Rich Club and reward network connectivity as endophenotypes for alcohol dependence: a DTI study

Nabi Zorlu¹, Necip Çapraz², Esra Oztekin¹, Başak Bagci¹, Maria A. Di Biase³, Andrew Zalesky³, Fazıl Gelal⁴, Emre Bora³,⁵, Ercan Durmaz¹, Lütfullah Besiroğlu¹, Aybala Sarıççek¹

¹ Department of Psychiatry, Katip Celebi University, Ataturk Training and Research Hospital, Izmir, Turkey
² Department of Psychiatry, Cizre State Hospital, Sirnak, Turkey
³ Melbourne Neuropsychiatry Centre, Department of Psychiatry, The University of Melbourne
⁴ Department of Radiodiagnostics, Katip Celebi University, Ataturk Training and Research Hospital, Izmir, Turkey
⁵ Department of Psychiatry, Dokuz Eylül University Medical School, Izmir, Turkey.

* Corresponding author at: Katip Celebi University Ataturk Training and Research Hospital, Department of Psychiatry, İzmir, Turkey. Tel.: +90 232 2444444x1581.
E-mail address: zorlunabi@hotmail.com (N. Zorlu).

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Abstract

We aimed to examine the whole-brain white matter connectivity and local topology of reward system nodes in patients with alcohol use disorder (AUD) and unaffected siblings, relative to healthy comparison individuals. Diffusion-weighted MRI scans were acquired from 18 patients with AUD, 15 unaffected siblings of AUD patients and 15 healthy controls. Structural networks were examined using network-based statistic (NBS) and connectomic analysis. Connectomic analysis showed a significant ordered difference in normalized rich club organization (AUD < Siblings < Controls). We also found rank ordered differences (Control > Sibling > AUD) for both nodal clustering coefficient and nodal local efficiency in reward system nodes, particularly left caudate, right putamen and left hippocampus. NBS analyses showed that AUD group had significantly weaker connectivity than controls in the right hemisphere, mostly in the edges connecting putamen and hippocampus with other brain regions. Our results suggest that reward system network abnormalities, especially in subcortical structures, and impairments in rich-club organization might be related to the familial predisposition for AUD.

Key words: Alcohol use disorder, endophenotype, rich club, structural connectivity,
INTRODUCTION

Alcohol use disorder (AUD) is one of the most common mental disorders (Grant et al. 2015) and associated with psychosocial dysfunction (Kendler et al. 2017), disability (Grant et al. 2015), elevated mortality (Kendler et al. 2016) and physical and psychiatric comorbidities (Grant et al. 2015; Rehm 2011). Nevertheless, despite the increasing prevalence and disease burden of AUD, the neurobiological mechanisms underlying the disorder remain unclear.

Diffusion-weighted magnetic resonance imaging (MRI) can be used to investigate white matter (WM) in vivo. Fractional anisotropy (FA) is a measure of the degree to which water diffusion is constrained in the brain and is widely used as a general index of WM integrity (Kubicki et al. 2002). Damage to WM or demyelination along neuronal axons results in more isotropic water movement and manifests in lower FA values. Previous diffusion studies have reported decreased FA in both recently detoxified and in long-term abstinent AUD patients, most consistently in the corpus callosum, centrum
semiovale, internal and external capsules, fornix, superior cingulate, and longitudinal fasciculi (Zahr & Pfefferbaum 2017; Zorlu et al. 2013, 2014).

