Determining Ethyl Glucuronide Cutoffs When Detecting Self-Reported Alcohol Use in Addiction Treatment Patients

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Background: Ethyl glucuronide (EtG) is an alcohol biomarker with potential utility as a clinical research and alcohol treatment outcome. Debate exists regarding the appropriate cutoff level for determining alcohol use, particularly with the EtG immunoassay. This study determined the EtG immunoassay cutoff levels that most closely correspond to self-reported drinking in alcohol-dependent outpatients.

Methods: Eighty adults with alcohol dependence and mental illness, taking part in an alcohol treatment study, provided urine samples 3 times per week for up to 16 weeks (1,589 samples). Self-reported drinking during 120 hours prior to each sample collection was assessed. Receiver operating characteristic analyses were conducted to assess the ability of the EtG immunoassay to detect self-reported alcohol use across 24- to 120-hour time periods. Sensitivity and specificity of EtG immunoassay cutoff levels was compared in 100 ng/ml increments (100 to 500 ng/ml) across 24 to 120 hours.

Results: Over half (57%) of the 1,589 samples indicated recent alcohol consumption. The EtG immunoassay closely corresponded to self-reported drinking from 24 (area under the curve [AUC] = 0.90, 95% confidence interval [CI]: 0.88, 0.92) to 120 hours (AUC = 0.88, 95% CI: 0.87, 0.90). When cutoff levels were compared across 24 to 120 hours, 100 ng/ml had the highest sensitivity (0.93 to 0.78) and lowest specificity (0.67 to 0.85). Relative to 100 ng/ml, the 200 ng/ml cutoff demonstrated a reduction in sensitivity (0.89 to 0.67), but improved specificity (0.78 to 0.94). The 300, 400, and 500 ng/ml cutoffs demonstrated the lowest sensitivity (0.86 to 0.33) and highest specificity (0.86 to 0.97) over 24 to 120 hours.

Conclusions: For detecting alcohol use for >24 hours, the 200 ng/ml cutoff level is recommended for use as a research and clinical outcome.

Key Words: Ethyl Glucuronide, Urine Testing, Alcohol Biomarkers, Alcohol Treatment.

A N ESTIMATED 17 million Americans suffer from alcohol use disorders (Grant et al., 2004) and nearly 80,000 people die from alcohol-related causes every year (Centers for Disease Control and Prevention, 2004). Accurate assessment of alcohol use is needed to identify alcohol use disorders and evaluate treatment effectiveness. Currently, self-report instruments such as the alcohol Timeline Followback (TLFB) and Alcohol Use Disorders Identification Test are considered the “gold standard” of alcohol assessment in clinical research and treatment. However, the validity of self-reported alcohol use can vary (Babor et al., 2000; Del Boca and Noll, 2000), particularly when respondents face social, treatment, or legal contingencies for alcohol use (Langenbucher and Merrill, 2001).

The combination of self-report and biological assessments, such as point-of-care drug immunoassay urine tests, is considered to be the best method for assessing illicit drug use (Donovan et al., 2012; Jatlow and O’Malley, 2010). These urine tests are frequently used in research and treatment settings because of their accuracy, rapid results, and detection period of at least 48 hours (Chermack et al., 2000; Donovan et al., 2012; Ries et al., 2002). Although many biomarkers of recent alcohol consumption exist, none to date, have demonstrated a detection period of more than 48 hours while still being feasible for use in a clinical research or treatment setting. Measures of blood alcohol content such as breath tests or urinary ethanol (EtOH) can only detect alcohol use during the preceding 12 hours, making them suitable for detecting current intoxication only (Helander et al., 1996). Transdermal alcohol monitors allow for continuous monitoring, but are relatively expensive and questions exist regarding their feasibility, convenience, and perceived stigma due to use in forensic settings (Gurvich et al., 2013). Enzymes such as gamma-glutamyltransferase can be measured at increased levels in people with alcohol dependence,
but have limited utility in detecting low levels of drinking, or infrequent, nonchronic binge drinking (Goldberg, 1980; Rosalki, 1975). Carbohydrate-deficient transferrin (CDT) is the most specific serum biomarker of heavy alcohol consumption; however, the sensitivity is somewhat limited, especially in people with severe liver disease, and point-of-care analysis of CDT is not yet feasible (Anton, 2001; Berthalet et al., 2014). Phosphatidylethanol has shown potential for detecting heavy drinking episodes for approximately 2 weeks but may have difficulty detecting lower levels of alcohol consumption and requires blood collection, which may not be feasible in addiction treatment settings (Helander et al., 2012). Finally, ethyl sulfate (EtS) has performed well as a biomarker for recent alcohol consumption (up to 36 hours); however, there is no commercially available EtS immunoassay (Helander and Beck, 2005; Wurst et al., 2006). Therefore these tests, while valuable in other contexts, have limited utility in clinical research or treatment settings.

