study was considered preliminary due to its small sample size (N=84) and focus on predicting outcome at 24 months. Taking advantage of the large and prospectively characterized samples collected through the Baby Sibling Research Consortium (BSRC), the present study is focused on identifying features in 18-month-old HR siblings predictive of ASD diagnosis at age 3. We focused on detection of behavioral features at 18 months, as this represents an age by which many parents begin to note concerns regarding their children’s development, but before most affected children receive diagnosis and enter treatment. Risk assessment at 18 months was based on coding of 29 individual items from the ADOS Module 1 with scores ranging from 0 to 3. The items provided a broad range of autism-relevant behaviors measured in a standardized fashion. To identify predictors of ASD, atypical (ATYP) and typical developmental (TD) diagnostic outcome at 3 years based on the ADOS items, we employed a CART approach. The advantage of this approach is that it allows for selection of the most predictive features from a multiplicity of behavioral symptoms and their interactions, resulting in a parsimonious mapping of different sets of predictors for later outcomes. It also allows for several distinct combinations of features to be related to a single diagnostic outcome, a characteristic that is particularly important given that ASD may arise through multiple etiological pathways. We examined (1) combinations of behavioral features at 18 months that are associated with ASD diagnosis at 3 years, and (2) the factors that affect accurate identification at 18 months of HR siblings who receive diagnosis of ASD at 3 years of age.

Method

Participants

Participants were 719 HR siblings: 413 males (57.4%) and 306 females (42.6%) from eight sites of the BSRC, a network of projects focused on studying the development of younger siblings of children with ASD: University of Toronto/Holland-Bloorview Kids Rehabilitation Hospital, Dalhousie/IWK Health Centre, University of Alberta/Glenrose Rehabilitation Hospital, Boston University/Boston Children’s Hospital, Kennedy Krieger Institute, University of California, Davis, University of California, Los Angeles, and Yale University. Infants were included in the study if they had an older biological sibling with ASD and had complete characterization data including ADOS and MSEL at 18 months and 3 years. All sites verified the older siblings’ diagnostic status based on the ADOS, Autism Diagnostic Interview –Revised, and/or Social Communication Questionnaire. Exclusion criteria were: identified neurological or genetic condition in the older or younger sibling (e.g., fragile X syndrome or tuberous sclerosis).

Measures

ADOS—The ADOS is a semi-structured, standardized assessment of social interaction, communication, and play skills as well as repetitive behaviors that are diagnostic of ASD. To quantify symptoms of ASD, at 18 months, all children were administered Module 1, which is designed to assess behavioral symptoms in nonverbal and minimally verbal young
children. At 3 years, either Module 1 (n=181) or 2 (for children with early phrase speech; n=538) was employed. To facilitate comparisons of symptom severity across modules and ages, total algorithm scores were converted into calibrated severity scores, which range from 1 to 10.

**MSEL**—The MSEL is a standardized developmental measure for children between birth and 68 months. Scores in four MSEL domains were considered in the current analysis: Fine Motor, Visual Reception, Expressive Language, and Receptive Language. Nonverbal Developmental Quotient (DQ; NVDQ) scores were computed as the average of Fine Motor and Visual Reception DQ scores, and verbal DQ (VDQ) scores as the average of Expressive Language and Receptive Language scores.

**Outcome Assessment**

Diagnostic groupings were based at 3 years on a combination of clinical best-estimate (CBE) and test scores on ADOS and MSEL (see Table S1, available online, for details). CBE was assigned at each site by an expert clinician based on a combination of measures including medical and developmental history as well as social, cognitive, verbal and adaptive functioning. Consistent with other BSRC reports, those in the ASD group had to meet CBE criteria for ASD and have an ADOS severity score in the clinical range. Toddlers with ATYP exhibited abnormal scores on either the ADOS or MSEL or both. The TD group had scores in the typical range on both instruments. Based on these criteria, the sample consisted of 157 (21.8%) siblings with ASD outcomes, 178 (24.8%) siblings with ATYP outcomes, and 384 (53.4%) siblings with TD outcomes.

