Longitudinal identification of clinically distinct neurophenotypes in young children with fragile X syndrome

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Fragile X syndrome (FXS), due to mutations of the \textit{FMR1} gene, is the most common known inherited cause of developmental disability. The cognitive, behavioral, and neurological phenotypes observed in affected individuals can vary considerably, making it difficult to predict outcomes and determine the need for interventions. We sought to examine early structural brain growth as a potential marker for identification of clinically meaningful subgroups. Participants included 42 very young boys with FXS who completed a T1-weighted anatomical MRI and cognitive/behavioral assessment at two longitudinal time points, with mean ages of 2.89 y and 4.91 y. Topological data analysis (TDA), an unsupervised approach to multivariate pattern analysis, was applied to the longitudinal anatomical data to identify coherent but heretofore unknown subgroups. TDA revealed two large subgroups within the study population based solely on longitudinal MRI data. Post hoc comparisons of cognition, adaptive functioning, and autism severity scores between these groups demonstrated that one group was consistently higher functioning on all measures at both time points, with pronounced and significant unidirectional differences ($P < 0.05$ for time point 1 and/or time point 2 for each measure). These results support the existence of two longitudinally defined, neuroanatomically distinct, and clinically relevant phenotypes among boys with FXS. If confirmed by additional analyses, such information may be used to predict outcomes and guide design of targeted therapies. Furthermore, TDA of longitudinal anatomical MRI data may represent a useful method for reliably and objectively defining subtypes within other neuropsychiatric disorders.

Over the past several decades, clinical neuroscience research has enhanced our understanding of the neurobiological and genetic bases for psychiatric disorders. This neurobiological and genetic information holds potential to inform disease-specific targeted interventions promising substantial improvement over current treatments (1). Fragile X syndrome (FXS) is the most common inherited cause for intellectual disability (2). The genetic etiology of FXS, a repeat expansion in the fragile X mental retardation-I gene (\textit{FMR1}) on the X chromosome, results in diminished levels of the fragile X mental retardation protein (\textit{FMRP}), which plays an essential regulatory role in synaptic plasticity and dendritic pruning, processes critical for neurodevelopment (3). The known biological basis of FXS allows for investigation of cognitive and behavioral (e.g., autism) symptoms in an etiologically homogeneous genetic disorder (4).

This well-characterized genetic disorder has also been the focus of targeted therapies aimed at correcting aberrant brain neurotransmitter systems such as up-regulation of metabotropic glutamate receptor 5 (mGluR5) signaling, which is disrupted in the \textit{Fmr1} knockout (KO) mouse model (5). Although several targeted therapies have demonstrated promising phenotype rescue in animal models (6–8), they have been unsuccessful thus far in human trials (9). One potentially important factor to consider within this context is neurobiological heterogeneity within full mutation FXS. Identification of specific subtypes within this disorder may inform the design of targeted therapies and aid in the prediction and testing of response to these therapies.

Despite sharing similar \textit{FMR1} mutation genotypes, our previous work suggests that there may be biologically defined subgroups within boys with FXS, which exhibit clinically distinct behavioral profiles at a young age (mean = 2.89 y). Specifically, classification based solely on anatomical MRI data revealed two large subgroups that differed in terms of neuroanatomical and cognitive/behavioral profiles (10). However, several important questions remain regarding development with respect to these subgroups. In particular, longitudinal research is critical in the case of FXS to increase our understanding of the developmental effects of \textit{FMRP} on the brain (11). In the present study, we report results of a 2-y follow-up using longitudinal neuroimaging data and topological data analysis (TDA) to understand variation in this well-characterized cohort of young boys with FXS. Specifically, we sought to understand if cognition and behavior developed differently depending on subgroup membership and if subgroup membership remained consistent over time.

**Significance**

We present a research study that uses early structural brain growth as a metric for defining subgroups within individuals with fragile X syndrome (FXS). Topological data analysis, a type of multivariate pattern classification, identified two large subgroups based solely on the longitudinal structural brain images. Post hoc analysis indicated significant, unidirectional differences in cognition, adaptive functioning, and autism severity scores. Our results support the use of longitudinal topological data analysis (TDA) as a putative tool for differentiating individuals with FXS based on neuroanatomical data. This information may be used to predict outcomes and guide design of targeted therapies for individuals with FXS. Longitudinal TDA may be a useful analysis tool to explore variation in other neuropsychiatric disorders.


