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Physiological Arousal in Autism and Fragile X Syndrome: Group Comparisons and Links

With Pragmatic Language

Jessica Klusek, Gary E. Martin, and Molly Losh

Abstract

This study tested the hypothesis that pragmatic (i.e., social) language impairment is linked to arousal dysregulation in autism spectrum disorder (ASD) and fragile X syndrome (FXS). Forty boys with ASD, 39 with FXS, and 27 with typical development (TD), aged 4–15 years, participated. Boys with FXS were hyperaroused compared to boys with TD but did not differ from boys with ASD. Dampened vagal tone predicted pragmatic impairment in ASD, and associations emerged between cardiac activity and receptive/expressive vocabulary across groups. Findings support autonomic dysfunction as a mechanism underlying pragmatic impairment in ASD and suggest that biophysiological profiles are shared in ASD and FXS, which has implications for understanding the role of fragile X mental retardation-1 (FMR1, the FXS gene) in the pathophysiology of ASD.

Key Words: autism; ASD; fragile X syndrome; arousal; vagal tone; heart rate; pragmatic language; social communication; endophenotype
Fragile X syndrome (FXS) is a monogenic disorder associated with significantly increased risk for autism spectrum disorder (ASD; Cohen, Pichard, & Tordjman, 2005). Pragmatic language impairment is seen in both ASD and FXS (Landa, 2000; Losh, Martin, Klusek, Hogan-Brown, & Sideris, 2012; Sudhalter & Belser, 2001), yet it is unknown whether such impairments arise from similar underlying impairments. Physiological dysregulation (e.g., elevated heart rate and differences in the functioning of the parasympathetic “rest and restore” system) has been documented in FXS and idiopathic ASD and, in FXS, has been hypothesized to be a causal factor in pragmatic language impairments (e.g., Belser & Sudhalter, 1995). The present study investigated the role of physiological dysregulation in pragmatic language deficits in ASD and FXS, examining physiological arousal in relation to clinically based and standardized assessments of pragmatic language across groups. Further understanding of arousal, a biophysiological marker for stress, as a mechanism linked with pragmatic language impairments of ASD and FXS may have implications for the development of targeted interventions and lend insight into shared biological pathways in ASD and FXS that can be traced back to the gene that causes FXS, *fragile X mental retardation-1* (*FMR1*).

**Genetic Basis of ASD and FXS**

ASD is characterized by atypical social and communicative development along with repetitive and restricted behavioral patterns (American Psychiatric Association, 2013). Affecting approximately 1 in 88 individuals, ASD is seen at high levels, and there is an urgent need to understand the etiological basis of the disorder (CDC, 2012). Evidence supports a strong genetic component in the etiology of ASD although the exact genetic underpinnings still remain undefined (Devlin & Scherer, 2012). The genetic basis of ASD is thought to be heterogeneous and complex with many different gene-gene and gene-environment interactions leading to the
common phenotypic endpoint of ASD (Abrahams & Geschwind, 2008). Single-gene disorders, such as FXS, are implicated in about 10% of cases of ASD (Betancur, 2011). The study of ASD within the context of associated genetic conditions provides a better-understood genetic paradigm for studying ASD, which may provide a starting point for pinning down pathophysiological mechanisms (Abrahams & Geschwind, 2008; Hagerman, Hoem, & Hagerman, 2010).

While the etiological basis of ASD is complex and heterogeneous, FXS can be traced back to a single genetic cause: a trinucleotide expansion on the *FMR1* gene (Pieretti et al., 1991). This expansion silences the gene and halts the production of fragile X mental retardation protein (FMRP), which is highly expressed in the brain and is thought to play a role in synaptic development (Hagerman & Hagerman, 2002). Deficiency in FMRP appears to underlie the neurobehavioral profile of FXS, which includes intellectual disability, language impairment, social difficulties, anxiety, and hyperactivity (Baumgardner, Reiss, Freund, & Abrams, 1995; Hagerman, 2002; Reiss & Dant, 2003).

Strikingly, 50%–75% of individuals with FXS meet criteria for ASD, and those who do not reach diagnostic thresholds nevertheless show symptoms consistent with ASD, such as reduced eye gaze and repetitive behaviors (Hagerman et al., 1986; Hall, Lightbody, & Reiss, 2008; Harris et al., 2008). This significantly elevated risk for ASD suggests that the *FMR1* mutation may play a role in the development of autistic symptoms, potentially through interactions with other genes that are involved in ASD (Hagerman, Au, & Hagerman, 2011). For example, FMRP assists in the translation of several proteins that are dysregulated in idiopathic ASD (e.g., neuroexin, CYFIP, PTEN), and the absence of FMRP in FXS has a detrimental effect on the normal expression of other genes (see Hagerman et al., 2010). In fact, a large number of
autism susceptibility genes are known interactors with \textit{FMR1} (Darnell & Klann, 2013; Darnell et al., 2011; Iossifov et al., 2012). The \textit{FMR1} mutation may therefore disrupt the normal function of a number of autism susceptibility genes, lowering the threshold of interacting genetic effects needed to produce ASD. In this way, FXS provides a simplified genetic model that may be useful in identifying genetic or molecular pathways implicated in ASD.

\textbf{Pragmatic Language in ASD and FXS}

Evidence suggests that individuals with idiopathic and FXS-associated ASD show similar symptom profiles on omnibus measures of autism symptomatology (Dissanayake, Bui, Bulhak-Paterson, Huggins, & Loesch, 2009; Rogers, Wehner, & Hagerman, 2001). Specific features of the language profiles associated with ASD, such as pragmatic language impairment, also appear to be shared in idiopathic ASD and ASD within the context of FXS (Losh, Martin, et al., 2012). Pragmatic language is defined as the use of language in social contexts to communicate meaning (Bates, 1976; McTear & Conti-Ramsden, 1992; Prutting, 1982). Pragmatic language difficulties are a universally observed feature of ASD (Landa, 2000; Tager-Flusberg, Paul, & Lord, 2005). For example, turn-taking, topic maintenance, and communicative repair skills are deficient in ASD (Adams, Green, Gilchrist, & Cox, 2002; Capps, Kehres, & Sigman, 1998; Geller, 1998; Paul et al., 1987; Tager-Flusberg & Anderson, 1991; Volden, 2004). Atypical pragmatic features such as echolalia, perseveration, and bizarre word choice are seen at increased rates in ASD (Ghaziuddin & Leonore, 1996; Ross, 2002; Schuler & Prizant, 1985), and narrative (storytelling) skills are impaired as well (Capps, Losh, & Thurber, 2000; Diehl, Bennetto, & Young, 2006; Losh & Capps, 2003; Loveland, McEvoy, & Tunali, 1990; Tager-Flusberg, 1995). Differences in pragmatic language use are thought to be a genetically meaningful feature of ASD, given that subclinical pragmatic differences present at increased rates among unaffected relatives of
individuals with ASD (Landa et al., 1992; Losh, Childress, Lam, & Piven, 2008) and show patterns suggestive of intrafamilial transmission (Klusek, Losh, & Martin, 2012).

