



Original Investigation | Infectious Diseases

Evaluation of Computerized Cognitive Training and Cognitive and Daily Function in Patients Living With HIV A Meta-analysis

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Abstract

IMPORTANCE In the era of antiretroviral therapy (ART), the incidence of HIV-associated neurocognitive disorder (HAND) has not yet been controlled. With the exception of ART, there is no beneficial pharmacologic treatment. However, some studies have reported that computerized cognitive training (CCT) programs may improve cognitive function among people living with HIV.

OBJECTIVE To examine the association between CCT programs and 8 domains measuring cognitive function (7 domains) and daily function (1 domain) among people living with HIV.

DATA SOURCES Records from the Cochrane Library, PsycINFO, PubMed, and Web of Science were searched from database inception to December 15, 2020. Supplementary searches to identify missing studies were conducted in Google Scholar using updated search terms from database inception to November 18, 2021.

STUDY SELECTION Studies that compared changes before and after a CCT intervention among people living with HIV were included. Search terms were a combination of words associated with HIV (eg, people living with HIV, HIV, and/or AIDS) and cognitive training (eg, cognitive intervention, nonpharmacology intervention, computer game, video game, computerized training, cognitive exercise, cognitive stimulation, and/or cognitive enhancement). Studies were included if they (1) used CCT as the primary intervention or combined CCT with other types of interventions; (2) used placebo, passive control conditions, traditional cognitive training, or single training tasks as control conditions; (3) reported changes between baseline and posttraining; (4) included participants 18 years or older; and (5) were randomized clinical trials (RCTs). Studies were excluded if they (1) were not associated with HIV, (2) were research protocols or feedback reports, (3) were case reports, or (4) did not report findings for domains of interest.

DATA EXTRACTION AND SYNTHESIS Two reviewers independently extracted data. This study followed the Preferred Reporting Items for Systematic Reviews and Meta-analyses (PRISMA) reporting guideline. Random-effects models were used to quantitatively synthesize the existing data.

MAIN OUTCOMES AND MEASURES Primary outcomes were the meta-synthesized changes in each domain after CCT.

RESULTS Among 1245 records identified, 1043 were screened after removal of duplicates. Of those, 1019 records were excluded based on titles and abstracts, and 24 full-text articles were assessed for eligibility. After exclusions, 12 eligible RCTs were selected for inclusion in the meta-analysis. These RCTs involved 596 total participants, with 320 individuals in the CCT group (mean age, 47.5-59.7

(continued)

Key Points

Question Are computerized cognitive training (CCT) programs associated with improvements in cognitive and daily function among people living with HIV?

Findings In this meta-analysis of 12 randomized clinical trials involving 596 participants living with HIV, CCT was associated with improved function in 6 of the 8 domains assessed, including abstraction and executive function, attention and working memory, memory, speed of information processing, motor skills, and daily function.

Meaning This meta-analysis found that CCT programs were associated with improvements in cognitive and daily function among people living with HIV.

Supplemental content

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Abstract (continued)

years; 0%-94% female; 8.3-14.2 years of education) and 276 individuals in the control group (mean age, 44.2-60.0 years; 19%-90% female; 9.0-14.9 years of education). The average HIV inhibition ratio (the proportion of participants who achieved virological suppression) ranged from 30% to 100%, and the CD4⁺ T-cell count ranged from 471 to 833 cells/ μ L. The time since training ranged from 3 to 24 weeks. After receipt of CCT, function significantly improved in 6 of the 8 domains: abstraction and executive function (standardized mean difference [SMD], 0.58; 95% CI, 0.26-0.91; P < .001), attention and working memory (SMD, 0.62; 95% CI, 0.33-0.91; P < .001), memory (SMD, 0.59; 95% CI, 0.20-0.97; P = .003), motor skills (SMD, 0.50; 95% CI, 0.24-0.77; P < .001), speed of information processing (SMD, 0.65; 95% CI, 0.37-0.94; P < .001), and daily function (SMD, 0.44; 95% CI, 0.02-0.86; P = .04). Sensory and perceptual skills (SMD, 0.06; 95% CI, -0.36 to 0.48; P = .78) and verbal and language skills (SMD, 0.46; 95% CI, -0.07 to 0.99; P = .09) did not significantly improve after CCT.

