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Acute and chronic effects of cannabinoids on human cognition-a systematic review

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**Abstract**

Cannabis use has been associated with impaired cognition during acute intoxication as well as in the unintoxicated state in long-term users. However, the evidence has been mixed and contested, and no systematic reviews of the literature on neuropsychological task-based measures of cognition have been conducted in an attempt to synthesize the findings. We systematically review the empirical research published in the past decade (from January 2004 to February 2015) on acute and chronic effects of cannabis and cannabinoids and on persistence or recovery after abstinence. We summarize the findings into the major categories of the cognitive domains investigated, considering sample characteristics and associations with various cannabis use parameters. Verbal learning and memory and attention are most consistently impaired by acute and chronic exposure to cannabis. Psychomotor function is most affected during acute intoxication, with some evidence for persistence in chronic users and after cessation of use. Impaired verbal memory, attention, and some executive functions may persist after prolonged abstinence, but persistence or recovery across all cognitive domains remains underresearched. Associations between poorer performance and a range of cannabis use parameters, including a younger age of onset, are frequently reported. Little further evidence has emerged for the development of tolerance to the acutely impairing effects of cannabis. Evidence for potential protection from harmful effects by cannabidiol continues to increase but is not definitive. In light of increasing trends toward legalization of cannabis, the knowledge gained from this body of research needs to be incorporated into strategies to minimize harm.

**Keywords**

review, systematic, cognition, chronic, human, acute, cannabinoids, effects

**Disciplines**

Education | Social and Behavioral Sciences

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Review

Biological Psychiatry

Acute and Chronic Effects of Cannabinoids on Human Cognition—A Systematic Review

# Samantha J. Broyd, Hendrika H. van Hell, Camilla Beale, Murat Yücel, and Nadia Solowij

## ABSTRACT

Cannabis use has been associated with impaired cognition during acute intoxication as well as in the unintoxicated state in long-term users. However, the evidence has been mixed and contested, and no systematic reviews of the literature on neuropsychological task-based measures of cognition have been conducted in an attempt to synthesize the ﬁndings. We systematically review the empirical research published in the past decade (from January 2004 to February 2015) on acute and chronic effects of cannabis and cannabinoids and on persistence or recovery after abstinence. We summarize the ﬁndings into the major categories of the cognitive domains investigated, considering sample characteristics and associations with various cannabis use parameters. Verbal learning and memory and attention are most consistently impaired by acute and chronic exposure to cannabis. Psychomotor function is most affected during acute intoxication, with some evidence for persistence in chronic users and after cessation of use. Impaired verbal memory, attention, and some executive functions may persist after prolonged abstinence, but persistence or recovery across all cognitive domains remains underresearched. Associations between poorer performance and a range of cannabis use parameters, including a younger age of onset, are frequently reported. Little further evidence has emerged for the development of tolerance to the acutely impairing effects of cannabis. Evidence for potential protection from harmful effects by cannabidiol continues to increase but is not deﬁnitive. In light of increasing trends toward legalization of cannabis, the knowledge gained from this body of research needs to be incorporated into strategies to minimize harm.

*Keywords:* Attention, Brain, Cannabis, Cognition, Executive function, Memory <http://dx.doi.org/10.1016/j.biopsych.2015.12.002>

### Shifts in public opinion and policies toward legalization of cannabis are poised to result in an increase in the prevalence of cannabis use beyond the 178 million users estimated to exist today ([1–3](#_bookmark6)). Although most individuals who try cannabis do not go on to use it regularly ([1](#_bookmark6),[2](#_bookmark7)), individuals who do so risk adverse effects to physical and mental health ([4](#_bookmark8)). Negative sequelae that have been attributed to regular and prolonged cannabis use include alterations to brain morphology ([5–7](#_bookmark10)) and function ([8–11](#_bookmark13)); psychosis risk ([12](#_bookmark19),[13](#_bookmark21)); poor psychosocial outcomes ([4](#_bookmark8),[14–17](#_bookmark24)); and impaired cognition, especially deﬁ- cits in attention, learning and memory ([18–21](#_bookmark29)), and executive functions ([9](#_bookmark15),[22](#_bookmark35)). Morphological and connectivity changes in brain structures with high amounts of cannabinoid receptors (e.g., hippocampus, prefrontal cortex, cerebellum) ([23](#_bookmark5)) may mediate observed cognitive deﬁcits in cannabis users ([5–7](#_bookmark10), [9–11](#_bookmark11),[24](#_bookmark12)), although direct structure/function relationships are not readily demonstrated.

A substantial number of studies have been published in recent years, prompted by renewed interest in understanding the effects of cannabis on the brain partly as a result of mounting evidence for links between cannabis use and psychosis ([25–28](#_bookmark9)) and recognition of similarities between cognitive impairment in cannabis users and deﬁcits observed

in patients with schizophrenia ([19](#_bookmark31)). Interest has emerged in examining the effects of different compounds within cannabis plant matter, speciﬁcally (2)-*trans*-Δ9-tetrahydrocannabinol (THC), the primary psychoactive constituent considered to be psychotogenic, and cannabidiol (CBD), the second most abundant cannabinoid, shown to have antipsychotic proper- ties ([29](#_bookmark14)) and to attenuate the psychotogenic effects of THC, with opposite effects on brain function ([30](#_bookmark16)). Recent critical reviews have focused on neuroimaging outcomes from acute cannabinoid challenge ([9](#_bookmark15),[31](#_bookmark17)) and on brain morphology in chronic users ([5](#_bookmark10),[7](#_bookmark12),[32](#_bookmark18)). However, to date, the literature on neuropsychological task-based measures of cognition has not been examined in the form of a systematic review. We systematically review the empirical research published in the past decade. We identify core themes that have emerged from the recent literature or continue to plague this ﬁeld and study limitations and future directions for this research area.

