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DOI
10.1016/j.neubiorev.2019.07.006

Publication date
2019

Document Version
Final published version

Published in
Neuroscience and Biobehavioral Reviews

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Link to publication

Citation for published version (APA):

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Is (poly-) substance use associated with impaired inhibitory control? A mega-analysis controlling for confounders


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https://doi.org/10.1016/j.neubiorev.2019.07.006

Available online 15 July 2019
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Many studies have reported that heavy substance use is associated with impaired response inhibition. Studies typically focused on associations with a single substance, while polysubstance use is common. Further, most studies compared heavy users with light/non-users, though substance use occurs along a continuum. The current mega-analysis accounted for these issues by aggregating individual data from 43 studies (3610 adult participants) that used the Go/No-Go (GNG) or Stop-signal task (SST) to assess inhibition among mostly “recreational” substance users (i.e., the rate of substance use disorders was low). Main and interaction effects of substance use, demographics, and task-characteristics were entered in a linear mixed model. Contrary to many studies and reviews in the field, we found that only lifetime cannabis use was associated with impaired response inhibition in the SST. An interaction effect was also observed: the relationship between tobacco use and response inhibition (in the SST) differed between cannabis users and non-users, with a negative association between tobacco use and inhibition in the cannabis non-users. In addition, participants’ age, education level, and some task characteristics influenced inhibition outcomes. Overall, we found limited support for impaired inhibition among substance users when controlling for demographics and task-characteristics.

1. Introduction

1.1. Substance use and response inhibition

1.1.1. What is response inhibition and how does it relate to substance use?

Inhibitory control, also known as response inhibition, has been defined as the ability to control one’s attention, behavior, thoughts, and/or emotions to override a strong internal predisposition or external lure, and instead do what is more appropriate or needed (Diamond, 2013). Loss of control over one’s behavior is a defining characteristic of addiction. The DSM-5 lists characteristics such as ‘taking larger amounts or over a longer period than was intended’ and ‘unsuccessful efforts to cut down or control alcohol use’ to define the loss of control over drinking (American Psychiatric Association, 2013). Moreover, inhibitory control has been proposed to play an important role at different stages of the addiction cycle, i.e., 1) initial use of substance; 2) transition from recreational use to heavier use and abuse; 3) continuation of use for those who get addicted; 4) relapse after abstinence (e.g., Garavan et al., 2015; Koob and Volkow, 2010). Furthermore, the dual process model on addiction proposes that an imbalance between a hyper-sensitized impulsive system, which is responsible for cue-reactivity, and a comprehensive reflective or control system (including inhibition of impulses) are important in the development of addiction (Bechara, 2005; Gladwin et al., 2011; Volkow et al., 2004, 2015).

Over the past two decades, multiple studies have focused on the relationship between chronic substance use and response inhibition, but findings have been equivocal. Inhibitory impairment has been associated with chronic use of some substances (e.g., cocaine, ecstasy, methamphetamine, tobacco, and alcohol) but not for others (e.g., opioids, cannabis, see for a meta-analysis, Smith et al., 2014). Results also vary in studies of single substances. For instance, heavy drinkers have been reported to make more commission errors than light drinkers on the Go/No-Go task (GNG, Kreusch, Quertemont et al., 2014), while alcohol-dependent and control participants did not differ significantly on the same measure (Kamarajan et al., 2005). Two main issues might explain these conflicting findings, namely the phenomenon of poly-substance use and the use of extreme group designs (i.e., comparing control participants and problematic or disordered substance users). In addition, sample demographics and task characteristics are often not taken into consideration. In order to address these issues in this mega-analysis, we aimed to investigate the relationship between inhibition and use of multiple substances by analyzing individual-level data, while taking demographics and task characteristics into account. In doing so, we did not exclusively focus on populations diagnosed with substance use disorders (SUD, American Psychiatric Association, 2013).

1.1.2. Experimental paradigms: the Go/No-Go task and the Stop-signal task

Successful suppression of motor responses can involve distinct behavioral processes such as “action restraint” or “action cancellation” (Schachar et al., 2007). Action restraint refers to stopping a prepared but not yet initiated response, which is commonly measured using the GNG and its variants, such as Conners’ continuous performance task (Conners and Sitarenios, 2011; Donders, 1969). These tasks focus on the ability to withhold responding if a no-go stimulus is presented. The main variables of interest are the rate of commission errors (i.e., failures to inhibit a response to no-go targets or false alarms), the rate of omission errors (i.e. failures to respond to go targets, or misses), and the response time (RT) to go stimuli. A relatively high rate of commission errors and a short go RT reflects suboptimal inhibition (Smith et al., 2014).
By contrast, action cancellation refers to stopping a response that is already underway. It is typically measured using the Stop-signal task (SST, Logan, 1994). In this paradigm, each trial starts with the presentation of a go signal that requires an overt response such as a button press. On a subset of trials (typically around 25%), the go signal is followed by a stop signal after a certain interval (stop-signal delay, SSD), upon which participants should inhibit their already initiated go response. Usually, an adaptive tracking algorithm controls the SSD, such that there is a 50% probability of inhibiting the response. A horse-race model, assuming an independent race between the ‘go’ and ‘stop’ processes, affords the estimation of the stop-signal reaction time (SSRT, Logan, 1994). Given that the response could not be withheld on n percent of all stop trials (usually around at 50%), SSRT is calculated by subtracting the mean SSD from the go RT that marks the nth percentile in the go RT distribution (Band et al., 2003).

In contrast to the GNG, the latency of the go response and the latency of the stop process are considered to be independent (Logan and Cowan, 1984). Thus, a longer SSRT reflects an inhibitory deficit, whereas a longer go RT is interpreted as a lack of attention among other influencing factors (preparation, choice, and speed-accuracy trade-off, Lifshitz et al., 2005).

In addition to the GNG and the SST, other experimental paradigms, such as the Stroop (Stroop, 1992) and Eriksen Flanker tasks (Eriksen and Eriksen, 1974) have been proposed to measure inhibitory capacities. However, these paradigms measure distractor inhibition rather than motor response inhibition (Nigg, 2000; Ridderinkhof et al., 2004). To keep the present review focused and allow for straightforward comparisons of results, we only included studies using the GNG and SST.

1.2. Research gaps and research needs

1.2.1. Previous meta-analyses and reviews

To date, there are at least nine published meta-analyses or review papers examining the relationship between inhibitory control and long-term substance use or behavioral addiction. In terms of scope, these studies can be classified into three categories. First, literature overviews focusing on a single substance (e.g., alcohol: Aragues et al., 2011; Stavro et al., 2013) or non-substance related disorder (e.g., gambling disorder: Chowdhury et al., 2017; Moccia et al., 2017). These reviews associated alcohol use with prolonged inhibition impairment, up to one month after abstinence (Stavro et al., 2013) and detoxified alcohol-dependent patients showed poor inhibition compared with healthy controls (Aragues et al., 2011). Polysubstance use was not systemically described or controlled for in either of the review studies on alcohol. Individuals with gambling disorder without comorbid SUD were reported to show large inhibition deficits (Chowdhury et al., 2017), which was attributed to impaired activity in prefrontal areas (Moccia et al., 2017). Second, other reviews focused on drawing general conclusions across multiple substances. For instance, Lipszyc and colleagues found that substance users generally did not differ significantly from controls in SST (Lipszyc and Schachar, 2010) and GNG performance (Wright et al., 2014). However, such a review does not provide a clear profile for the effects of these substances in isolation or of specific interactions (i.e., greater than additive or compensation effects). A third category of literature reviews included multiple substances and the results were specified by the substance. Examples include a recent systematic review focused on neuroimaging findings (Luijten et al., 2014) and a meta-analysis focused on behavior (Smith et al., 2014). The latter meta-analysis indicated that inhibitory deficits were apparent for heavy use/disorders related to cocaine, ecstasy, methamphetamine, tobacco, alcohol, and gambling but not for opioids or cannabis, without testing the interaction effect of using multiple substances (Smith et al., 2014). In sum, the current findings and conclusions of reviews and meta-analyses are rather inconsistent. If a conclusion can be drawn, it appears to be the counterintuitive conclusion that reviews and meta-analyses that focused on a specific addictive substance or behavior are more likely to report a significant association with inhibitory control compared to those reporting on multiple substance use. Importantly, none of these reports have considered several key variables that might bias the results, which will be highlighted in the next section.