CONCLUSIONS AND RELEVANCE This meta-analysis of RCTs found that CCT programs were associated with improvements in cognitive and daily function among people living with HIV. Future studies are needed to design optimal specific training programs and use implementation science to enable the transformation of CCT from a scientific research tool to a real-world clinical intervention.

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Introduction

People living with HIV experience cognitive deterioration, also known as HIV-associated neurocognitive disorder (HAND), despite the widespread use of antiretroviral therapy (ART). 1-4 Based on commonly used criteria, ⁵ HAND is diagnosed by assessing 7 cognitive domains, including speed of information processing, sensory and perceptual (sensory/perceptual) skills, memory (learning and recall), attention and working memory (attention/working memory), motor skills, verbal and language (verbal/language) skills, and abstraction and executive function (abstraction/ executive function). ^{6,7} A diagnosis of HAND can have substantial consequences for an individual's daily function. 8,9 In addition, HAND has been associated with reduced life satisfaction 10-12 and social isolation, ¹³ producing public health and economic burdens worldwide. ^{14,15} With the exception of ART, there is no specific pharmacologic treatment for HAND. 16,17 However, studies have reported potential benefits from nonpharmacologic interventions that may ameliorate cognitive decline and reduce the odds of developing HAND among people living with HIV. 18,19 These interventions have attracted the attention of many researchers. Chan et al 18 focused on different types of cognitive and neurologic rehabilitation strategies among people living with HIV. Their results highlighted the importance of ART and suggested that nonpharmacologic strategies might improve cognitive function, either as stand-alone interventions or as part of a multidisciplinary approach.

Because of its safety, relevance, low cost, scalability, and convenience, computerized cognitive training (CCT) has been one of the most commonly used nonpharmacologic interventions. ²⁰⁻²² Computerized cognitive training programs aim to incorporate guided drills and practice for single or multiple cognitive domains through specific standardized procedures, differentiating CCT from other cognitive remediation approaches. ²³ The theoretical premise behind CCT is that it can stimulate neuroplasticity. ²⁴ From a neuropsychological perspective, ²⁵ CCT has the potential to shape brain structure and reorganize function among cognitively healthy older adults ²⁶⁻²⁸ and individuals with Alzheimer disease, ^{29,30} Parkinson disease, ^{31,32} attention deficits, ³³ and acquired brain injury. ³⁴ Furthermore, physiological parameters, including brain metabolism ³⁵ and inflammatory, ³⁶ hormonal, ³⁷ and sleep-related factors, ³⁸ may also be transformed through brain plasticity. ²⁵ In addition, many studies involving people living with HIV have reported that CCT may reduce the risk of cognitive deterioration in several domains, ³⁹ such as working memory, ⁴⁰⁻⁴³ speed of information

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processing, ^{40,44} executive function and attention, ⁴⁵⁻⁴⁷ and memory (learning and recall). ⁴² However, the benefits reported in original articles using the same CCT program (Posit Science software) have been inconsistent. Pope et al⁴⁸ found that this software could improve abstraction/executive function, whereas Fazeli et al⁴⁹ reported that the software could not only enhance abstraction/executive function but also improve attention, working memory, and speed of information processing among people living with HIV.

Because of the mixed results reported in original studies, Vance et al³⁹ investigated the findings in a systematic review. After identifying 13 items that fit their selection criteria, they found that most of the CCT programs were associated with improvements in cognitive function that translated into better daily function, improved mood, more substantial locus of control, and enhanced quality of life. Nevertheless, Vance et al³⁹ did not calculate the effect size for each subdomain or perform sensitivity or moderator analyses. In addition, to our knowledge, no previous meta-analysis has confirmed the association of CCT with daily function and with each cognitive domain categorized by the Frascati criteria. ⁵ The potential factors associated with CCT outcomes for each cognitive domain among people living with HIV have also not been confirmed.

Given these knowledge gaps, we conducted a meta-analysis to assess the associations of CCT programs with cognitive and daily function among people living with HIV. The present study aimed to (1) assess the extent of improvement in each domain after CCT among people living with HIV and (2) explore the consistency of the domain results among all of the potential factors. Furthermore, the study aimed to provide suggestions for future implementation of CCT interventions among people living with HIV.