METHOD

This systematic review was conducted in accordance with Preferred Reporting Items for Systematic Reviews and Meta- Analyses guidelines ([Figure 1](#_bookmark0)) ([33](#_bookmark20)). The search strategy and

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Figure 1. Preferred Reporting Items for Systematic Reviews and Meta- Analyses ﬂow diagram for systematic search and identiﬁcation of studies meeting inclusion criteria for systema- tic review. Selection criteria were as follows: 1) neuropsychological or cog- nitive experimental tasks administered to regular or former cannabis users or after acute administration of cannabis or synthetic or phytocannabinoid compounds; 2) cannabis (or cannabi- noids) as the primary drug of interest; and 3) human participants. Exclusion criteria were as follows: 1) studies in which cannabis was not the primary drug of concern; 2) questionnaire (trait) measures of cognition; 3) major psychopathology or neurologic condi- tions within assessed sample; 4) animal research; 5) neuroimaging, electrophysiologic, or autonomic measures as primary outcome vari- ables; 6) treatment (e.g., cognitive- behavioral therapy) as the primary focus; 7) real-world multiplex tasks requiring simultaneous use and inte- gration of multiple aspects of cogni- tion, such as driving; and 8) case studies.

Records identified through PubMed (n = 3021)

Records identified through Scopus (n = 3220)

Records after titles assessed in PubMed (n = 316)

Records after titles assessed in Scopus (n = 225)

Records after duplicates removed (n = 356)

Full text records screened (n = 356)

Records excluded (n = 260)

Studies included in qualitative synthesis (n = 105)

data extraction are detailed in [Supplement 1](#_bookmark4), and selection criteria, resulting in 105 studies included for review, are provided in the legend of [Figure 1](#_bookmark0). From each study, we extracted participant demographics (age, sex, IQ), cannabis use metrics (e.g., age of onset and duration, frequency, and quantity of use), period of abstinence before testing, extent of other substance use (including alcohol and tobacco), dosing details for acute administration studies, cognitive domains investigated, experimental tasks employed, and key cognitive ﬁndings. The primary results of interest for qualitative syn- thesis of ﬁndings were group differences in performance and associations with cannabis use metrics.

RESULTS

We provide a summary appraisal of ﬁndings organized by primary cognitive domain in order of evidential strength from most to least consistently impaired. A more detailed account of ﬁndings from all studies is provided in [Supplement 1](#_bookmark4), and detailed data extracted from each article are tabulated in [Table](#_bookmark4) [S1](#_bookmark4) in [Supplement 2](#_bookmark4). Within each cognitive domain, we address ﬁrst acute administration studies, then studies of chronic exposure to cannabis, followed brieﬂy by absti- nence studies. Further consideration is given to the important issue of persistence or recovery of function separately in the

Discussion. [Table 1](#_bookmark2) provides a qualitative summary of ﬁndings across all cognitive domains examined.

Memory

Memory function has been the most consistently impaired cognitive domain affected by cannabis, and studies from the past 10 years continue to extend the evidence base. The most extensive evidence for impairment is within verbal learning and memory.

Verbal Learning and Memory. Most often measured using word list learning tasks, with several immediate and delayed recall trials and a recognition trial, verbal learning and memory tasks have been identiﬁed as particularly sensitive to the acute ([20](#_bookmark33),[34](#_bookmark22),[35](#_bookmark23)) and chronic ([18](#_bookmark29)) effects of cannabis. Further clear evidence has emerged for impairing effects of acute intravenous (IV) THC ([36–40](#_bookmark25)), vaporized cannabis ([41](#_bookmark34),[42](#_bookmark36)) and oral nabilone ([43](#_bookmark37)) on immediate and delayed recall and sometimes recognition accuracy. Predosing with CBD or greater CBD content in cannabis may protect against some THC-induced verbal learning and memory deﬁcits ([40](#_bookmark32),[44](#_bookmark38)). Impaired verbal learning and memory continues to be con- sistently observed in chronic cannabis users, including ado- lescents ([45–47](#_bookmark40)) and young adults ([48–52](#_bookmark46)) [with some

#### Table 1. Strength and Consistency of Evidence for Impairment Associated With Acute and Chronic Cannabis Use and for Recovery of Function With Abstinence From Research Published in the Past Decade[*a*](#_bookmark1)