Pragmatic language deficits are also seen in FXS. For example, conversation in FXS is characterized by impaired topic maintenance (Roberts et al., 2007; Sudhalter & Belser, 2001; Sudhalter, Cohen, Silverman, & Wolf-Schein, 1990; Wolf-Schein et al., 1987), poor ability to repair communicative breakdowns (Abbeduto et al., 2008), and stereotyped and perseverative language (Belser & Sudhalter, 2001; Martin et al., 2012; McDuffie et al., 2010; Roberts et al., 2007; Sudhalter et al., 1990; Wolf-Schein et al., 1987). Importantly, children with idiopathic ASD and FXS with comorbid ASD perform comparably on standardized pragmatic language assessments, and pragmatic language abilities are associated with FMR1-related genetic variation in FXS (Losh, Martin, et al., 2012). Furthermore, recent research suggests that pragmatic language profiles overlap in the broad autism phenotype and the FMR1 premutation. Female carriers of the FMR1 premutation and mothers of individuals with ASD exhibit conversational pragmatic language difficulties that are elevated in comparison to controls and which are not only similar in severity but also show qualitative overlap as evidenced by similar performance on pragmatic subdomains (see Losh, Klusek, et al., 2012). Together, these studies suggest that FMR1 may be involved in pragmatic language impairments associated with ASD.

Although significant overlap in pragmatic language deficits in ASD and FXS has been documented (along with overlapping pragmatic language profiles in unaffected genetic carriers in FXS and ASD, i.e., parents), it is unclear whether the pragmatic impairments in these groups stem from common underlying factors; only a handful of studies have directly compared neurobiological characteristics in these disorders, and very few have related those features to pragmatic language. While arousal dysregulation has also been proposed as a mechanism
underlying social deficits in idiopathic ASD (e.g., Dawson & Lewy, 1989), most theories of ASD have focused on neurocognitive models to account for pragmatic language impairment, such as impaired social cognition or central coherence (Martin & McDonald, 2003). On the other hand, several studies have proposed arousal dysregulation as a cause of pragmatic language difficulties in FXS (Belser & Sudhalter, 1995; Cohen, 1995). In this model, arousal dysmodulation causes an individual to remain “on edge” during social situations, over time leading to social withdrawal and limiting opportunities to learn skills through interaction with others (Rubin & Burgess, 1991). Given that physiological regulatory deficits have also been documented in ASD (Bal et al., 2010; Ming, Julu, Brimacombe, Connor, & Daniels, 2005), this study explored physiological dysregulation as a predictor of pragmatic language deficits in both disorders.

**Physiological Arousal in ASD and FXS**

The present study focused on cardiac indices of physiological arousal: heart rate (a measure of general arousal) and respiratory sinus arrhythmia (RSA), an index of parasympathetic control of the heart via the vagal nerve (Porges, 2007). These indices of physiological functioning are thought to mark one’s capacity for engagement with the social environment (Porges & Furman, 2011).

**Heart rate.** Increased general arousal, indexed by elevated heart rate, is a hallmark feature of FXS (Heilman, Harden, Zageris, Berry-Kravitz, & Porges, 2011; Roberts, Boccia, Bailey, Hatton, & Skinner, 2001; Roberts, Tonnsen, Robinson, & Shinkareva, 2012). It is unclear whether heart rate is also elevated in ASD as there is great inconsistency across prior reports, possibly due to differences in sample characteristics and the heterogeneity that is characteristic of ASD. Increased heart rate has been detected in ASD in comparison to typical and
developmentally delayed comparison groups (Bal et al., 2010; Goodwin et al., 2006; Mathewson et al., 2011; Woodard et al., 2012) although a number of other reports have found heart rate in ASD to be similar to that of controls (Althaus, Mulder, Mulder, Aarnoudse, & Minderaa, 1999; Corona, Dissanayake, Arbelle, Wellington, & Sigman, 1998; Sigman, Dissanayake, Corona, & Espinosa, 2003). Hyperarousal is hypothesized to underlie communication deficits (e.g., Belser & Sudhalter, 1995; Belser & Sudhalter, 2001) although few studies have directly examined this hypothesis in ASD or FXS. One study of children with ASD found that elevated heart rate was associated with reduced use of communicative gestures (Patriquin, Scarpa, Friedman, & Porges, 2011). Another preliminary report examining two males with FXS found that perseverative and tangential language was associated with increased arousal as indexed by skin conductance responses (Belser & Sudhalter, 1995).

**Vagal tone.** The vagal nerve works as part of the parasympathetic “rest and restore” system, counteracting sympathetic “fight or flight” excitation to promote a physiological state that facilitates social engagement (see Porges & Furman, 2011). Numerous studies of typical development link high vagal tone with enhanced social skills (e.g., Blair & Peters, 2003; Calkins & Keane, 2004; Kok & Fredrickson, 2010). Reduced vagal tone is well documented in FXS (Hall, Lightbody, Huffman, Lazzeroni, & Reiss, 2009; Heilman et al., 2011; Roberts et al., 2001; Roberts et al., 2012). Some investigations of ASD have also detected dampened vagal tone (Bal et al., 2010; Mathewson et al., 2011; Ming et al., 2005; Porges et al., 2013; Van Hecke et al., 2009) although reports are inconsistent (Althaus et al., 1999; Levine et al., 2012; Toichi & Kamio, 2003). Diminished vagal tone has been shown to relate to poorer receptive vocabulary and social skills in children with ASD (Patriquin et al., 2011; Van Hecke et al., 2009) and with more severe autistic traits in toddlers with FXS (Roberts et al., 2012).
In addition to examining vagal activity at rest, the measurement of vagal tone specifically during social-communicative interactions can shed light on physiological adaption to social demands. In typical development (TD), children who show the greatest increases in vagal tone in response to social contexts have better receptive and expressive language abilities, and children who exhibit vagal withdrawal are more likely to present with anxiety, depression, and internalizing problems (Heilman et al., 2008; Suess & Bornstein, 2000). Few studies have investigated physiological profiles of ASD and FXS during social-communicative contexts in order to determine how these profiles might relate to atypical pragmatic language features. In one study of children with FXS, Hall et al. (2009) found that vagal tone during conversation with an examiner did not relate to the extent of gaze avoidance exhibited by the children although the authors did not measure other pragmatic features. In another study of young children with ASD, vagal tone during a social listening task (child-directed speech) prospectively predicted later parent-reported social-communication abilities, and baseline vagal tone was predictive of later skills (Watson, Baranek, Roberts, David, & Perryman, 2010). In sum, little is known about the role of the physiological system in pragmatic language deficits in ASD and FXS, and no studies, to our knowledge, have directly compared cardiac indicators of physiological profiles in ASD and FXS, limiting our understanding of biophysiological pathways that may be shared in ASD and FXS and could be linked to similar behavioral endpoints (such as pragmatic language).

**Study Rationale and Hypotheses**

This study addressed the hypothesis that physiological dysregulation in ASD and FXS is linked with pragmatic language impairment. The study addressed two primary questions:

1. *Do cardiac indicators of physiological activity differ in boys with ASD, FXS, and TD during resting and conversational contexts?* Prior to examining associations between arousal and
pragmatic language in each group, it was important to establish profiles of physiological activity and determine any differences and similarities across groups. To do so, we examined group differences in heart rate and vagal tone measured at rest and during unstructured conversation with an examiner.

2. *Do cardiac indicators of physiological activity predict pragmatic language impairment in ASD, FXS, and TD?* We examined within-group associations between physiological activity (at rest and during conversation) and pragmatic language ability, measured in separate contexts. We also examined associations with behavioral symptoms of anxiety, which could importantly relate to both cardiac activity and pragmatic language impairment.