Methods

This study was registered in the International Prospective Register of Systematic Reviews (PROSPERO; registration No.: CRD42020210805). The study followed the Preferred Reporting Items for Systematic Reviews and Meta-analyses (PRISMA) reporting guideline for meta-analyses.⁵⁰

Search Strategy

We performed a record search of electronic databases, including the Cochrane Library, PsycINFO, PubMed, and Web of Science, with no limitations on publication type, from database inception to December 15, 2020. We also conducted a supplementary search using additional search terms from a previous meta-analysis. ⁵¹ The search terms were a combination of words associated with HIV (eg, people living with HIV, HIV, and/or AIDS) and cognitive training (eg, cognitive training, cognitive intervention, cognitive rehabilitation, nonpharmacology intervention, mnemonic training, processing speed training, working memory training, N-back training, attention training, reasoning training, computer game, video game, computerized training, computerized intervention, cognitive exercise, brain exercise, cognitive stimulation, and/or cognitive enhancement). Additional searches to identify missing studies were also conducted from database inception to November 18, 2021, using Google Scholar and the reference lists of reviews and included studies. These supplementary searches did not identify any missing studies or studies published after the conclusion of the primary search (December 15, 2020). A full description of the initial and supplementary search strategies is available in eTable 1 and eTable 2 in the Supplement.

Selection Criteria

Studies were eligible for inclusion if they (1) used CCT as the primary intervention or combined CCT with other types of interventions; (2) used a placebo, passive control conditions, traditional cognitive training, or single training tasks as control conditions; (3) reported changes between baseline and posttraining; (4) included participants 18 years or older and (5) were randomized clinical trials (RCTs). Studies were excluded if they (1) were not associated with HIV, (2) were research protocols or feedback reports, (3) were case reports, or (4) did not report findings for domains of interest.

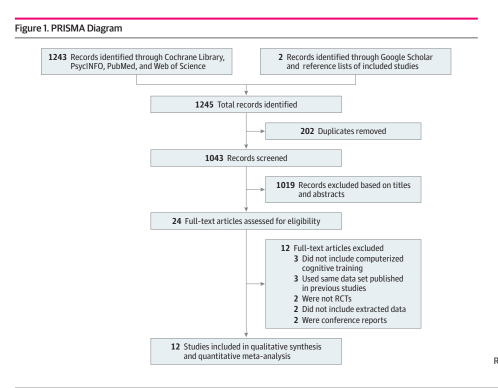
Duplicate studies were removed using EndNote X9 software (Clarivate), and 2 reviewers (J.W. and J.H.) separately selected search results based on titles and abstracts. Assessment of full-text articles to determine eligibility of the remaining studies was conducted by the same 2 reviewers. Disagreements about eligibility between reviewers were resolved through discussion with 2 investigators (B.S. and T.Z.). A detailed description of reasons for exclusion is available in eTable 3 in the Supplement, and the study selection process is shown in **Figure 1**.

Data Extraction

Relevant data were independently extracted and cross-checked by 2 researchers (J.W. and J.H.) using an Excel spreadsheet (Microsoft Corporation). The summary statistics collected for each outcome included number of participants, means, and SDs. Training outcomes included daily function (1 domain) and cognitive tasks (7 domains, which included speed of information processing, sensory/ perceptual skills, memory [learning and recall], attention/working memory, motor skills, verbal/ language skills, and abstraction/executive function). The 7 cognitive domains were categorized using Frascati criteria⁵ and selected based on criteria from a previous study⁵² and work by Lezak et al.⁵³ Details about the categorization process are shown in eTable 4 in the Supplement. Training dose was defined by the total number of training sessions, duration of each session, total training hours, training frequency, and time since training. Other information extracted from each RCT included the name of the first author, study location, year of publication, sample size, sex distribution, mean age and educational level of participants, current CD4* T-cell counts, and current HIV inhibition ratio (the proportion of participants who achieved virological suppression).