First-degree relatives of patients with AUD are at greater risk of developing the disorder and heritability of AUD is estimated at 50–85% (Long et al. 2017; Verhulst, Neale & Kendler 2016), suggesting a large familial component to the disorder. Therefore, the investigation of unaffected first-degree relatives, in addition to AUD patients and healthy controls, may help to distinguish biomarkers of genetic risk without the confounding effects of the burden of illness, medication or clinical state. Previous studies examining WM integrity in genetically high-risk groups compared to controls found lower FA in frontal and parietal regions (Acheson et al. 2015; Herting, Fair & Nagel 2011), no difference (Wetherill et al. 2012) or higher FA values in several regions throughout the brain (Squeglia et al. 2014). However, most of these studies were conducted in adolescents or young adults with a family history of alcoholism, and therefore, who were at risk of developing AUD in the future. Furthermore, previous studies (Acheson et al. 2015; Herting, Fair & Nagel 2011; Wetherill et al. 2012; Squeglia et al. 2014) did not include a patient group in the analysis, which limits endophenotypic inference regarding WM abnormalities. Additionally, the degree of penetrance for candidate genes and environmental factors might also affect structural changes associated with genetic risk. Therefore, siblings of individuals with AUD provide a unique opportunity to investigate shared genetic and environmental risk factors of the disorder.
Although previous diffusion studies have improved our understanding of WM alterations in AUD, they have typically applied a regional perspective, which limits interpretations at a whole-brain or network level. To overcome this limitation, recent studies have applied connectomics to examine alterations in the brain network topology of patients with psychiatric disorders (Lord et al. 2017). In connectomic analyses, a structural network of the brain is mapped that comprises cortical and subcortical regions (nodes) and WM connections between nodes (edges). Small-worldness is a key feature of brain network topology, which reflects an optimal balance between local segregation and global integration (Bullmore & Sporns 2012). The most commonly used measures to characterize network topology are clustering coefficient (CC) and modularity (Q) for network segregation and characteristic path length (CPL) and global efficiency (GE) for network integration. Briefly, higher levels of CC and Q suggest the presence of locally connected clusters or modules implicated in specialized information processing. In addition, low CPL and high GE levels suggest a more globally integrated network. Another important feature of the brain network topology is rich-club organization, which is thought to represent a backbone facilitating network integration (van den Heuvel & Sporns 2013). In short, some brain regions are connected to a large number of other regions. Moreover, some of these so-called hub regions are also more likely to be interconnected among each other than expected by chance. This densely interconnected group of hub regions is called a rich club. Importantly, previous studies
reported genetic influences on structural network topology (Bohlken et al. 2014; Fulcher & Fornito 2015). Furthermore, two previous studies reported intermediate levels of rich-club organization, clustering and efficiency between patients and controls in siblings of patients with schizophrenia (Collin et al. 2014; Lo et al. 2015). Therefore, given the heritability of AUD, we hypothesize that the same patterns are evident in the siblings of patients with AUD.

The first connectomic analysis of AUD did not identify any abnormalities in AUD individuals in both global functional network topology and local topology of the striatal nodes, using functional MRI (Sjoerds et al. 2017). In contrast, the second study found higher CC, indicating higher clustered and segregated network topology and information processing in a functional network of AUD patients compared to controls (Morris et al. 2017). A third study was conducted in relatives of patients with AUD on functional networks (Holla et al. 2017), which found reduced CC, small-worldness, and local network efficiency in substance-naive high-risk male offspring with a family history of AUD compared to controls without a family history of AUD. However, it is not known whether abnormalities in functional networks in AUD reflect underlying changes in structural topology. Given the evidence for WM abnormalities in AUD, connectomic analysis using structural networks might provide further insight about the mechanisms underlying AUD.
Reward processing abnormalities are one of the major models for addiction (Volkow & Morales 2015) and thought to be mediated by cortico-limbic-striatal brain networks, particularly medial orbitofrontal cortex (mOFC) (Peters & D’Esposito 2016; Zald et al. 2014), striatum (Luijten et al. 2017; Richter et al. 2017), hippocampus (Richter et al. 2017) and amygdala (Sescousse et al. 2013). Furthermore, reward processing abnormalities seem to be an important risk for developing AUD (Hill & Brien 2015). However, whether abnormalities in nodes of the reward system are premorbid conditions or consequences of alcohol use remains unclear. Given the heritability of AUD and brain network topology (Bohlken et al. 2014; Fulcher & Fornito 2015), examining reward network topology in unaffected siblings of patients with AUD might provide insight into whether reward system abnormalities might be a candidate endophenotype.

Here, we applied the network-based statistic (NBS) (Zalesky, Fornito & Bullmore 2010) to identify white matter connections that were significantly altered in AUD individuals and unaffected siblings, relative to healthy comparison individuals. We also compared global and local graph measures between these groups to determine whether white matter connectivity alterations are related to familial liability. We hypothesized that siblings have intermediate levels of rich-club organization and local topology of reward system nodes between patients with AUD and controls.
MATERIALS AND METHODS

Participants

A total of 18 patients with AUD who had been abstinent for at least 2 weeks before scanning, 15 siblings of AUD patients and 15 healthy controls were enrolled in the study. All subjects were male and right-handed. The groups were matched for age and years of education. The advantage of studying subjects abusing various substances in Turkey is that there are groups of subjects with single substance abuse, particularly alcohol. Hence, we recruited patients with AUD who abused only alcohol to date.