Ethyl glucuronide (EtG) is a minor nonoxidative hepatic metabolite of EtOH. It can be detected in a variety of bodily tissues (e.g., hair and nails) and in urine. A commercially available immunoassay test of this metabolite can be conducted by nontechnical staff using a relatively small bench-top analyzer that delivers results of a semi quantitative assessment of EtG concentration in urine within 20 minutes (Leickly et al., 2015). This semiquantitative assessment allows for the use of varying cutoff levels above which results are considered to be positive for alcohol consumption. Preliminary studies have shown EtG immunoassay to have detection rates similar to the more established liquid chromatography–tandem mass spectrometry (LC–MS/MS) method of EtG testing (Böttcher et al., 2008; Leickly et al., 2015; McDonell et al., 2011), which has limited utility in clinical settings as it requires transportation of samples to commercial laboratories and carries a relatively high cost.

Despite its promise as an alcohol biomarker, widespread use of EtG remains limited outside of forensic settings. Due to concerns about overdetection of alcohol use based on incidental nonbeverage alcohol exposure, the Substance Abuse and Mental Health Services Administration (SAMHSA) Advisory (2011) recommended a relatively conservative cutoff level of 500 ng/ml. However, few studies have examined this issue in alcohol-dependent populations (Jatlow and O’Malley, 2010). Furthermore, when EtG is used as a clinical research or treatment outcome, the 500 ng/ml cutoff level may underdetect drinking (Anton, 2014; Jatlow and O’Malley, 2010; Jatlow et al., 2014). Therefore, research is needed to build consensus regarding an acceptable EtG cutoff level, as commercial laboratories, immunoassay manufacturers, and foreign regulatory authorities use cutoff levels ranging from 100 to 1,000 ng/ml (Rohrig et al., 2006; Thierauf et al., 2009, 2010).

In a recently published study, Jatlow and colleagues (2014) used an alcohol challenge paradigm and data gathered from 2 clinical trials to investigate appropriate EtG cutoff levels. In the alcohol challenge study, standardized low, medium, and high doses of alcohol were administered to a small sample (n = 18) of non-alcohol-dependent participants, and their EtG levels were monitored via LC/MS–MS analysis over 12, 24, 48, and 72 hours. At 12 hours postadministration, detection rates of low-dose alcohol use by EtG at the 100 and 200 ng/ml cutoffs were both 100%. In contrast, the 500 ng/ml cutoff only detected 50% of cases of low-dose alcohol use. Results from the clinical trials mirrored those of the challenge study, and saw slight increases in sensitivity at lower cutoffs. Data from the 3 phases of this study suggest that EtG cutoff levels much lower than 500 ng/ml are needed to detect moderate to low levels of drinking, particularly when detecting alcohol use for more than 48 hours. In addition, results of this study provide initial evidence that relatively low cutoff levels might be appropriate to detect alcohol use in clinical research.

Despite results of Jatlow and colleagues (2014) and similar studies, no previous research has investigated optimal cutoff levels for detecting any recent drinking using the EtG immunoassay. Further, most previous studies of EtG rely on small samples sizes (Albermann et al., 2012; Dahl et al., 2011), few within subject observations (Jatlow et al., 2014; Wurst et al., 2008), and include participants with relatively low levels of alcohol consumption (Anton, 2014; Wurst et al., 2004). This study compares the sensitivity and specificity of varying EtG immunoassay cutoff levels, ranging from 100 to 500 ng/ml, when detecting any self-reported alcohol use across 24 to 120 hours in a total of 1,589 samples submitted by 80 alcohol-dependent adults.