1.2.2. Important factors to consider

1.2.2.1. Polysubstance use. Polysubstance use broadly refers to the consumption of more than one drug over a defined period, either simultaneously or at different times (Connor et al., 2014; Subbaraman and Kerr, 2015). This involves different sub-categories, namely using different substances, the dependence of one substance and co-use of other substances or dependence on multiple substances. For instance, tobacco smoking is strongly associated with alcohol and marijuana use (Connor et al., 2014), opioids and benzodiazepines are often prescribed simultaneously (Jones et al., 2012), and stimulants users are more likely to be heavy drinkers (McCabe et al., 2005). Note that there is some evidence indicating that concurrent use of substances can lead to additionally toxic effects because of a toxic metabolite, as was reported for alcohol and cocaine (Pennings et al., 2002). It is also possible that the use of one substance decreases the negative effect of another substance, as found with alcohol and cannabis (Schweinsburg et al., 2011). Hence, studying interactions between drugs on neurocognitive functions is important, given the frequent occurrence and possible interaction effects. However, studies comparing substance users versus non-users or light users have typically focused on the primary substance of concern, while ignoring secondary substances. Up to now, only a few studies have investigated the relationship between polysubstance use and inhibition (Gamma et al., 2005; Moallem and Ray, 2012; Verdejo-Garcia et al., 2007). Heavy drinking smokers did not show poorer SST response inhibition than smokers only and heavy drinkers only (Moallem and Ray, 2012). Similarly, ecstasy polysubstance users did not show more strongly inhibited inhibitory brain mechanisms compared with controls (Gamma et al., 2005), and cocaine and heroin polysubstance users showed similar commission error rates as controls in the GNG (Verdejo-Garcia et al., 2007). A limitation of the latter two studies is that the greater-than-additive effect could not be examined without a group of single substance users. The lack of studies calls for a synthesis of research that does take polysubstance use into account.

1.2.2.2. Substance use as a continuous variable. All the above-mentioned reviews and meta-analyses included comparisons between a control or light user group and a heavy or problematic user group. Scores retained as a result of such extreme group designs are often coded and analyzed in terms of low versus high, reducing individual differences into a binary code. This practice involves ignoring individual-differences of substance use in favor of creating quasi-arbitrary groups assumed to be homogeneous on the variable of interest (MacCallum et al., 2002; Royston et al., 2006; Preacher et al., 2005). In the current study, we aimed to quantify substance use as a continuous variable.

1.2.2.3. Abstinence. Studies on long-lasting effects of substance use have generally been conducted by testing recently abstinent users. With respect to response inhibition, some studies have found that abstinence from cocaine, methamphetamine and heroin normalized inhibitory function (Morie et al., 2014; Schulte et al., 2014), however, one study found sustained suboptimal performance after heroin abstinence (e.g., Pu et al., 2008). In addition, the duration of abstinence appears to moderate the return to normal functioning, which may explain these conflicting findings (Schulte et al., 2014). In order to preclude this as a confounder, we did not include studies on abstinence in (formerly) dependent users. All participants indicated substance use in everyday life, but were requested to refrain from using all substances (in most cases excluding tobacco) 24 h to one week before testing.
1.2.2.4. Individual-level and task-level variables. Some individual-level and task-level factors are known to affect inhibitory control and are therefore included in this mega-analysis, including the demographic variables age, sex, and education years. For GNG, six task parameters were controlled for: no-go percentage, number of experimental trials, working memory load (taxed or not), substance-related stimuli (used or not), cued GNG or not, and task complexity. For the SST, five task parameters were controlled for: number of experimental trials, stop-trial percentage, SSD settings, stop-signal modality, and SSRT calculation method. Reasons for controlling these confounders are based on a large primary literature on these tasks and are summarized in Supplementary Materials S1. Except for sex, for which the interaction with substance use was considered, all other factors were only controlled for regarding their main effect.

1.3. Why a mega-analysis rather than a meta-analysis?

A meta-analysis combines the summary statistics (i.e., effect sizes of included studies), while a mega-analysis combines the raw individual data from different studies. The latter method allows studying the combined effect of individual characteristics (cf. Price et al., 2016) and examining the interaction effect of multiple substances used with enhanced statistical power (Riley et al., 2010). Therefore, we implemented a mega-analysis with individual-level data.

1.4. The goal of the current study

Our primary goal was to examine the main and interaction effects of various kinds of long-term substance use on response inhibition. As the interaction effects of substance use on inhibition are rarely investigated and reported, we explore these interactions in the current study. We do so while controlling for demographics (e.g., age, sex, education years) and task-related factors (e.g., no-go percentage, number of trials, whether stimuli are substance-related) that likely explain performance variance between studies and individuals. Interactions between substance use and sex were also included. Based on the literature reviewed above, we tested the following hypotheses: 1) According to Smith et al. (2014) and other findings (Colzato et al., 2007; Fillmore and Rush, 2002; Quednow et al., 2007), we assumed that the inhibitory deficit would be more pronounced in users of psychostimulants (e.g., cocaine, ecstasy, methamphetamine, tobacco, and alcohol), especially for cocaine and amphetamines, given the known neuropsychopharmacology of the cortical and subcortical networks underlying impulse control (i.e., the right dorsolateral and inferior frontal cortices, Koob and Volkow, 2010; Smith et al., 2014); 2) Given the literature, and as a validation of our individual-level mega-analysis, we expect some demographics (e.g., age and sex) and task characteristics (e.g., no-go percentage, whether stimuli are substance-related) to be associated with inhibition performance (see for expected directions of effects, Supplementary Materials S1).

2. Method

2.1. Study identification and selection

PsycINFO, Medline, EMBASE, Web of Science, CINAHL, and Cochrane Library were searched until 01/03/2016. Search terms and synonyms indicating substance use (alcohol, amphetamine, cocaine, cannabis, heroin, ketamine, methamphetamine, benzodiazepines, gambling, gamer, and internet addiction) were combined with terms indicative of inhibition (go/no-go, inhibitory control, inhibitory process, response inhibition, stop task, etc.). Published meta-analyses and reviews were also checked for additional studies (Horsley et al., 2011). Although behavioral addictions (e.g., gambling, internet addiction) were initially included, there were too few relevant studies to allow further analyses.

2.1.1. Eligibility criteria

The first author (YL) assessed the eligibility of all records using the following initial inclusion criteria: (a) presented in English; (b) conducted on human participants; (c) reported at least one measure from the following: no-go commission errors or go RT in the GNG; SSRT or go RT in the SST; (d) reported use of at least one kind of substance (e.g., alcohol, tobacco, cannabis, amphetamine, cocaine, ecstasy). Note that we included behavioral data from fMRI/EEG studies if available. In addition, we ran supplementary analyses to investigate whether inhibition performance varied with study type (behavioral/EEG/fMRI). It turned out that study type did not systematically influence behavioral performance (see Supplementary Materials S2). We excluded studies (a) that presented stop signals using a single SSD, as this is known to invalidate a performance strategy of delayed responding (Logan, 1994); (b) in which the percentage of no-go or stop trials was higher than 50%, as this is known to invalidate the task (Nieuwenhuis et al., 2004; Randall and Smith, 2011); (c) that focused on the acute effects of substances on inhibition; (d) that recruited participants with a family history of substance dependence; (e) that included polysubstance users; (f) with participants that already received treatment for SUD or abstained from substance use; (g) with participants younger than 18. The exclusion of both intoxicated and abstinent consumers may have kept heavily affected/addicted participants from being included in the sample.