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TDA is one of a general class of approaches to analyzing high-dimensional data known in the literature as multivariate pattern analysis (MVPA). Multivariate approaches are designed to detect effects that may be discernible within the relationships (patterns) among variables but which may elude detection when variables are examined in isolation. TDA is an unsupervised method of classification and it is used to identify coherent but potentially heretofore unknown groups within the study population. Other examples of unsupervised approaches using MVPA include independent component analysis (12) and clustering via correlation matrices (13).

A distinct advantage of TDA is the resulting visual representation called a Reeb graph. A Reeb graph is a compressed and easily readable visual representation of the data, which preserves its underlying geometric structure and facilitates identification of salient features (14). These features can encode not only information about clusters that may exist within the data but also information about spectra (i.e., variation in the data due to underlying continuous parameters). Thus, TDA is “a methodology intermediate between modeling by algebraic equations, which is continuous but not very flexible, and cluster analysis, which is discrete and therefore misses continuous phenomena” (ref. 15, p. 109). Finally, another advantage of TDA is that it can be applied directly to very high-dimensional data and so is able to preserve finer characteristics of the population structure that may be obscured through the process of dimensionality reduction required by other approaches to MVPA.

We hypothesized that TDA of longitudinal neuroimaging data would reveal clinically distinct subgroups similar to our cross-sectional analysis reported earlier (10). This hypothesis was tested by examining variables that were not included in the Reeb graph construction process. Specifically, we examined longitudinal changes in cognition and behavior within the groups defined by the TDA. We also examined neuroanatomical differences between the groups using a univariate analysis. Finally, we compared subgroup membership based on longitudinal analysis to subgroup membership based on the cross-sectional analysis previously reported (10).

Participants included 52 boys with FXS who were part of an ongoing longitudinal brain imaging study (10, 11, 16–24). The mean age at enrollment was 2.89 y, SD = 0.63, and 41 of the original 52 individuals were followed up an average of 2.01 y later (SD = 0.28). Demographic information is presented in Table S1.

**Results**

**Multivariate Pattern Analysis.** The primary outcome of the TDA is a Reeb graph (Fig. 1) that displays connected nodes representing candidate subgroups within the data. Two large and distinct subgroups were present in the data: group A appearing on the left (n = 13) and group B on the right (n = 24). There are also three small, disconnected nodes (two containing a single participant each, one green and one purple, and one containing two participants, in yellow). Due to the limited sample size in these smaller groups, remaining analyses focus on the two larger groups, A and B. Cognitive and behavioral scores are presented for individuals in the smaller groups in Table S2.

**Cognitive/Behavioral Results.** There were no differences in age between groups A and B at time 1 or time 2, and the follow-up duration was not different between groups (all p > 0.10; Fig. 2 and Table S3). Longitudinal analyses of cognitive/behavioral scores include a main effect of the anatomically defined subgroup, indicating higher performance of group B on the Vineland Adaptive Behavior Scales (25) (P = 0.016) and trends for the Mullen Scales of Early Learning (26) (P = 0.090) and the Autism Diagnostic Observation Schedule–Generic (ADOS) (27) (P = 0.056) composite scores. Note that higher scores on the ADOS indicate higher severity of autism symptoms, whereas higher scores on the Vineland and Mullen indicate higher levels of adaptive function and cognitive ability, respectively. Significant main effects of time indicate decreasing Vineland (P < 0.001) and increasing ADOS (P = 0.015) scores for all individuals regardless of group and a trend for decreasing Mullen scores (P = 0.053) over time. Group-by-time interactions were not significant for any of the cognitive/behavioral measures global scales (all p > 0.10; see Table S4 for subscale results).

Follow-up cross-sectional t tests at each time point indicated significantly higher scores for group B on the Vineland at both time points (time 1 P = 0.009 and time 2 P = 0.001) and on the Mullen at time 1 (P = 0.015) and lower scores for group B on the ADOS at time 2 (P = 0.033). We also observed a floor effect on Mullen scores: at time 1, 85% of individuals in group A and 42% in group B received a standard score of 49 (the lowest possible standard score). At time 2, 75% of individuals in group A and 67% in group B received a score of 49. Analysis of Mullen raw scores is presented in Table S5.

Post hoc analyses showed a significant group-by-time interaction for the Vineland daily living subscale (P = 0.004) indicating a more rapid decline in standard scores for group A (Table S4). There was no significant group difference in FMRP percentages (P > 0.10).