Exclusion criteria for the alcohol dependence group were as follows: (1) any lifetime substance use other than alcohol (except nicotine), (2) current or past history of any serious psychiatric illness, including any psychotic disorder, bipolar disorder, (3) current use of psychotropic medication (4) current or past history of any significant neurological disorders, (5) history of loss of consciousness for more than 30 min, (6) any severe hepatic, endocrine, renal disease, and (7) any contraindications for MRI scanning (metal implants, pacemakers, etc.). Siblings and control subjects met the same criteria as patients, except for the history of alcohol dependence. All subjects were interviewed using the Structured Clinical Interview for DSM-IV Axis I Disorders (First et al. 1999) to exclude participants with past or current comorbid Axis I diagnoses and to confirm the diagnosis of alcohol dependence in the clinical group.
AUD patients were interviewed in order to determine the age at which they started drinking, the length of time they had alcohol dependence and duration from the last alcohol consumption. Monitoring of blood and urine for the presence of alcohol, amphetamines, barbiturates, benzodiazepines, cocaine, cannabis, and opiates was performed to assure sobriety at the day of scanning for only patients with AUD.

All subjects gave written informed consent to participate in the study. The study was approved by local research and ethics committees.

**Imaging protocol**

MRI was performed using a 1.5T MR system (GE SignaHDxt, General Electric Medical Systems, Milwaukee, WI, USA). Diffusion imaging data were acquired in 41 diffusion gradient directions (b-value=1000 s/mm²) and a single non-diffusion weighted reference image using a sequence optimized to collect diffusion-weighted images (repetition time=6500 ms, echo time=90 ms, voxel size=1x1x2 mm³).

**Data pre-processing and network construction**

DTI data were analyzed using FMRIB's (Oxford Centre for Functional MRI of the Brain) Diffusion Toolbox, which is part of FSL (FMRIB Software Library) (Smith et al. 2004).
Motion and eddy current artifacts were corrected using FSL’s “eddy-correct” command. A brain mask of the non-diffusion-weighted image was created using FSL’s Brain Extraction Tool (Smith 2002). Diffusion tensors were then calculated with FSL DTIFIT for whole brain volumes, and the resulting FA volume of each subject was then registered to the FMRIB58_FA template using the FSL nonlinear registration tool FNIRT to obtain the warp field from native to standard space. The nonlinear warp was initialized with an affine registration generated with the FSL linear registration tool FLIRT.

Deterministic whole-brain streamline counts were generated using the fiber assignment by continuous tracking (FACT) algorithm using the TrackVis Diffusion Toolkit (trackvis.org) in native diffusion space for each individual after rotation corrections were applied (Leemans & Jones 2009); parameters included a 35° degree angle threshold, twenty seeds were placed in every voxel and a minimum streamline length of 20 mm was used.

Network nodes were based on the 116 cortical, subcortical, and cerebellar regions comprising the automated anatomical labeling (AAL) atlas (Tzourio-Mazoyer et al. 2002). AAL labels were mapped to each individual’s native space using the inverse of the warp field previously computed. Each individual’s network was represented with a symmetric N × N connectivity matrix, where N denoted the number of nodes. Each element of the connectivity matrix was populated with the number of streamlines...
between the corresponding pair of regions (streamline count), which served as a measure of inter-regional connection strength.

**NBS analyses**

AUD patients and siblings were separately compared with controls. For each of the comparisons (AUD vs Controls and Siblings vs Controls), all connectivity matrices were subject to a group threshold, applied across all subjects, to eliminate spurious connections (de Reus & van den Heuvel 2013). This involved eliminating any pair of regions that was not interconnected by one or more streamlines in at least 25% of all individuals. In performing the NBS calculations, to localize differences in the connection strengths (streamline count) to specific networks, while controlling the family-wise error (FWE), the primary threshold for each inter-regional connection was set to a t value = 2.3 (10000 permutations).