MATERIALS AND METHODS

Participants

Participants were 80 adults with DSM-IV (American Psychiatric Association, 2000) diagnoses of alcohol dependence and co-occurring mood (72.4%, n = 55) or psychotic (27.6%, n = 21) disorder. Their average age was 47.2 (SD = 11.2), and 71.3% (n = 57) of participants were male. Reported ethnicities were 53.8% (n = 43) Caucasian, 30% (n = 24) African American, 7.6% (n = 6) Hispanic, 2.5% (n = 2) American Indian, 1.3% (n = 1) Asian/Pacific, 3.8% (n = 3) multi racial, and 1.3% (n = 1) other ethnicities. At baseline, participants reported drinking on average 16.2 days (SD = 8.1) of the previous 30 days.

Study Procedures

All study procedures were approved by the Institutional Review Board of the University of Washington. Participants were enrolled in a randomized controlled trial of a contingency management intervention for alcohol dependence. More information about the parent grant can be found at clinicaltrials.gov (identifier: NCT01567943). All participants participated in a 4-week baseline observation period where they submitted urine samples and provided self-reported alcohol use 3 times per week. They received reimbursement in the form of prizes for submitting these data. Participants were then randomized to 12 weeks of a contingency management group where they received prizes for submitting urine samples negative for alcohol and gift cards for attending intensive outpatient addiction treatment groups or a
noncontingent control group where they received prizes for submitting urine samples and self-report data. All participants received treatment as usual, which included intensive outpatient addiction treatment located in an urban community mental health center in Seattle, WA. As part of their involvement in this study, participants submitted up to 51 urine samples (M = 19.0, SD = 16.0) for EtG immunoassay testing. These urine samples were collected 3 times per week across a 16-week assessment period, and monthly during a 3-month follow-up period. A total of 1,589 urine samples were collected.

EtG immunoassays were conducted onsite at an outpatient addiction clinic by clinical research staff using spectrophotometry on a commercially available ThermoFisher Indiko analyzer (Fremont, CA). Bachelor’s degree-level research staff with no formal laboratory or technical training participated in a 1-day training on the operations and maintenance of the analyzer before performing analyses. Diagnostic Reagents Incorporated EtG enzyme immunoassay tests were conducted using EtG 100, 500, 1,000, 2,000 ng/ml, and negative calibrators and EtG 100 and 375 ng/ml controls. Antibody/substrate and enzyme conjugate reagents were used and the analyzer was calibrated weekly. To prevent bacterial hydrolysis, a reported source of exogenous EtG in urine samples (Helandet al., 2007), all samples were analyzed on the day of collection and stored until analysis in a 4°C refrigerator with all calibrators, controls, and reagents. The DRI EtG immunoassay is linear up to 2,000 ng/ml, with a reportable range of 0 to 2,000 ng/ml (the range of the lowest and highest calibrators). As per manufacturer recommendation, dilution procedures were conducted when EtG immunoassay concentrations displayed an error message indicating high absorbance. Only 10 samples (0.6%) required dilution due to high absorbance.

To avoid positive EtG immunoassay results due to inadvertent alcohol exposure, participants were advised at the beginning of the study and reminded at each appointment to avoid using nonbeverage sources of EtOH, including hand sanitizers and mouthwashes. Use of chloral hydrate, the only medication known to interfere with EtG immunoassay analysis (Arndt et al., 2009), was not reported by study participants when asked monthly about prescription medications used.

A measure was created for the purposes of this study to assess the hours since the last drinking episode, as well as the number of standard drinks consumed at the last drinking episode. These variables are known to have the greatest impact on EtG test results. These data were collected using a calendar method, similar to the alcohol TLFB (Sobell and Sobell, 2000). Self-reported hours since last alcohol use (up to 120 hours) were assessed as a continuous integer when each urine sample was collected. Standard drinks consumed at the most recent drinking episode were also assessed as a continuous integer. Using these data, a summary variable was created that indicated whether or not any alcohol use occurred in the prior 24-, 48-, 72-, 96-, and 120-hour period (e.g., the 48-hour detection period includes drinking that occurred between 1 to 48 hours).

