Impact of Adolescent Alcohol and Drug Use on Neuropsychological Functioning in Young Adulthood: 10-Year Outcomes

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Because of ongoing neuromaturation, youth with chronic alcohol/substance use disorders (AUD/SUD) are at risk for cognitive decrements during young adulthood. We prospectively examined cognition over 10 years based on AUD/SUD history. Youth (N = 51) with no AUD/SUD history (n = 14), persisting AUD/SUD (n = 18), or remitted AUD/SUD (n = 19) were followed over 10 years with neuropsychological assessments. Groups were compared at baseline and 10-year follow-up. Both AUD/SUD groups declined in visuospatial construction at year 10 (p = .001). Further, cumulative alcohol use (p < .01) and drug withdrawal (p < .05) predicted year-10 visuospatial function. Alcohol use predicted verbal learning/memory (p < .05), while stimulant use predicted visual...
learning/memory ($p = .01$). More recent substance use predicted poorer executive function ($p < .05$). In conclusion, heavy alcohol and other substance use from adolescence through young adulthood may produce cognitive disadvantages, including visuospatial and memory decline. Youth with heavy, chronic alcohol use and/or drug withdrawal symptoms may be at particular risk.

KEYWORDS adolescence, alcohol, executive function, memory; neurocognition, substance use disorders, visuospatial function, withdrawal, young adulthood

INTRODUCTION

Alcohol and other substance use disorders (AUD/SUD) are common among youths and often continue into adulthood. In 2008, 72% of high school seniors reported having at least tried alcohol, and 25% reported heavy drinking (five or more drinks in a row) in the past two weeks (Johnston, O’Malley, Bachman, & Schulenberg, 2009). Use tends to increase in young adulthood, with 40% of college students reporting recent heavy drinking (Johnston et al., 2009). Because neuromaturation continues through adolescence and young adulthood (Toga, Thompson, & Sowell, 2006), the neurocognitive effects of alcohol and other drug use are of great concern, and youths with AUD/SUD may be at particular risk for cognitive dysfunction.

In adult populations, chronic heavy alcohol use leads to neurocognitive deficits. Specifically, broad cognitive impairment is common among adult alcoholics during the acute detoxification period (Fein, Bachman, Fisher, & Davenport, 1990), including visuospatial, memory, and executive deficits (Reed, Grant, & Rourke, 1992; Sher, Martin, Wood, & Rutledge, 1997; Sullivan, Rosenbloom, & Pfefferbaum, 2000) as well as gait and balance abnormalities (Sullivan et al., 2000). After several weeks of abstinence, attention and concentration, reaction time, and memory tend to improve (Bates, Voelbel, & Buckman, 2005; Fein et al., 1990). Executive function, processing speed, and non-verbal skills appear more resistant to recovery, while the improvement of verbal abilities is less clear (Bates et al., 2005; Fein et al., 1990). After years of abstinence, adults appear to broadly recover neurocognitive abilities (Reed et al., 1992), although spatial processing may remain affected (Fein, Torres, Price, & Di Scalfani, 2006). The degree of neurocognitive recovery in adults depends on age, premorbid cognitive ability, duration of abstinence, and amount of drinking in the intervening time (Bates et al., 2004; Fein et al., 1990; Rourke & Grant, 1999).

Neuropsychological function among adolescents with heavy alcohol use has not been as thoroughly examined. However, adolescents with alcohol...
use disorders (AUD) have shown decrements in visuospatial and attention performance (Tapert, Granholm, Leedy, & Brown, 2002); retention of verbal and non-verbal material (Brown, Tapert, Granholm, & Delis, 2000); language and executive function (Moss, Kirisci, Gordon, & Tarter, 1994); and intellectual function, academic achievement, speed of processing, and sustained attention (Tapert & Brown, 2000; Tarter, Mezzich, Hsieh, & Parks, 1995). However, because intellectual abilities differed between groups in some cases (Moss et al., 1994; Tarter et al., 1995), the causes of the neuropsychological abnormalities are less clear.

In addition to cognitive deficits, adolescents with AUD have shown altered patterns of brain activation in response to a spatial working memory task during functional magnetic resonance imaging (fMRI), despite performance that is similar to controls (Caldwell et al., 2005; Tapert et al., 2004). In particular, the degree of abnormal activation was greater in adolescents who had more extensive histories of withdrawal/hangover symptoms and alcohol consumption. The results suggest possible reorganization of neuronal circuits that are detectable during the early stages of AUD and may suggest that certain brain areas compensate for decreased activation in other areas (Tapert et al., 2004). Although the groups performed similarly on neuropsychological tests, it is unclear whether continued alcohol use would result in behavioral differences (Caldwell et al., 2005).