**Statistical Analysis**

We employed CART analysis to identify the individual items of the ADOS at 18 months of age that best predicted diagnostic outcomes at 3 years of age. CART analysis is a decision-tree technique that uses recursive partitions of the data to predict a categorical or continuous response variable. A decision tree is a flow-chart-like structure, where each internal (non-leaf) node denotes a test on an attribute (e.g., ADOS item X), each branch represents the outcome of a test (e.g., the score of item X), and each leaf represents a class label (e.g., ASD, ATYP, or TD). At each step, the model selects the best variable and cutoff score among all available predictor variables to make a partition. The nested structure of partitions within CART analysis naturally incorporates interactions among variables in the model, and the option to stop the growth of the tree at any partition (i.e., “pruning” the tree) provides a method of variable selection by predictive importance.

A validation sample comprising 20% of the participants (n = 154), selected randomly from each outcome group (35 ASD, 40 ATYP, and 79 TD) and site, was initially set aside and used only for assessment of the final CART model on out-of-sample data. The remaining participants (n=565; 122 ASD, 138 ATYP, and 305 TD) were retained in the training set and used to construct the tree. CART models were built using the “tree” package in R. To prevent over-fitting the model to the training data, an optimal tree size was identified by examining misclassifications over all three groups using 10-fold cross-validation on the training set for tree sizes varying from 2 to 20 leaves. The final tree was pruned to this...
optimal misclassification-minimizing size, with the fitted label for each leaf in the final tree assigned by a ‘majority vote’ (i.e., defined as the most prevalent diagnostic group in the leaf, with ties broken by giving higher precedence to groups having a higher proportion of their membership within that leaf). Comparisons of cases that were correctly classified and misclassified by CART were conducted using linear mixed models with Tukey-Kramer correction for multiple post hoc comparisons. Effects of site and gender were included in the models as indicated.

Results

Sample description

ASD, ATYP, and TD groups were similar with regard to racial (Caucasian vs. non-Caucasian) and ethnic (Hispanic vs. non-Hispanic) composition (Table 1). Boys were more likely to be diagnosed with ASD than girls (29.3% versus 11.8%) and to have an ATYP outcome (28.1% versus 20.3%). The groups did not differ in the age of recruitment, or in maternal and paternal age. They had similar proportions of college-educated mothers, but fathers of children with TD outcomes were more likely to hold a college degree than fathers of children with ASD or ATYP. There were no differences among groups in age at the 18 month visit, but the groups differed in autism severity scores (ASD>ATYP>TD) and verbal and nonverbal (ASD<ATYP=TD) scores. There were no significant differences among sites in the proportion of ASD outcomes (X[^2][5]=5.39, p=.369), ADOS-calibrated severity score, verbal DQ, and nonverbal DQ at either of the ages (all p-values >.07).

Predicting ASD diagnosis based on behavioral features at 18 months

CART cross-validation analysis indicated that the optimal misclassification error rate was obtained for a tree size of 8 leaves (Figure 1, Table 2). Three leaves received a label of “ASD,” capturing 57% of ASD cases (H, F, and D). A majority (93%) of TD siblings was classified correctly into leaves A, C, E, and G, thus these leaves received a label of “TD.” Very few (4%) children with ATYP outcomes were assigned their own leaf (B); instead, the ATYP siblings were distributed largely between TD (77%) and ASD (19%) leaves. Inclusion of site in the model did not alter the tree structure, suggesting that site was not an important predictor of classification outcome under CART analysis. Taken together, at 18 months, CART analysis predicted ASD versus non-ASD (ATYP and TD combined) diagnostic status at 36 months with an overall accuracy of 82.7%, 56.6% sensitivity, 89.8% specificity, positive predictive value (PPV) of 60.5%, and negative predictive value (NPV) of 88.2%. Analogous prediction rates on the validation subsample were 77.3% overall accuracy, 45.7% sensitivity, 86.6% specificity, 50.0% PPV, and 84.4% NPV.