**Graph theoretical analyses**

For each dataset, a threshold was applied to each weighted undirected structural connectivity matrix (edges with less than 3 streamlines were set to zero) to reduce false positive connections and graph theory metrics were computed with the brain connectivity toolbox (Rubinov & Sporns 2010).
First, we examined global network measures including clustering coefficient (CC), modularity (Q), characteristic path length (CPL), global efficiency (GE), and small-worldness. CC is defined as the fraction of triangles around a node and is equivalent to the fraction of node’s neighbors that are neighbors of each other (Rubinov & Sporns 2010). Q quantifies the degree to which the network may be subdivided into such clearly delineated groups (Rubinov & Sporns 2010). CPL is the average shortest path length in the network and the GE is the average inverse shortest path length in the network (Rubinov & Sporns 2010). CC and CPL were normalized by 1,000 randomized null network preserving the number of nodes, edges, and degree distribution (CC\_norm and CPL\_norm). Small-worldness was calculated as; CC\_norm/CPL\_norm. Additionally, we also examined the nodal network measures in the a priori selected ROIs, including bilateral putamen and caudate nucleus, hippocampus, amygdala and medial OFC, based on previous research as mentioned above, including the nodal local efficiency (E\_nodal) and clustering coefficient (CC\_nodal). E\_nodal is the GE computed on the neighborhood of the node (Rubinov & Sporns 2010).

Second, we calculated the rich club coefficient as the connection density of the subnetwork defined by the set of nodes with degrees exceeding $k$. The degree threshold, $k$, was systematically varied to test rich club organization at different levels. The rich club coefficient was normalized (RC\_norm) with respect to an ensemble of 1000 surrogate networks with the same number of nodes, edges, and degree distribution as
the empirical data, but which was randomized in all other respects. $RC_{norm}>1$ over a range of $k$ implies the presence of a rich-club in a network (van den Heuvel & Sporns 2011).

Nonparametric Jonckheere-Terpstra (JT) permutation analysis (10000 permutations) was applied to test whether graph metrics have a familial pattern (AUD > Siblings > Controls or Controls > Siblings > AUD). False discovery rate (FDR) correction for multiple comparisons (Benjamini & Hochberg 1995) was applied to the global (5 statistical tests), nodal (10 statistical tests, one per region) and rich-club organization (16 statistical tests, one per $k$ levels) analyses. We used additional post-hoc independent sample $t$ test to compare graph metrics between patients with AUD versus controls and siblings versus controls if JT test indicated significant differences after FDR correction.

**Robustness of main results**

The stability of results for the global, nodal and rich-club organization was checked using a variety of thresholds on the streamline count. We utilized five additional thresholds of streamline counts $\geq 0, 1, 2, 4$ and $5$. We also normalized edge values to remove the possible influence of the streamline counts connecting pairs of regions across individuals.
To check the stability of NBS results, we applied different thresholds across a range from 5% to 30% in steps of 5%.

RESULTS

Demographic and alcohol use variables

Groups were comparable for age, educational level and smoking status. Table 1 shows the demographic and alcohol use variables for the 3 groups.

Table 1 here

Global Metrics

JT testing did not show ordered differences for CCnorm, CPLnorm, small-worldness and GE. We found ordered difference for Q, such that Patients > Siblings > Controls at an uncorrected level but not after FDR correction (Table 2).

Table 2 here

Reward System Node-Specific Metrics

There were significant ordered differences for both CCnodal and Enodal (Control > Sibling > AUD) in the left hippocampus and caudate and right putamen that survived
FDR correction. Post-hoc t tests demonstrated significantly reduced CC\textit{nodal} and Enodal in AUD compared with controls in all nodes. Siblings showed a trend for both CC\textit{nodal} (p=0.70) and Enodal (p=0.054) in right putamen (Figure 1) (see Supporting Information Table S1 and Table S2 for further information).

**Figure 1 here**

**Rich-Club Organization**

There was a significant ordered difference in normalized rich club organization after FDR correction (AUD < Siblings < Controls) over the range of k from 13 to 17 (Figure 2). Post-hoc t tests demonstrated significantly reduced normalized rich club organization in AUD compared with controls in all significant k values. Siblings showed significantly reduced RC\textit{norm} in k=17 (p=0.046) and a trend for k=16 (p=0.071).