Certain brain areas may be especially vulnerable to the effects of heavy alcohol consumption. Animal studies have shown decreased neurogenesis in the adolescent forebrain and hippocampal areas subsequent to ethanol exposure (Crews, Mdzinarishvili, Kim, He, & Nixon, 2006). In humans, the volume of the prefrontal cortex appears reduced in adolescent heavy drinkers (De Bellis et al., 2005; Medina et al., 2008), as does the hippocampus (De Bellis et al., 2000; Medina, Schweinsburg, Cohen-Zion, Nagel, & Tapert, 2007; Nagel, Schweinsburg, Phan, & Tapert, 2005). Alcohol-related prefrontal and hippocampal dysmorphology may help explain the cognitive and memory disadvantages among AUD adolescents.

Most of the studies above are cross-sectional, and considerably less is known about the neurocognitive effects of continued heavy alcohol use by youths as they transition into adulthood. To examine the long-term effects of AUD and SUDs, our group is conducting a longitudinal study of adolescents who received treatment for alcohol and other substance use problems. Previous examinations of the data showed that continued substance use following treatment was associated with poorer visuospatial and attention functioning four years post-treatment (Tapert & Brown, 1999), and lower visuospatial, attention, learning, and retention function eight years post-treatment (Tapert et al., 2002). Withdrawal symptoms (especially from alcohol) were associated with poorer visuospatial functioning (Brown et al., 2000; Tapert & Brown, 1999; Tapert et al., 2002), as well as verbal and non-verbal retrieval (Brown et al., 2000).
The current study prospectively examines neuropsychological functioning of youths after 10 years as a function of cumulative alcohol and drug involvement. In addition to examining differences between healthy young adults and those with persisting AUD/SUD, we were interested in whether individuals with remitted AUD/SUD would show long-term improvements in neurocognitive performance. Based on previous findings, we hypothesized that, relative to controls, young adults with persisting AUD/SUD would show decrements in visuospatial skills, attention, executive function, learning, and retention, while those with remitted AUD/SUD would show similar performance to controls by the 10-year follow-up. We also hypothesized that greater cumulative use of alcohol and other drugs as well as substance withdrawal symptoms over follow-up would be linked to poorer neurocognition at the 10-year follow-up, particularly in visuospatial functioning and retention.

METHODS

Participants

Participants were part of a longitudinal research project following adolescents (N=187; ages 13 to 18 years) with and without alcohol and other substance use disorders who received neuropsychological examinations at baseline and semiannually thereafter up to 10 years (Brown, Myers, Mott, & Vik, 1994; Brown, Vik, & Creamer, 1989; Tapert et al., 2002). Clinical participants were consecutive admissions to inpatient adolescent substance abuse treatment centers who met diagnostic criteria for alcohol abuse or dependence at project intake. The majority of youths also met criteria for at least one other SUD. The comparison sample was solicited through advertisements in the same communities from which clinical participants originated. Community adolescents were selected to be similar to clinical participants on age (±1 year), gender, socioeconomic status, ethnic background, years of education (±1 year), and prevalence of family history of AUD and SUD. Community adolescents were excluded if they had a history of AUD, SUD, or alcohol or drug use problems at project intake.

Recruitment procedures excluded youths who

1. did not have a resource person (typically a parent) to corroborate biographical, health, and substance involvement information;
2. lived >50 miles from the research facility;
3. had an Axis I psychiatric disorder other than conduct disorder predating the onset of regular substance use;
4. did not speak English fluently;
5. had a history of head trauma with loss of consciousness >2 minutes;
6. had any medical or physical condition that could compromise neuropsychological (NP) performance (e.g., epilepsy, migraine); or
7. had prenatal exposure to alcohol (mother drank 3 or more drinks on an occasion or >3 times per month during pregnancy).

Informed assent and consent, approved by University of California, San Diego Institutional Review Board and clinical agencies, was independently obtained from each participant and parent.

Intake assessments were conducted in the treatment facility approximately three weeks after admission for clinical participants, and at the research site for community youths. Follow-up neuropsychological assessments were conducted 2, 4, 6, 8, and 10 years after intake evaluations. Substance involvement information was obtained on over 90% of the cohort through in-person or phone interviews at each follow-up time point (Anderson, Ramo, Cummins, & Brown, 2010). At year 10, individuals who did not complete the study did not differ from included cases on the basis of demographics (age, ethnicity, handedness, education) or substance use patterns at intake. However, individuals who did not complete the 10-year substance use assessment were slightly more likely to be male and have lower parental socioeconomic status (SES) at intake relative to participants who completed the 10-year assessment.

NP testing rates were not as high, due to participant relocations beyond the 50-mile limit. Of the 187 participants who received a NP assessment at intake, 79 participants (42.2%; 51 clinical and 28 community youths) had a 10-year follow-up NP assessment. Reasons for non-completion of the NP at year 10 included the following:

1. participant relocation ($N = 68$);
2. not able to be contacted ($N = 31$);
3. study dropout ($N = 5$);
4. death ($N = 3$); and
5. incarceration ($N = 1$).