The analysis identified three combinations of behavioral features predictive of diagnostic outcome at 36 months (Figure 1, Table 2). The largest proportion of affected siblings (41%, leaf H) was classified based on a combination of poorly modulated eye contact to initiate, terminate, or regulate social interactions (item B1 of Module 1, score 2), the absence of giving objects to others to share (B8, score 1 or 2), and limited use of emotional or descriptive communicative gestures other than pointing (A8, score 1 or 2). Relative risk for ASD in this leaf was 3.17, i.e., compared to all siblings in our sample, siblings in leaf H
were over 3 times more likely to be diagnosed with ASD. Toddlers in this leaf had ADOS-calibrated severity scores that were high and stable from 18 to 36 months, very low verbal scores, and low-to-average nonverbal scores (Table 3). An additional 10% of the children with ASD were classified by their impaired eye contact (B1, score 2), and the lack of ability to spontaneously engage in pretend play (C2) (leaf F, score 2 or 3). The relative risk for ASD in this leaf was 1.79, suggesting an almost two-fold increased likelihood of ASD in this leaf. Toddlers in leaf F had moderate calibrated symptom severity scores that remained stable over time, and displayed below-average verbal and average nonverbal skills (Table 3). Finally, a third group of siblings with ASD classified into leaf D (6%) displayed appropriate eye contact meshed with other communicative behaviors (B1, score 0), but presented with repetitive and stereotyped behaviors ranging from relatively mild to severe (D4, score 1, 2, or 3) and rarely or never gave objects to get help or to share (B8, score 2) during the ADOS assessment. Relative risk for ASD in this leaf was 3.24. Children in this leaf had calibrated severity scores at 18 months in the borderline range; however, their symptoms worsened over time (Table 3). They also exhibited below-average verbal skills and average nonverbal skills at both time points.

**Comparison of HR siblings with ASD classified correctly versus misclassified**

Subsequently, we compared siblings with ASD outcomes who were at 18 months “Classified Correctly” (ASD:n=69) or “Misclassified” (i.e., classified as ATYP or “TD:n=53). There were no significant differences between correctly classified and misclassified participants with ASD with regard to gender, age at recruitment or assessment, race or ethnicity, and maternal or paternal age and education (Table S2, available online). However, there was a significant effect of age (F (1,119) = 112.7, p < .001), classification group (F (1,113) = 69.7, p < .001), and age × classification interaction (F (1,119) = 24.96, p< .001) on autism severity score (Figure 2). The misclassified children had significantly lower severity scores than those classified correctly at both time points and severity scores increased over time in both groups (all post hoc comparisons are significant at least at p<.01 level with Tukey-Kramer correction). Neither the effect of site (p = .28) nor gender (p = .23) was significant. For verbal DQ there were significant effects of age (F [1, 77] = 9.84, p = .003) and classification (F [1,113] = 32.49, p < .001), but no interaction effect (p = .10). The misclassified siblings had higher VDQs at both time points than those classified correctly, and in both groups the VDQ scores increased over time. There was no effect of site (p = .15) or gender (p = .94) on VDQ. Finally, for NVDQ there was a significant effect of age (F [1, 78] = 11.91, p < .001) and classification (F [1,113] = 19.22, p < .001), but no interaction effect (p = .55). Misclassified children had higher NVDQ scores than those who were classified correctly, and in both groups, the scores declined slightly over time. There was no site effect (p = .17) or gender effect (p = .54) for NVDQ.

**Discussion**

This is the first large-scale, multisite study aimed at identifying specific behavioral features that distinguish HR siblings with ASD from their typically and atypically developing high-risk peers as early as 18 months of age. Almost half of the HR siblings exhibited clinically-relevant concerns at 36 months in the form of ASD (21.8%) or other atypical developmental
outcomes (24.8%). Compared to recent population prevalence rates of ASD, the HR siblings in our study were 14.7 times more likely to develop ASD. Boys were 2.5 times more likely to be affected by ASD, and 1.5 more likely to have ATYP outcomes than girls. Despite these differences in gender distribution with regard to outcome, boys and girls were equally likely to be classified correctly by the CART analysis suggesting that this analytic approach did not favor detection of signs of ASD in one gender over another.