Acute and Chronic Effects of Cannabinoids on Cognition

|  |  |  |
| --- | --- | --- |
|  | Number of Studies |  |
|  |  | Persistence |  |  |  |  |
| Cognitive Domain | Acute[*b*](#_bookmark1) | With Chronic[*b*](#_bookmark1)Abstinence[*b*](#_bookmark1) | Pertinent Cannabis Use Parameters | Acute | Chronic | Abstinence |
| Memory |  |  |  |  |  |  |
| Verbal learning and memory | 111 | 111 12 | Frequency; lifetime use; duration; age of onset; sex | 11 ([36–44](#_bookmark25),[77](#_bookmark26),[87](#_bookmark27)) | 20 ([44–58](#_bookmark38),[60–62](#_bookmark40),[66](#_bookmark42),[142](#_bookmark44)) | 9 ([55](#_bookmark56),[57](#_bookmark60),[60](#_bookmark64),[63–67](#_bookmark37),[95](#_bookmark39)) |
| Working memory | 12 | 12 12 | Frequency; lifetime use; recency; sex | 20 ([36–40](#_bookmark25),[42](#_bookmark26),[43](#_bookmark27),[68–78](#_bookmark30),[88](#_bookmark32),[143](#_bookmark36)) | 16 ([46](#_bookmark42),[48](#_bookmark46),[49](#_bookmark48),[51–53](#_bookmark50),[55](#_bookmark52),[57](#_bookmark54),[79–84](#_bookmark56),[115](#_bookmark60),[119](#_bookmark66)) | 7 ([55](#_bookmark56),[57](#_bookmark60),[64](#_bookmark39),[65](#_bookmark41),[67](#_bookmark45),[79](#_bookmark66),[85](#_bookmark75)) |
| Other memory function | 1 | 12 2 | Age of onset; frequency; recency | 2 ([42](#_bookmark36),[144](#_bookmark109)) | 8 ([45](#_bookmark40),[49](#_bookmark48),[78–80](#_bookmark65),[119](#_bookmark66),[144](#_bookmark68),[145](#_bookmark98)) | 4 ([63](#_bookmark37),[65](#_bookmark41),[79](#_bookmark66),[95](#_bookmark95)) |
| Attention |  |  |  |  |  |  |
| Attention | 111 | 111 12 | Dose; age of onset; length | 16 ([36](#_bookmark25),[37](#_bookmark26),[39](#_bookmark30),[42](#_bookmark36),[43](#_bookmark37),[68](#_bookmark47),[70](#_bookmark51),[71](#_bookmark53),[76](#_bookmark61),[77](#_bookmark62), | 14 ([45](#_bookmark40),[46](#_bookmark42),[54](#_bookmark55),[55](#_bookmark56),[57](#_bookmark60),[61](#_bookmark67),[79–81](#_bookmark66),[84](#_bookmark68), | 10 ([55](#_bookmark56),[57](#_bookmark60),[63](#_bookmark37),[64](#_bookmark39),[67](#_bookmark45),[79](#_bookmark66),[91–](#_bookmark86) |

of abstinence; withdrawal effects

[86–90](#_bookmark77),[143](#_bookmark79))

[91–94](#_bookmark70))

[93](#_bookmark86),[95](#_bookmark89))

Attentional bias 1 111 NA Craving; dependence;

frequency; CBD

1 ([102](#_bookmark71)) 7 ([96–102](#_bookmark96)) None

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|  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- |
| Psychomotor Function | 111 | 1 | 1 |  | 18 ([37](#_bookmark26),[42](#_bookmark36),[43](#_bookmark37),[68](#_bookmark47),[70](#_bookmark51),[73](#_bookmark57),[74](#_bookmark58),[76](#_bookmark61),[77](#_bookmark62),[89](#_bookmark82),[90](#_bookmark84),[103–107](#_bookmark73),[143](#_bookmark76),[146](#_bookmark78)) | 10 ([46](#_bookmark42),[48](#_bookmark46),[51](#_bookmark50),[54](#_bookmark55),[57](#_bookmark60),[66](#_bookmark43),[78](#_bookmark65),[80](#_bookmark68),[91](#_bookmark86),[108](#_bookmark83)) | 6 ([57](#_bookmark60),[63–65](#_bookmark37),[67](#_bookmark39),[91](#_bookmark41)) |
| Executive Function |  |  |  |  |  |  |  |
| Planning, reasoning, interference control, and problem solving | 12 | 12 | 12 | Neurodevelopmental stage; age of onset; frequency | 12 ([37–41](#_bookmark26),[77](#_bookmark27),[86](#_bookmark30),[89](#_bookmark32),[103](#_bookmark34),[104](#_bookmark62),[106](#_bookmark77),[109](#_bookmark82)) | 23 ([46](#_bookmark42),[48](#_bookmark46),[52–54](#_bookmark52),[57](#_bookmark54),[60](#_bookmark55),[61](#_bookmark60),[66](#_bookmark64),[78](#_bookmark67),[81–84](#_bookmark43),[93](#_bookmark65),[98](#_bookmark70),[110–115](#_bookmark71),[147](#_bookmark72)) | 9 ([57](#_bookmark60),[60](#_bookmark64),[63–](#_bookmark37)[65](#_bookmark37),[67](#_bookmark39),[85](#_bookmark41),[93](#_bookmark45),[95](#_bookmark75)) |
| Inhibition | 11 | 12 | NA | Frequency; task complexity | 5 ([42](#_bookmark36),[89](#_bookmark82),[103](#_bookmark73),[104](#_bookmark76),[109)](#_bookmark85) | 9 ([45](#_bookmark40),[50](#_bookmark49),[54](#_bookmark55),[56](#_bookmark59),[82](#_bookmark71),[110](#_bookmark87),[116–118](#_bookmark95)) | None |
| Verbal ﬂuency | 2 | 12 | 12 |  | 3 ([36](#_bookmark25),[38](#_bookmark27),[44](#_bookmark38)) | 6 ([44](#_bookmark38),[48](#_bookmark46),[51](#_bookmark50),[53](#_bookmark54),[54](#_bookmark55),[61](#_bookmark67)) | 4 ([65](#_bookmark41),[67](#_bookmark45),[93](#_bookmark91),[119](#_bookmark98)) |
| Time estimation | 12 | 2 | 2 |  | 6 ([73](#_bookmark57),[74](#_bookmark58),[77](#_bookmark62),[86](#_bookmark77),[148](#_bookmark113),[149](#_bookmark115)) | 1 ([55](#_bookmark56)) | 1 ([55](#_bookmark56)) |
| Decision Making | 1– | 1– | 2 | Age of onset; lifetime exposure; frequency; cannabis use disorder | 7 ([103](#_bookmark73),[105](#_bookmark78),[106](#_bookmark80),[109](#_bookmark85),[120–122](#_bookmark99)) | 17 ([45](#_bookmark40),[48](#_bookmark46),[50](#_bookmark49),[56](#_bookmark59),[78](#_bookmark65),[82](#_bookmark71),[84](#_bookmark74),[94](#_bookmark93),[113](#_bookmark92),[116](#_bookmark95),[117](#_bookmark96),[123–126](#_bookmark101),[128](#_bookmark103),[150](#_bookmark105)) | 3 ([85](#_bookmark75),[127](#_bookmark108),[128](#_bookmark110)) |