No statistically significant differences were found between youths who did and did not receive a NP assessment at 10-year follow-up on (1) intake demographics (age, gender, ethnicity, handedness, education, parental SES), substance involvement, or NP performance; or (2) 10-year demographics or substance involvement, although participants who did not complete a NP assessment at year 10 were slightly more likely to use stimulants more frequently at year 10 (intake-only NP: mean = 1.8 ± 6.1 days per month; 10-year NP: mean = 1.2 ± 5.6). In addition, 10 community controls developed substance-related problems by the 10-year follow-up point and were excluded from the current study because (1) of the small size of this group, and (2) they did not logically fit into the other groups.

For the current evaluation, participants were excluded for recent substance use at the 10-year assessment (<48 hours; $N = 18$) to avoid the effects of intoxication or hangover. This resulted in a sample with at least 2 days
since last alcohol or other substance use (range = 2 to >1000). Participants in this sample reported the last substance used as (1) alcohol ($N=35$), (2) marijuana ($N=4$), (3) stimulants ($N=2$), (4) hallucinogens ($N=1$), or (5) opiates ($N=1$), and 8 participants denied recent use of any substances (i.e., within 1,000 days of assessment). Please refer to Table 1 for more detailed recent and cumulative substance use patterns.

Within this sample ($N=51$), three groups of individuals were studied:

1. community controls with no history of AUD/SUD at intake through follow-up (controls; $N=14$);
2. participants who met AUD/SUD criteria at intake, and had (a) alcohol or drug dependence symptoms or problems in the past two years or (b) past 3-month drug or heavy alcohol use (5 or more drinks on one occasion) at year 10 (persisting AUD/SUD; $N=18$); and
3. participants who met AUD/SUD criteria at intake but reported (a) no alcohol or drug dependence symptoms or problems in the past two

<table>
<thead>
<tr>
<th>TABLE 1</th>
<th>Substance Use Characteristics of Youths Followed over 10 Years</th>
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<tbody>
<tr>
<td></td>
<td>Controls</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Substance Use at Intake</strong></td>
<td></td>
</tr>
<tr>
<td># Alcohol dependence sxs ever$^{***}$</td>
<td>0.1 (0.3)</td>
</tr>
<tr>
<td># Drug dependence sxs ever$^{***}$</td>
<td>0.3 (1.1)</td>
</tr>
<tr>
<td><strong>Substance Use in Past 2 Years</strong></td>
<td></td>
</tr>
<tr>
<td>Days since last substance use</td>
<td>162.0 (356)</td>
</tr>
<tr>
<td>Alcohol, drinks per month, past 3 months</td>
<td>5.1 (5.5)</td>
</tr>
<tr>
<td>Alcohol dependence sxs, past 2 yrs$^{***}$</td>
<td>0.0 (0.0)</td>
</tr>
<tr>
<td>Alcohol withdrawal sxs, past 2 yrs$^{***}$</td>
<td>0.0 (0.0)</td>
</tr>
<tr>
<td>Drug withdrawal sxs, past 2 yrs$^{***}$</td>
<td>0.0 (0.0)</td>
</tr>
<tr>
<td>Drug dependence sxs, past 2 yrs$^{***}$</td>
<td>0.0 (0.0)</td>
</tr>
<tr>
<td><strong>Average Use over Follow-Up Period: Intake to Year 10</strong></td>
<td></td>
</tr>
<tr>
<td>Drug dependence sxs$^{a,d,***}$</td>
<td>0.1 (0.3)</td>
</tr>
<tr>
<td>Drug withdrawal sxs$^{b,d,***}$</td>
<td>0.0 (0.0)</td>
</tr>
<tr>
<td>Alcohol dependence sxs$^{c,d,***}$</td>
<td>0.2 (0.3)</td>
</tr>
<tr>
<td>Alcohol withdrawal sxs$^{d,***}$</td>
<td>0.0 (0.2)</td>
</tr>
<tr>
<td>Alcohol, avg. drinks per month$^{c,d,***}$</td>
<td>5.4 (5.1)</td>
</tr>
<tr>
<td>Marijuana, avg. use days per month$^{c,d,***}$</td>
<td>0.1 (0.2)</td>
</tr>
<tr>
<td>Stimulants, avg. use days per month$^{a,***}$</td>
<td>0.1 (0.2)</td>
</tr>
</tbody>
</table>

*Note: Values represent means (and standard deviations). Refer to methods section for description of composite variables summarizing substance use over the 10-year follow-up. Sxs = symptoms.

Mann-Whitney post hoc tests:

$^{a}$Persisting and Remitted AUD/SUD > Controls;

$^{b}$Persisting AUD/SUD > Remitted AUD/SUD and Controls;

$^{c}$Persisting AUD/SUD > Controls;

$^{d}$Persisting AUD/SUD > Remitted AUD/SUD;

$^{**} p < .01$. $^{***} p < .001$ between three groups.
years and (b) no recent drug or heavy alcohol use at year 10 (remitted AUD/SUD; N = 19).