To identify indices of diagnostic outcome at the age of 3 years, we applied a nonparametric decision-tree learning algorithm to an array of behavioral ratings collected during a standardized social interaction at 18 months. By considering combinations of only six behavioral features at 18 months, it was possible to identify ASD cases with approximately 83% overall accuracy, 57% sensitivity, and 90% specificity in the training sample. This is a relatively high level of accuracy, given that a large proportion of HR siblings showed broader phenotype features already at 18 months, and that siblings missed by the classification procedure were higher-functioning with an increase in symptom severity after 18 months and therefore were unlikely to trigger clinical concerns at this age. Classification based on social-communicative features and other autism-specific behaviors at 18 months was not particularly successful at isolating siblings with non-ASD atypical outcomes. Almost 20% of this group showed characteristics seen in the ASD leaves, whereas over 70% were classified together with TD siblings at 18 months. Follow-up work will be necessary to better understand the distribution of ASD-related traits at 18 months in siblings without ASD and their association with outcomes.

Although the search for invariant biological markers of complex developmental disorders including ASD is ongoing, diagnosis of ASD still relies on the analysis of the behavioral phenotype, a phenotype that is not only highly heterogeneous, but also undergoes important transformations in the first years of life. These factors motivated our focus on a very narrow developmental epoch along with the examination of a high number of combinations of behaviors known to be impaired in the early stages of ASD as potential prognostic indicators. The strength of this analytic approach lies in its ability to enhance the predictive value of individual behaviors by taking under consideration how they interact with other behaviors. Specifically, at 18 months, atypical eye contact was noted in 34% of siblings. Yet, only 40% of them had ASD by 3 years, suggesting that at this age, atypical eye contact by itself is a poor prognostic indicator of ASD amongst HR siblings. However, if at 18 months the atypical eye contact co-occurred with a paucity of communicative gestures and limited use of giving objects to share, then the likelihood of ASD outcome was over three times higher than in HR siblings in general and almost all children who displayed these characteristics had clinically significant (ASD or ATYP) outcomes at 3 years. In a small proportion of siblings with ASD, eye contact was not impaired at 18 months, but these siblings displayed emerging repetitive behaviors and limited use of giving objects to others for any purpose. Children with this combination of features were over three times more likely to have ASD than the HR siblings in general, and almost all siblings in this group had ASD or other atypical outcome at 3 years. Finally, poor eye contact paired with a limited ability to spontaneously engage in pretend play, although not very specific to ASD (relative risk: 1.79), appeared to more generally signal the presence of developmental challenges, as almost 2/3 of siblings in this group had outcomes of either ASD or ATYP at 3 years of age.
Importantly, the groups of siblings identified by the three combinations of features displayed distinct developmental paths between 18 and 36 months with regard to autism severity, verbal, and nonverbal skills. Taken together, these findings suggest that different constellations of individual behaviors observed at 18 months are associated with a common diagnostic outcome. While all HR siblings should be monitored throughout the first three years of life, the presence of these combinations of characteristics at 18 months—the first characterized by drastically limited nonverbal communication and the second by the presence of repetitive behaviors—should trigger consideration of a comprehensive diagnostic assessment and, if indicated, targeted intervention.

One of the key challenges in the field of autism research is the identification of more phenotypically homogeneous subgroups amongst individuals affected by the disorder. Identification of such subgroups may facilitate gene finding and discovery of novel treatment targets and strategies. Amongst the otherwise highly heterogeneous sample of HR siblings with ASD, our analysis identified a large (41%) subgroup of infants who were highly symptomatic and exhibited marked language delays by 18 months. Importantly, they continued to have pronounced autism symptoms and significant language delays at the age of 3 years. An earlier study identified a group of siblings with similar characteristics around the first birthday using an analogous analytic approach, albeit in a much smaller sample. Taken together, these findings suggest that in a large minority of HR siblings, clinical symptoms of ASD paired with significant language delays may emerge around 18 months or before and remain relatively stable over time. Although a more long-term follow up would be necessary, this group may represent toddlers at risk for less optimal outcomes amongst children affected by ASD.