After applying the inclusion and exclusion criteria by YL, a second rater (author YG) assessed the eligibility of a random subset (20%) of the records and obtained 100% agreement. Authors of eligible studies were invited via email to contribute raw data. Repeated attempts were made (i.e., four reminders were sent) if no response was received. Corresponding authors of the identified studies were asked to share their raw individual data, following our instructions on data requirements. The ‘essential variables’ included a set of pre-identified variables, including sociodemographic characteristics (e.g., age, sex, and education), typical alcohol and tobacco use (as alcohol and tobacco are two most commonly used substances), and task performance (Table S1a, S1b). ‘Optional variables’ (Supplementary Materials S3) included other demographic information recorded (e.g., race), other substance use (e.g., cocaine, cannabis) and questionnaires administered (e.g., Alcohol Use Disorder Identification Test (AUDIT), Saunders et al., 1993). The ‘optional variables’ were defined in a more flexible format with open questions. A study was included in our mega-analysis only if information about all ‘essential variables’ could be provided.

2.1.2. Quality assessment and data extraction

As the quality of included studies can influence mega-analysis in unpredictable ways (i.e., shortcomings in original studies will be carried over to the mega-analysis and thus weaken its conclusions, Müller et al., 2019), a quality assessment of original studies was conducted. The methodological quality of studies was assessed by two authors (YL and YG) separately. We used the National Heart, Lung, and Blood Institute (NHLBI) Quality Assessment Tool for Observational Cohort and Cross-Sectional Studies, which is widely used and recommended by Cochrane for quality assessment of observational and cross-sectional studies (Table S2, National Heart, Lung and Blood Institute, 2014). The total agreement (Good/Fair/Suboptimal) between assessors was high (GNG: 20/24 = 83%, SST: 16/20 = 80%). Inter-rater reliability, measured using Spearman’s rank correlation coefficient was high for GNG ($r = 0.84, p < 0.001$) and moderate for SST ($r = 0.56, p = 0.01$, Kendall, 1938).

All provided data, including predictors (i.e., substance use, demographics, task characteristics) and dependent variables were merged into four datasets separated based on the four dependent variables (i.e., the commission error rate in GNG, go RT in GNG, SSRT in SST, and go RT in SST. As speed-accuracy trade-off is a potential issue in GNG (Zhao et al., 2017), a balanced integration score was calculated (Liesefeld and Janczyk, 2019). Main results applying this score as the outcome are presented in Supplementary Materials S4. The first author (YL)
| Study | Sample size (n) | Age (mean) | MA (%) | Education level (%) | Brain structure (mean) | Substance use intensity | Study phase | Duration (mm) | Other substances use info provided | Task number | Sex ratio (♂/♀) | Task complexity | Task performance | Behavioral findings reported in the original publications | Number of cases excluded | Groups excluded |
|-------|----------------|------------|--------|--------------------|------------------------|------------------------|-------------|-------------|----------------|----------------|----------------|----------------|----------------|----------------|------------------------------------------------|----------------------|----------------|
| Y. Liu et al. (2019) | 41 | 20.65 (1.27) | 41 | NA | Alcohol | >1 daily drinker with a history of binge drinking | Male dominant | 20 | Yes | No | No | No | 10 (5/5) | 408 (4) | There were no differences in commission error rate and go RT, and no significant correlation between commission error rate and go RT. | 1 | 2 |
| Glass et al. (2013) | 106 | 22.64 (4.08) | 49 | 3.2 (2.13) | Adult | N/A | Male dominant | 20 | No | No | Yes | Yes | No | 19 (16/3) | 33 (3/3) | There were no significant correlations between the two variables, and there were no significant differences in the between-group analyses. | 1 | 2 |
| Holzinger et al. (2018) | 83 | 19.86 (3.91) | 40 | 22.9 (2.47) | Adult | All participants took part in both groups in the last part of the study. | Male dominant | 20 | Yes | No | No | Yes | 7 (7/0) | 3 (3/0) | There were no differences in commission error rate and go RT, and there were no significant correlations between the two variables. | 1 | 2 |
| Rommerts et al. (2013) | 29 | 22.7 (7.16) | 35 | 2.4 (2.88) | Adult | Male dominant | Male dominant | 20 | Yes | No | Yes | No | 5 (1/4) | 20 (5/3) | There were no differences in commission error rate and go RT. | 1 | 2 |
| Rommerts et al. (2012) | 30 | 21.42 (1.43) | 47 | 3.5 (2.37) | Adult | Male dominant | Male dominant | 20 | Yes | No | No | No | 4 (4/0) | 3 (3/0) | There were no differences in commission error rate and go RT. | 1 | 2 |
| Lind et al. (2012) | 96 | 21.91 (4.17) | 61 | NA | None | Male dominant | Male dominant | 20 | No | Yes | Yes | No | 41 (19/0) | 30 (0/0) | Executive functions were more strongly correlated with commission errors. | 26 | 26 |
| Legaré-Claude et al. (2014) | 57 | 16.74 (0.55) | 46 | 14.1 (2.85) | Adult | Male dominant | Male dominant | 20 | No | Yes | No | Yes | 4 (4/0) | 0 (0/0) | There were no differences in commission error rate and go RT. | 1 | 2 |
| Leijon et al. (2011) | 39 | 24.48 (1.58) | 72 | 24.4 (1.11) | Tobacco | Male dominant | Male dominant | 20 | No | Yes | No | No | 30 (15/15) | 20 (10/10) | There were no differences in commission error rate and go RT. | 1 | 2 |
| Leijon, O. (2014) | 32 | 25.25 (3.13) | 60 | 15.5 (2.01) | Tobacco | Male dominant | Male dominant | 20 | No | Yes | No | No | 21 (11/10) | 40 (20/20) | There were no differences in commission error rate and go RT. | 1 | 2 |
| Leijon, O. (2013) | 40 | 20.15 (2.02) | 67 | 14.9 (0.19) | Tobacco | Male dominant | Male dominant | 20 | Yes | No | No | No | 39 (16/23) | 30 (0/30) | There were no differences in commission error rate and go RT. | 26 | 26 |
| Leijon et al. (2010) | 36 | 20.58 (1.02) | 100 | 14.9 (1.02) | GA | Male dominant | Male dominant | 20 | No | Yes | No | Yes | 43 (24/19) | 40 (20/20) | There were no differences in commission error rate and go RT. | 1 | 2 |
| Mahieddine et al. (2010) | 39 | 30.64 (1.94) | 72 | 14.3 (2.91) | None | Male dominant | Male dominant | 20 | No | Yes | No | Yes | 14 (6/8) | 11 (6/5) | There were no differences in commission error rate and go RT. | 1 | 2 |
| Patel et al. (2012) | 35 | 21.25 (1.99) | 51 | 14.5 (2.79) | Alcohol | Male dominant | Male dominant | 20 | No | Yes | No | Yes | 15 (7/8) | 20 (10/10) | There were no differences in commission error rate and go RT. | 1 | 2 |
| Patel et al. (2010) | 200 | 21.06 (1.63) | 40 | 17.6 (1.01) | Alcohol | Male dominant | Male dominant | 20 | No | Yes | No | No | 14 (6/8) | 30 (15/15) | There were no differences in commission error rate and go RT. | 1 | 2 |
| Plé et al. (2007) | 81 | 30.03 (1.58) | 66 | 11.9 (1.13) | Cocaine | Male dominant | Male dominant | 20 | Yes | Yes | Yes | Yes | 10 (12/13) | 30 (0/0) | There were no differences in commission error rate and go RT. | 1 | 2 |
| Quattrone et al. (2007) | 71 | 21.36 (1.78) | 100 | 13.9 (1.68) | Ecstasy & cannabis | Male dominant | Male dominant | 20 | No | Yes | Yes | No | 25 (12/13) | 11 (6/5) | Ecstasy group performed better on commission error rate and go RT, but there was no significant difference in other variables. | 1 | 2 |
| Marc et al. (2004) | 82 | 25.59 (5.88) | 40 | 15.8 (2.19) | Tobacco | Male dominant | Male dominant | 20 | No | Yes | No | Yes | 29 (12/13) | 25 (9/16) | There were no differences in commission error rate and go RT. | 1 | 2 |
| Bobeck et al. (2000) | 39 | 22.38 (3.31) | 31 | 16.3 (2.43) | Ecstasy | Male dominant | Male dominant | 20 | No | Yes | No | Yes | 31 (15/16) | 30 (15/15) | There were no differences in commission error rate and go RT. | 1 | 2 |

(continued on next page)
performed the data merging, which was verified by two authors (RW and WW).