Measures

STRUCTURED CLINICAL INTERVIEW

A 90-minute confidential structured clinical interview (Brown et al., 1994; Brown et al., 1989) administered to youths and parents assessed demographic information, as well as social, academic, and occupational functioning, family history of AUD/SUD, maternal drinking during pregnancy, physical and emotional health problems, and SES (higher scores reflect lower SES) (Hollingshead, 1965).

CUSTOMARY DRINKING AND DRUG USE RECORD (CDDR)

The CDDR (Anderson, Ramo, Schulte, Cummins, & Brown, 2007; Brown et al., 1998) gathered self-report information on lifetime and current alcohol and drug involvement. The follow-up version obtained information on the use of alcohol (beer, wine, liquor) and drugs (marijuana, amphetamines, barbiturates, hallucinogens, cocaine, inhalants, opiates, and prescription medications or other drugs not previously specified), life problems related to alcohol and drug use, DSM-III-R and DSM-IV alcohol- and drug-dependence criteria (American Psychiatric Association, 1987, 1994), and alcohol and drug withdrawal symptomatology. The CDDR incorporates the Cahalan drinking classification procedure (Cahalan, 1970), Drug Indulgence Index (Lu, 1974), and Alcohol Dependence Scale (Horn, Skinner, Wanberg, & Foster, 1984) items. At each follow-up assessment, alcohol and drug information was collected for the past three months and since the previous assessment (e.g., two years). Good internal consistency, test-retest reliability, and inter-rater reliability have been demonstrated in adolescent and young adult samples using these time periods (Brown et al., 1998; Stewart & Brown, 1995).

NEUROPSYCHOLOGICAL TEST BATTERY

The NP battery at project intake included the following: Wechsler Intelligence Scale for Children–Revised (WISC–R; Wechsler, 1974) Vocabulary, Similarities, Block Design, Arithmetic, and Digit Span (digits forward, digits backward) subtests; Controlled Oral Word Association Test (COWAT; Benton, Hamsher, Varney, & Spreen, 1983); Wechsler Memory Scale (WMS) Visual Reproduction subtest (Wechsler, 1945); California Verbal Learning Test (CVLT; Delis, Kramer, Kaplan, & Ober, 1987); and Trail Making Test (Reitan & Wolfson, 1993). The year-10 battery included: Wechsler Adult Intelligence Scale–Revised
(WAIS–R; Wechsler, 1981) Vocabulary, Similarities, Block Design, Arithmetic, and Digit Span subtests; COWAT; CVLT-II (Delis, Kramer, Kaplan, & Ober, 2000); and Trail Making Test. In addition, WMS Visual Reproduction was replaced with the Rey-Osterrieth Complex Figure test (ROCF; Osterrieth, 1944) copy and 30-minute delay, employing the Taylor system for scoring accuracy (Lezak, 1995). To reduce the influence of practice effects, alternate forms of tests were administered when possible. Testing was approximately two hours at each time point. Finally, each NP measure was normed using T-scores based on the control group's raw scores at intake and year 10. Therefore, the mean of the control group at both time points is $T = 50$.

Procedures

Clinical and community youths were administered the NP test battery and structured interview by a trained psychometrist at project intake and at each follow-up. Adolescents recruited from treatment programs received the initial assessment during inpatient stay after a three-week detoxification period. The other assessments were conducted at the research facility or the participant’s residence. Youths and parents were each interviewed privately by different interviewers. At each follow-up, 10% of youths were randomly selected for urine drug toxicology screening to confirm self-reported substance use. Discrepancies in self-report and toxicology results (<1% of cases) were scored for use on toxicology-identified substances.

Data Analyses

NEUROPSYCHOLOGICAL DATA REDUCTION

To reduce the number of comparisons and Type I error risk, composite scores were generated to summarize the cognitive domains tested. We used a hybrid method that relied on established cognitive domains (Lezak, Howieson, & Loring, 2004) and reliability analyses (Delis, Jacobson, Bondi, Hamilton, & Salmon, 2003). After performing necessary $\log_{10}$ transformations to meet the assumptions of parametric analysis (intake and year-10 Trails A and B time-to-completion), standard scores were created for each individual cognitive measure for the entire sample ($N = 51$) at both intake and year 10. For each measure, higher standard scores represented superior performance. Next, tests were grouped to form distinct cognitive domains derived from a theoretical background, and standard scores were averaged within each domain. To ensure internal consistency of the individual measures in each composite score, we calculated standardized Cronbach’s $\alpha$ coefficients. We reexamined composites with Cronbach’s $\alpha$ coefficients less than .60 until good reliability was obtained. In two cases, the neuropsychological measure was best
represented by standing alone (i.e., Trails A: time-to-completion and WISC-R or WAIS-R Block Design).