**Method**

**Participants**

Participants included 40 school-aged boys with idiopathic ASD, 39 boys with full-mutation FXS, and 28 boys with TD, aged 4–15 years. Participants were drawn from a larger pool of children participating in ongoing longitudinal studies of speech/language in FXS and ASD, which have been described previously (see Losh et al, 2012; Zajac, Harris, Roberts, & Martun, 2009). Given the longitudinal design of the larger studies, in some instances an individual participant had available data from several different time points. In these cases, the time point was selected that best facilitated group-level matching on chronological age given that age has a known impact on cardiac activity (Alkon et al., 2003; Bar-Haim, Marshall, & Fox, 2000). All participants spoke English as their primary language and were regularly using sentences of three or more words. Only boys participated in the study because girls with FXS are generally less severely affected and show more heterogeneous profiles (Hagerman, 2004). Recruitment was focused in the Eastern and Midwestern regions of the United States through local advertisement and through
the Research Participant Registry Core of the Carolina Institute for Developmental Disabilities at the University of North Carolina at Chapel Hill.

Group characteristics are presented in Table 1. On average, the FXS group was older than the ASD and TD groups, and the mean age of the ASD and TD groups were similar; chronological age was controlled for statistically in group comparisons. Nonverbal mental age, as measured with the Leiter International Performance Scale-Revised (Leiter-R; Roid & Miller, 1997), was similar in the ASD and TD groups while the boys with FXS had a significantly lower mean mental age than the other groups. Receptive and expressive vocabulary differed across groups as measured with the Peabody Picture Vocabulary Test-III (PPVT; Dunn & Dunn, 1997) and Expressive Vocabulary Test (EVT; Williams, 1997) with FXS showing the lowest vocabulary skills, followed by the ASD group. As expected, the TD group had higher vocabulary skills than the ASD and FXS groups. The groups did not differ in race, household income, or maternal education (ps > .135). Thirty percent (12/40) of the boys with ASD and 41% (16/39) of the boys with FXS were reported by their caregivers to be using psychoactive medications at the time of assessment (see Table 1).

**Procedures**

All procedures were approved by the Institutional Review Boards of the University of North Carolina at Chapel Hill and Northwestern University. Testing took place in a research laboratory or the participant’s home or school. Assessments were conducted within the context of a broader research protocol, which generally lasted 4–6 hr. In general, structured assessments that required simple, nonverbal responses (i.e., pointing), such as the PPVT, were administered first to allow some “warm up” time, followed by more interactive tasks, such as the physiological assessment. However, the protocol was flexible, and the order of assessments and number of breaks were
adapted to the needs of the participant. The assessments were conducted by two trained examiners who were experienced in working with children with developmental disabilities.

Characterization of ASD. The Autism Diagnostic Observation Schedule (ADOS; Lord, Rutter, DeLavore, & Risi, 2001) was used to confirm ASD in the ASD group, to rule out ASD in the TD group, and to determine ASD status in the FXS group. The ADOS was also used as a continuous measure of autism symptom severity as described by Gotham, Pickles, and Lord (2009). The ADOS was administered concurrently for 36 participants with ASD, 30 with FXS, and 20 with TD. The remaining participants were administered the ADOS within one year of the time of the main assessment. Nonconcurrent ADOS data were used to confirm diagnostic status but were not included in correlational analyses in order to prevent confounds related to potential changes in symptom severity over time. The “autism spectrum” cutoffs of the revised diagnostic algorithms were used to determine the presence of ASD (Gotham et al., 2008; Gotham, Risi, Pickles, & Lord, 2007). The ADOS was scored by examiners who met standard requirements for research reliability. Mean inter-rater reliability was calculated on a randomly selected subset (13%) of participants, and reliability was 86% for individual items, 85% for diagnostic algorithm items, and 93% for diagnostic classification. Thirty-two of the boys with FXS met criteria for ASD.

Cognitive assessment. Nonverbal cognition was assessed with the Brief IQ Composite of the Leiter-R (Roid & Miller, 1997). Due to time constraints, Leiter-R data were unable to be collected for seven boys with ASD, and scores from the Performance IQ scale of the Wechsler Abbreviated Scale of Intelligence (Wechsler, 1999) were used as a substitute measure of nonverbal intelligence.
Receptive and expressive language assessment. The PPVT (Dunn & Dunn, 1997) was employed to assess receptive vocabulary. Two boys with ASD received the Peabody Picture Vocabulary Test-IV (Dunn & Dunn, 2007). Expressive vocabulary was measured with the EVT (Williams, 1997). Age-equivalent scores for the PPVT and EVT were used in analysis. Two boys with ASD and one boy with TD were missing PPVT data; the EVT was missing for four boys with ASD and one boy with TD.

Pragmatic language assessment. The Pragmatic Judgment subtest of the Comprehensive Assessment of Spoken Language (CASL-PJ; Carrow-Woolfolk, 1999) was administered as a measure of the knowledge and use of language in social contexts. In this standardized assessment, participants answer questions about what should be said or done in various social situations. CASL-PJ scores were not collected for the TD group because this was not an aim of the larger study from which the boys were recruited. CASL-PJ scores were missing for seven participants with ASD and two participants with FXS due to time constraints; one participant with ASD obtained a raw score of “0,” and scores for this participant were considered missing. Age-equivalent scores were used in analysis.

The Pragmatic Rating Scale-School Age (PRS-SA; Landa, 2011) was administered as a semi-naturalistic measure of conversational pragmatic language abilities. The PRS-SA consists of 33 items that sample a range of pragmatic features, such as the ability to initiate topics, the provision of necessary background information, or the use of appropriate rate and volume of speech. Items are tallied to obtain a total score ranging from 0 to 66 with a higher score indicating greater impairment. The PRS-SA was scored from video, based on interaction that occurred during the ADOS. The PRS-SA was only available for those participants who had been administered the ADOS concurrently (i.e., the PRS-SA was missing for four participants with
ASD, nine with FXS, and eight with TD). Five children from the ASD group and six from the FXS group were administered the ADOS module 2 (for use with individuals who have phase speech), and the remaining children were administered the ADOS module 3 (for verbally fluent individuals); there were no significant differences between the PRS-SA scores of children who had been administered module 2 versus module 3 within each group. The first author (JK), who had achieved reliability with the developer of the PRS-SA, coded all of the samples. The coder was blind to the diagnosis of 46% of the participants with ASD, 70% with FXS, and 90% with TD (the coder had assisted in participant assessment, and it was not possible to maintain blinding to all participants). Twenty percent of the sample was randomly selected and second-scored by a blind, independent rater who had also achieved reliability with the developer of the PRS-SA. Intraclass correlations were computed and inter-rater reliability (ICC [3, 2]) was 0.97 (0.91 for ASD, 0.91 for FXS, and 0.94 for TD).