**Univariate Neuroanatomical Results.** Group A demonstrated significantly enlarged gray and white matter volumes relative to group B across widespread regions including caudate, thalamus, frontal lobe, temporal lobe, and cerebellum (Fig. 3). There were no regions for which group B demonstrated enlarged gray or white matter volume relative to group A. No significant group-by-time interaction was found with and without including age at

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Fig. 1. Reeb graph depicting topological organization of the longitudinal anatomical data. The longitudinal TDA, as visualized with Ayasdi Core software, divided the data into two large and distinct subgroups based solely on the anatomical images. Each dot represents a node, the size of which corresponds to the number of participants that were clustered to form that node. Lines or edges between nodes indicate they have participants in common. The subgroups include a node collection that appears on the left (group A, n = 13) and a node collection on the right (group B, n = 24). There are also three small, disconnected nodes (two containing single participants, one green and one purple, and one yellow group containing two participants, on the bottom left). The Ayasdi Core software allows for coloring the Reeb graph based on any measure present in the dataset. Each node is colored based on the value of principle component 1 across participants in each node to illustrate how the data vary along that component; however, the subgroupings are based on the separation of the graph structure into two primary collections of connected nodes.
time 1 as a covariate ($P > 0.10$, FDR corrected). Longitudinal analyses of total brain volume include a main effect of the anatomically defined subgroup, indicating larger volumes for group A relative to group B and a significant main effect of time indicating larger volumes at time 2 (both $P < 0.001$, FDR corrected) but no group-by-time interaction ($P > 0.10$, FDR corrected). Follow-up $t$ tests indicated that brain volumes were larger for group A at time 1 and time 2 (both $p < 0.001$, FDR corrected).

Comparison Between Longitudinal and Cross-Sectional TDA. We also compared classification of subgroups based on longitudinal data to classification based on cross-sectional data that had been previously published (10). The longitudinally defined higher-functioning group B was consistent with the higher-functioning group classified at mean age 2.89. Of the 16 individuals from the original higher-functioning group who were followed longitudinally, all 16 were classified in the higher-functioning group based on their longitudinal data. The longitudinally based lower-functioning cross-sectional group was not as stable as the higher-functioning group. In fact, four of the individuals who were originally in the lower-functioning group were classified in the higher-functioning group based on their longitudinal data. In other words, these individuals initially shared the neuroanatomical characteristics of the lower-functioning cross-sectional group but at time 2 follow-up were observed to have neuroanatomical trajectories that led to their inclusion in the higher-functioning longitudinal group. The change in Vineland scores for these individuals from time 1 to time 2 (which fall between $−3.98$ and $−4.73$) demonstrates a smaller decline in standard scores compared with the change in means of group A ($−5.49$) and group B ($−5.05$; Fig. S1).

Discussion

In this study, we hypothesized that TDA-based analysis of longitudinal neuroanatomical data would identify clinically meaningful subgroups within a cohort of males with the FMR1 full mutation. Indeed, TDA classified the longitudinal anatomical data into two subgroups that were clinically distinct in terms of cognitive/behavioral performance. Group B demonstrated higher levels of adaptive function and cognition along with lower levels of autism symptoms and smaller total brain volumes relative to group A. These results suggest TDA applied to longitudinal neuroimaging data can differentiate clinically meaningful subgroups that are not based on an a priori hypothesis. This information could have significant utility in characterizing subgroups within the larger FXS population by advancing understanding of different neurobiological mechanisms and/or unique brain-environment interactions within each subgroup. Further, subgroup-specific information might be used in the future to inform targeted therapies and aid in predicting response to treatment.

The results demonstrated a higher level of adaptive functioning (Vineland) for group B relative to group A, and this difference was stable over time (no interaction with time). The Mullen and ADOS data revealed trends for higher performance and fewer autism symptoms for group B, but group differences were only significant at time 1 for Mullen and at time 2 for ADOS. We observed a floor effect on the Mullen scores, which may have prevented us from detecting effects of group or time. Analysis of raw Mullen scores revealed significantly lower scores on the expressive language subscale for group A at time 2 and a trend for a group-by-time interaction. Lack of significant difference in the other raw Mullen scores may be due to generally low performance in both groups.
Group differences for ADOS scores were not seen at time point 1, perhaps because autism symptoms were not pronounced enough or the ADOS version we used lacked sensitivity. We used the original ADOS assessment for this study (27). However, since completion of our data collection, toddler (28) and infant (29) versions of the ADOS have been introduced. These age-appropriate assessments may be more sensitive to quantifying early autism symptoms. Nevertheless, the significant main effect of time indicates increasing scores in both subgroups suggesting that autism symptoms increase with age in young boys with FXS and that early intervention should be considered for these individuals.