Six summary scores representing performance at intake and year 10 were derived:

1. Language (WISC-R or WAIS-R Vocabulary and Similarities subtests);
2. Psychomotor Speed (Trails A: time-to-completion);
3. Visuospatial Construction (WISC-R or WAIS-R Block Design);
4. Visuospatial Learning & Memory (WMS Visual Reproduction Immediate and Delayed Recall [intake]; ROCF copy and delayed recall [year 10]);
5. Verbal Learning & Memory (CVLT or CVLT-II: total of trials 1–5, short-delay free recall, long-delay free recall, percent savings [(total words correctly recalled on long-delay free recall/total words recalled on trial 5) × 100]); and
6. Executive Function (Trails B: time-to-completion, COWAT, WISC-R or WAIS-R Arithmetic and Digit Span subtests [total number of digits recalled in forward and backward sequence]).

Cronbach’s \( \alpha \) for each composite score ranged from .72 to .83 with two exceptions: Language at intake (\( \alpha = .66 \)) and Executive Function at year 10 (\( \alpha = .65 \)).

SUBSTANCE USE, DEPENDENCE, AND WITHDRAWAL SCORES

To examine the effects of specific substances on neurocognition, indices were created to represent cumulative consumption of the three most commonly used substances in this sample (Tapert et al., 2002). Alcohol consumption was calculated as the average number of standard ethanol consumption units per month (over the preceding 3 months) at each follow-up time point over the 10-year period. Marijuana use was quantified as the average days per month of marijuana use at each follow-up time point. Stimulant use was calculated as the average days per month on which amphetamine or cocaine use occurred at each follow-up. Alcohol withdrawal was computed as the average number of different types of alcohol withdrawal symptoms endorsed for the preceding two years at 4-, 6-, 8-, and 10-year follow-ups, plus the number of different alcohol withdrawal symptoms endorsed for the preceding 3 months at the 6-, 12-, and 24-month follow-up time points. Drug withdrawal and alcohol- and drug-dependence symptoms were estimated in the same manner. The substance use assessment portion of the follow-up includes more than 90% of the sample at each time point over the 10-year period, and all 51 participants had substance use data from at least four follow-up time points to facilitate computing these summary scores. Thus, we can be confident of the composite substance use scores derived herein. Distributions of variables were examined, and all substance use, dependence, and withdrawal scores were log_{10} transformed to meet the assumptions for parametric analysis (Tabachnick & Fidell, 2007).
DEMOGRAPHIC COMPARISON

One-way ANOVAs and Fisher’s exact tests compared groups on demographic variables, including age, gender, ethnicity, handedness, SES, education, and the number of neuropsychological testing sessions completed. The non-parametric Kruskal-Wallis procedure examined group differences in alcohol and other substance use. As groups differed on age at intake (1.3 years) and years of education at year 10 (2.5 years), these were entered as covariates for subsequent group comparisons. SES also differed between groups, but since year-10 participant SES (Hollingshead score) and education were highly correlated (Spearman’s $\rho = -0.66$, $p < .001$), education was selected as the most appropriate covariate for this age group. Gender, education at year 10, and participant SES were not linked to year-10 NP scores. Analyses used SPSS 14.0 (Chicago, Illinois).

LONGITUDINAL ANALYSES

Repeated measures ANCOVAs compared groups on each of the six NP composite scores at intake and year-10 time points ($\alpha = .05$) controlling for age at intake and years of education at year 10. Hierarchical regression analyses were conducted to ascertain which composite substance involvement variables most strongly predicted performance at year 10. For the first set of regressions, Block 1 entered intake age, intake NP composite score, and year-10 education; Block 2 entered (stepwise) the cumulative alcohol, marijuana, and stimulant scores. In the second set of regressions, Block 1 entered intake age, intake NP composite score, and year-10 education; Block 2 entered (stepwise) cumulative alcohol withdrawal symptoms, cumulative drug withdrawal symptoms, and days since last substance use at year 10. Dependent variables were the six 10-year NP composite scores.

RESULTS

Demography and Substance Use

The three groups were similar in gender, ethnicity, and handedness distributions, and parent SES at baseline (see Table 2). Remitted AUD/SUD youths were about a year older than controls at intake, but this age difference was non-significant at 10-year follow-up. On average, the remitted and persisting AUD/SUD groups had lower levels of education and SES at year 10 than controls (note that higher Hollingshead SES scores reflect lower SES). Because the effects of repeated testing can confound the interpretation of NP performance (Matarazzo & Herman, 1984), the number of times the NP battery was administered to each participant was totaled. Importantly, groups were statistically equivalent with regard to the number of times tested. A repeated measures ANOVA showed that groups were similar in their estimated IQ
Neuropsychological 10-Year Outcomes