**Physiological assessment.** Heart activity data were collected during a baseline resting condition and a conversational condition. During baseline, participants watched 10 min of an animated children’s film. Movie watching was chosen as the baseline task because it was thought that the disability groups would be unable to complete a more traditional baseline task (sitting alone quietly); this approach is consistent with several other physiology studies of developmental disabilities (e.g., Hall et al., 2009; Roberts et al., 2011; Roberts et al., 2001). Immediately following the baseline period, a conversation condition took place in which participants conversed with an examiner about any topic of interest for 10 min. In order to obtain a naturalistic sample, examiners maintained the conversation by commenting, asking questions, and bringing up topics of possible interest to the participant. No specific demands were placed on the participants other than that they remained seated and did not touch the electrodes (i.e.,
participants were not specifically directed to maintain eye contact or instructed to act in a certain way. Heart rate was not measured during the ADOS (the conversational sample that served as the basis for the PRS-SA) because of concerns that wearing the electrodes might influence participants’ behavior in ways that could lead to invalid ASD characterization. Conversely, it was not possible to rate the PRS-SA directly from the physiology conversational condition due to the brevity of the sample (the PRS-SA was developed for use with a 30–60 min conversational sample).

Electrocardiogram data was collected with an Alive Wireless Heart Monitor (Alive Technologies, Copyright 2005–2009), either via two electrodes that were placed on the participant’s chest or with an elastic Polar belt that contained electrode receptors (the belt was presented as an option for participants who refused the electrodes). Data were sampled at a rate of 300 times per second. Twelve boys with FXS and three with ASD (not included in sample ns) were recruited for the study, but data collection was unable to be completed due to random equipment malfunction, excessive movement/recording artifacts (>20%), or uncooperativeness. Electrocardiogram data from the last 5.5 min of baseline and the first 8 min of conversation were analyzed. This portion of the baseline was selected in order to obtain a “pure” resting physiological estimate by allowing time for the participant to acclimate to the sensation of the electrodes. Only the first 8 min of the conversational condition were analyzed to ensure equal sample time across participants (occasionally, the sample ended premature of the full 10 min due to participant uncooperativeness). Data were edited with CardioEdit software (Brain-Body Center, University of Illinois at Chicago) by a reliable research assistant. Editing consisted of visual inspection of the electrocardiogram signal to identify invalid heart periods (e.g., faulty R wave detection) and manual adjustment of artifacts using integer arithmetic. Estimates for RSA
and interbeat interval (IBI) were extracted. Briefly, IBI is measured as the time in milliseconds between successive R waves. To extract RSA, sequential heart periods are sampled at 250-ms epochs, and data is detrended with a 21-point moving polynomial algorithm (Porges & Bohrer, 1990). Data is then bandpass filtered to extract variance associated with spontaneous breathing parameters (0.24–1.04 Hz), and the variance is transformed to its natural logarithm to yield an estimate of RSA. The average RSA and IBI for each condition, measured from 30-s epochs, were used in analyses. Change scores, measuring reactivity, were computed by subtracting conversational estimates from baseline.

**Anxiety.** Symptoms of anxiety were assessed with the Child Behavior Checklist-1½–5 years (CBCL; Achenbach & Rescorla, 2000), which was completed by the primary caregiver. The Diagnostic and Statistical Manual of Mental Health Disorders-IV (DSM-IV)–oriented anxiety subscale was used, and it corresponds to the diagnostic criteria for anxiety outlined in the DSM-IV (American Psychiatric Association, 1994). The preschool version of this assessment was administered as the younger items were judged to be more closely aligned with the mental ages of the study participants. Raw scores were used in analysis in accordance with the recommendations of the test publishers for use in research (Achenbach & Rescorla, 2000). These data were missing for eight participants with ASD, 13 participants with FXS, and seven participants with TD.

**Data Analysis**

Group differences in IBI and RSA were tested with repeated measures analysis of covariance (ANCOVA) with condition (baseline, conversation) as a within-participants factor, controlling for chronological age. ANCOVA was also conducted to test group differences in the magnitude of IBI and RSA change, controlling for chronological age. Next, the impact of autism symptom
severity on cardiac activity was explored by (a) examining the association between the cardiac indices and continuously distributed autism symptom severity (ADOS severity score) with Spearman correlations, and (b) by examining means descriptively among ASD subgroups of the FXS group. Given the limited sample size of children with FXS without comorbid ASD (n = 7), group differences were not tested statistically. Spearman correlations were used to examine the relationship between the six cardiac variables (IBI and RSA estimates for baseline, conversation, and change) and clinical symptoms of anxiety, and group differences in the severity of anxiety were tested with ANOVA.

Analyses were then conducted to test cardiac activity as a predictor of pragmatic language ability across groups. First, Spearman correlations were used to explore relationships between the six cardiac variables, receptive vocabulary (PPVT), expressive vocabulary (EVT), and pragmatic language (CASL-PJ and PRS-SA) within each group. These exploratory analyses were conducted in order to guide more focused analyses examining cardiac activity as a predictor of pragmatic language, given that pragmatic language ability may be influenced by receptive/expressive language skills (McTear & Conti-Ramsden, 1992). Linear regression models were used to test cardiac activity as a predictor of pragmatic language after accounting for receptive/expressive vocabulary (PPVT and EVT).

Results

Group Differences in Cardiac Activity and the Impact of Autism

Descriptive statistics for the physiological variables are presented in Table 2. Examination of individual patterns of physiological activity showed that 21/40 (52.5%) of the participants with ASD, 28/39 (71.8%) with FXS, and 15/28 (53.6%) with TD increased RSA from baseline to conversation. Most participants decreased IBI (i.e., increased heart rate) in conversation; 82.5%
(33/40) of the participants with ASD, 82.1% (32/39) with FXS, and 92.9% (26/28) with TD increased heart rate during conversation.

**Group differences in heart rate.** Group membership had a significant main effect on IBI ($F[2, 103] = 3.42, p = .036, \eta^2 = .06$). Follow-up Bonferroni-corrected pair-wise comparisons showed that the FXS group had significantly lower IBI (i.e., faster heart rate) than the TD group ($p = .040$) but did not differ from the ASD group ($p = .173$). The mean IBIs of the ASD and TD groups did not differ ($p = .349$). A main effect was detected for condition with lower IBI (i.e., faster heart rate) during conversation than at baseline ($F[1, 103] = 4.93, p = .029, \eta^2 = .05$). The interaction between group and condition was not significant ($p = .855$), indicating that all groups showed similar increases in heart rate in response to the conversational task. The magnitude of IBI change did not differ across groups ($F[3, 107] = 0.67, p = .562, \eta^2 = .01$).

**Group differences in vagal tone.** The effect of the group and its interaction with condition had nonsignificant effects on RSA ($ps > .690$). The main effect of condition approached significance ($F[1, 103] = 2.87, p = .093, \eta^2 = .03$) with a trend for higher vagal tone during the conversational condition. The magnitude of RSA change did not differ across groups ($F[3, 107] = 1.26, p = .161, \eta^2 = .04$).

**Cardiac activity in ASD subgroups of FXS.** Although the small number of participants with FXS without comorbid ASD (FXS-only, FXS-O) prevented group differences from being tested statistically, cardiac activity was examined descriptively across the FXS-O and FXS with comorbid ASD (FXS-ASD) groups, controlling for chronological age. Age-adjusted group means for IBI are depicted in Figure 1. Overall, IBI in the FXS-ASD subgroup was lower than that of the FXS-O group (indicating faster heart rate in the FXS-ASD group). The FXS-ASD
group had the lowest mean IBI of all of the groups, and the IBI estimates of the FXS-O group were more similar to that of the ASD group.

Age-adjusted group estimates for RSA are presented in Figure 2. The boys with FXS-O had higher mean RSA in comparison with all other groups across both conditions. In contrast, the RSA of the boys with FXS-ASD was the lowest of all of the groups at baseline with a sharp increase in response to the conversation condition. During conversation, RSA of the FXS-ASD group was similar to that of the TD group.