**Data Analysis**

Receiver operator characteristic (ROC) analyses were conducted to investigate the sensitivity and specificity of EtG immunoassay in terms of predicting self-reported alcohol use. Hours since last drink, recorded as a continuous integer at each urine sample collection, were coded into a binary measure for positive/negative self-reported alcohol use at 24-, 48-, 72-, 96-, and 120-hour detection periods. ROC analyses were conducted separately for each of the detection periods and the area under the curve (AUC) and 95% confidence intervals (CIs) were calculated. For each of the 5 assessment periods, the sensitivity and specificity of EtG immunoassay cutoff was reported in 100 ng/ml intervals, ranging from 100 to 500 ng/ml. Statistical analyses were conducted in SPSS version 19.0 (SPSS, 2010).

**RESULTS**

Participants self-reported alcohol use within the previous 5 days on 57.2% (909/1,589) of assessments. The mean number of standard drinks reported during the last drinking episode was 4.12 (SD = 6.37, range 0 to 96). EtG immunoassay results were positive in 50.7% (805/1,589) and 34.8% (553/1,589) of the samples at the 100 and 500 ng/ml cutoff levels, respectively. EtG immunoassay results suggested a bimodal distribution, with values clustering near the lower (0 ng/ml) and upper limits of the immunoassay (2,000 ng/ml). Therefore, while the mean EtG immunoassay value was 682 ng/ml (SD = 865 ng/ml), the median value was 104 ng/ml.

**Sensitivity and Specificity**

Overall, the EtG immunoassay was able to correctly identify self-reported drinking from 24 (AUC = 0.90, 95% CI: 0.88, 0.92) to 120 hours (AUC = 0.88, 95% CI: 0.87, 0.90). When balancing sensitivity and specificity, optimal EtG immunoassay cutoff levels for different times periods since last drink were as follows: 420 ng/ml (24 hours, sensitivity = 0.86, specificity = 0.85), 150 ng/ml (48 hours, sensitivity = 0.84, specificity = 0.83), 100 ng/ml (72 hours, sensitivity = 0.83, specificity = 0.81), 86 ng/ml (96 hours, sensitivity = 0.81, specificity = 0.80), and 80 ng/ml (120 hours, sensitivity = 0.81, specificity = 0.78). Figure 1 displays the sensitivity and specificity of the EtG immunoassay cutoff levels from 100 to 500 ng/ml in 100 ng/ml increments across the 24- to 120-hour assessment periods. As Fig. 1 demonstrates, the 100 ng/ml cutoff level had the highest sensitivity, relative to higher cutoff levels across all time periods ranging from 0.93 (24 hours) to 0.78 (120 hours). The 100 ng/ml cutoff level also had the lowest level of specificity across all time periods, relative to other cutoff levels. Across all time periods, the 200 ng/ml cutoff level provided higher sensitivity than 300 ng/ml and higher cutoff levels, while providing improved specificity, relative to the 100 ng/ml. Cutoff levels above 200 ng/ml performed similarly across all assessment periods, with relatively low sensitivity and high specificity.

**DISCUSSION**

The current study builds upon the growing literature suggesting that relatively low EtG cutoff levels are needed to detect alcohol use for more than 24 hours (Dahl et al., 2011; Hegstad et al., 2013; Jatlow et al., 2014; Stewart et al., 2013). Results of this study suggest that the 100 ng/ml cutoff level had the highest level of self-reported drinking detection, relative to higher cutoff levels throughout the 120-hour assessment period. The relatively lower specificity of the 100 ng/ml cutoff level (particularly during the 24- and 48-hour assessment periods) may be reflective of the limitations of assay specificity (Leickly et al., 2015) or may be due to the fact that this low cutoff level may be
detecting use that occurred beyond the 24- to 48-hour assessment period, rather than nonbeverage alcohol use.

Consistent with previous literature (Wurst et al., 2004), cutoff levels of 300 ng/ml and higher appear to be suitable for detection of drinking during the first 24 hours after alcohol use. However, they appeared to be less effective at detecting drinking for more than 24 hours. Importantly, the 500 ng/ml cutoff level used by most commercial laboratories did not demonstrate a benefit in terms of specificity, relative to the 300 or 400 ng/ml cutoff levels. The 200 ng/ml cutoff level offered specificity that is nearly comparable to higher cutoffs while offering increased sensitivity; therefore, it may be an ideal cutoff for those wishing to balance sensitivity and specificity when detecting drinking in clinical and research settings.