TABLE 2 Demographic Characteristics of Youths Followed over 10 Years

<table>
<thead>
<tr>
<th></th>
<th>Controls</th>
<th>Persisting AUD/SUD</th>
<th>Remitted AUD/SUD</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td>14</td>
<td>18</td>
<td>19</td>
</tr>
<tr>
<td>Age at intakea</td>
<td>15.2 (1.6)</td>
<td>15.8 (1.2)</td>
<td>16.5 (0.8)</td>
</tr>
<tr>
<td>Age at year-10 testing</td>
<td>25.4 (1.7)</td>
<td>26.7 (2.2)</td>
<td>26.8 (1.1)</td>
</tr>
<tr>
<td>Gender (males:females)</td>
<td>6.8</td>
<td>14:4</td>
<td>9:10</td>
</tr>
<tr>
<td>Caucasian (%)</td>
<td>64%</td>
<td>67%</td>
<td>79%</td>
</tr>
<tr>
<td>Right-handed (%)</td>
<td>86%</td>
<td>89%</td>
<td>84%</td>
</tr>
<tr>
<td>Parent SES at intake</td>
<td>31.8 (13.4)</td>
<td>31.7 (15.1)</td>
<td>25.8 (9.3)</td>
</tr>
<tr>
<td>Participant SES at year 10b**</td>
<td>32.4 (13.9)</td>
<td>47.8 (14.2)</td>
<td>44.3 (12)</td>
</tr>
<tr>
<td>Years of education at year 10b***</td>
<td>15.1 (2.1)</td>
<td>12.6 (1.9)</td>
<td>12.7 (1.9)</td>
</tr>
<tr>
<td># of NP testings by year 10</td>
<td>7.43 (1.3)</td>
<td>6.94 (1.1)</td>
<td>7.16 (1.3)</td>
</tr>
</tbody>
</table>

Note: Values represent means (and standard deviations) unless otherwise noted. SES determined by Hollingshead scores (higher scores reflect lower SES).

Post hoc tests:
a Remitted AUD/SUD > Controls;
b Controls > Persisting and Remitted AUD/SUD;
*p < .05. **p < .01. ***p < .001 between three groups.

levels at intake and year 10 [WISC-R and WAIS-R Vocabulary scores: group: $F(2,48) = 1.19$, $\text{NS}$, $\eta^2_p = .05$; group $\times$ time: $F(2,48) = 2.03$, $\text{NS}$, $\eta^2_p = .08$]. Substance use was comparable among persisting AUD/SUD and remitted AUD/SUD participants at intake, and generally higher among persisting AUD/SUD throughout follow-up (see Table 1).

Neuropsychological Functioning

REPEATED MEASURES ANALYSIS

After controlling for age at intake and education at year 10, a main effect of group [$F(2,46) = 3.67$, $p < .05$, $\eta^2_p = .14$] was found for Visuospatial Construction (i.e., Block Design), with controls performing worse than the remitted and persisting AUD/SUD groups at intake. A group $\times$ time interaction for Visuospatial Construction [$F(2,46) = 7.73$, $p = .001$, $\eta^2_p = .25$] indicated that, despite performing better than controls at baseline, both persisting and remitted AUD/SUD groups showed greater declines in visuospatial construction abilities from intake to 10-year follow-up (see Figure 1). A marginally significant group $\times$ time effect was also detected for the Language composite score [$F(2,46) = 2.78$, $p = .073$, $\eta^2_p = .11$]. Again, both abuse groups performed better than controls at intake but showed greater declines than controls over the 10-year follow-up period.

SUBSTANCE USE AND WITHDRAWAL REGRESSIONS

Hierarchical regressions ascertained whether alcohol and other substance use over the 10-year follow-up period predicted NP performance at year 10.
Greater cumulative alcohol use significantly predicted poorer year-10 Visuospatial Construction performance \(F(4,46) = 12.09, p < .001; R^2_\Delta = 10.3\%, \beta = -.33, p < .01\) and Verbal Learning & Memory \(F(4,44) = 8.27, p < .001; R^2_\Delta = 7.4\%, \beta = -.28, p < .05\) above and beyond the effects accounted for by age, intake Visuospatial Construction or Verbal Learning & Memory performance, and year-10 education, while cumulative marijuana and stimulant use were not predictive in either case. However, greater stimulant use over the 10-year period significantly predicted poorer Visual Learning & Memory performance at year 10 \(F(4,46) = 6.64, p < .001; R^2_\Delta = 10.1\%, \beta = -.33, p = .01\) above and beyond the effects accounted for by age, intake Visual Learning & Memory functioning, and education at year 10 (see Table 3).

A second set of regressions examined whether alcohol and other drug withdrawal symptoms over the 10-year follow-up period or recency of substance use at year 10 predicted NP performance at year 10. After controlling for age and Visuospatial Construction performance at intake and year-10 education, having more drug withdrawal symptoms over the follow-up period was predictive of poorer year-10 Visuospatial Construction performance \(F(4,46) = 10.85, p < .001; R^2_\Delta = 7.6\%, \beta = -.31, p < .05\), while alcohol withdrawal over follow-up and days since last substance use at year 10 were not predictive of year-10 performance. In addition, more recent use of alcohol

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**FIGURE 1** Persisting and remitted AUD/SUD groups show reduced block design T-scores from intake to 10-year follow-up, relative to control participants. Note: Each NP measure was normed using T-scores based on the control group’s raw scores at intake and year 10. Therefore, the mean of the control group at both time points is \(T = 50\). Error bars depict standard error of the mean.
or other drugs predicted poorer Executive Function at year 10 \( F(4,42) = 13.54, p < .001; R^2_D = 7\% \), \( \beta = .28, p < .05 \) above and beyond the effects accounted for by age, intake Executive Function, and education at year 10.