Almost half of siblings later diagnosed with ASD did not trigger an ASD classification at 18 months based on the CART analysis. Neither gender distribution nor parental characteristics differed between the correctly classified and misclassified ASD cases. Importantly, the misclassified siblings appeared to have few developmental delays at 18 months, and their behavioral response to the ADOS probes were largely in the non-clinical range. However, while they maintained average verbal and nonverbal skills over the subsequent months, they evidenced a rapid increase in autism symptom severity between 18 and 36 months. Several, not necessarily mutually exclusive, hypotheses can be advanced to explain this finding. First, it is plausible that the siblings were either asymptomatic or mildly symptomatic at 18 months and in the subsequent months withdrew socially and either failed to advance or showed a loss of previously acquired social-communicative skills. Second, it is possible that their higher verbal and nonverbal skills masked their social disability in the context of the playful interaction with a highly supportive adult, but as their speech and representational skills advanced, their atypical interactive and communicative behaviors became apparent (e.g., stereotyped speech, all-encompassing interest, and limited conversational skills). Finally, it is possible that we missed these cases because we focused our investigation too narrowly on autism-specific behaviors elicited in a particular context. Considering additional information derived from other social contexts (e.g., peer interactions), developmental domains (e.g., emotion regulation) or levels of analysis (e.g., social perception and attention) may help identify those higher-functioning siblings with ASD who were missed at 18 months. Direct investigation of these hypotheses will markedly enhance our understanding.
of both methodological and theoretical issues surrounding the developmental dynamics of HR siblings with ASD.

It is important to note that 18 months is a developmental time point marked by important transitions in the emergence of representational thought, language, and social interactions. Given the developmental nature of ASD, it is possible that the predictive signs of ASD identified at this time point may not generalize to other developmental epochs. The identification of combinations of features specific to, for example, 6-, 12-, 18- and 24-month age levels will be crucial for furthering our understanding of processing underlying emergence of social disability and for the advancement of developmentally-sensitive ASD screening instruments for high-risk siblings in the general population. Our analytic approach was designed to identify diagnostic features that differentiate toddlers with ASD who have an older sibling on the autism spectrum from HR siblings who do not develop the disorder but may display broader phenotype features. It remains an empirical question whether these findings generalize to a broader population of affected toddlers. Finally, although we have focused our analysis on minimizing misclassification error, alternative analytic strategies could be used if the purpose of the analysis was to develop screening instruments where greater emphasis is placed on maximizing sensitivity over specificity. The exploration of these variations and techniques, as well as the incorporation of additional data sources in classification and prediction, may result in advancing development of early screening and diagnostic methods.

This study has several important clinical and theoretical implications. A large minority of HR siblings with ASD display marked symptoms at 18 months, whereas in others, symptoms become pronounced after 18 months, suggesting at least two distinct windows of opportunity for identification of the affected cases. Moreover, several combinations of clinical features at 18 months were predictive of ASD outcome, each associated with a different developmental course and clinical profile by the age of 3 years. The clinical implications of these results are amplified as the combinations of predictive features were derived from the ADOS, a well-validated, standardized, and therefore readily replicable assessment of ASD symptoms. Combined, these findings suggest the presence of different developmental pathways to the common diagnostic ASD outcome, pathways characterized by distinct combinations of early markers. The findings also reinforce the need for repeated diagnostic screening in the first three years of life to identify individual cases of ASD as soon as behavioral symptoms become apparent. Such rapid detection will enhance our ability to ameliorate the impact of primary symptoms on the development of social and nonsocial cognition as well as to prevent secondary symptoms from emerging. Better understanding of the neurobiology of the various underlying pathways to ASD may advance both the identification of novel treatment targets and the design of interventions tailored to specific clinical profiles and their developmental dynamics.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.
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References


Figure 1.
Classification and Regression Tree (CART) diagram predicting groups with autism spectrum disorder (ASD), atypical development (ATYP), and typical development (TD) based on 18-month Autism Diagnostic Observation Schedule (ADOS) item data. Note: The bars reflect the proportion of toddlers from each diagnostic group assigned into each leaf based on their presentation during ADOS evaluation. ADOS items used in the tree are B1 (Unusual Eye Contact), D4 (Unusually Repetitive Interests or Stereotyped Behaviors [RSB]), A3 (Intonation of Vocalizations or Verbalizations), B8 (Giving), C2 (Imagination/Creativity), and A8 (Gestures). The numbers displayed to the left and to the right of the item label reflect the actual scores on the given item of the ADOS.
Figure 2.
Marginal means (+/− 2 standard errors) of severity scores, verbal developmental quotient (DQ) scores, and nonverbal DQ scores in children with autism spectrum disorder (ASD) classified correctly as ASD and misclassified as typically or atypically developing at 18 months.
Table 1

