Exposure and Loss of Environmental Enrichment Mediates Ethanol Consumption in Adolescent Female Rats

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https://doi.org/10.33697/ajur.2019.032

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ABSTRACT
Alcohol use among adolescent females has significantly increased in the United States with young women drinking alcohol at the same rate as young men. One potential treatment strategy that could help sustain alcohol abstinence is Environmental Enrichment (EE). Environmental enrichment is a process concerning the stimulation of the brain by one's physical and social surrounding, which promotes non-drug reinforcement alternatives (e.g. voluntary exercise) supporting drug abstinence. Thus, the primary focus of this study was to investigate the effect of EE on ethanol (ETOH) abstinence in adolescent female rats. All adolescent female rats, starting on postnatal day 30, had 24-h access to 2%, then 4%, and then 6% ethanol concentrations. At the end of the four weeks, the environmental conditions were switched (EE → NEE and NEE → EE) and the 6% ethanol measure was repeated. We found that EE significantly reduced ethanol consumption for adolescent female rats compared to controls. Further, the removal of EE opportunities resulted in a significant increase in ethanol consumption. Collectively, the results suggest that access to enriched life conditions are important in facilitating alcohol abstinence in adolescent female rats.

KEYWORDS
Adolescent Females; Alcohol Consumption; Environmental Enrichment; Alcohol Use Disorder; Treatment Strategy; Alcohol Abstinence; Ethanol; Adolescent Female Rats

INTRODUCTION
Alcohol use among young women has significantly increased in the United States within recent years. Young women are now drinking alcohol at similar rates as young men, suggesting that differences in consumption of alcohol for males and females has dramatically narrowed. For example, thirty-seven percent of ninth grade girls—averaging at about 14 years old—report drinking in the past month, surpassing the percentage rate reported for ninth grade boys. Unfortunately, the increased use of alcohol in young women can set the stage for development of an alcohol use disorder (AUD) later in life. According to the National Institute of Health, AUD is a chronic relapsing brain disease characterized by compulsive alcohol use, loss of control over alcohol intake, and a negative emotional state when not using. Given the recent upward trend of AUDs among adolescent females, it is important that substance abuse animal research examine possible treatment strategies that may be effective for treating adolescent female alcohol abuse.

One promising treatment strategy that has shown to support drug abstinence is environmental enrichment. Environmental Enrichment (EE) can be defined as the non-contingent delivery of alternative non-drug rewards such as food, social interaction, novelty objects and voluntary physical activity either in the presence of drug (concurrent) or in the absence of drug (non-concurrent). Access to nondrug alternatives can impede or prevent acquisition and decrease drug-maintained responding. For example, given the choice between drug and other types of rewards (e.g., toys or social interaction) they will typically prefer the alternative rewards over drug. Further, the removal of such non-drug alternatives may also result in increased drug taking.

To date, research has provided promising evidence that EE may indeed support drug abstinence in male animal populations. Access to EE can prevent the acquisition of drug taking and decrease drug responding in male rats. For example, male animal studies have shown that EE reduces cocaine and heroin’s reinforcing effects when concurrently available with the drug. However, there is a paucity of research concerning EE as a possible treatment strategy for substance abuse in female animal populations. To our knowledge, there have been no studies examining EE as a possible treatment method for alcohol consumption in adolescent female rats. Therefore, the current study examined a novel approach by implementing an EE treatment strategy that may help support alcohol abstinence in adolescent female rats.
Our primary focus was to investigate the effects of exposure and loss of EE on ethanol (ETOH) consumption in adolescent female rats. During phase one, adolescent female rats were either placed in the EE or in standard non-enriched cages (NEE) with both groups having access to ETOH. For both groups, adolescent female rats were given 2% ETOH for one week, then increased to 4% for another week and finished with 6% over the span of two weeks. At the end of the fourth week, we switched the groups (phase two) where female rats that were originally in the EE were now placed into the NEE and rats that were first in the NEE were given access to the enriched environment. We hypothesized that both groups of adolescent female rats would show significantly less ETOH consumption while in the EE condition, than in the NEE condition. If these results were to be observed then this would lend support for the implementation of EE as a strategy to facilitate alcohol abstinence in adolescent female rats.