We noted a significant effect of time for Vineland and a trend for Mullen scores in the overall study sample with the change in scores indicating worse performance at later ages. Decreases in Vineland and Mullen standard scores indicate a slowing of the growth of adaptive and cognitive functioning. In other words, these individuals are falling farther behind their same-age typically developing peers, but this does not represent a loss of function per se. Decreasing trajectories are consistent with longitudinal analyses of adaptive behavior (30) and cognitive functioning (31) in individuals with FXS.

The lack of significant group-by-time interactions within any of the three assessment global scales suggests that the cognitive/behavioral differences were consistent over time. It is possible that the tests were not sufficiently sensitive to capture differential trajectories of cognitive/behavioral function within subgroups. This lack of sensitivity is most apparent in the range of scores in the Mullen (for which we noted a floor effect) and ADOS severity scores, which range from 1 to 10 for this measure. The Vineland scores demonstrated a greater range, and we observed a group-by-time interaction for the daily living subscale. This result indicates a steeper slope or more rapid decline in standard scores for group A. The daily living subscale measures an individual’s skills in personal, domestic, and community domains. Recent longitudinal research in FXS indicates a weakness for the daily living domain relative to the socialization domain across development (30). Our results suggest that for a subset of individuals with FXS, daily living may be particularly vulnerable to a more rapid widening of the gap in scores relative to those of typically developing peers. Identification of individual strengths and weaknesses and knowledge of time-dependent changes in adaptive behavior profiles are critical for tailoring medical and behavioral interventions. Clearly, further longitudinal follow-up beyond age 5 will be required to bear out strengths and weaknesses of individual subgroups.

The longitudinal TDA revealed a complex multivariate pattern of subgroup differences in brain structure, and the results indicated significant, widespread group differences that were stable over time (no group-by-time interaction). The univariate results indicated enlarged volumes in group A for gray matter and white matter regions including caudate, thalamus, frontal lobe, temporal lobe, and cerebellum. These univariate results are consistent with what our group has found when comparing this entire cohort of individuals with FXS to typically developing groups and clinical comparison groups (11, 16, 17). Other studies have confirmed enlarged subcortical and cortical volume and increased cortical thickness in older individuals with FXS (32, 33). These neuroanatomical differences may result from aberrant synaptic activity and altered dendritic pruning, which are downstream effects of FMR1 gene mutations (3). Our results demonstrate that the aforementioned subcortical and cortical enlargement is not homogeneous across males with the FMR1 full mutation. Within our study population, the subgroup with widespread enlarged brain regions (i.e., generally larger brain volume) presented with lower adaptive, behavioral, and cognitive scores.

In the longitudinal analysis, two major groups accounted for over 90% of the data, whereas in the single-time point analysis (10), two groups accounted for only 73% of the data ($\chi^2 = 4.522$, df = 1, $P = 0.033$). This difference suggests that subgroup identification based on longitudinal histories may lead to a clearer separation of the data than identification based on cross-sectional analysis. This finding is consistent with neuroimaging research demonstrating that longitudinal analysis is more powerful than cross-sectional analysis at detecting gene–brain relationships (34). The enhanced classification in the present study cannot be explained by attrition alone: 4 of the 11 individuals lost to attrition were unclassified in the cross-sectional analysis, and 7 were classified into one of the two cross-sectional subgroups. Longer longitudinal studies are required to determine the stability of these groups across development, and replication in independent samples will be necessary to establish the generalizability of these results to other populations. Further, studies with larger sample sizes will be important for examining outcomes within individuals who were not classified into one of the larger subgroups.

There was no difference in FMRP levels between the lower-functioning and higher-functioning groups, a finding that is consistent with our cross-sectional results (10). Thus, subgroups were identified among young children with FXS using factors other than FMRP. This discrimination is important: within males with FXS, FMRP levels are skewed to the lower end of the spectrum within a small range (in the present study FMRP levels range from 1 to 20%). However, FMRP levels may prove more meaningful for differentiation within more diverse samples of individuals with FXS. For example, previous studies have reported associations between FMRP and brain function in females (35, 36) and between FMRP and anatomy in mixed sex samples (18) where a larger range in FMRP levels is observed. Although FMRP is essential for regulating synaptic plasticity and dendritic pruning (3) it also interacts with several other proteins that have unique expression profiles during development (37). Follow-up studies examining other genetic factors, including relevant functional polymorphisms for other loci may be informative for understanding complex gene–brain–behavior relationships in individuals with FXS.