2.1.3. Publication bias check

To examine whether significant findings in the original papers are indicative of evidential value, a \( p \)-curve was calculated and plotted (Simonsohn et al., 2015). In a \( p \)-curve, the x-axis represents \( p \)-values below 0.05, and the y-axis represents the percentage of studies yielding such a \( p \)-value. A right-skewed \( p \)-curve indicates evidential value, whereas a left-skewed \( p \)-curve, many \( p \)-values just below 0.05, may be indicative of flexibility in data analysis (Simonsohn et al., 2015). If the data did not indicate evidential value, a 33% power test is performed to examine whether the absence of evidential value is due to insufficient power. A \( p \)-curve disclosure table was added in Supplementary Materials (Table S3) according to Simonsohn et al. (2015). \( p \)-curves and corresponding analyses were conducted using the \( p \)-curve app 4.06 (http://www.p-curve.com/app4, 2018).

2.2. Individual participant data meta-analysis

The analysis was conducted in the following steps: 1) apply additional exclusion criteria to the merged datasets; 2) standardize all continuous independent variables; 3) determine substance-related one-way variables; 4) dummy code all discrete variables; 5) determine and generate substance-related interaction variables; 6) multiple imputations of the missing values using all main and interaction variables; 7) build the linear mixed regression model with fixed effects of all predictors and a random intercept; 8) variable selection by stepwise backward elimination. These eight steps are outlined in more detail below.

2.2.1. Construction of the database

2.2.1.1. Individual and group exclusion criteria. The data from the included studies were stacked into a single data file for each dependent variable, with unique identifiers for each study and for each participant. We further applied some minimal exclusion criteria to the individuals. That is, we excluded a participant if (1) he/she was younger than 18 years old; (2) he/she had missing data on all indices of substance use; (3) the dependent variable of current analysis (e.g., commission error rate) was missing; (4) SSRT was negative. A group of substance users from a certain study was excluded if the substance was not included as a predictor in the model. This happened when there was limited data provided for that substance (see criteria in 2.2.1.3.1). For example, if it was concluded that opiate use was assessed insufficiently across all studies, we did not add opiate as a predictor. Consequently, opiate users were excluded from the analysis. The excluded cases and groups from each study are listed in Tables 1 and 2.

2.2.1.2. Standardisation of independent variables

Demographics like age and education level were transformed respectively into continuous variables years and years of education according to the education system in the country where the study was conducted. Task characteristics such as no-go RT were reported. Alcohol consumption was converted into the continuous variable grams of ethanol per month. Data on alcohol consumption were provided in two different ways. Most researchers provided data based on timeline follow-back (TLFB). These data were either already in grams per month or could be transformed by making use of standard drinks adjusted for country (Cooper, 1999). Some studies only had data from more general questionnaires. For instance, three studies (de Ruiter et al., 2012; Luijten et al., 2013a; Rossiter et al., 2012) provided the raw data of the AUDIT (Saunders et al., 1993). In that case, we multiplied midpoints of item 1 (frequency), midpoints of item 2 (drinking days per week) and standard drinks in the country where the study took place. Similarly, four studies (Littel et al., 2012; Luijten et al., 2011; Luijten, Meerkerk et al., 2015; Luijten et al., 2013b) provided Quantity

Note: go-RT: reaction time for correct go trials; M: Mean; SD: Standard Deviation; NA: Not Available; AUDIT: Alcohol Use Disorder Identification Test; VAT: Videogame Addiction Test; AUQ: Alcohol Use Questionnaire; FTND: Fagerström Test for Nicotine Dependence; SDS: Severity of Dependence Scale.

*Unpublished dataset at time of searching literature.

Why comparisons between substance users and controls could not be obtained from the original paper

interested in the difference between the increasing and decreasing limb of BAC but we only used baseline data when participants were sober.

the correlation between commission error rate and binge score was not reported.

focused on the experimental effect (different kinds of cued GNG) instead of the individual difference.
Table 2
Description of included SST studies (dependent variable is SSRT).