**Relationship between cardiac activity and autism symptom severity.** Autism severity was not associated with any of the physiological variables in the ASD ($p_s > .286$) or FXS ($p_s > .228$) groups. In the TD group, continuously distributed symptoms of autism were significantly associated with RSA at baseline ($\rho = -.58, p = .007$) and during conversation ($\rho = -.57, p = .009$) with lower vagal tone associated with increased autism symptom severity.

**Relationship between cardiac activity and symptoms of anxiety.** ANOVA indicated a significant group effect for the CBCL-anxiety subscale ($F[2, 76] = 5.49, p = .006$). Bonferroni-corrected post-hoc comparisons showed that, whereas both disability groups exhibited more anxiety than the boys with TD ($p_s < .027$), the FXS and ASD groups did not differ from each other ($p = 1.00$). No significant associations were detected between anxiety and any of the cardiac variables across groups ($p_s > .136$). Anxiety symptoms were also not associated with either pragmatic language measures across the groups ($p_s > .092$).

**Heart Rate and Pragmatic Language Ability**

**Associations between heart rate and pragmatic language.** Exploratory correlations between the IBI variables and pragmatic language are presented in Table 3. Among the boys with ASD, performance on the CASL-PJ was associated with baseline and change in IBI,
indicating that pragmatic language skills got worse with higher heart rate and less increase in arousal in response to conversation. In the FXS group, a marginal association between IBI change and pragmatic language performance on the PRS-SA was detected in the opposite direction \((p = .052)\) with more substantial arousal change (i.e., greater arousal in response to the conversational condition) associated with worse pragmatic language skills. In the boys with TD, no significant correlations were observed between the IBI and pragmatic language variables although a trend-level association between conversational IBI and pragmatic performance on the PRS-SA was detected \((p = .073)\) with greater arousal during conversation linked with worse pragmatic skills.

Exploratory correlations between IBI and expressive and receptive vocabulary are presented in Table 3 in order to inform specific associations with pragmatics language versus general language skills. In the ASD group, IBI change was significantly associated with performance on the EVT with less change in arousal corresponding to worse expressive vocabulary skills. No other significant associations were detected with receptive/expressive vocabulary.

**Heart rate as a predictor of pragmatic language.** Significant correlations were followed up with more specific analyses testing heart rate as a predictor of pragmatic language skills after controlling for receptive/expressive vocabulary. First, baseline IBI was tested as a unique predictor of CASL-PJ performance in the ASD group; after accounting for the variance associated with receptive and expressive vocabulary, baseline IBI did not account for significant additional variance in the CASL-PJ \((R^2 = .88, R^2\Delta = .01, F\Delta [1, 26] = 0.87, p = .360)\). IBI change was also not a significant predictor of CASL-PJ performance after accounting for receptive/expressive vocabulary in ASD \((R^2 = .87, R^2\Delta = .01, F\Delta [1, 24] = 0.01, p = .971)\).
Vagal Tone and Pragmatic Language Ability

**Associations between vagal tone and pragmatic language.** In ASD, baseline RSA was associated with the CASL-PJ with lower vagal tone linked with poorer pragmatic performance. A significant association between dampened RSA during conversation and worse pragmatic skills on the PRS-SA was also detected in ASD (along with a trend-level association in the same direction with pragmatics on the CASL-PJ, $p = .061$). In the FXS group, RSA change was negatively associated with CASL-PJ performance, indicating that the boys with FXS who showed less vagal change from the baseline to the conversational task had worse performance on the standardized measure of pragmatic language. No other significant correlations were detected between the RSA variables and pragmatics although a marginal association was detected in the TD group between RSA change and the PRS-SA ($p = .097$). Correlations are presented in Table 3.

Exploratory associations between RSA and receptive and expressive vocabulary showed a negative relationship between RSA change and performance on the EVT and PPVT in the FXS group (consistent with worse vocabulary skills as the boys showed less vagal reactivity). No other significant associations were observed with receptive/expressive vocabulary.

**Vagal activity as a predictor of pragmatic language.** Linear regression was used to test conversational RSA as a predictor of PRS-SA performance in ASD; results indicated that RSA during conversation accounted for a significant proportion of variance in PRS-SA scores (11%), even after accounting for receptive and expressive vocabulary ($R^2 = .36$, $R^2\Delta = .11$, $F\Delta [1, 30] = 5.04, p = .032$). A second regression was run to test baseline RSA as a predictor of CASL-PJ scores in ASD; after accounting for receptive/expressive vocabulary skills, baseline RSA was not
a significant predictor of pragmatic performance on the CASL-PJ in ASD ($R^2 = .88$, $R^2\Delta < .01$, $F\Delta [1, 26] = 0.16$, $p = .697$).

Next, RSA change was tested as a predictor of pragmatic language on the CASL-PJ among the boys with FXS. Results showed that the combined influence of receptive/expressive vocabulary accounted for significant variance in the CASL-PJ scores in FXS ($R^2 = .69$, $F\Delta [2, 34] = 37.97$, $p < .001$), but the addition of RSA change to the model did not account for significant variance above and beyond receptive/expressive vocabulary ($R^2 = .69$, $R^2\Delta = .01$, $F\Delta [1, 33] = 0.01$, $p = .925$).

**Discussion**

This study examined cardiac arousal and parasympathetic activity as potential neurobiological markers of pragmatic language difficulties in boys with ASD, FXS, and TD. Boys with ASD and FXS showed similar physiological profiles, including similar levels of general arousal and vagal tone at rest and during conversion with an examiner. The boys with FXS were hyperaroused in comparison to the boys with TD, but their general arousal level did not differ from the boys with ASD. The arousal level of the ASD group was moderately elevated but did not differ significantly from either the FXS or TD groups. Vagal tone did not differ across groups, suggesting that parasympathetic nervous system functioning was similar across groups. Furthermore, boys with ASD and FXS showed physiological responses that were similar to those of boys with TD in response to the conversational task, indicating that conversation did not elicit atypically elevated levels of arousal in either the ASD or FXS groups. Dampened vagal tone during conversation predicted impaired pragmatic language performance in ASD even after accounting for variation in receptive and expressive vocabulary skills. Together, these findings add to emerging evidence for a role of autonomic nervous system dysfunction in pragmatic
language impairment in ASD. Results also have implications for understanding biophysiological pathways that may be shared in ASD and FXS and which may be linked to FMR1-related genetic effects.

**Group Differences in Physiological Profiles**

To our knowledge, this is the first report to directly compare cardiac indices of arousal in ASD and FXS. As a single-gene disorder that is associated with significantly elevated risk for ASD, FXS may be a useful model for reducing “genetic noise” to speed the identification of pathophysiological mechanisms implicated in ASD. In the present study, the physiological profiles of FXS did not differ from those of ASD, suggesting that biophysiological pathways that are disrupted in FXS may also be common to idiopathic ASD and, providing support for FMR1, the FXS gene as a potential candidate gene for ASD. While a body of research supports phenotypic overlap in idiopathic ASD and FXS-associated ASD (Bailey et al., 1998; Dissanayake et al., 2009; Rogers et al., 2001), the results of the present study contribute new evidence to support shared biophysiological characteristics in these disorders. Specifically, the boys with ASD and the boys with FXS did not differ on measures of general arousal or vagal tone in either condition examined. The study of the autonomic nervous system as an intermediate biological process that may be dysregulated in both FXS and ASD may help bridge the gap between behavior and genes; current genetic models suggest that FMR1 may lead to the ASD phenotype through its interaction with other “background” genes, thereby lowering the genetic threshold needed to produce ASD (Bailey, Hatton, Skinner, & Mesibov, 2001; Harris et al., 2008; Rogers et al., 2001). Given that biological mechanisms are more directly linked with underlying genetic effects than are behavioral symptoms, identifying biophysiological pathways
that are common to ASD and FXS may help uncover molecular/genetic mechanisms that interact with \textit{FMR1} to produce the ASD phenotype.