METHODS AND PROCEDURES

Subjects
Twenty Sprague-Dawley adolescent female rats weighing between 300 and 350 g were used as subjects for both phases of the experiment. Estrous levels in free cycling females were not assessed. Each rat was housed in a climate-controlled environment ranging from 70.0-72.0 degrees Fahrenheit with constant access to water and Lab Rat Chow was provided during all phases of the experiment. Each rat was individually housed under a reversed 12-hour light: dark cycle (lights on at 19:00 h). This study complied with the guidelines of the National Institute of Health guide for the care and use of laboratory animals and was conducted in accordance with the SUNY Cortland’s Institutional Animal Care and Use Committee ethics protocol (IACUC Protocol #43).

Environmental Enrichment
Every female adolescent rat was provided a standard cage, 25 × 20 × 15 cm³ for the NEE and 60 × 38 × 20 cm³ for the EE (depending upon experimental phase). The EE contained running wheels, a 10-cm diameter tunnel, ladders, treats, and two additional objects that were rotated daily, such as a jingly ball, paper roll, and a dog chew toy. The components and procedure of rotating objects daily within enrichment cages, are similar to those used in other enrichment studies that have shown effects of the treatment.8,20-22

Procedure
The procedure consisted of two four week phases. During phase 1, starting on postnatal day 30, adolescent female rats were assigned to either the EE (n = 10) or NEE (n = 10) with a 14-oz drinking bottle containing an ascending series of ethanol concentrations placed on their home cages 24 hours per day (continuous access). Twenty female rats were exposed to different concentrations of ETOH by a fade-in procedure (Table 1). This procedure gradually introduces the rats to the ETOH solutions that provide a measure of volitional intake under restrictive conditions.21 This fade-in series started with 2% ETOH for the first seven days, followed by 4% for the next seven days and then 6% for 14 days. Bottles were weighed to the nearest 0.1g at the same time daily. At the end of the fourth week, we switched the groups (phase 2) where female rats that were in the EE were now placed into the NEE (EE-NEE) and rats that were in the NEE were given access to the enriched environment (NEE-EE). During phase 2, all rats had continued access to 6% for 14 days.

<table>
<thead>
<tr>
<th>Experimental Groups: (EE → NEE vs. NEE → EE)</th>
<th>Week 1</th>
<th>Week 2</th>
<th>Weeks 3 &amp; 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phase 1: Ethanol Exposure</td>
<td></td>
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<tr>
<td>Duration</td>
<td>7 Days</td>
<td>7 Days</td>
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<tr>
<td>Phase 2: Switched</td>
<td></td>
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<tr>
<td>Ethanol Exposure</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Duration</td>
<td>7 Days</td>
<td>7 Days</td>
<td>N/A</td>
</tr>
</tbody>
</table>

Table 1. Ascending series of ETOH concentrations administered during phases 1 and 2.
Statistical Analysis
The dependent variable consisted of the amount of ethanol consumption (6%) by the female rats across both phases. The independent variables were environmental history (NEE or EE) and groups (EE→NEE vs. NEE→EE). A two-way ANOVA with group (between-subjects) and environmental history (repeated measures) was conducted comparing the mean 6% ETOH consumption amounts in the EE and NEE conditions for both EE→NEE and NEE→EE groups. Differences were considered significant at the p<0.05 level. Standard error bars are indicated in Figure 1. All statistical analyses were conducted using IBM Statistical Package for Social Sciences (SPSS) 26 software.