<table>
<thead>
<tr>
<th>Study</th>
<th>Demographic information</th>
<th>Substance use</th>
<th>Task Characteristics</th>
<th>Task performance</th>
<th>Between-group findings reported in the original publication</th>
<th>Number of cases excluded</th>
<th>Groups excluded</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Sample size (male/total)</td>
<td>Death</td>
<td>Education years</td>
<td>Other substance</td>
<td>Task</td>
<td>No-go hit rate (%)</td>
<td>Signal-to-noise ratio</td>
</tr>
<tr>
<td>Bélanger et al. (2017)</td>
<td>159</td>
<td>21.56 (5.16)</td>
<td>64</td>
<td>NA</td>
<td>Cannabis</td>
<td>All participants used cannabis at least once a week in the past month and at least 10 times in the past 6 months.</td>
<td>NA</td>
</tr>
<tr>
<td>Boot et al. (2016)</td>
<td>119</td>
<td>21.73 (3.12)</td>
<td>5</td>
<td>1445 (1.56)</td>
<td>Alcohol</td>
<td>All participants use alcohol on a regular basis. Hinge score was calculated based on the first item of ASSQ.</td>
<td>NA</td>
</tr>
<tr>
<td>Boot et al. (2017)</td>
<td>186</td>
<td>36.77 (12.8)</td>
<td>32</td>
<td>16.45 (2.7)</td>
<td>Depression</td>
<td>No special requirement for substance use.</td>
<td>Cannabis, ecstasy</td>
</tr>
<tr>
<td>Cohen et al. (2007)</td>
<td>24</td>
<td>20.33 (8.3)</td>
<td>83</td>
<td>NA</td>
<td>Cannabis</td>
<td>Recreational cannabis users should consume cannabis 1 to 6 times per month by creating a score for a minimum of 2 years.</td>
<td>NA</td>
</tr>
<tr>
<td>Courney et al. (2012, 2017)</td>
<td>304</td>
<td>37.15 (10.0)</td>
<td>7</td>
<td>12.59 (3.25)</td>
<td>Alcohol</td>
<td>All participants were problem-drinkers, with a minimum of 49 standard drinks per month.</td>
<td>NA</td>
</tr>
<tr>
<td>de Ruiter et al. (2012)</td>
<td>38</td>
<td>36.2 (4.25)</td>
<td>1</td>
<td>11.86 (1.63)</td>
<td>Gambling &amp; Tobacco</td>
<td>Problem gamblers were diagnosed by DSM-5. Heavy cannabis use (at least 15 cigarettes per day).</td>
<td>NA</td>
</tr>
<tr>
<td>Filby et al. (2014)</td>
<td>714</td>
<td>21.14 (7.2)</td>
<td>74</td>
<td>13.52 (5.8)</td>
<td>Cannabis</td>
<td>All participants were cannabis users with at least 4 uses per week (or at least 6 months prior). Among these, 34 were diagnosed with cannabis dependence according to DSM-5. Participants in the cocaine use group scored 34 on DAST, himbiously used cocaine for a minimum of 6 months and used it in the past week.</td>
<td>Cannabis, ecstasy</td>
</tr>
<tr>
<td>Fillmore et al. (2002)</td>
<td>344</td>
<td>48.27 (8.8)</td>
<td>68</td>
<td>12.18 (1.4)</td>
<td>Cocaine</td>
<td>All participants were cocaine users with at least 4 uses per week (or at least 6 months prior). Among these, 34 were diagnosed with cannabis dependence according to DSM-5. Participants in the cocaine use group scored 34 on DAST, himbiously used cocaine for a minimum of 6 months and used it in the past week.</td>
<td>NA</td>
</tr>
<tr>
<td>Galderisi et al. (2001)</td>
<td>59</td>
<td>18.69 (1.1)</td>
<td>64</td>
<td>12.75 (1.37)</td>
<td>Tobacco</td>
<td>Daily smokers should smoke cigarettes daily for at least 6 months.</td>
<td>NA</td>
</tr>
<tr>
<td>Glass et al. (2000)</td>
<td>400</td>
<td>44.1 (6.7)</td>
<td>47</td>
<td>13.22 (2.7)</td>
<td>Alcohol &amp; Tobacco</td>
<td>A self-developed variable of alcohol severity was used, with participants categorized as alcohol abusers, 35 as alcohol dependence without physical dependence, 35 as alcohol dependence with physical dependence.</td>
<td>Cannabis, ecstasy</td>
</tr>
<tr>
<td>Kariri et al. (2014)</td>
<td>53</td>
<td>28.3 (4.1)</td>
<td>47</td>
<td>15.55 (1.5)</td>
<td>Alcohol</td>
<td>All participants were categorized as heavy drinkers with at least 2 drinks a day for 6 months per week. Among them, 12 participants were with AUDIT score &lt;16.</td>
<td>Cannabis</td>
</tr>
</tbody>
</table>
| Kiguel et al. (2015) | 75 | 28 (3.2) | 30 | 17.75 (2.56) | Gambling & Tobacco | pathological gambling (%) and cocaine dependence (DSM-IV-TR) were diagnosed with DSM-IV-TR. | Cannabis | 205 | 20 | Visual | Staircase | Integration | 268.63 | 571.18 | NA | 44 | Gambling disorder | (continued on next page)
Frequency Variability (QFV) score (Lemmens et al., 1992). Again, items of quantity, frequency, and standard drinks were multiplied together. Smoking was coded as cigarettes per day. Two studies (Moallem and Ray, 2012; Rossiter et al., 2012) only had data from the Fagerström Test for Nicotine Dependence (FTND, Heatherton et al., 1991). In these cases, the midpoint of the answer to item “How many cigarettes a day do you smoke” was used for daily cigarette use. One study used a self-developed 7-point Likert scale for the past 6 months tobacco consumption, for which we estimated daily cigarette use with the midpoint scores (Ames et al., 2014). Alcohol and tobacco use were standardized across the full dataset. All the other substance use variables had to be treated as dichotomous variables, as insufficient information was provided for treating it as a continuous variable in the model (see details below).

### Dichotomous variables

For interpretability, dichotomous variables were effect-coded with value +1 or -1. Except for alcohol and tobacco use, other substances were coded as ‘lifetime use (yes = 1/no = -1)’.

Four dummy task-characteristics were defined to classify the GNG studies: ‘working memory load (low/high)’, ‘substance-related (yes/no)’ ‘cued GNG (yes/no)’, and ‘task complexity (low/high)’. High working memory load, substance-related, cued GNG versions and complicated tasks were assigned the value of 1 (otherwise -1). Tasks with high working memory load were also assigned a value of 1 for task complexity as the association between stimuli and response was more complicated in these tasks.

Similarly, for the SST, three dummy task characteristics were extracted, including ‘stop-signal modality (visual/auditory)’, ‘SSD (fixed/staircase-tracking)’ and ‘SSRT calculation (integration/others)’. These variables were assigned a value of 1 if auditory stop signals were used; staircase-tracking procedure for SSD; and integration method for SSRT calculation (otherwise -1).

### Identification and generation of substance-related variables

Except for alcohol use and tobacco use, other kinds of substances had missing data as not all studies provided information. Data provided varied in the level of detail, the way questions were asked, and the substances of main interest. For instance, depending on...
the primary substance of interest, some studies provided detailed information for cannabis use but no information on cocaine use (Bidwell et al., 2013), with an opposite pattern for others (Colzato et al., 2007). In the following section, we explain the criteria for including substance-related variables in the model.

2.2.1.3.1 One-way variables. Due to missing data, a criterion was needed to include a variable in the model. We decided on a minimum of 100 participants per cell for a substance (which comes down to a power of 0.94 for the effect size of 0.5). As a result, final models for the GNG (both commission error rate and go RT) included cannabis, cocaine, amphetamine, ecstasy, and hallucinogens, in addition to alcohol and tobacco. For the SST (both SSRT and go RT), the final models included cannabis, cocaine, and ecstasy in addition to alcohol and tobacco.

2.2.1.3.2 Two-way variables. There were two types of two-way variables; the interaction of sex × substance and substance1 × substance2. Variables of sex × substance were created by multiplying sex with substance directly. For the second type, in order to evaluate whether there was sufficient data to assess these interactions, we again applied a criterion for inclusion. For example, dummy coding cannabis and cocaine use yielded a two by two table cannabis (yes/no) × cocaine (yes/no). The corresponding interaction was only entered into the model if all four cells had more than 20 entries. For alcohol and tobacco use, we dichotomized the data by a median split for table construction only. We performed an additional analysis to test whether the number of substances used was a predictor of inhibition performance, and this was not the case (see Supplementary Materials S5). The list of included two-way variables can also be found in Supplementary Materials (Table S4a-S4d). Demographics (in addition to sex) and task parameters could further moderate the relationship between substance use and inhibition. This, however, was not the focus of the current paper. In order to explore this potential issue, we analyzed interactions between alcohol on the one hand and demographics and task parameters on the other (see Supplementary Materials S6).

2.2.1.3.3 Three-way variables. Three-way variables were generated based on the substance1 × substance2 variables combined with sex. The corresponding variables were entered into the model only when all the eight cells in the three-way table sex (male/female) × substance1 (yes/no) × substance2 (yes/no) consisted of at least 10 entries. The list of three-way variables can be found in Supplementary Materials (Table S4a-4d).

2.2.2 Missing data for independent variables and their interactions

In the analysis of GNG commission error rate, the percentage of missing values ranged from 0 to 68.2% (highest: alcohol × hallucinogens × sex) and in the GNG go RT analysis, it ranged from 0 to 69.6% (highest: alcohol × hallucinogens × sex). For the SST, the percentage of missing values ranged from 0 to 84% for the SSRT (highest: tobacco × ecstasy × sex) and from 0 to 83.2% for the go RT (highest: tobacco × ecstasy × sex, a full list of missing data per variable can be found in Table S4a-4d).

In order to deal with these missing data, we used multiple imputations (Rubin, 2004). The default imputation option in SPSS was chosen. It first scans the data and determines the suitable method for imputation (Monotone or Fully Conditional Specification, FCS; Dong and Peng, 2013). All variables in the mixed regression model, including the main and interactive predictors and the dependent variable, were used for imputation. Apart from that, the discrete variable ‘tobacco lifetime use’ was also used, as some studies assessed tobacco use dichotomously (smokers/non-smokers). It has been suggested that the number of imputations should be similar to the percentage of cases that are incomplete (White et al., 2011) and the precision improves by increasing the number of imputations (Bodner, 2008). Therefore, 100 complete data sets were generated, which were combined into a pooled

---

Fig. 1. PRISMA for the mega-analysis detailing our search and selection decisions.
result using the method proposed by Rubin (Rubin, 2004) and Schafer (Schafer, 1997).