Sample Characterization Based on 36-Month Outcome

<table>
<thead>
<tr>
<th>Variable</th>
<th>ASD</th>
<th>ATYP</th>
<th>TD</th>
<th>p-value</th>
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</thead>
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<tr>
<td>n (% of total sample)</td>
<td>157 (21.8%)</td>
<td>178 (24.8%)</td>
<td>384 (53.4%)</td>
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</tr>
<tr>
<td>Age of Recruitment, months</td>
<td>8.2 (3.8)</td>
<td>7.7 (3.7)</td>
<td>7.5 (4.4)</td>
<td>p = .25</td>
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<td>Males (%)</td>
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<td>116 (65.2%)</td>
<td>176 (46.0%)</td>
<td>p &lt; .001</td>
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<td>Relative Male to Female Risk Ratio</td>
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<td>1.5 : 1</td>
<td>0.63 : 1</td>
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</tr>
<tr>
<td>% Caucasian</td>
<td>81.3</td>
<td>83.7</td>
<td>84.7</td>
<td>p = .66</td>
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<tr>
<td>% Hispanic</td>
<td>13.7</td>
<td>3.8</td>
<td>8.1</td>
<td>p = .09</td>
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<tr>
<td>Maternal Age, years</td>
<td>34.0 (4.6)</td>
<td>35.2 (4.6)</td>
<td>34.4 (4.4)</td>
<td>p = .06</td>
</tr>
<tr>
<td>Paternal Age, years</td>
<td>36.3 (5.5)</td>
<td>37.3 (5.3)</td>
<td>36.9 (5.2)</td>
<td>p = .26</td>
</tr>
<tr>
<td>% of mothers with college degree or higher</td>
<td>71.0</td>
<td>72.9</td>
<td>78.4</td>
<td>p = .22</td>
</tr>
<tr>
<td>% fathers with college degree or higher</td>
<td>68.0&lt;sup&gt;a&lt;/sup&gt;</td>
<td>63.2&lt;sup&gt;a&lt;/sup&gt;</td>
<td>75.7&lt;sup&gt;b&lt;/sup&gt;</td>
<td>p = .02</td>
</tr>
</tbody>
</table>

18 Months

| Age, months                                                  | 18.4 (0.6)   | 18.4 (0.6)   | 18.3 (0.5)    | p = .21 |
| ADOS Severity Score                                          | 4.6 (2.8)<sup>a</sup> | 2.8 (1.9)<sup>b</sup> | 2.1 (1.5)<sup>c</sup> | p < .001|
| MSEL Verbal DQ                                               | 73.5 (23.5)<sup>a</sup> | 91.4 (22.5)<sup>b</sup> | 97.2 (20.6)<sup>b</sup> | p < .001|
| MSEL Nonverbal DQ                                            | 95.5 (12.3)<sup>a</sup> | 103.8 (12.5)<sup>b</sup> | 105.9 (11.3)<sup>b</sup> | p < .001|

36 Months

| Age, months                                                  | 37.5 (2.4)<sup>a</sup> | 38.3 (2.8)<sup>b</sup> | 37.7 (2.8)<sup>c</sup> | p = .02 |
| ADOS severity score                                          | 7.0 (1.7)<sup>a</sup> | 4.1 (1.8)<sup>b</sup> | 1.3 (0.5)<sup>c</sup> | p < .001|
| MSEL Verbal DQ                                               | 81.3 (27.0)<sup>a</sup> | 98.1 (16.4)<sup>b</sup> | 105.9 (14.6)<sup>c</sup> | p < .001|
| MSEL Nonverbal DQ                                            | 86.4 (21.8)<sup>a</sup> | 100.7 (16.5)<sup>b</sup> | 109.9 (13.7)<sup>c</sup> | p < .001|

Note: Means are reported with standard deviations in parentheses: M (SD), except where variable is labeled as a percentage (%). ADOS = Autism Diagnostic Observation Schedule; ASD = autism spectrum disorder; ATYP = atypical development; DQ = Developmental Quotient; MSEL = Mullen Scales of Early Learning; TD = typical development.