While the arousal levels of the boys with ASD and FXS did not differ from each other, only the boys with FXS showed significantly elevated arousal in comparison to the boys with TD. Descriptively, the boys with ASD showed arousal levels that were lower on average than that of the FXS group but higher on average than those of the TD boys (with neither difference statistically significant). There is great inconsistency across prior investigations of arousal in ASD with a number of reports indicating that heart rate in ASD is similar to controls (Althaus et al., 1999; Corona et al., 1998; MacCulloch & Williams, 1971; Sigman et al., 2003) and other reports suggesting that heart rate is atypically elevated (Bal et al., 2010; Goodwin et al., 2006; Mathewson et al., 2011; Woodard et al., 2012). The inconsistency across studies might suggest that elevated heart rate is only characteristic of particular subgroups of individuals with ASD, which is consistent with the “middle of the road” arousal level detected in this study. Even if arousal modulation dysfunction is evident only in a subgroup of individuals with ASD, physiological regulatory difficulties may serve as a useful marker for identifying homogeneous subgroups of ASD, helping to uncover biological systems that may respond to targeted pharmaceutical or behavioral treatments. Additional research investigating arousal modulation as a potential endophenotype (i.e., a marker of genetic susceptibility) for ASD might prove useful as measures of cardiovascular activity are noninvasive, reliable, quantitative indices that tap biologically meaningful functions that may be more closely tied to underlying genetic variation than clinical phenotypes.

The finding of elevated heart rate in FXS is consistent with a body of prior reports documenting hyperarousal in FXS (Hall et al., 2009; Heilman et al., 2011; Roberts et al., 2001;
Roberts et al., 2012). Furthermore, the results of this study suggest that hyperarousal is a chronic physiological condition in FXS as heart rate was consistently elevated across both experimental conditions. Given that the parasympathetic vagal tone of the FXS group was similar to the TD boys, it may be that the elevated heart rate in FXS was driven by increased sympathetic tone. This interpretation is consistent with evidence from skin conductance studies showing elevated sympathetic tone in FXS (Miller et al., 1999; Roberts, Mazzocco, Murphy, & Hoehn-Saric, 2008). While this study cannot directly address this hypothesis, future research might include measures of both parasympathetic and sympathetic tone to pinpoint specific areas of breakdown in the autonomic system in FXS.

Contrary to hypotheses, vagal tone in the FXS and ASD groups did not differ from that of the boys with TD. A number of prior reports have documented dampened vagal tone in FXS (Boccia & Roberts, 2000; Hall et al., 2009; Heilman et al., 2011; Roberts et al., 2001; Roberts, Boccia, Hatton, Skinner, & Sideris, 2006) whereas findings from previous studies of ASD are mixed with reports that vagal tone is either reduced (Bal et al., 2010; Ming et al., 2005; Van Hecke et al., 2009) or similar to controls (Althaus et al., 1999; Toichi & Kamio, 2003; Watson, Roberts, Baranek, Mandulak, & Dalton, 2012). The mean vagal estimates detected in this study are consistent with prior reports: At baseline, the boys with FXS showed vagal tone that was lower on average than the ASD and TD groups, who were similar to each other. It is possible that this study was underpowered to detect group effects. The sample characteristics of the present study may have also influenced findings; nonverbal children were excluded from the study, and perhaps this subgroup displays a distinct physiological profile from higher functioning verbal children. Additionally, there exists considerable variation in the methods employed in prior studies of physiology in ASD and FXS. In this study, unlike others (e.g., Heilman et al., 2011),
cardiac activity was sampled after several hours of working with the examiner, possibly allowing the participants to become more comfortable (and less physiologically defensive) than if cardiac activity had been sampled earlier. It will be important to study directly the influence of familiarity with conversational partners on physiological arousal in future studies.

**Physiological Activity as a Predictor of Pragmatic Language**

A main finding of this work is that parasympathetic tone during conversation predicted pragmatic language skill in ASD even after accounting for receptive and expressive vocabulary. It is interesting that pragmatic language was specifically related to vagal tone in the conversational context but not at baseline. This finding is similar to that of Watson et al. (2010), who found that vagal tone during a social context (but not while watching a nonsocial video) accounted for significant variance in parent-reported social-communication outcomes of young children with ASD. As suggested by Watson et al. (2010), children with ASD who show higher vagal tone in social contexts may present with a physiological state that is more optimal for engaging with social stimuli. Consistent with transactional theories of social learning, increased social engagement is thought to lead to greater opportunities for social learning over time, including the learning of pragmatic conversational rules (Chapman, 2000; Dickinson & McCabe, 1991). Perhaps resting vagal tone was not related to pragmatic language outcomes because social learning might depend specifically on the ability to make physiological adjustments to adapt to social demands.

The identification of neurophysiological markers associated with social dysfunction in ASD has implications for the development of interventions targeted at correcting breakdowns at the physiological level. Vagal tone appears to respond to noninvasive treatments, such as massage therapy or breathing exercises, in typical development (Chambers & Allen, 2002;
Feldman & Eidelman, 2003; Lee, 2005; Miu, Heilman, & Miclea, 2009), and some evidence suggests that pharmaceutical intervention may improve arousal modulation in FXS (Roberts et al., 2011). The results of this study support the investigation of treatments targeting autonomic dysfunction as a means to improve communication outcomes in ASD.

It is unclear why physiological profiles predicted pragmatic language outcomes in the ASD group but not in the FXS group. When considering marginal associations, some differential patterns begin to emerge across groups with more significant heart-rate change associated with better pragmatic skills in the ASD group and worse pragmatic skills in the FXS group. Given that baseline arousal of the groups differed, it is possible that the groups relied on different regulatory processes to achieve an optimal arousal level; while the ability to increase arousal assisted the children with ASD in meeting task demands, it appears that additional increases in arousal did not improve the performance of the already hyperaroused FXS group. Although cardiac activity did not independently predict pragmatic language skills in the FXS group, several associations were detected with receptive and expressive vocabulary skills, suggesting that physiological modulation does play a role in communication abilities in FXS. Future research investigating potential interacting factors, such as cognition, anxiety, and executive abilities, might clarify the role of physiological activity in the pragmatic language performance of individuals with FXS.

**Conversation as an Elicitor of Arousal in ASD and FXS**

A notable finding was that unstructured conversation with an examiner did not result in atypically elevated arousal in either ASD or FXS. While the boys with ASD and FXS did increase arousal in response to conversation, this appeared to be a typical response given that increases of a similar magnitude were observed in the TD group. Hall et al. (2009) also found
that boys with FXS showed increased heart rate in response to a conversational task, which was similar to the increases that were observed in their TD siblings. Evidence that conversation does not elicit hyperarousal in ASD and FXS might imply that atypical social behaviors of these disorders are not related to socially induced hyperarousal—at least during unstructured conversation with an examiner. Clinically, these findings suggest that clinicians working with children with ASD or FXS need not be wary of engaging with the child in fear of causing the child to become overly aroused. This finding does not, however, preclude a relationship between general physiological health and social engagement; it merely implies that social interaction itself does not appear to be a catalyst for hyperarousal.