CBD, cannabidiol; NA, not available (not investigated).

*a*The prevalence of studies focused on acute vs. chronic effects is unequal, as is the focus on individual cognitive domains; strength metrics are based on qualitative interpretation of the literature, subjectively weighed on greater or lesser evidence for impairment across the published studies, considering the number of studies conducted and their quality (e.g., design, sample size), reached by consensus between the authors of this review.

*b*111, strong and largely consistent evidence for impairment; 11, moderate evidence for impairment; 1, weak evidence for impairment, being based on only a small number of studies;

12, mixed evidence; 2, little or no evidence for impairment.

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exceptions ([53](#_bookmark54), [54](#_bookmark55))] and even in only occasional users ([55](#_bookmark56)). Signiﬁcant associations between poorer performance in reg- ular users and frequency, quantity, duration, and age of onset of cannabis use have been reported ([47](#_bookmark44),[49](#_bookmark48),[56–58](#_bookmark59)). Consistent with previous ﬁndings ([59](#_bookmark63)), long-term users appear to be more affected than short-term users ([60](#_bookmark64),[61](#_bookmark67)). One study reported greater impairment associated with higher THC compared with CBD exposure ([62](#_bookmark69)). Improvement ([63](#_bookmark37)) or recovery ([55](#_bookmark56),[57](#_bookmark60), [64–66](#_bookmark39)) with abstinence has been observed in some studies or indices, but not others ([64](#_bookmark39),[65](#_bookmark41),[67](#_bookmark45)) (see [Supplement 1](#_bookmark4) and further on).

Working Memory. Whether working memory is impaired by cannabis is less clear, possibly because of the wide range of different working memory tasks employed. Acute administra- tion of THC, dronabinol, or nabilone affected working memory inconsistently across Sternberg, delayed matching to sample, spatial or numeric working memory, *n*-back, digit recall, and digit span tasks ([36–40](#_bookmark25),[42](#_bookmark26),[43](#_bookmark27),[68–78](#_bookmark30)). Similarly, chronic can- nabis use was shown to impair working memory in young adults on immediate recall ([79](#_bookmark66)), verbal reasoning ([80](#_bookmark68)), and verbal *n*-back ([81](#_bookmark70)) working memory tasks, but not on spatial working memory ([48](#_bookmark46),[82](#_bookmark71)) or digit span ([52](#_bookmark52),[53](#_bookmark54)), whereas spatial working memory was impaired in adolescent users ([46](#_bookmark42)), suggestive of differential effects in the developing brain. In older users, two studies reported no impairment on letter- number sequence and spatial span tests of the Wechsler Memory Scale ([83](#_bookmark72)) or on an *n*-back task ([84](#_bookmark74)), whereas recent or heavy users were impaired on a range of Wechsler Adult Intelligence Scale working memory tasks, with greater fre- quency and quantity of use correlated with poorer perform- ance ([57](#_bookmark60)). Impaired working memory persisted for a few weeks in some studies ([65](#_bookmark41)) but appears to mostly resolve with longer periods of abstinence ([35](#_bookmark23),[55](#_bookmark56),[57](#_bookmark60),[67](#_bookmark45),[85](#_bookmark75)).

For other memory function, see [Supplement 1](#_bookmark4).

Attention

Impaired attention has been considered a hallmark of the intoxicating effects of cannabis. Further evidence has accumu- lated in support of acute exposure to cannabinoids impairing focused, divided, or sustained attention, often in a dose- dependent manner ([36](#_bookmark25),[37](#_bookmark26),[42](#_bookmark36),[43](#_bookmark37),[68](#_bookmark47),[70](#_bookmark51),[71](#_bookmark53),[86–88](#_bookmark77)). In cases where lesser impairments were observed, this may be due to the development of tolerance among daily users ([76](#_bookmark61),[89](#_bookmark82),[90](#_bookmark84)). Previous evidence for deﬁcits in attention after chronic cannabis exposure was mixed, but more recent studies provide some clarity. Numerous studies report impairment in adolescent and adult cannabis users with a wide range of exposure as well as former users abstinent for several weeks on measures of sustained and divided attention, processing speed, rapid visual information processing, visual search, tracking, trail making, and paced serial addition ([46](#_bookmark42),[54](#_bookmark55),[55](#_bookmark56),[57](#_bookmark60),[61](#_bookmark67),[91–94](#_bookmark86)). Users absti- nent for 23 days remained impaired relative to control subjects despite improvements in sustained and divided attention with increasing abstinence ([91](#_bookmark86)) with poor attentional performance associated with younger age of onset in this study ([91](#_bookmark86)) and another study of adolescents abstinent for 30 days ([95](#_bookmark95)). However, no difference between abstinent former users and

control subjects on broader measures of attention was also reported ([57](#_bookmark60),[63](#_bookmark37),[64](#_bookmark39),[79](#_bookmark66),[95](#_bookmark95)). Therefore, cannabis-related atten- tional impairment may reﬂect residual effects that dissipate gradually as cannabinoids are cleared from the body ([Supplement 1](#_bookmark4)).