This mirrors Jatlow and colleagues’ (2014) recommendation that a cutoff of 200 ng/ml be used in clinical research.

There are several limitations to the present study. First, this study compares EtG immunoassay results to self-reported alcohol consumption data rather than data collected from an alcohol challenge or controlled drinking experiment. Self-reported alcohol use data have been shown to be vulnerable to inaccurate reporting, particularly underreporting of alcohol use when drinking carries real or perceived negative consequences (Langenbucher and Merrill, 2001). More severe drinking problems, higher levels of pre-treatment drinking, and greater levels of cognitive impairment have all been shown to be correlated with less accurate self-report in clinical trials (Babor et al., 2000). It is possible...
that the underreporting of alcohol use could have contributed to decreased specificity in this study.

During the 120-hour assessment period, 100 (6.3%) false positives were recorded at the 100 ng/ml cutoff level, 42 (2.6%) at the 200 ng/ml cutoff, 22 (1.4%) at the 300 ng/ml cutoff, 21 (1.3%) at the 400 ng/ml cutoff, and 18 (1.1%) at the 500 ng/ml cutoff. The data were collected in the context of a contingency management treatment study. When reinforcers were contingent on EtG negative results, 8% of the time participants submitted an EtG positive (>100 ng/ml) urine sample they denied drinking during the previous 120 hours. When reinforcers were not contingent on EtG test results, 5% of the time participants who submitted an EtG-positive (100 ng/ml) urine sample denied drinking during the previous 120 hours. This was a small but statistically significant difference, $\chi^2(1) = 4.5, p = 0.033$. Regardless of study condition, the levels of agreement between EtG immunoassay and self-report in the present study are similar to agreement between urine tests of illicit drugs and self-report in previous research (Chermack et al., 2000; Decker et al., 2015; Hilario et al., 2014). Although self-report has limitations as a validity outcome, it nevertheless provides valuable information in terms of evaluating the accuracy of EtG, particularly in samples where alcohol challenge experiments might not be appropriate, such as those receiving alcohol treatment.

A second limitation is that participants in this study were suffering from co-occurring mental illness in addition to alcohol dependence. Therefore, results may not generalize to other alcohol-dependent populations. Third, regular dilution was not performed, and EtG/creatinine ratios were not calculated to account for varying urine dilution. Recent research (Jatlow et al., 2014; Stewart et al., 2013) has shown these adjustments to be unnecessary, and tests were conducted in this manner to emulate the way in which they would be performed in an actual outpatient addiction clinic. Additionally, confirmatory testing of EtG immunoassay results by EtG-LC-MS/MS was conducted on a random selection of urine samples, rather than all EtG immunoassay positive samples. However, this was likely unnecessary as there appears to be a high level of agreement between EtG immunoassay and EtG-LC-MS/MS (Böttcher et al., 2008; Leickly et al., 2015).

Despite these limitations, results of this study suggest that EtG immunoassay can be used to accurately assess recent alcohol use in a clinical setting. When used as a clinical research or treatment outcome measure, EtG immunoassay can be a relatively low-cost alternative to the more expensive EtG-LC-MS/MS testing. The ability to conduct tests onsite at an outpatient addiction clinic and receive results rapidly using an easily operable analyzer adds to the utility of EtG immunoassay for researchers and clinicians interested in monitoring client alcohol consumption. Additionally, a point-of-care EtG immunoassay dipstick test was recently released (Premier Biotech, Excelsior, MN). While the dipstick test utilizes a cutoff level of 500 ng/ml and little independent information is available about its accuracy, this technology further increases the feasibility of onsite EtG testing. Like all other alcohol and drug biomarkers, sensitivity declines with increasing time since alcohol consumption. Therefore, rates of detection are improved with shorter testing intervals, which advances in technology continue to make more feasible. In conclusion, study results suggest that a cutoff level of 200 ng/ml provides the best balance between sensitivity and specificity in detecting alcohol use within the past 24 to 120 hours. When used in conjunction with self-report, EtG is a valuable tool that is likely to improve the accuracy of alcohol use assessment in clinical research and addiction treatment settings.

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