**Autism Severity and Physiological Activity**

A remaining question is why autism severity was associated with physiological activity in TD but not within ASD or FXS as the parasympathetic nervous system is thought to set the stage for social engagement, which is a core deficit in ASD and FXS. The robust correlations detected in the TD group are somewhat unexpected given the limited range of ADOS severity scores in the TD group (by definition, participants with TD had subclinical ADOS severity scores that ranged from one to four points). Other studies have also failed to detect a relationship between heart rate/vagal tone and autism severity among children with idiopathic ASD (Jansen, Gispen-de Wied, van der Gaag, & van Engeland, 2003; Patriquin et al., 2011; Van Hecke et al., 2009). With regards to FXS, a recent report by Roberts et al. (2012) did find that vagal tone and autism severity were associated in young toddlers with FXS. In the present study of older children with FXS, such a relationship was not observed although associations followed the same direction. More research is needed to clarify developmental changes in vagal activity over time and whether associations between vagal activity and behavior are most prominent during specific
developmental stages. Although this study did not detect a relationship between cardiac activity and general symptoms of ASD, physiological modulation was linked to communication difficulties that represent a subdomain of the ASD phenotype. This area of research warrants further attention as understanding how independent biophysiological mechanisms may lead to the presentation of ASD (or to the presentation of subdomains of ASD) will be informative for the development of interventions targeting these pathways.

**Links With Anxiety**

In studies of FXS, anxiety is often hypothesized to be rooted in hyperarousal (e.g., Sudhalter & Belser, 2001). However, the few studies that have tested this hypothesis empirically have failed to detect a relationship between anxiety and cardiac arousal in FXS (Keysor, Mazzocco, McLeod, & Hoehn-Saric, 2002) and ASD (Jansen et al., 2006; Mathewson et al., 2011) as well as in individuals with anxiety disorders (Kelly, Brown, & Shaffer, 1970; McLeod, Hoehn-Saric, Zimmerli, de Souza, & Oliver, 1990; Tyrer & Alexander, 1980). The results of this study are consistent with these prior reports; cardiac activity was not associated with parent-reported symptoms of anxiety in any group. It is notable, however, that the ASD and FXS groups did not differ on parent-reported symptoms of anxiety, suggesting that the syndrome-specific associations between physiological activity and pragmatic language were not driven by group differences in anxiety level. Given that anxiety affects a significant proportion of individuals with ASD and FXS (Cordeiro, Ballinger, Hagerman, & Hessl, 2011; Merenstein et al., 1996; van Steensel, Bögels, & Perrin, 2011; White, Oswald, Ollendick, & Scahill, 2009), further exploration of the role of anxiety (and its interaction with physiological arousal) in behavioral outcomes may be useful for honing treatments in these populations.

**Directions and Limitations**
Several limitations of the study should be considered in interpreting results. First, the mean chronological ages of the groups differed (a confound for which we controlled statistically). Second, although the sample was relatively large in comparison to other studies of FXS, we may nonetheless lack statistical power to detect group differences. Third, the physiological responses observed in this study may be the most representative of those boys with FXS who have comorbid ASD given that the majority of the boys with FXS in this study met diagnostic criteria for ASD. The small participant numbers of children with FXS without ASD also prevented formal examination of the impact of ASD status on the physiological responses of individuals with FXS, which would be informative in understanding the biophysiological underpinnings of ASD in FXS. The sample of this study also consisted of children who were able to use phrase speech (so that participants could complete the conversational task of interest), which may limit generalizability to nonverbal individuals. Additionally, a number of children from the FXS and ASD groups were taking psychoactive medications, which may influence heart activity, depending on individual dosage and physical characteristics, such as weight and metabolic functioning (O’Brien & Oyebode, 2003; Rechlin, 1995; Silke, Campbell, & King, 2002). However, we felt it was important to include children who were taking medications to improve external validity as 50%–75% of children with ASD and FXS use psychotropic medications (Mandell et al., 2008; Morgan, Roy, & Chance, 2003; Valdovinos, Parsa, & Alexander, 2009). Finally, the design of the current work can only shed light on how cardiac activity during one conversational condition predicts pragmatic performance in a different social-communicative context (it was not possible to collect pragmatic and physiological data from the same conversational sample, see Methods). Future work might sample cardiac activity and pragmatic language concurrently in order to clarify context-specific relationships. Future studies would also
benefit from assessments at different developmental periods and from following children longitudinally to understand how physiological characteristics may be related to language outcomes across time, thus enhancing understanding of physiological underpinnings of social language in ASD and FXS and providing insights into developmental periods that could be most responsive to intervention.
Table 1

**Group Characteristics**

<table>
<thead>
<tr>
<th></th>
<th>ASD $n = 40$</th>
<th>FXS $n = 39$</th>
<th>TD $n = 28$</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Chronological age in years</strong></td>
<td>10.11 (2.96)$^a$</td>
<td>11.90 (2.46)$^b$</td>
<td>8.84 (2.41)$^a$</td>
</tr>
<tr>
<td>$M (SD)$, range</td>
<td>4.08–14.56</td>
<td>6.47–15.89</td>
<td>4.02–13.68</td>
</tr>
<tr>
<td><strong>Nonverbal mental age$^1$</strong></td>
<td>8.67 (3.96)$^a$</td>
<td>5.14 (0.60)$^b$</td>
<td>10.17 (3.91)$^a$</td>
</tr>
<tr>
<td>$M (SD)$, range</td>
<td>4.67–19.67</td>
<td>3.50–6.67</td>
<td>5.17–18.83</td>
</tr>
<tr>
<td><strong>Receptive vocabulary age$^2$</strong></td>
<td>8.58 (4.50)$^a$</td>
<td>6.29 (1.45)$^b$</td>
<td>12.63 (5.21)$^c$</td>
</tr>
<tr>
<td>$M (SD)$, range</td>
<td>2.58–22.00</td>
<td>3.92–9.33</td>
<td>5.67–22.00</td>
</tr>
<tr>
<td><strong>Expressive vocabulary age$^3$</strong></td>
<td>7.79 (3.73)$^a$</td>
<td>5.43 (1.22)$^b$</td>
<td>10.57 (4.16)$^c$</td>
</tr>
<tr>
<td>$M (SD)$, range</td>
<td>3.17–19.75</td>
<td>3.58–8.25</td>
<td>5.42–22.00</td>
</tr>
</tbody>
</table>

**Medication %**

<table>
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<tr>
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<th>FXS</th>
<th>TD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antidepressant</td>
<td>2.5</td>
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<tr>
<td>Stimulant</td>
<td>7.5</td>
<td>10.3</td>
<td>0</td>
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<tr>
<td>Antipsychotic</td>
<td>0</td>
<td>5.1</td>
<td>0</td>
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<tr>
<td>Antianxiety</td>
<td>0</td>
<td>2.6</td>
<td>0</td>
</tr>
<tr>
<td>Alpha adrenergic agonist</td>
<td>2.5</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>More than one</td>
<td>17.5</td>
<td>15.4</td>
<td>0</td>
</tr>
</tbody>
</table>