**Figure 2 here**

**Structural Connectivity**

NBS analyses identified a single network of significantly decreased structural connectivity in AUD group compared to controls (t=2.3, p=0.014; Figure 3). The network was on the right hemisphere and consisted of ten nodes connected by nine edges. These connections linked the putamen with the inferior frontal gyrus (both triangular and
opercular parts), inferior temporal gyrus and supramarginal gyrus. Another three edges linked the hippocampus with inferior temporal gyrus, middle occipital gyrus and superior occipital gyrus. Two further edges linked the superior frontal gyrus with caudate and inferior frontal gyrus, opercular part (Figure 3). We further examined the nine edges showing decreased streamline count in AUD group to see whether streamline counts also show significant ordered differences similar to topological analyses. All edges except edge between the right putamen and inferior frontal gyrus triangular part showed significant ordered differences (AUD < Siblings < Controls) after FDR correction (see Supporting Information Table S3 for further information). There were no differences between AUD group and controls at higher supra-thresholds (t=2.5, t=3.0).

There was a trend for a reduced structural connectivity in siblings compared to controls (t=2.3, p=0.066). The network was on the left hemisphere and connections linked the middle temporal gyrus with inferior temporal gyrus and inferior occipital gyrus.

There were no edges showing increased structural connectivity in AUD group or siblings compared with controls with NBS.

Figure 3 here
We did not find any significant correlations between the alcohol use variables and rich-club organization, reward system metrics and streamline counts in the AUD group.

**Reproducibility of the findings**

Reward system node-specific metrics were generally robust to the choice of streamline threshold. In particular, CCnodal and Enodal in the left caudate and right putamen remained statistically significant for all streamline thresholds. Enodal in the left hippocampus was statistically significant for 5 of 7 thresholds and CCnodal was statistically significant for 4 of 7 thresholds (see Supporting Information Table S4 and S5 for further information). Global metrics remained unchanged across all streamline thresholds. On the other hand, rich-club organization was only significant for streamline count threshold of $\geq 1$ (k=19, p=0.032; k=20, p=0.046; k=21, p=0.046).

All nine edges that we found significantly decreased in AUD group compared to controls remained stable for the various thresholds (5%, p = 0.006; 10%, p = 0.003; 15%, p = 0.012; 20%, p = 0.014). Only for the 30% (p = 0.017), edge between putamen with the supramarginalgyrus did not remain significant.

**DISCUSSION**
We examined AUD patients, unaffected siblings of AUD patients and healthy controls in order to identify alterations in anatomical connectivity that may be associated with the familial risk for AUD. We found rank ordered differences in network topology in reward system nodes (Control>Siblings>AUD), particularly caudate, putamen and hippocampus and abnormal rich club organization in AUD patients and unaffected siblings of AUD patients. Our results suggest that rich club organization and subcortical connectivity abnormalities might be imaging endophenotypes.

We found significant rank ordered differences of CC\textit{nodal} and E\textit{nodal}, such that values were lowest in AUD group, intermediate in sibling group and highest in controls, in the left caudate and right putamen and left hippocampus. The lower levels of CC\textit{nodal} observed in these nodes reflect reduced connectivity between the neighbors of these nodes that suggest weaker specialized information processing and lower levels of E\textit{nodal} suggest less efficient (with more edges) connection of these nodes with all other nodes in the network. Striatum as well as hippocampus has been implicated in reward-related and motivated behaviors which are thought to be a core mechanism of addiction (Volkow & Morales 2015). A recent PET study found increased reward-induced dopamine release in the bilateral caudate nucleus, putamen and ventral striatum and extensive reward-induced dopamine release in dorsal striatum was associated with better reinforcement learning task performance in the healthy volunteers (Kasanova et al. 2017). Furthermore, another recent study found that genetic variations associated
with dopamine D2 receptor expression affect modulation of striatal and hippocampal brain responses during encoding of reward-predicting items and healthy controls with lower dopamine D2 receptor expression exhibited better reward-related memory (Richter et al. 2017). Striatal and hippocampal abnormalities were also found in the relatives of substance dependent subjects with different neuroimaging methods. A PET study in younger drinkers with short drinking histories found an approximately fourfold striatal dopamine increase after tasting a preferred alcoholic beverage in subjects with a first-degree relative with AUD compared to family history-negative subjects (Oberlin et al. 2013). Additionally, a voxel-based morphometry study found larger medial temporal lobe and putamen volumes in the siblings of stimulant dependent patients (Ersche et al. 2012). Thus, it is possible that reward system alterations in subcortical structures reflect a genetic predisposition rather than a consequence of alcohol dependence. Speculatively, these abnormalities might lead to increased reward sensitivity, which may confer greater risk for initiation and maintenance of alcohol use.