A subset of participants was taking medications that are known to affect neurological functioning (Table S6). Higher medication use in the clinical in comparison to longitudinal groups is consistent with the higher levels of cognitive and behavioral dysfunction and lower adaptive functioning we observed in children comprising this group.

It is unlikely that medication use significantly influenced brain development and the resulting subgroup clustering we obtained in our analysis of longitudinal data. Specifically, very few children were taking medication at time 1 (only three in group A), at which time the clustering was very similar to that obtained from the longitudinal data. Of the individuals who switched from the low- to the high-functioning subgroup, none were taking medications at the first time point, and only one participant was taking medication at the second time point. Accordingly, it is unlikely that medication use affected the differences in clustering between single- and multi-time point data. Future studies with larger sample sizes and longer follow-up durations will be important for examining how medication influences both brain development and developmentally based subgrouping within FXS.

In summary, our results support longitudinal TDA as a putative tool for differentiating individuals with FXS based on neuroanatomical data. These results confirm the presence of two longitudinally defined, neuroanatomically distinct subgroups that are associated with clinically relevant phenotypes among young boys with fragile X syndrome. The subgroup differences were relatively stable across time in terms of adaptive function, cognitive function, and autism symptoms. Furthermore, classification based on longitudinal data may be superior to classification based on single-time
point data, consistent with improved performance using additional data. This information may be used to predict outcomes and guide design of targeted therapies for individuals with FXS. In addition, longitudinal TDA may represent an informative tool for exploration of neurodevelopmental variation in other specific and idiopathic disorders such as sex chromosome aneuploidy conditions, idiopathic autism spectrum disorder, and attention deficit hyperactivity disorder.

Materials and Methods

Participants and Data Collection. Participants were recruited by collaborating research teams at Stanford University and the University of North Carolina at Chapel Hill. The study protocols were approved by the human subjects committees at Stanford University and the University of North Carolina at Chapel Hill, and informed consent was obtained from all parents. Children with FXS were recruited through registry databases maintained by both institutions, through postings to the National Fragile X Foundation website and quarterly newsletter, and through mailings to other regional FXS organizations. FMR1 full mutation was confirmed via Southern blot DNA analysis, and FMRP percentage was estimated based on the percentage of peripheral lymphocytes containing FMRP (38). Participants using medication were included in the study (Table S6).

Participants completed the same battery of assessments at each time point, and all assessments were administered and scored under the supervision of a doctoral-level clinician. The Mullen Scales of Early Learning (26) assessed cognitive abilities. The Vineland Adaptive Behavior Scales, Interview Edition, Survey Form (25), was used to measure age-appropriate skills required for navigating daily life. The same parent completed the interview for all four participants (one in group A, two in group B, and one in that was not assigned to either of the subgroups). When these participants were removed from the repeated measures analysis of Vineland composite scores, the pattern of results did not change. The main effect of group remained significant (higher scores for group B, P = 0.024), the main effect of time remained significant (decreasing scores for all individuals, P < 0.001), and the group-by-time interaction remained not significant (P > 0.10). The group-by-time interaction for the daily living subscale also remained significant (P = 0.005). The Autism Diagnostic Observation Schedule—Generic (ADOS) (27) was administered and scored by a research-certified clinician or by individuals who were directly trained by a research-certified administrator. The ADOS was administered by a different clinician at each time point with the exception of six cases. Reliability among administrators was verified within and across sites via consensus scoring for 20% of the assessments. ADOS-G scores were converted using the ADOS-2 summary scores (39) and calibrated severity scores (40). The stability of the calibrated severity scores has been demonstrated over time and when administered by different clinicians (41). Thus, having a different administrator at time 1 and time 2 likely did not bias our longitudinal results. There were no significant site differences on any of the cognitive/behavioral variables (all P > 0.05).

Whole brain T1-weighted MRI data were collected from participants at each time point (GE 1.5 T Signa, corona orientation; inversion recovery preparation pulse = 300 ms; repetition time = 12 ms; echo time = 5 ms; flip angle = 20°; slice thickness = 1.5 mm; excitations = 1; field of view = 20 cm; matrix = 256 × 192; voxels = 0.78 mm3). Plastic geometric phantoms were scanned at regular intervals during the study to ensure calibration of MRI scanners across sites. Results of calibration procedures have been reported (17).