EXPLORATORY ANALYSES

We examined potential gender differences between groups for each of the six NP domains using repeated measure ANCOVAs, controlling for age at intake and education at year 10. No significant gender or group \( \times \) gender effects were detected.

DISCUSSION

We studied the long-term NP consequences of protracted AUD/SUD over a 10-year period from mid-adolescence to young adulthood. Previous studies have reported visuospatial, memory, and attention deficits among individuals with continued heavy drinking or drug use (Brown et al., 2000; Tapert & Brown, 1999; Tapert et al., 2002). The primary aim of the current study was to examine whether youths with remitted AUD/SUD would show improvements in NP functioning relative to youths with persisting AUD/SUD. Community controls with no history of AUD/SUD were also examined. The three groups performed similarly in visuospatial learning and memory, verbal learning and memory, executive function, and psychomotor speed at project intake and at a 10-year follow-up assessment. However, young adults with a history of AUD/SUD, whether persisting or remitted, showed

<table>
<thead>
<tr>
<th>Dependent variable</th>
<th>Alcohol use</th>
<th>Marijuana use</th>
<th>Stimulant use</th>
<th>Alcohol withdrawal</th>
<th>Drug withdrawal</th>
<th>Days since last substance use</th>
</tr>
</thead>
<tbody>
<tr>
<td>Language</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
</tr>
<tr>
<td>Psychomotor Speed</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
</tr>
<tr>
<td>Visuospatial</td>
<td>-.33**</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>- .31**</td>
<td>n.s.</td>
</tr>
<tr>
<td>Construction</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
</tr>
<tr>
<td>Visual Learning &amp;</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
</tr>
<tr>
<td>Memory</td>
<td></td>
<td>-.33**</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
</tr>
<tr>
<td>Verbal Learning &amp;</td>
<td>-.28*</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
</tr>
<tr>
<td>Memory</td>
<td></td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
</tr>
<tr>
<td>Executive Function</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>.28**</td>
</tr>
</tbody>
</table>

Note: All regressions controlled for intake age, intake neuropsychological performance, and years of education at year 10.

*p < .05. **p ≤ .01.
a relative decline in visuospatial construction performance from intake to the 10-year follow-up relative to controls without a history of substance-related problems, even after controlling for age and education. A similar pattern was found for language abilities, although this finding was only marginally significant. These findings suggest that an individual’s lifetime history of substance-related problems may negatively influence his or her cognitive function long after he or she has discontinued heavy or disordered use of alcohol or other drugs.

When examining multiple substance use indicators simultaneously, average alcohol use and average drug withdrawal symptoms over the 10-year follow-up appeared most predictive of visuospatial construction skills in young adulthood. Alcohol use during the follow-up period also predicted verbal learning and memory, while stimulant use significantly predicted visual learning and memory performance. In all cases, more use or withdrawal symptoms over the follow-up period was associated with poorer NP outcomes. Furthermore, having used alcohol or other drugs more recently at the 10-year assessment predicted poorer executive function. Overall, this pattern of findings suggests that youths with remitted AUD/SUD may not perform substantially better than youths who continue to have alcohol- or drug-related problems. Rather, the cumulative effects of protracted substance use and withdrawal during mid- to late adolescence and into young adulthood appear related to long-term cognitive functioning in the mid-twenties, particularly visuospatial construction and visual and verbal memory. In addition, recent substance use may influence executive function in young adulthood.

The association of poorer visuospatial and memory performance with protracted substance use is in concordance with previous reports from this longitudinal study (Tapert & Brown, 1999; Tapert et al., 2002), as well as other investigators (Fein et al., 1990; Fein et al., 2006; Reed et al., 1992; Sher et al., 1997; Sullivan et al., 2000). Consistent with our previous reports on NP functioning at four and eight years after substance use treatment during adolescence (Tapert & Brown, 1999; Tapert et al., 2002), more drug withdrawal symptoms over the follow-up period were associated with poorer visuospatial function. However, in the current study, alcohol withdrawal symptoms were not related to visuospatial performance, and we did not find decrements in attention (Brown et al., 2000; Tapert & Brown, 1999; Tapert et al., 2002) perhaps due to limited power. This could suggest that adolescent substance-use-related visuospatial and memory problems may be particularly enduring, or vulnerable to the impact of chronic use.