2.3. Statistical analysis

Backward elimination was used for variable selection. Initially, each imputed dataset was analyzed with a linear mixed model including all the above-mentioned main, second order, and third order effects as fixed effects and a random intercept (for which a model summary can be found in Tables S4a–S4d). We did not include random slopes and thus assumed that predictors had similar effects in each study. The fixed effects that were least significant (i.e., the one with the largest p-value) were removed and the model was refitted. Each subsequent step removed the least significant variable in the model until all remaining variables or its higher order variables had p-values smaller than 0.05 (Draper and Smith, 2014). For instance, if the variable alcohol × tobacco was significant, then variables of alcohol and tobacco would also be included in the model, irrespective of their independent significance.

3. Results

3.1. Study selection

3.1.1. Summary of authors’ responsiveness

Applying the inclusion and exclusion criteria resulted in a sample of 153 potentially eligible studies (Fig. 1). Out of these targeted papers, 4 researchers responded that they no longer had access to the datasets, 21 declined to participate, 52 did not respond to our invitation and 11 did not have all the basic information we asked for. In total, we obtained raw data from 65 studies. Out of these, 22 had to be excluded because the authors could not provide all the ‘essential variables’, such as data on monthly alcohol use in grams was unavailable (9 studies), missing data of tobacco use (5 studies), participants were abstaining from cocaine and 1 pathological gambling), 1 (5%) reported the opposite suboptimal outcomes (1 study, provided stop latency instead of SSRT). The full list can be found in Supplementary Materials S7. The final dataset for the GNG comprised of 23 independent datasets from 24 papers (in some cases, more than one paper was published with the same dataset).

Table 3a

<table>
<thead>
<tr>
<th>Study</th>
<th>Q1</th>
<th>Q2</th>
<th>Q3</th>
<th>Q4</th>
<th>Q5</th>
<th>Q6</th>
<th>Q7</th>
<th>Q8</th>
<th>Q9</th>
<th>Q10</th>
<th>Q11</th>
<th>Q12</th>
<th>Q13</th>
<th>Q14</th>
<th>Quality Rating</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ames et al. (2014)</td>
<td>yes</td>
<td>yes</td>
<td>NR</td>
<td>no</td>
<td>no</td>
<td>no</td>
<td>no</td>
<td>yes</td>
<td>yes</td>
<td>no</td>
<td>yes</td>
<td>NR</td>
<td>NA</td>
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<td>fair</td>
</tr>
<tr>
<td>Claus et al. (2013)</td>
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<td>yes</td>
<td>NR</td>
<td>no</td>
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<td>no</td>
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<td>yes</td>
<td>yes</td>
<td>no</td>
<td>yes</td>
<td>NR</td>
<td>NA</td>
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</tr>
<tr>
<td>Hendershot et al. (2015)</td>
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<td>yes</td>
<td>NR</td>
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<td>no</td>
<td>no</td>
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<td>yes</td>
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</tr>
<tr>
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</tr>
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<td>no</td>
<td>no</td>
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<td>NR</td>
<td>NA</td>
<td>yes</td>
<td>good</td>
</tr>
<tr>
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<td>yes</td>
<td>NR</td>
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<td>NR</td>
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<td>NR</td>
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</tr>
<tr>
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<td>NR</td>
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<td>no</td>
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</tr>
<tr>
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<td>NR</td>
<td>CD</td>
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<td>NR</td>
<td>NA</td>
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<td>fair</td>
</tr>
<tr>
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<td>yes</td>
<td>NR</td>
<td>CD</td>
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<td>yes</td>
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<td>NR</td>
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<td>no</td>
<td>yes</td>
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<td>NR</td>
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</tr>
<tr>
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<td>NR</td>
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</tr>
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</tr>
<tr>
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</tr>
<tr>
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<td>NR</td>
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</tr>
<tr>
<td>Roberts and Garavan (2010)</td>
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<td>NR</td>
<td>no</td>
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<td>NR</td>
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</tr>
<tr>
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<td>Takagi et al. (2014)</td>
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<td>yes</td>
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<td>no</td>
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<td>NR</td>
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<td>yes</td>
<td>good</td>
</tr>
<tr>
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<td>NR</td>
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<td>no</td>
<td>yes</td>
<td>NR</td>
<td>yes</td>
<td>yes</td>
<td>good</td>
</tr>
</tbody>
</table>

Note: CD: cannot determine; NA: not applicable; NR: not reported; Meanings of criteria Q1-Q14 can be found in Table S2.
NHBLI assessment tool (see Tables 3a and 3b). For the GNG, most (58.3%) of the studies were of intermediate quality, 37.5% of high quality and 4.2% of suboptimal quality. For the SST, 40% of studies were of high quality and another 60% of intermediate quality. The main limitations were small sample size, especially for the studies focused on neuroimaging findings, and insufficient control of confounders such as the history of other kinds of drug use. For a few studies, the population was not fully described, lacking information of where and when the participants were recruited. To explore whether different study types differ in methodological quality, we did a chi-square test based on Tables 3a and 3b. The results indicate that the percentages of studies of good, fair and suboptimal quality did not differ between behavioral (10/23, 13/23, 0/23), EEG (4/8, 3/8, 1/8) and fMRI (3/12, 9/12, 0/12) studies ($\chi^2 (4, N = 44) = 6.51, p = 0.15$).

3.3. Publication bias check

To examine evidential value in the original studies, a $p$-curve was calculated (Supplementary Materials Fig. S1). Out of the 31 effect sizes (unavailable for some studies), 11 were statistically significant ($p < 0.05$), with 8 $p < 0.025$. The $p$-curve analysis on the association between substance use and response inhibition indicated no evidential value (full $p$-curve $z = −0.98, p = 0.16$; half $p$-curve $z = 0.58, p = 0.72$). However, this was likely due to a lack of power (33% power test, full $p$-curve $z = −0.95, p = 0.17$).

3.4. Main outcomes

3.4.1. GNG: no-go commission errors

None of the substance-related variables or their interactions had a significant effect on the commission error rate. Among all other variables, two demographic variables and three task characteristics significantly predicted commission error rates. Age significantly predicted commission error rate ($\beta = −0.01, p < 0.01, 95\% CI [−0.02, 0.00])$, indicating that older participants showed decreased commission error rates. Education years also significantly predicted commission error rate ($\beta = −0.01, p = 0.03, 95\% CI [−0.02, 0.00]$), indicating the higher the educational level, the lower the commission error rates. The nominal variable working memory load had a significant effect on commission error rate ($\beta = 0.10, p < 0.01, 95\% CI [0.07, 0.14]$), indicating that when working memory load was high, participants made more commission errors. The no-go percentage had a significant effect on commission error rate ($\beta = −0.04, p < 0.01, 95\% CI [−0.07, −0.02]$), such that the higher the no-go percentage, the lower the rate of commission errors. The number of trials also had a significant effect on commission error rate ($\beta = 0.04, p < 0.01, 95\% CI [0.02, 0.07]$), indicating higher commission error rates when there were more trials.