<sup>a, b, c</sup> Within each row, means with different superscripts differ significantly, at least at p<.05 based on post hoc differences corrected for multiple tests using Tukey’s honest significant difference test for continuous variables and Bonferroni correction for categorical variables.
Table 2

Composition of the Subgroups Based on Classification and Regression Tree (CART) Analysis at 18 Months

<table>
<thead>
<tr>
<th>ADOS-G items predictive of ASD</th>
<th>Leaf Label</th>
<th>ASD (n, %)</th>
<th>ATYP (n, %)</th>
<th>TH (n, %)</th>
<th>Total Leaf Size</th>
<th>Relative Risk for ASD (95% Confidence Interval)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B1: Appropriate eye contact</td>
<td>D4: No RSB</td>
<td>A: Normal intonation</td>
<td>15 (12%) [7%]</td>
<td>36 (26%) [16%]</td>
<td>169 (55%) [77%]</td>
<td>220 [100%]</td>
</tr>
<tr>
<td></td>
<td></td>
<td>A: Somewhat odd intonation</td>
<td>2 (2%) [15%]</td>
<td>6 (4%) [32%]</td>
<td>2 (1%) [18%]</td>
<td>11 [100%]</td>
</tr>
<tr>
<td>D4: Some RSB</td>
<td>B8: Good or limited giving</td>
<td>C: Normal giving</td>
<td>20 (16%) [15%]</td>
<td>43 (31%) [32%]</td>
<td>71 (23%) [53%]</td>
<td>134 [100%]</td>
</tr>
<tr>
<td>B1: Poor eye contact</td>
<td>B8: No giving</td>
<td>D: ASD</td>
<td>7 (6%) [70%]</td>
<td>2 (1%) [20%]</td>
<td>1 (0%) [10%]</td>
<td>10 [100%]</td>
</tr>
<tr>
<td>B8: Good giving</td>
<td>C2: Spontaneous pretend play</td>
<td>E: TD</td>
<td>2 (2%) [4%]</td>
<td>15 (11%) [33%]</td>
<td>29 (10%) [63%]</td>
<td>46 [100%]</td>
</tr>
<tr>
<td>C2: No spontaneous pretend play</td>
<td>F: ASD</td>
<td>12 (10%) [39%]</td>
<td>7 (5%) [23%]</td>
<td>12 (4%) [39%]</td>
<td>31 [100%]</td>
<td>1.79 [1.12 – 2.87]</td>
</tr>
<tr>
<td>B8: Limited Giving</td>
<td>A8: Variety of gestures</td>
<td>G: TD</td>
<td>13 (11%) [33%]</td>
<td>12 (9%) [30%]</td>
<td>15 (5%) [38%]</td>
<td>40 [100%]</td>
</tr>
<tr>
<td>A8: Limited use of gestures</td>
<td>H: ASD</td>
<td>50 (41%) [69%]</td>
<td>17 (12%) [23%]</td>
<td>6 (2%) [8%]</td>
<td>73 [100%]</td>
<td>3.17 [2.54 – 3.96]</td>
</tr>
</tbody>
</table>

Total Outcome Group           | 122 (100%) | 138 (100%) | 305 (100%) | 565 | 565 | 1.00 [0.96 – 1.04] |

Note: n = 565; 20% of original sample removed for the validation set. Column marginals (proportion of all children of a particular outcome at 36 months, as given by the column header, found in each leaf) are given in parentheses (i.e., while row marginals (proportion of children within that leaf with an ultimate diagnosis given by the column header) are given in square brackets []). Relative risk estimates for autism spectrum disorder (ASD) are based on prevalence of ASD amongst all high-risk (HR) siblings in this sample (risk compared to the general population is 14.7 times greater). ADOS = Autism Diagnostic Observation Schedule; ATYP = atypical development; RSB = repetitive and stereotyped behaviors; TD = typical development.
Table 3
Characterization of Subgroups of High-Risk Siblings Based on Classification and Regression Tree (CART) Analysis at 18 Months