**Race %**

<table>
<thead>
<tr>
<th></th>
<th>ASD</th>
<th>FXS</th>
<th>TD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Caucasian</td>
<td>87.2</td>
<td>88.9</td>
<td>78.6</td>
</tr>
<tr>
<td>African American</td>
<td>7.7</td>
<td>0</td>
<td>7.1</td>
</tr>
<tr>
<td>Asian</td>
<td>0</td>
<td>8.3</td>
<td>0</td>
</tr>
<tr>
<td>Multiracial</td>
<td>2.6</td>
<td>2.8</td>
<td>10.7</td>
</tr>
<tr>
<td>Other</td>
<td>2.6</td>
<td>0</td>
<td>3.6</td>
</tr>
</tbody>
</table>

**Income %**

<table>
<thead>
<tr>
<th></th>
<th>ASD</th>
<th>FXS</th>
<th>TD</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt; 20k</td>
<td>6.9</td>
<td>0</td>
<td>5.6</td>
</tr>
<tr>
<td>20k–39k</td>
<td>20.7</td>
<td>4.2</td>
<td>5.6</td>
</tr>
<tr>
<td>40k–59k</td>
<td>6.9</td>
<td>12.5</td>
<td>22.2</td>
</tr>
<tr>
<td>60k–79k</td>
<td>24.1</td>
<td>16.7</td>
<td>16.7</td>
</tr>
<tr>
<td>&gt; 80k</td>
<td>41.4</td>
<td>66.7</td>
<td>50</td>
</tr>
</tbody>
</table>

**Maternal education level %**

<table>
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<tr>
<th></th>
<th>ASD</th>
<th>FXS</th>
<th>TD</th>
</tr>
</thead>
<tbody>
<tr>
<td>High school</td>
<td>25.7</td>
<td>16.7</td>
<td>13.6</td>
</tr>
<tr>
<td>Associate</td>
<td>17.1</td>
<td>11.1</td>
<td>9.1</td>
</tr>
<tr>
<td>Bachelor</td>
<td>37.1</td>
<td>44.4</td>
<td>31.8</td>
</tr>
<tr>
<td>Master</td>
<td>17.1</td>
<td>19.4</td>
<td>36.4</td>
</tr>
<tr>
<td>Doctorate</td>
<td>2.9</td>
<td>8.3</td>
<td>9.1</td>
</tr>
</tbody>
</table>

*Note. $^1$Leiter International Performance Scale-Revised, $^2$Peabody Picture Vocabulary Test, $^3$Expressive Vocabulary Test. ASD = autism spectrum disorder; FXS = fragile X syndrome; TD*
= typical development. Means in the same row with different superscripts differ significantly at \( p < .05 \).
Table 2

Descriptive Statistics of Cardiac Indices Across Conditions and Groups

<table>
<thead>
<tr>
<th>Cardiac index</th>
<th>Group</th>
<th>Estimate</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Observed M (SE)</td>
<td>Adjusted M (SE)</td>
</tr>
<tr>
<td>Baseline IBI</td>
<td>ASD</td>
<td>690.373 (128.06)</td>
<td>693.32 (17.95)</td>
</tr>
<tr>
<td></td>
<td>FXS</td>
<td>654.18 (100.77)</td>
<td>642.29 (19.15)</td>
</tr>
<tr>
<td></td>
<td>TD</td>
<td>702.00 (122.42)</td>
<td>715.84 (22.41)</td>
</tr>
<tr>
<td>Conversation IBI</td>
<td>ASD</td>
<td>641.29 (78.74)</td>
<td>644.45 (12.82)</td>
</tr>
<tr>
<td></td>
<td>FXS</td>
<td>627.21 (84.12)</td>
<td>612.75 (13.68)</td>
</tr>
<tr>
<td></td>
<td>TD</td>
<td>649.34 (90.40)</td>
<td>665.27 (16.01)</td>
</tr>
<tr>
<td>Change IBI (baseline – conversation)</td>
<td>ASD</td>
<td>49.43 (76.34)</td>
<td>49.11 (10.56)</td>
</tr>
<tr>
<td></td>
<td>FXS</td>
<td>33.13 (44.20)</td>
<td>34.60 (11.27)</td>
</tr>
<tr>
<td></td>
<td>TD</td>
<td>53.37 (75.97)</td>
<td>51.79 (13.18)</td>
</tr>
<tr>
<td>Baseline RSA</td>
<td>ASD</td>
<td>4.35 (1.38)</td>
<td>4.36 (0.22)</td>
</tr>
<tr>
<td></td>
<td>FXS</td>
<td>4.01 (1.34)</td>
<td>3.99 (0.23)</td>
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<td></td>
<td>TD</td>
<td>4.35 (1.42)</td>
<td>4.37 (0.27)</td>
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<tr>
<td>Conversation RSA</td>
<td>ASD</td>
<td>4.69 (1.18)</td>
<td>4.68 (0.18)</td>
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<tr>
<td></td>
<td>FXS</td>
<td>4.55 (1.10)</td>
<td>4.59 (0.19)</td>
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<tr>
<td></td>
<td>TD</td>
<td>4.43 (1.05)</td>
<td>4.39 (0.22)</td>
</tr>
<tr>
<td>Change RSA (baseline – conversation)</td>
<td>ASD</td>
<td>−0.40 (1.12)</td>
<td>−0.39 (0.17)</td>
</tr>
<tr>
<td></td>
<td>FXS</td>
<td>−0.50 (1.01)</td>
<td>−0.56 (0.18)</td>
</tr>
<tr>
<td></td>
<td>TD</td>
<td>−0.08 (0.94)</td>
<td>−0.02 (0.21)</td>
</tr>
</tbody>
</table>

Note. Adjusted means depict estimates controlling for chronological age. ASD = autism spectrum disorder; FXS = fragile X syndrome; TD = typical development.
### Table 3

**Correlations Between Cardiac Activity and Language Ability**

<table>
<thead>
<tr>
<th>Cardiac index</th>
<th>Group</th>
<th>Language Measure</th>
<th>PPVT</th>
<th>EVT</th>
<th>CASL-PJ</th>
<th>PRS-SA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline IBI</td>
<td>ASD</td>
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<td>.30</td>
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**Note.** IBI = interbeat interval; RSA = respiratory sinus arrhythmia; PPVT = Peabody Picture Vocabulary Test; EVT = Expressive Vocabulary Test; CASL-JP = Comprehensive Assessment of Spoken Language Pragmatic Judgment subtest; PRS-SA = Pragmatic Rating Scale-School Age; higher scores on the PRS-SA indicate greater impairment. ASD = autism spectrum disorder; FXS = fragile X syndrome; TD = typical development.

*p < .05, **p < .01, †p < .10.
Figure 1. Mean Inter-beat-interval Estimates across Groups

Note. Means adjusted for chronological age. ASD = autism spectrum disorder; FXS-ASD = fragile X syndrome with autism spectrum disorder; FXS-O = fragile X syndrome without autism spectrum disorder; TD = typical development.
Figure 2. Mean Respiratory Sinus Arrhythmia Estimates across Groups

Note. Means adjusted for chronological age. ASD = autism spectrum disorder; FXS-ASD = fragile X syndrome with autism spectrum disorder; FXS-O = fragile X syndrome without autism spectrum disorder; TD = typical development.
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