Attentional Bias. Greater attentional bias to cannabis- related stimuli has been noted in chronic cannabis users ([96–102](#_bookmark96)) and during acute intoxication in individuals with lower CBD/THC ratios in hair ([102](#_bookmark71)). There are no studies of abstinent users after cessation of use, which may be a particularly important target for future research to strengthen the efﬁcacy of treatment programs aimed at maintaining abstinence.

Psychomotor Function

Finger tapping, critical tracking, choice reaction time tasks, and digit-symbol substitution tasks have been used to meas- ure psychomotor function. In infrequent users, smoked or vaporized cannabis impaired critical tracking ([42](#_bookmark36),[89](#_bookmark82),[103](#_bookmark73),[104](#_bookmark76)), affected reaction time and motor control in a dose-dependent manner ([70](#_bookmark51)), and disrupted motor function in a task with a motivational component ([105](#_bookmark78)). In heavy users, high-dose smoked cannabis resulted in more collisions in a virtual maze task ([106](#_bookmark80)) but did not affect critical tracking ([89](#_bookmark82),[90](#_bookmark84)). Oral administration of THC [or IV ([37](#_bookmark26))], nabilone, or dronabinol impaired psychomotor function in seven of eight studies ([37](#_bookmark26),[43](#_bookmark37),[68](#_bookmark47),[74](#_bookmark58),[76](#_bookmark61),[77](#_bookmark62),[107](#_bookmark81)), with only one study ﬁnding no sig- niﬁcant impairment ([73](#_bookmark57)). Findings regarding the chronic effects of cannabis on psychomotor function are mixed, being reported as impaired ([51](#_bookmark50),[54](#_bookmark55),[80](#_bookmark68),[108](#_bookmark83)), improved ([48](#_bookmark46)), and unaf- fected ([46](#_bookmark42),[78](#_bookmark65)). Psychomotor function was impaired in users abstinent for 23–35 days ([57](#_bookmark60),[65](#_bookmark41),[67](#_bookmark45),[91](#_bookmark86)), with a trend also after 12 months of abstinence (ﬁnger tapping) ([64](#_bookmark39)). The weight of evidence suggests that psychomotor function is affected by acute intoxication and that this likely persists for some time after chronic cannabis exposure.

Executive Function

Planning, Reasoning, Interference Control, and Prob- lem Solving. On similar tasks of planning, reasoning, inter- ference control, and problem solving, THC administration was found to impair performance in some studies ([38](#_bookmark27),[41](#_bookmark34),[103](#_bookmark73), [104](#_bookmark76),[106](#_bookmark80),[109](#_bookmark85)), but not others ([39](#_bookmark30),[86](#_bookmark77),[89](#_bookmark82)), equally across samples of occasional, moderate, and heavy users. Impaired perform- ance may depend on extent of prior exposure, route of administration, dose delivered, and blood cannabinoid concen- trations at baseline and after dosing ([89](#_bookmark82)). With regard to the chronic effects of cannabis, numerous studies reported null ﬁndings in case-control comparisons ([46](#_bookmark42),[54](#_bookmark55),[66](#_bookmark43),[81](#_bookmark70),[95](#_bookmark95),[98](#_bookmark97), [110](#_bookmark87),[111](#_bookmark88)), whereas several others found cannabis-related deﬁ- cits in heavy users ([53](#_bookmark54),[78](#_bookmark65),[83](#_bookmark72),[112](#_bookmark90)), including adolescents ([113](#_bookmark92)), early-onset but not late-onset adult users ([114](#_bookmark94)), and older users ([57](#_bookmark60)), and associated with persistent use in a longitudinal study ([60](#_bookmark64)). Impaired executive function was evident despite intact performance on other cognitive tasks ([48](#_bookmark46),[82](#_bookmark71)) and associated with self-reported problems related to cannabis use ([115](#_bookmark93)). Studies in which executive dysfunction was detected tended

to have older samples than the studies in which no impairments were observed, which included predominantly adolescent and young adult users. It may be that executive dysfunction becomes more evident beyond the period of maturation of the frontal lobes, perhaps reﬂecting perturbed neurodevelop- ment. Consistent with this interpretation, three abstinence studies reported no group differences relative to control sub- jects in younger samples ([65](#_bookmark41),[85](#_bookmark75),[95](#_bookmark95)), whereas persistent exec- utive dysfunction was observed in users abstinent for $28 days aged 35–50 years ([57](#_bookmark60)) and in users abstinent for 12 months aged 38–51 years ([64](#_bookmark39)), where impaired performance on block design tests was the only signiﬁcant difference detected between abstinent users and their nonuser twins. Older users have also been exposed to far more cannabis use over the lifetime.