3.4.2. SST: SSRT

Lifetime cannabis use significantly predicted SSRT, with users showing longer SSRT than non-users ($\beta = 5.59, p = 0.03, 95\% CI [0.41, 10.77]$). Tobacco use was positively, although not significantly, associated with SSRT ($\beta = 3.21, p = 0.06, 95\% CI [−0.13, 6.55]$), indicating that the more tobacco was consumed, the longer SSRT. The tobacco × cannabis interaction also had a significant effect on SSRT ($\beta = −4.19, p = 0.03, 95\% CI [−8.03, −0.37]$, Fig. 2). Post-hoc analyses were performed by splitting the imputed data sets and fitting the same restricted model without the interaction term. These analyses revealed that for the cannabis non-users, higher tobacco use was associated with longer SSRT ($\beta = 6.44, t = 2.70, p < 0.01$). For cannabis users, no effect of tobacco use on SSRT was observed ($\beta = −0.15, t = −0.05, p = 0.96$). When split based on cigarette smoking (median-split of $z$-score), the following effects were obtained: for low tobacco users, cannabis lifetime users did not differ significantly from cannabis non-users in SSRT ($\beta = 7.62, t = 1.90, p = 0.06$). A similar finding was observed among high tobacco users ($\beta = 4.80, t = 1.74, p = 0.08$).

Education years also significantly predicted SSRT ($\beta = −9.33, p < 0.01, 95\% CI [−12.88, −5.80]$), indicating that the higher the education level, the shorter the SSRT. Age significantly predicted SSRT ($\beta = 13.46, p < 0.01, 95\% CI [9.29, 17.63]$), with an increase in SSRT along with an increase in age. The number of trials also significantly predicted SSRT ($\beta = −17.44, p < 0.01, 95\% CI [−30.60, −4.28]$), indicating a decrease in SSRT when there were more trials. In addition, stop-signal modality had an effect on SSRT ($\beta = −28.58, p = 0.01, 95\% CI [−50.61, −6.56]$), indicating that auditory stop signals induced shorter SSRT compared to visual stop signals. SSD also had a significant effect on SSRT ($\beta = −33.29, p = 0.04, 95\% CI [−64.61, −1.96]$), indicating that the staircase-tracking procedure resulted in shorter SSRT compared to the fixed SSD procedure.

For both SSRT and commission error rate, models including the interaction between alcohol use on the one hand and demographics and task parameters on the other resulted in largely comparable findings as more.
In the model including interactions with demographics and task-parameters, presented here\(^2\). Only in the GNG, an interaction between alcohol use and age appeared ($\beta = 0.01$, $p = 0.02$, 95% CI [0.001, 0.02]). For light drinkers, older people made less commission errors ($\beta = -0.02$, $t = -2.56$, $p = 0.01$), which was in line with the main effect of age. Whereas for heavy drinkers, this relationship was absent ($\beta = -0.01$, $t = -1.50$, $p = 0.14$). All other interactions with alcohol were found to be non-significant (Supplementary Materials S6).

Outcomes for go RT in GNG and SST can be found in Supplementary Materials S8. Briefly, older people had longer go RT in both GNG and SST. Higher educated people had shorter go RT in SST. Although the interaction between cocaine and tobacco had an effect on go RT in SST, post-hoc analysis revealed no significant simple effect.

4. Discussion

Previous individual studies, reviews, and meta-analyses investigating inhibitory control deficits in relation to long-term substance use and SUD have provided mixed results (Luijten et al., 2014; Smith et al., 2014; Wright et al., 2014). These inconsistent findings might at least partly be due to insufficient control of frequently occurring polysubstance use. In addition, studies differed in sample demographics and task-related variables and used extreme group designs. The current mega-analysis aggregated data of 3610 individuals, from 43 studies, in which polysubstance use, demographics, and task parameters were included in the prediction of inhibition performance by means of an imputed multilevel analysis. Most of the included studies were of medium to high quality, which validates the overall conclusions drawn. Surprisingly, our overall pattern of results indicated that most types of substance use did not show an association with response inhibition. While for most substances no effects were found, lifetime cannabis use was found to be associated with impaired inhibition, as indexed by an increased SSRT in the SST. Tobacco use was also associated with impaired inhibition as indexed by the same variable. In addition, an interaction between lifetime cannabis and tobacco use was found on SSRT, which indicated a strong positive relationship between daily tobacco use and SSRT in participants who did not use cannabis (indicating poorer inhibition), and the absence of such a relationship in users smoking cannabis. In addition, demographic factors such as age and years of education and task characteristics such as no-go percentage, affected inhibition performance in the expected direction, strengthening the credibility of the other results.

4.1. Response inhibition and substance use

The main significant finding of our mega-analysis was that lifetime cannabis use was associated with prolonged response inhibition in the SST. One possible explanation is that this could (partly) involve subacute effects of cannabis use (i.e., lasting 7h to 4 weeks after last cannabis use, Gruber and Yurgelun-Todd, 2005; Pope and Yurgelun-Todd, 1996; Schulte et al., 2014). Acute cannabis use (i.e., 0–6 hours after last cannabis use) has been consistently reported to impair response inhibition in the SST (Metrik et al., 2012; Ramaekers et al., 2006). In contrast, findings of its long-term effect (i.e., 3 weeks or longer after last cannabis use) were mixed (Craen et al., 2011), with some confirming an impairing effect (Moreno et al., 2012), while others did not (Tapert et al., 2007). To have a closer look at the effect of cannabis, we compared cannabis daily users with less frequent users. A linear mixed regression model was built with the fixed effect of ‘cannabis daily users (yes/no)’ and a random intercept. It indicated that cannabis daily users did not differ from less frequent users on their stopping latency (i.e., SSRT, $\beta = -6.42$, $p = 0.90$, 95% CI [$-114.27, 127.10$]), which does not support the hypothesis of subacute cannabis effects. Despite conflicting behavioral findings of the relationship between cannabis use and response inhibition, abnormalities in neural activation have often and more consistently been reported in relation to acute as well as chronic cannabis use compared with non-users (systematic review: Wrege et al., 2014). Age of onset may have a moderating effect on the neural effects of cannabis (Hester et al., 2009), but we did not have sufficient data to test this hypothesis.

In line with previous findings, tobacco use tended to impair inhibition. Participants with a higher level of tobacco dependence demonstrated a lower level of response inhibition capacities (Billieux et al., 2010), and smokers performed worse than non-smokers in a smoking-related GNG (Luijten et al., 2011). However, it should be noted that the main effect of tobacco use was qualified by a significant interaction with cannabis use, indicating a negative effect of tobacco use only in non-cannabis users. Another study reported that co-administration of cannabis and tobacco attenuated the impairment in delayed recall memory caused by cannabis alone (Hindocha et al., 2017), and other reports have indicated weaker impairment on some measures after polysubstance use (e.g., alcohol and cannabis, Schweinsburg et al., 2011). One possible interpretation of these findings is that cannabis has a protective effect when used together with other substances such as alcohol and tobacco (cf., Viveros et al., 2006). Due to the high co-occurrence of cannabis and tobacco use (Badiani et al., 2015; Leatherdale et al., 2006), and the fact that concurrent tobacco use contributes to cannabis dependence symptoms (Ream et al., 2008), further studies of the combined and single effects on response inhibition are warranted to elucidate these findings.

What could explain the low evidence for a relationship between (most) long-term substance use and inhibition? On closer inspection, only 30% of studies included reported evidence for negative associations between substance use (or gambling) and response inhibition (Tables 1 and 2). In contrast, other studies reported evidence for positive associations between substance use and inhibition performance in GNG and SST (significant: Glass et al., 2009; nonsignificant: Galván et al., 2011; Papachristou et al., 2012b; Vonmoos et al., 2013). In light of this, it is less surprising that the integrated results indicated overall largely null findings (most of the confidence intervals ranged around zero). Similarly, only one out of the five studies included in a recent review (Carbia et al., 2018) reported impaired response inhibition—as measured by SST and GNG tasks—in binge drinkers compared with controls (Czapla et al., 2015).