Voxel-Based Morphometry Preprocessing. Voxel-based morphometry (VBM) preprocessing of MRI data was performed using Statistical Parametric Mapping 5 (SPM5) (www.fil.ion.ucl.ac.uk/spm) and VBM5.1 (dbm.neuro.uni-jena.de/vbmm). Images were bias-field corrected and segmented to gray matter, white matter, and cerebrospinal fluid. A Hidden Markov Random Field (prior probability weight 0.3) incorporated spatial constraints from neighboring voxels. Images were normalized with a 12-parameter affine transformation with a spatial frequency cutoff of 25 in all three (x, y, z) directions and resampled to 1 × 1 × 1 mm voxels. Linear and nonlinear Jacobian modulation parameters were applied. For each time point, customized gray matter, white matter, and cerebrospinal fluid templates created using all participants in our previous studies (11) were used for VBM preprocessing. Segmentation and normalization accuracy were manually inspected for each image.

Multivariate Pattern Analysis Using Topological Data Analysis. Fig. 52 summarizes these procedures. The normalized images were smoothed to 4 mm to reduce noise and enhance further computation. Next, gray and white matter images for all participants for both time points were combined into a single matrix, M, with the data for each individual entered as a row and voxel intensity as the columns, M = (# participants) × (# gray voxels time 1 + # white voxels time 1 + # gray voxels time 2 + # white voxels time 2). Ayasdi Core software, a more recent version of iris (42), was used to construct a Reeb graph to identify topological (i.e., shape) features. Construction of a Reeb graph includes (i) column selection, (ii) metric selection, (iii) filter selection, and (iv) selection of gain and resolution values. Column selection defines the subject space, S, a high-dimensional space in which each participant’s data represent a point. A subset of columns of M was selected corresponding to voxels whose variance was at least 0.03 across participants. (In modulated and segmented images voxel intensities correspond to proportions of volume and thus range in value from 0 to 1.) Each selected column was mean centered and scaled to have a variance of 1 such that all voxels were weighted equally. The next step, metric selection, defined the distances within S yielding a measure of similarity between participants. The metric chosen for the present study was Euclidean distance using all of the columns in M, which is a generalization of the usual distance between points P1 = (x1, y1) and P2 = (x2, y2) in the 2D coordinate plane:

\[ d(P1, P2) = \sqrt{(x1 - x2)^2 + (y1 - y2)^2}. \]

Filter selection transforms the similarity information into a visual representation, a 2D rendering of a 3D graph. The first two principal component scores (PC1 and PC2) of the variance-normalized data were used as filters, such that two participants were judged to be similar (i.e., shared the same bin) if both their PC1 and PC2 scores were sufficiently similar. Next, Euclidian distance was used to further cluster the participants within each bin, such that two participants within the same bin were merged into the same cluster if the distance between them was sufficiently small; each cluster obtained in this way was then viewed as a node of the Reeb graph. Any two clusters (i.e., nodes) that shared a participant was then joined by an edge. Although bin size was governed by the resolution parameter, bins were allowed to overlap (i.e., share participants) to an extent governed by the gain parameter, and this overlap may lead clusters from different bins to share participants. Thus, resolution and gain determine the nodes and edges of the Reeb graph, which was projected onto a 2D plane for visualization. We may think of the resulting graph as a view of the data through a microscope, where the TDA gain and resolution parameters play roles analogous to level of light and level of magnification. The parameter values that clearly decomposed the data into subgroups were a gain of 4 for PC1 and PC2 and a resolution of 20 for PC1 and PC2.

Analysis of Cognitive/Behavioral Data. Repeated measures ANOVA was used to assess longitudinal change with longitudinally based TDA group as the independent variable, Mullen, Vineland, or ADOS score as the dependent variable and time as the repeated factor. Standard t tests were used to compare cross-sectional cognitive/behavioral scores between subgroups at each time point.

Univariate Analyses of Magnetic Resonance Images. Post hoc descriptive comparisons of gray and white matter volume, using age at time 1 as a covariate, were performed to illustrate cross-sectional structural brain differences between subgroups at time 1 and at time 2. Longitudinal gray and white matter images representing change in volume from time 1 to time 2 were also compared between groups statistically to illustrate differential changes between the groups over time. For both cross-sectional and longitudinal comparisons FSL’s randomize tool (https://fsl.fmrib.ox.ac.uk/fsl/fslwiki/Randomise) was implemented with 5,000 permutations and threshold-old-free cluster enhancement (43). Clusters corresponding to P < 0.005 (FWE corrected) were retained. We also compared total brain volumes longitudinally with repeated measures ANOVA and examined cross-sectional differences using t tests.

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