We also found impaired rich-club organization in patients with AUD and to a lesser degree, in siblings of AUD patients compared to controls (AUD < Sibling < Controls) that suggest reduced structural connectivity among the central hubs in the AUD and sibling groups. These findings are in line with previous studies reporting heritability of structural brain connectivity features (Bohlken et al. 2014; Fulcher and Fornito 2015). Rich-club organization is thought to mediate the integration of information between segregated
parts of the brain network (van den Heuvel & Sporns 2013). However, the clinical importance of rich-club organization is still an open question. A recent study reported stronger rich club organization is associated with better cognitive performance in healthy volunteers (Baggio et al. 2015). Therefore, our results may partly explain cognitive abnormalities reported in both alcoholics (Le Berre, Fama & Sullivan 2017) and first-degree relatives of patients with AUD (Cservenka 2016).

NBS analyses showed that AUD group had significantly weaker connectivity than controls in the right hemisphere, mostly in the edges connecting putamen and hippocampus with other brain regions. There were no differences between sibling and control groups but sibling group showed intermediate streamline numbers in the edges that was found lower in AUD group compared to controls. We found four decreased edges included in putamen. Animal studies have shown that dorsolateral striatum (putamen in humans) is essential for habitual alcohol seeking (Corbit & Janak 2016) and according to contemporary theories of addiction, putamen is one of the core regions for the transition from goal-directed reward related behavior into habitual consumption of alcohol (Everitt & Robbins 2016). However, habitual alcohol seeking is not enough to explain addiction process and compulsive alcohol seeking despite negative consequences is essential for addiction. Compulsive drug seeking is thought to be due to disrupted cortical control, especially prefrontal cortex, over behavior with the long-term drug use (Everitt & Robbins 2016). Considering that the right inferior frontal gyrus
known to mediate response inhibition (Fuentes-Claramonte et al. 2016), our finding of reduced connectivity between right putamen and right inferior frontal gyrus could be interpreted as evidence of reduced fronto-striatal control due to the long-term alcohol use. However, due to cross-sectional nature of our study, we are unable to establish a clear temporal relationship between alcohol use and reduced connectivity between cortical and subcortical regions.

It is also important to note that middle aged sibling group in our study had not developed AUD despite the fact that they shared reward system alterations with AUD group. This might suggest that more risk factors are involved in the development of AUD and/or that protective factors might also prevent siblings from developing AUD. For example siblings showed intermediate values between AUD group and controls while siblings were not significantly different from controls on these values except rich-club organization. Lower number or expression of high-risk genes in siblings compared to patients with AUD might be a possible explanation for intermediate levels that was shown in our study. Another possible explanation might be the environmental factors differentiated between siblings and patients might contribute to AUD in patients but not in siblings on the basis of distributed structural network topology in both groups. A longitudinal study would be required to elucidate these relationships.
The current study has several potential limitations. The most obvious is the cross-sectional nature of the study, so that potential differential changes in WM measurements over the course of illness in AUD remain to be directly established. Second, our sample was relatively small, which might reduce the power of the group comparison especially for the unaffected sibling group as not all individuals in this group are likely to be carrying the susceptibility gene to AUD. Third, the field strength of 1.5 Tesla should be noted. Fourth, DTI-based deterministic tractography has been shown to have a limited capacity for resolving crossing, converging or diverging fibers which might lead to unreliable connectivity matrices that comprise false positive connections (Jbabdi & Johansen-Berg 2011, Zalesky et al. 2016, Maier-Hein et al. 2017). Fifth, due to an absence of standardized methodological criteria, the choice of streamline thresholds is rather arbitrary, although the majority of our findings were robust to the choice of threshold. Rich-club organization was most sensitive to the choice of threshold and thus these findings should be interpreted cautiously with respect to the contribution of weak and potentially spurious connections. Finally, we cannot extend our findings to females with AUD and their siblings as our sample included only men.

In conclusion, our results suggest that reward system network abnormalities, especially in subcortical structures, and impairments in rich-club organization might be related to the familial predisposition for AUD.
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Declarations of interest
All authors declare that they have no conflicts of interest.

AUTHORS CONTRIBUTION

NZ, NC, AS, and LB were responsible for the study concept and design. NC, EO, BB, ED, and FG contributed to the acquisition of imaging data. NZ analyzed the data. NZ drafted the manuscript. AZ, EB, and MAD provided critical revision of the manuscript for important intellectual content. All authors critically reviewed content and approved final version for publication.