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\(^2\) In the model including interactions with demographics and task-parameters, tobacco and cannabis use were both positively associated with SSRT. However, their interaction was not significant, but the three-way interaction with sex was. Post-hoc tests indicated that, only for male non-cannabis users, tobacco use was positively associated with SSRT (see in Supplementary Materials S6)
One explanation is that chronic recreational substance use without a diagnosis of SUD is not associated with response inhibition impairment. In other words, a threshold effect rather than a linear effect might exist between substance use and response inhibition performance. Alternatively, there might be a linear relationship, albeit shallow and we only see the effects when comparing very extreme groups (e.g., healthy controls vs. SUD in clinical samples). As a result of our exclusion criteria, Fig. S2a and S3a indicate that only a minority of the participants reached the level of SUD (either reported in individual paper or categorized based on questionnaire score), and most others were still within the normal range of use. It is conceivable that inhibition is only impaired in SUD (Björk et al., 2004; Fernández-Serrano et al., 2011; Noël et al., 2007; Petit et al., 2014). Alternatively, inhibition problems may play a role in the transition from heavy use to SUD. In the SST sample, there were more people diagnosed with tobacco dependence (about 10%, Fig. S3a), which might explain why a positive (although not significant) association of SSRT and tobacco use was found.

A second possibility is that substance use is actually associated with impaired inhibition, but we were unable to detect this. Possible reasons include: sample characteristics (as was discussed in the last paragraph), the type of tasks included, outcome measures (i.e., effects may only be visible in biological markers but not in behavior), and statistical power. Regarding tasks included, there is the possibility that (heavy) use of psychoactive substances does not lead to a general inhibition problem, but only to a specific problem in the domain of substance use (hence an interaction between an appetitive process and suboptimal control, Jones et al., 2018). A related explanation can be that self-control failures like maladaptive substance use may reflect a reduced mobilization of inhibitory control in substance-related contexts rather than generally impaired inhibitory control competencies (Kronke et al., 2018, 2015; Wolff et al., 2016). However, in a secondary analysis, we did not find that substance-related GNG moderated the relationship between alcohol and commission error rate (see details in 4.2). Furthermore, the SST and GNG measure stimulus-driven (exogenous) inhibition, which may not closely match real-world ‘loss of control’ behavior related to substance use (e.g., an initial intention to have one drink escalating into a binge-drinking session, failed suppression of craving, etc.). These examples reflect a different type of inhibition, namely endogenous or intentional rather than exogenous inhibition. Intentional inhibition paradigms such as the Marble task (Schel et al., 2014) could be considered in future research. Regarding outcome measures, it is possible that biological but not behavioral markers might be more sensitive to inhibition impairments among substance users (Garrison and Potenza, 2014). Relatedly, some of the included MRI studies reported specific group-related abnormalities in brain activation but not in behavioral outcomes (e.g., Claus, Ewing et al., 2013; de Ruiter et al., 2012; Galván et al., 2011; Karoly, Weiland et al., 2014; Luijten et al., 2013a; Roberts and Garavan, 2010). In addition, a recent study indicated that resting state fMRI connectivity might serve as a promising biomarker of alcohol use disorder severity (Fede et al., 2019; see further, Steele et al., 2019 for additional recent approaches to identifying biomarkers for addiction). Alternatively, Kwako et al. (2018) suggested a dimensional approach to biomarkers in terms of executive functions (inhibitory control, working memory, etc.), which includes measuring neuropsychological tests and epigenetic changes in relevant genes (e.g., COMT). With respect to statistical power, polysubstance use was coarsely defined, such that substances other than alcohol and tobacco had to be coded in a binary lifetime use variable. It is still possible that (heavy) use of a specific combinations of substances at the same time (e.g., cocaine and alcohol, Schulte et al., 2014) does have a negative impact, which did not emerge from our analysis here using binary variables. In addition, the total author response rate was low, which we discuss as a limitation. Currently, it remains an open question whether substance use is not associated with a motor inhibition impairment or if we were incapable of detecting such an impairment.

4.2. Demographics and task parameters

Our results indicate that age is a significant predictor of performance. In the GNG-task, the age-related increase in accuracy is most likely due to the strategic slowing of responses (confirmed by longer go RTs). In the SST, SSRT increased with age. Education was positively correlated with inhibition capability in both tasks. There was not a significant effect of sex on inhibition, nor any interactions between sex and substance use. In the GNG, higher working memory load, lower no-go percentages, and a higher number of experimental trials resulted in more commission errors. These effects are in line with the primary literature on these tasks and are further discussed in Supplementary Materials SI. Somewhat surprisingly, we did not obtain an effect of substance-related GNG on performance measures compared to classical task versions. This is in line with a recent meta-analysis, where the main effect of appetitive cues was not observed after correction for publication bias, and where drinking status (light vs. heavy drinkers) also did not moderate this effect (Jones et al., 2018). In a small exploratory analysis, we examined the alcohol × substance-related task interaction effect, which was not a significant predictor of commission error rates in GNG (Supplementary Materials S6). Still, since our conclusion is based on only 5 out of 23 included studies, future research should address this question. In the SST, visual (vs. auditory) stop signals, fewer number of trials and fixed SSDs (vs. staircase-tracking procedure) induced prolonged SSRT (elaboration in Supplementary Materials S1).

4.3. Implications

Our results showed no relationship between the use of most substances and impaired response inhibition, except for a relationship between cannabis use and impaired inhibition, and in non-cannabis users an association between cigarette use and impaired inhibition. What are the theoretical implications? First, these findings could be of relevance for the current debate on the question whether addiction should be considered a chronic brain disease or not (Heather et al., 2017; Lesnner, 1997; Lewis, 2015; Volkow et al., 2015). The current findings do not support the idea that long-term recreational substance use leads to irreparable problems in inhibition, although it cannot be excluded that inhibition problems are present in a (subgroup of) people diagnosed with SUD. Second, in many dual process models of addiction, suboptimal inhibition of stimulus-driven appetitive processes (cue-reactivity) plays an important role in the escalation of use (e.g., Baler and Volkow, 2006; Wiers et al., 2007). An alternative perspective does not emphasize the competition between stimulus-driven and goal-directed processes, but rather between different goal-directed processes (Moors et al., 2017). Individuals learn to mobilize and allocate resources strategically according to goal saliency and importance (Köpetz et al., 2013). In this way, the inhibition capability of substance users is expected to fluctuate moment-to-moment (i.e., state-like) based on the external and internal context. Note again that the current findings do not exclude the possibility that in severe addiction(s), chronic inhibition problems of stimulus-driven processes do play a role. It merely underscores the goal-directed nature of (heavy) substance use. Third, impaired response inhibition as an immediate consequence of substance consumption may be more important than general inhibitory impairments in the long term. Compared with long-term (non-dependent) substance use, acute use is more consistently related to impaired inhibitory control that enhances further consumption (Gan et al., 2014).

4.4. Limitations and suggestions for future study

There are several limitations of the current study worth considering. First, the response rate was rather low. Although more than 100 studies met our inclusion and exclusion criteria, authors of only 65 studies provided raw data. The reasons for this include inaccessibility of the data, data could not be shared due to regulations, and a lack of success
We thank Dr. Rebecca L. Ashare, Dr. Elliott T. Berkman, Dr. Craig R. Colden, Dr. Piike Erika, Dr. Mark T. Fillmore, Dr. Rual Gonzalez, Dr. Bernice Porjesz, Dr. Olgia Rass, and Dr. Craig R. Rush who contributed raw data without co-authorship.

Yang Liu thanks the China scholarship council (CSC) (No. 201506990019) for fellowship support.

HMH is supported by a VICI grant awarded by the Netherlands Organization of Scientific Research (NWO) [grant number 453-12-005]

MY was supported by a National Health and Medical Research Council of Australia Fellowship (#APP1117188) and the David Winston Turner Endowment Fund.

Appendix A. Supplementary data

Supplementary material related to this article can be found, in the online version, at doi:https://doi.org/10.1016/j.neubiorev.2019.07.006.

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