"'

Inhibition. Measures of inhibition are derived from para- digms such as go/no-go or stop-signal reaction time tasks. Acute administration of THC has consistently been reported to increase stop-signal reaction time in both occasional and heavier cannabis users ([42](#_bookmark36),[89](#_bookmark82),[104](#_bookmark76),[109](#_bookmark85)). Findings in chronic users are more mixed ([45](#_bookmark40),[50](#_bookmark49),[54](#_bookmark55),[56](#_bookmark59),[82](#_bookmark71),[116–118](#_bookmark95)) ([Supplement 1](#_bookmark4)).

Verbal Fluency. Three studies reported no effect on verbal ﬂuency during acute intoxication with THC or cannabis ([36](#_bookmark25),[38](#_bookmark27),[44](#_bookmark38)), but there has been little consistency regarding chronic or abstinence effects ([44](#_bookmark38),[48](#_bookmark46),[51](#_bookmark50),[53](#_bookmark54),[54](#_bookmark55),[61](#_bookmark67),[65](#_bookmark41),[67](#_bookmark45),[85](#_bookmark75), [93](#_bookmark91),[110](#_bookmark87),[119](#_bookmark98)) ([Supplement 1](#_bookmark4)). The ﬁndings suggest that if verbal ﬂuency is impaired in cannabis users, it is more likely to be in older individuals with longer durations of exposure, whereas in younger users, impairment in verbal ﬂuency may depend on intellectual functioning and on the task employed.

Time Estimation. The subjective effect of cannabis dis- torting time is well known, but objective evidence from new studies of time estimation is limited ([Supplement 1](#_bookmark4)).

In summary, executive function subdomains are differentially affected by acute administration and chronic exposure to cannabis. There are clear acutely impairing effects on inhibition, whereas planning, problem solving, reasoning, and interference control are inconsistently impaired, and the moderators of impaired performance require further investigation. The latter subdomains may be more affected in older chronic users or with greater exposure to cannabis. Literature assessing recov- ery of executive functions with abstinence is very sparse, and this is an important area for optimizing treatment programs for cannabis dependence. For example, one study found that poor neurocognitive performance was associated with relapse to cannabis use at 1-year follow-up in adolescents ([95](#_bookmark95)).

Decision Making, Reward Processing, and Delay Discounting

Measures of risky and impulsive decision making include performance on the Iowa Gambling Task, delay discounting tasks, and behavioral risk-taking tasks. Some studies found that acute administration of THC adversely affected decision making by altering sensitivity to reward and punishment

and increasing risk taking in infrequent ([120](#_bookmark99)) and regular users ([106](#_bookmark80),[121](#_bookmark100),[122](#_bookmark102)), but not all studies found impaired decision making ([103](#_bookmark73),[109](#_bookmark85)). Evidence for effects of chronic cannabis use on decision-making is mixed. Although several studies reported poorer decision-making performance across a range of tasks ([48](#_bookmark46),[78](#_bookmark65),[82](#_bookmark71),[117](#_bookmark96),[123](#_bookmark101),[124](#_bookmark103)), especially decreased sensitivity to loss and greater sensitivity to gains ([123](#_bookmark101)), clear group differences were not found in other studies ([45](#_bookmark40),[50](#_bookmark49),[94](#_bookmark93),[125–127](#_bookmark105)). Cognitive ﬂexibility in decision making is also affected by chronic cannabis exposure ([45](#_bookmark40),[113](#_bookmark92),[116](#_bookmark95)). Behav- ioral risk taking was greater in adolescents with a mean period of abstinence of 53 days ([85](#_bookmark75)) and associated with extent of prior exposure in users with an abstinence period of 25 days ([127](#_bookmark108)), but was unaffected in other studies of current users ([50](#_bookmark49),[56](#_bookmark59),[125](#_bookmark105)). No delay discounting performance differences were found between current and abstinent users and control subjects ([128](#_bookmark110)). Thus, risky decision making and sensitivity to reward are increased during acute intoxication. Despite a large number of studies of chronic users in the past decade, the extent to which these effects persist in chronic or abstinent users remains unclear.

DISCUSSION

In general, the literature on the cognitive effects of cannabis exposure continues to be plagued with complexity in terms of heterogeneity of both the extent of cannabis exposure in the samples assessed and the means of assessing cognitive function ([18–21](#_bookmark29)). The nature of the samples recruited and the way in which prior exposure to cannabis is quantiﬁed affect interpretation of ﬁndings for acute and chronic effects. In studies of chronic users, associations with frequency of use are more likely to reﬂect the residual effects of acute or subchronic intoxication, associations with dose of exposure may reﬂect residual or cumulative effects, and associations with the duration of use are likely to indicate more persistent cannabis-related alterations ([20](#_bookmark33)). These parameters may also interact with the neurodevelopmental period during which regular cannabis use is started, and increasing consideration has been given to age of onset effects and assessing adolescent users. Potential sex differences are insufﬁciently addressed in studies of predominantly male cannabis users ( 75% of studies reviewed), although many argue that this overrepresentation matches the sex distribution of cannabis users in the general population ([129](#_bookmark111)). Where sex effects were tested ( 12% of studies), very few were found (see [Supplements 1](#_bookmark4) and [2](#_bookmark4)). Evidence for sex differences in cannabinoid metabolism, action, and brain morphology in cannabis users ([130–132](#_bookmark112)) suggests that future research should further investigate these differences in relation to cognition, both following acute administration and with chronic expo- sure. Several key issues are critical to consider further.