References


Table 1 Demographics and clinical characteristics of in the patients with alcohol use disorder (AUD), siblings and controls.

<table>
<thead>
<tr>
<th></th>
<th>AUD (n= 18)</th>
<th>Siblings (n=15)</th>
<th>Controls (n= 15)</th>
<th>Statistics</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>40.8± 6.4</td>
<td>42.5± 8.1</td>
<td>42.8± 7.2</td>
<td>F=0.391, p=0.679</td>
</tr>
<tr>
<td>Education (years)</td>
<td>8.9± 3.2</td>
<td>8.6± 3.1</td>
<td>8.7± 4.0</td>
<td>F=0.240, p=0.788</td>
</tr>
<tr>
<td>Smokers</td>
<td>14/18</td>
<td>9/15</td>
<td>10/15</td>
<td>χ=1.248, p=0.536</td>
</tr>
<tr>
<td>Duration of</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>dependence (years)</td>
<td>12.0± 6.5</td>
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<tr>
<td>Duration of</td>
<td>38.9± 24.9</td>
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<tr>
<td>abstinence (days)</td>
<td></td>
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</table>
Age at first use 17.6± 5.2 (years)

Data are presented as mean ± standard deviation unless otherwise indicated.

Table 2 Global metrics of brain WM structural networks in the patients with alcohol use disorder (AUD), siblings and controls.

<table>
<thead>
<tr>
<th></th>
<th>AUD (n=18)</th>
<th>Siblings (n=15)</th>
<th>Controls (n=15)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>CCnorm</td>
<td>4.713± 0.593</td>
<td>4.561± 0.554</td>
<td>4.564± 0.494</td>
<td>0.40</td>
</tr>
<tr>
<td>CPLnorm</td>
<td>2.515± 1.333</td>
<td>2.777± 1.179</td>
<td>2.419± 0.466</td>
<td>0.38</td>
</tr>
<tr>
<td>Small-worldness</td>
<td>2.138± 0.676</td>
<td>1.834± 0.518</td>
<td>1.965± 0.485</td>
<td>0.24</td>
</tr>
<tr>
<td></td>
<td>GE</td>
<td>Modularity (Q)</td>
<td></td>
<td></td>
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<tr>
<td>----------------</td>
<td>-----------</td>
<td>----------------</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>0.704± 0.070</td>
<td>0.746± 0.109</td>
<td></td>
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<tr>
<td></td>
<td>0.695± 0.105</td>
<td>0.746± 0.109</td>
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<tr>
<td></td>
<td>0.695± 0.105</td>
<td>0.604± 0.026</td>
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<tr>
<td></td>
<td>0.84</td>
<td>0.02</td>
<td></td>
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<tr>
<td></td>
<td>0.604± 0.017</td>
<td>0.604± 0.026</td>
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</tbody>
</table>

Data are presented as mean± standard deviation unless otherwise indicated. CCnorm= normalized clustering coefficient; CPLnorm; normalized characteristic path length.
Nodes showed significant rank ordered differences, such that patients with alcohol use disorder (AUD) < Siblings < Controls for both nodal clustering coefficient (CC\textit{nodal}) and nodal local efficiency (Enodal) after FDR correction.

* Post-hoc t tests demonstrated significantly reduced CC\textit{nodal} and Enodal in AUD group compared with controls in all nodes.
Figure 2 Groups showed rich club organization over a range of k from 7 to 22. 
* Normalized rich club coefficient (RCnorm) showed significant rank ordered differences, such that patients with alcohol use disorder (AUD) < Siblings < Controls over a range of k from 13 to 17 after FDR correction. Post-hoc t tests demonstrated significantly reduced RCnorm in AUD group compared with controls in all significant k levels. Siblings showed significantly reduced RCnorm in k=17 (p=0.046).
Figure 3 Decreased structural connectivity in patients with alcohol use disorder compared to controls determined by network-based statistic analysis. R, right; SFGdor, superior frontal gyrus dorsolateral; IFGoperc, inferior frontal gyrus opercular part; IFGtriang, Inferior frontal gyrus triangular part; CAU, caudate; PUT, putamen; HIP, hippocampus; ITG, inferior temporal gyrus; SMG, supramarginal gyrus; SOG, superior occipital gyrus; MOG, middle occipital gyrus.