RESULTS
A two-way ANOVA with group (between-subjects) and environmental history (repeated measures) as factors was conducted comparing the mean 6% ETOH consumption amounts in the EE and NEE conditions for both EE→NEE and NEE→EE groups. The 6% ETOH consumption over 14 days for the EE→NEE female adolescent rats during the NEE condition yielded a mean of 17.73 ± 2.16 grams (g), while the 6% ETOH consumption over 14 days for the EE→NEE group during the EE condition led to a mean of 5.40 ± 0.93(g). The 6% ETOH consumption over 14 days for the NEE→EE rats during the NEE condition yielded a mean of 19.71 ± 2.45(g), while the 6% ETOH consumption over 14 days for this same group led to a mean of 7.54 ± 1.23(g). (Figure 1). For both EE→NEE and NEE→EE groups, there was a significant effect of environmental history \(F(1, 39) =16.59, p<.01\]. However, there were no significant effects for group \(F = 3.59, p>.05\] or group by environmental history interaction \(F = 0.89, p>.05\].

DISCUSSION
The current study examined a possible treatment method for supporting alcohol abstinence in adolescent female rats. We found that EE exposure led to significantly less consumption of ethanol for both female groups, while loss of EE led to a significant increase in ethanol intake. The results suggest that having access to EE opportunities while alcohol is readily available may be an effective treatment strategy for promoting decreased ethanol consumption in adolescent female rats. A possible explanation for this finding is that the introduction of rewarding stimulation experienced in the enriched environment might have reduced the reinforcing effects of the ethanol through a behavioral contrast mechanism. Behavioral contrast refers to a change in the rate of reinforcement on one component of a concurrent schedule produces an opposite change in the rate of response on another component creating an inverse relationship.\textsuperscript{23-25} For example, a change to a high reinforcement rate in one schedule component typically results in a lower response rate in the other schedule component. A relevant example for the current experiment is when environmental enrichment studies provide concurrent access to alternative reinforcement (e.g. toys and treats) while the drug is
The current findings may have important implications concerning treatment strategies for alcohol abuse in adolescent females. Reinforcement (e.g., wheel running) it leads to a reduction in the reinforcing effects of the drug, thereby facilitating abstinence. (largely with animal male populations) using EE by demonstrating that when having concurrent access to alternative activity in the presence of drug found that access to nondrug alternatives led to a decrease in drug-maintained responding. 

An important aspect of the current study was to examine the changes in adolescent female ethanol consumption after loss of EE stimulation. Previous research has shown that the removal of non-drug alternatives results in the behavioral resurgence of drug responding or an increase in drug taking. Furthermore, in humans, researchers have suggested a link between the removal of alternative, reinforcing events and increases in drug intake or instances of behavioral resurgence after periods of abstinence. For example, researchers examined data from a Health and Retirement study in order to explore the relationship between involuntary job loss and smoking intensity, as well as, relapse in abstinent smokers. They found that involuntary job loss contributed significantly to elevated levels of smoking in individuals who already smoked. Furthermore, risk of relapse doubled after job loss in ex-smokers. Similarly, we found that loss of access to non-drug alternatives (EE) led to a significant increase in ethanol consumption for adolescent female rats. Thus, the common feature that both the current and previous studies all demonstrate is that when stimulation or reward is derived from a source other than the drug itself (enrichment), there is a reduction in the reinforcing effects of the drug(s), thereby supporting abstinence. Further, when non-drug alternative rewards are no longer being delivered, a behavioral resurgence of drug use could be the result.

One possible neural mechanism whereby EE may produce its effect on ethanol consumption in adolescent female rats is by disrupting neural circuits in areas involved in ethanol taking. For example, researchers have found that EE housing reduced the incentive value of novelty and the reinforcing properties of ETOH that are mediated by the mesocorticolimbic dopaminergic reward system. The disruption of neural circuitry by EE is supported by other studies that found EE rats previously trained to self-administer cocaine and after cue-induced relapse tests for cocaine had activated cFos protein in the mesocorticolimbic system to a lesser extent than in NEE animals. Brain alterations in neural pathways associated with alcohol abuse suggests that EE may play a disruptive role in the neural mechanisms associated with alcohol consumption. Further, this may help explain how the concurrent introduction of rewarding stimulation (EE), may have the effect of reducing the reinforcing effects of ethanol through the contrast mechanism described above.