<table>
<thead>
<tr>
<th>Leaf (Label)</th>
<th>n</th>
<th>% Boys</th>
<th>Age of recruitment (months)</th>
<th>Age of assessment (months)</th>
<th>Severity 18m</th>
<th>Severity 36m</th>
<th>VDQ 18m</th>
<th>VDQ 36m</th>
<th>NVDQ 18m</th>
<th>NVDQ 36m</th>
</tr>
</thead>
<tbody>
<tr>
<td>A (TD)</td>
<td>220</td>
<td>44.1</td>
<td>7.4 (4.6)</td>
<td>18.3 (.6)</td>
<td>1.5 (9)</td>
<td>2.1*** (1.7)</td>
<td>100 (28)</td>
<td>104 (16)</td>
<td>105 (11)</td>
<td>109 (16)</td>
</tr>
<tr>
<td>B (ATYP)</td>
<td>11</td>
<td>72.7</td>
<td>7.2 (2.3)</td>
<td>18.2 (.4)</td>
<td>1.6 (1.2)</td>
<td>4.6** (1.9)</td>
<td>108 (7)</td>
<td>106 (15)</td>
<td>111 (12)</td>
<td>105 (13)</td>
</tr>
<tr>
<td>C (TD)</td>
<td>134</td>
<td>63.9</td>
<td>7.5 (3.7)</td>
<td>18.3 (.5)</td>
<td>1.9 (1.1)</td>
<td>2.9*** (2.4)</td>
<td>97 (20)</td>
<td>101 (17)</td>
<td>105 (12)</td>
<td>103 (16)</td>
</tr>
<tr>
<td>D (ASD)</td>
<td>10</td>
<td>90.0</td>
<td>9.2 (4.7)</td>
<td>18.2 (.6)</td>
<td>3.2 (1.5)</td>
<td>6.4** (2.8)</td>
<td>83 (25)</td>
<td>84 (28)</td>
<td>93 (19)</td>
<td>90 (29)</td>
</tr>
<tr>
<td>E (TD)</td>
<td>46</td>
<td>58.7</td>
<td>7.8 (3.4)</td>
<td>18.4 (.5)</td>
<td>3.7 (1.4)</td>
<td>2.7 (2.2)</td>
<td>85 (17)</td>
<td>104*** (14)</td>
<td>102 (8)</td>
<td>105 (16)</td>
</tr>
<tr>
<td>F (ASD)</td>
<td>31</td>
<td>71.0</td>
<td>7.4 (3.0)</td>
<td>18.5 (.6)</td>
<td>4.8 (1.7)</td>
<td>4.1 (2.7)</td>
<td>85 (24)</td>
<td>93 (28)</td>
<td>102 (10)</td>
<td>97 (19)</td>
</tr>
<tr>
<td>G (TD)</td>
<td>40</td>
<td>57.9</td>
<td>7.9 (3.3)</td>
<td>18.4 (.5)</td>
<td>4.6 (1.3)</td>
<td>4.1 (2.4)</td>
<td>84 (16)</td>
<td>99** (19)</td>
<td>105 (12)</td>
<td>98 (17)</td>
</tr>
<tr>
<td>H (ASD)</td>
<td>73</td>
<td>74.0</td>
<td>8.1 (3.8)</td>
<td>18.5 (.7)</td>
<td>6.1 (1.9)</td>
<td>6.1 (2.6)</td>
<td>62 (21)</td>
<td>77** (26)</td>
<td>92 (13)</td>
<td>84 (21)</td>
</tr>
</tbody>
</table>

Note: Post hoc comparisons between scores at 18 and 36 months with Bonferroni correction for multiple comparisons. Variables associated with significantly different comparisons are boldface within each row. Excepting n and % Boys, variables are reported as means and standard deviation: m(SD). ASD = autism spectrum disorder; ATYP = atypical development; NVDQ = nonverbal developmental quotient; TD = typical development; VDQ = verbal developmental quotient.

*** p<.0001,
** p<.001