Possible reasons for discrepancies with previous reports include different statistical approaches and sample characteristics. For example, due to upgrades in the neuropsychological battery over the follow-up period, the current study included measures that were similar at intake and 10-year follow-up, while earlier reports included a wider range of NP tests. Furthermore, measures included in each of the composite scores varied between
studies. The study inclusion/exclusion criteria at project intake were the same for all participants, but previous reports may have excluded participants from analyses for different reasons. For example, while the current study excluded participants for recent substance use prior to NP testing, previous reports used even more stringent substance exposure exclusions due to larger sample sizes (Tapert & Brown, 1999; Tapert et al., 2002). In addition, a unique aspect of the current study was the examination of individuals with remitted AUD/SUD relative to those with persisting AUD/SUD and healthy controls. Previous reports from this sample examined only two groups or defined the groups slightly differently. In the current study, we did not detect deficits in intellectual function or processing speed (Tarter et al., 1995). Possible reasons for this discrepancy include use of different recruitment sources, distributions of male and female participants, neuropsychological batteries, inclusion/exclusion criteria, length of abstinence, and approaches to data analysis.

Although our sample size is modest, the current findings suggest that heavy, chronic alcohol and other substance use during late adolescence and early adulthood may produce long-term cognitive disadvantages, primarily in visuospatial and memory abilities. Furthermore, by including a group of remitted users, this investigation demonstrated that having a lifetime history of heavy alcohol or other substance use during adolescence and young adulthood appears to be more important than whether or not individuals continued to have substance-related problems into their mid-twenties. Youths who consume heavy quantities of alcohol and/or consistently experience drug withdrawal symptoms over several years may especially be at risk. Chronic stimulant use may also influence visual memory functioning, and more recent substance use may have a specific impact on executive functions, which in the current study focused primarily on verbal working memory. Alternatively, some youths may have had a disadvantageous developmental course predetermined by factors other than substance use, which adversely influenced both cognitive performance and substance use outcomes.

Alcohol-related cognitive deficits may have important negative consequences for adolescents and young adults. For example, in the early phases of abstinence, recall deficits may disadvantage youths in academic environments (Brown et al., 2000). Cognitive ability also appears to moderate the relationship between coping skills and substance use after treatment (Tapert, Brown, Myers, & Granholm, 1999). That is, among adolescents with poor cognitive abilities, those with fewer coping skills had worse substance use outcomes one year after treatment, while having a larger repertoire of coping skills appeared to protect these individuals from subsequent drug and alcohol use. For those with average or better intellectual abilities, other factors such as temperament, family, community, and work environments may account for future substance exposure (Brown et al., 2008). Thus, treatment for substance use problems may be improved with the integration of coping skills training, as well as cognitive habilitation (Moss et al., 1994). The cognitive decline evident
in this and prior longitudinal studies of AUD/SUD youths is subtle and seldom obvious to the individual substance abuser or those in the academic environments. Consequently, youths in treatment may benefit from personal feedback regarding their neurocognitive functioning and potential consequences of continued use (Ramo, Myers, & Brown, 2007). It is likely that substance-related cognitive deficits have other consequences for adolescents and young adults, but more research is needed to examine such characteristics.

Some limitations with the current study should be considered. First, our sample size is relatively small due to the stringent inclusion/exclusion criteria and participant relocations over a 10-year study period. However, participants who received a NP assessment at year 10 were generally similar to those who did not on baseline demographic, substance use, and NP variables, and on 10-year demographic and substance use characteristics. While it is possible that heavier substance users were lost during follow-up, our data suggest that the majority of the original participants at least completed the substance use assessment at year 10. Second, practice effects could influence results given the repeated testing over the 10-year follow-up period, although groups did not differ in the number of NP assessments received during the study. Third, while participants were excluded for using substances before testing to avoid effects of acute intoxication, this did not rule out potential effects of protracted hangover or withdrawal symptoms on NP functioning. Nevertheless, this abstinence duration generalizes readily to substance users with frequent use patterns. Fourth, this clinical sample consists of individuals who have a lifetime history of AUD and most are polysubstance users; consequently, the effects of any single substance cannot be isolated. Furthermore, the community sample was matched to the clinical sample on demographics and family history of AUD/SUD, and it could be that the family history and SES factors modify what would be expected in a true representative community sample. Therefore, the current findings suggest that heavy, chronic alcohol use in the context of other substance involvement may be associated with visuospatial and memory decrements relative to individuals of similar demographics.

Finally, for this study we summarized substance use patterns by creating composite scores that reflected average use of substances, dependence symptoms, and withdrawal symptoms over the 10-year period. While this approach enabled us to retain more participants in this longitudinal study, important information about the long-term effects of alcohol and drugs on cognition may be gained by examining the fluctuating patterns of substance use in more detail. Given the low base rate of alcohol withdrawal symptoms and the high prevalence and intensity of use, consideration of alternative means to assess alcohol withdrawal symptoms may be merited. Future studies might examine the neurocognitive outcomes of various substance use trajectories particularly during critical periods of brain maturation (Anderson et al., 2010) as well as factors that either contribute to cognitive decline or are neuroprotective during the transition to young adulthood. The clinical and
occupational consequences, as well as the neural correlates (e.g., structural and functional neuroimaging studies) of these substance-related cognitive deficits may have important ramifications.

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