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Dose, Route of Administration, and Tolerance Effects

Acute administration studies often, but inconsistently, report dose-dependent effects ([Supplements 1](#_bookmark4) and [2](#_bookmark4) contain further details of doses administered). Discrepancies may be due to differing routes of administration, with IV, smoked, and vaporized cannabinoids exerting more immediate but

shorter-lasting effects than effects following oral or sublingual administration. The duration of impaired cognition after acute administration has not been further investigated; most studies conducted a single assessment immediately after or within 2–3 hours of dosing. The development of tolerance is often assumed to occur in frequent or dependent users and is inferred to explain modest effects in some studies, with supportive reference to preclinical literature and some prior, but limited, human studies. Most studies of acute adminis- tration examined effects in a single sample of control subjects or occasional or infrequent users (23 studies), or a single sample of regular, frequent, or daily users (11 studies), whereas only 4 studies directly compared effects in cannabis users versus control subjects (2 studies; [37](#_bookmark26),[87](#_bookmark79)) or in frequent versus infrequent users (2 studies; [89](#_bookmark82),[149](#_bookmark115)) ([Supplement 2](#_bookmark4)). These latter studies reported blunted impairing effects in regular or frequent users of acute smoked cannabis on critical tracking and divided attention or of IV THC on spatial working memory, verbal memory (inconsistently), and time estimation. Tolerance may occur in other cognitive domains, but the research over the past decade has not signiﬁcantly advanced knowledge of tolerance effects beyond the limited previous evidence in humans ([87](#_bookmark79)). Although cognitive impairment after acute canna- binoid exposure may be blunted in regular users, substantial further evidence has accumulated that it nevertheless still exists across multiple domains (psychomotor, attention, and, incon- sistently, memory) and with potential real-world effects on complex tasks involving these domains, such as driving ([133](#_bookmark114)).

Speciﬁcity to Cannabis and Potential Confounds

The literature continues to be riddled with a range of potential confounds that may affect speciﬁc attribution of impairment to cannabinoids, including largely premorbid functionality and other substance use. The strongest design to address these is a large-sample prospective study that controls for cognitive ability assessed before initiation of cannabis use, years of education, and other substance use. One such study examined neuropsychological change from childhood to age 38 and found that persistent cannabis use or dependence was associated with an IQ decline of 6 points; this reﬂected broad decline in functioning not speciﬁc to any particular cognitive domain; cessation of use did not restore IQ in adolescent-onset users; and speciﬁcity to persistent cannabis use was determined by ruling out effects caused by recent cannabis use and persistent tobacco, alcohol, or other drug use ([60](#_bookmark64)). Two further studies controlling for premorbid function showed that adolescent cannabis users showed impaired verbal learning and memory

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([47](#_bookmark44)) and immediate and delayed memory and processing speed ([79](#_bookmark66)), whereas former users did not differ from control subjects ([79](#_bookmark66)). Numerous studies have not controlled for premorbid functioning or ascertained current intellectual functioning;

.50% of studies did not report IQ or years of education. Where groups were matched on IQ or IQ was controlled for in analyses, persistent impairments to cognition were identiﬁed for immedi- ate and delayed memory ([79](#_bookmark66)), verbal learning and memory ([39](#_bookmark30),[61](#_bookmark67)), verbal ﬂuency, attention, and executive function ([61](#_bookmark67)) [although not in other studies ([84](#_bookmark74),[128](#_bookmark110))].

Alcohol and tobacco use as well as other illicit substance use are common in cannabis users, and a disconcerting

number of studies continue to fail to report or account for this ([Supplements 1](#_bookmark4) and [2](#_bookmark4)). Means of dealing with these potential confounds include applying a heterogeneous range of exclusion criteria, often poorly speciﬁed; matching compar- ison groups on levels of use; and covarying for other substance use but with wide-ranging metrics (e.g., quantity/ frequency measures, lifetime use, dependence/abuse scores). Mostly, the cognitively impairing effects of chronic cannabis use have held after controlling for other substance use, but not in all studies ([Supplement 1](#_bookmark4)). Greater standardization of not only cannabis use metrics across studies but also tobacco, alcohol, and other drug use would enable better character- ization of the speciﬁcity of cannabis effects. Further research to examine additive, interactive, or synergistic effects of cannabinoids with other substances is required.

Persistence or Recovery of Function With Abstinence

Recovery of function after prolonged abstinence remains con- tentious. The persistence of impairments or recovery of cogni- tive function after a period of abstinence .24 hours, most often for several weeks (21–35 days; only a few went beyond this period) was examined in 18 new studies. In 11 studies, cross- sectional comparisons were performed of former users (absti- nent 7 days to .1 year) and current users or nonuser control subjects; 9 of these studies assessed adolescents or young adults. Seven were prospective studies, assessing cannabis users at baseline and at follow-up (with some intervening measures) over periods ranging from 21 days to 4–8 years of abstinence. Although this is a stronger design, there remains a conspicuous paucity of these studies, as two were confounded by concomitant pharmacologic treatments ([63](#_bookmark37),[93](#_bookmark91)).