Previous male animal studies that delivered alternative non-drug rewards such as, food, novelty objects, and voluntary physical activity in the presence of drug found that access to nondrug alternatives led to a decrease in drug-maintained responding. Similarly, the current study found that when adolescent female rats were provided access to EE alternatives there was a reduction in ethanol consumption. Therefore, EE may also be an effective treatment strategy for substance abuse when implemented in female populations.

One limitation to the present study is that we did not assess estrous levels in freely cycling adolescent female rats, while in the EE or NEE conditions. Increased estrogen levels have been shown to be related to increased alcohol use in females. However speculative, it may be that EE disrupts the estrogenic effects that contribute to female’s increased sensitivity to the rewarding effects of alcohol. In part, this possible explanation could account for significantly less ETOH consumption rates by female rats during the EE condition compared to the NEE condition. Therefore, further studies should examine the possible role that EE may play in mediating the estrogenic effects on alcohol use in adolescent female rats. Another limitation to the present study involves the restricted age cohort of investigating the possible EE preventative effects on ETOH in adolescent female rats. Future studies should examine the heterogeneity of treatment effects concerning EE on ETOH use by observing other female age cohorts.

CONCLUSIONS

The primary focus of this study was to investigate a potential treatment strategy that would support ETOH abstinence in adolescent female rats. We found that exposure to EE led to significantly less consumption of ETOH for both female groups, while loss of EE led to a significant increase in ethanol volitional intake. The present results support previous research findings (largely with animal male populations) using EE by demonstrating that when having concurrent access to alternative reinforcement (e.g., wheel running) it leads to a reduction in the reinforcing effects of the drug, thereby facilitating abstinence. The current findings may have important implications concerning treatment strategies for alcohol abuse in adolescent females.
ACKNOWLEDGEMENTS
The authors thank the State University of New York at Cortland for their generous funding that made this work possible.

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ABOUT STUDENT AUTHORS
Natalie Lipari will graduate in 2020 with a Bachelor of Science degree in Psychology and plans to continue her research focus in the area of substance abuse as she pursues her Ph.D. in Behavioral Neuroscience. Max Baron will graduate in 2021 from the University of Michigan with a Bachelor of Science degree in Biopsychology, Cognition and Neuroscience. He plans to continue research in the area of Behavioral Neuroscience upon graduate school acceptance.

PRESS SUMMARY
Recent research has shown that female adolescent alcohol abuse is on the rise in the United States. In fact, the number of adolescent females (ages 12-17) with an alcohol use disorder (AUD) actually surpassed the number of adolescent males with an AUD. The consequences of underage drinking can be devastating and may result in truancy, motor vehicle injuries, sexual assault cases, and even death. Thus, with the rise of adolescent female alcohol consumption and increased risk of AUD, it is essential to determine a treatment method that supports long-term alcohol abstinence in females. One potential treatment is the implementation of environmental enrichment (EE). Environmental enrichment is a process concerning the stimulation of the brain by one’s physical and social surrounding that promotes non-drug reinforcement alternatives supporting drug abstinence. Further, accumulating evidence indicates that exposure to an EE during the earlier stages of life reduces the effects of abused drugs and decreases the vulnerability to develop a substance abuse disorder. To our knowledge, there have been no studies that have examined EE as a possible treatment strategy to support alcohol abstinence in adolescent female rats. Therefore, the current research sought to examine this novel approach by implementing EE as a viable treatment option to support abstinence in female alcohol abuse. The results suggest that access to enriched life conditions are important in facilitating alcohol abstinence in adolescent female rats.