Cross-sectional studies report persistent impairing effects on some aspects of attention, verbal and working memory, and psychomotor speed in adolescents abstinent for 28 days

([65](#_bookmark41)) and 35 days ([67](#_bookmark45)), but not on other aspects of these and other cognitive domains. Poor performance was associated with lifetime cannabis exposure ([65](#_bookmark41)) or an earlier age of use onset in adolescents with 30 days of abstinence ([95](#_bookmark95)) and predicted relapse to cannabis use during a 1-year follow-up ([95](#_bookmark95)). Even after 53 days of abstinence, adolescents showed impaired working memory and risk taking ([85](#_bookmark75)). Young adults abstinent for up to 4 weeks showed poorer verbal ﬂuency relative to control subjects ([119](#_bookmark98)) and nonsigniﬁcantly poorer performance on a gambling task, associated with prior quan- tity of weekly cannabis use ([127](#_bookmark108)). Older adults abstinent for

1. days were impaired on executive function and information processing but not attention and working memory ([57](#_bookmark60)). Lyons *et al.* ([64](#_bookmark39)) argued that their study of monozygotic twins, discordant for cannabis use with .12 months abstinence, provides evidence against long-term effects of cannabis on cognition, despite ﬁnding impaired performance on block design tests and trends toward poorer long delay and cued verbal recall and poorer ﬁnger tapping performance in the former users. Such studies offer a rare opportunity to control for underlying vulnerability to cannabis use, which may never- theless interact with effects of persistent use.

In the prospective studies, only two controlled for premor- bid function; one found that adolescent former users did not

Table 2. Key Findings for Cognitive Impairment in Cannabis Users

Acute Effects of Cannabis on Cognition Impaired verbal learning and memory

Impaired working memory and other memory functions Impaired attention, task and dose dependent

Impaired inhibition, less so for other executive functions Impaired psychomotor function

Chronic Effects of Cannabis on Cognition Impaired verbal learning and memory Impaired attention and attentional bias Possible impaired psychomotor function

Mixed evidence for executive function and decision making

Most associated with cannabis use parameters, particularly frequency of use and age of onset

Recovery of Function With Abstinence

Likely persistent effects on attention and psychomotor function Possible persistent effects on verbal learning and memory Evidence insufﬁcient and mixed

### differ from control subjects on any cognitive measures ([79](#_bookmark66)), whereas another large and well-controlled prospective study showed that cannabis use before the age of 18 resulted in greater decline in IQ by age 38 persisting even after cessation or reduction of use in the past year ([60](#_bookmark64)). Poor verbal learning improved in adolescents by 2 weeks of abstinence, but visual search remained impaired at 3 weeks relative to baseline ([55](#_bookmark56)). An adult sample showed improvements in critical tracking and divided attention with increasing abstinence periods over 8–23 days, but the sample subjects remained impaired relative to control subjects; withdrawal and age of onset effects were observed ([91](#_bookmark86)). In a large prospective study following a sample across three waves of data collection spaced 4 years apart, former heavy users abstinent for $12 months improved rela- tive to ongoing heavy users on immediate verbal memory and did not differ from nonusers on any cognitive measure ([66](#_bookmark43)).

Given the plasticity of the human brain, recovery of function might be anticipated, and consistent with previous ﬁndings ([134](#_bookmark116),[135](#_bookmark117)), the evidence from some (but not all) of these recent studies suggests that verbal learning and memory impairment may recover with prolonged abstinence. Yet evidence for persistent cognitive dysfunction continues to emerge. Neither the precise cannabis use metrics required for the persistence of cognitive deﬁcits nor the neural mecha- nisms underlying the persistence of deﬁcits have been elucidated. The latter likely include neurotransmitter system dysfunction ([136–138](#_bookmark102)) and regional brain structural and con- nectivity alterations ([32](#_bookmark18),[139](#_bookmark104)), which may take signiﬁcant time to be restored to pre–cannabis use integrity and functionality [however, see Yücel *et al.* ([140](#_bookmark106))]. Previous studies using sensitive measures of brain function detected duration of exposure-related impaired attentional processes in 2-year abstinent users ([141](#_bookmark107)). Further well-controlled prospective studies monitoring restoration of brain function and structure from current use through cessation of use and over prolonged abstinence are urgently needed.

CONCLUSIONS

**Further signiﬁcant evidence has emerged supporting the ﬁnding that acute and chronic exposure to cannabinoids impairs cognition, especially in the domains of verbal learning, memory, and attention (**[**Table 2**](#_bookmark3)**). Mixed evidence across the range of other cognitive domains is likely due to ongoing heterogeneity in the cognitive tests employed, prior cannabis use histories, and the assessment of cannabis use metrics as well as the neurodevelopmental stage at both onset and cessation of cannabis use. Nevertheless, it is clear from the literature reviewed that cognitive impairment on a range of domains can persist beyond the period of acute intoxication and potentially affect daily functioning in cannabis users and hence the range of adverse educational and other psychoso- cial outcomes identiﬁed as associated with frequent use, in particular for adolescent users (**[**17**](#_bookmark28)**). Multiple potential moder- ators of cannabinoid effects remain underinvestigated, includ**- ing premorbid and other individual differences, genetic factors, sex, psychopathology, and polydrug use, and few studies consider the range of compounds in cannabis that interac- tively moderate the effects of THC, despite increasing interest in CBD. Further prospective and mechanistic studies are required to understand the impact of cannabinoids during brain maturation in adolescence through young adulthood and interacting with normal and abnormal aging processes in later years. The time course and moderators of potential recovery of cognitive function necessitate more precise delineation. In light of increasing trends toward legalization or medicalization of cannabis, it is imperative to research further the parameters of cannabis use that result in impairment and the potential for protection from cognitive harm by CBD (evidence is growing but is not deﬁnitive) such that harm minimization strategies may be implemented, and to understand the therapeutic parameters of any of the cannabinoids to enable the beneﬁts of medications without concomitant brain and cognitive harm.

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