Statistical Analysis

Data analysis was performed using Comprehensive Meta-Analysis software, version 3 (Biostat, Inc), and the funnel plot was constructed using Review Manager software, version 5.4 (Cochrane Training). Because of the inherent heterogeneity across studies, we used random-effects models to estimate pooled effect sizes. Standardized mean differences (SMDs) were calculated as the mean change from pretraining to posttraining in the intervention group minus the mean change from pretraining to posttraining in the control group divided by the combined pretest SD (adjusted for



 $RCT\ indicates\ randomized\ clinical\ trial.$

bias). The inverse variance method was used to connect the SMDs of each study. We used Q tests (ie, χ^2 tests) and the I^2 statistic to assess the statistical and proportional significance of heterogeneity. We also used the Egger regression intercept test to estimate publication bias. The threshold for statistical significance was 2-tailed P < .05.

Study quality was evaluated using the Cochrane risk of bias tool, which divided risk of various biases into 3 grades: low, high, and unclear (**Figure 2**; eFigure in the Supplement). We used GRADEpro software, version 3.6 (McMaster University and Evidence Prime, Inc.), to assess the methodological quality of included evidence (eTable 5 in the Supplement).

Sensitivity analyses were performed for significant results. The moderators included participant age (<18 years vs \geq 18 years), proportion of women (<50% vs \geq 50%), years of education (\leq 12 years vs >12 years), current CD4⁺ T-cell counts (<500 cells/µL vs \geq 500 cells/µL), current HIV inhibition ratio (<100% vs 100%), total sessions (<22 vs \geq 22), session duration (<60 minutes vs \geq 60 minutes), session frequency (<3 sessions per week vs \geq 3 sessions per week), total training time (<10 hours vs \geq 10 hours), and time since training (<10 weeks vs \geq 10 weeks). Study participants were also divided into those with normal vs impaired cognitive status.

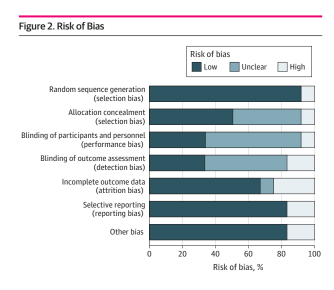
Results

Search Results

Because the methods and results of supplementary research covered the initial research results, we used the flowchart from our supplementary research to describe study selection in this article (Figure 1). We included a total of 1245 records. After removing all duplicates (n = 202), 1043 records were screened. Of those, 1019 records were excluded based on titles and abstracts, and 24 full-text articles were assessed for eligibility. After exclusions, 12 eligible RCTs^{40-42,44-49,54-56} were selected for inclusion in the meta-analysis.

Study Characteristics

Studies included data from the US, $^{40,41,44,46-49,54-56}$ Uganda, 42 and Italy, 45 comprising 596 total participants (320 participants in the CCT group and 276 participants in the control group). Participant ages ranged from a mean of 47.5 years 45 to 59.7 years 42 in the CCT group and 44.2 years 46 to 60.0 years 42 in the control group. The proportion of women ranged from $0\%^{54}$ to $94\%^{40}$ in the CCT group and $19\%^{45}$ to $90\%^{40}$ in the control group. Years of education ranged from 8.3 years 42 to 14.2 years 40 in the CCT group and 9.0 years 45 to 14.9 years 40 in the control group. Only 9 studies $^{40,42,44-49,54}$ reported etiologic data from CCT groups; in those studies, CD4 $^+$ T-cell counts



ranged from 471 cells/ μ L⁴⁴ to 833 cells/ μ L,⁴⁹ and the HIV inhibition ratio ranged from 30%⁴⁴ to 100%.⁴⁵ Additional characteristics of the included RCTs and participants are shown in the **Table**.

The total number of training sessions ranged from 6^{54} to 48, 47 and session duration ranged from 20 minutes 54 to 90 minutes. 56 Training frequency ranged from 2 sessions per week 41,42,46 to 4 sessions per week, 48 total number of training hours ranged from 1.7^{54} to 20.0, 55,56 and time since training ranged from 3 weeks 54 to 24 weeks. 45

Although all 12 studies^{40-42,44-49,54-56} used CCT programs, the intervention tools were different. Cody et al,⁴¹ Fazeli et al,⁴⁹ and Pope et al⁴⁸ used the same cognitive training program (BrainHQ.com; Posit Science) along with transcranial direct current stimulation (tDCS). Ownby et al⁵⁴ also used tDCS combined with a video game (GT Racing 2; Gameloft). Towe et al^{46,47} used an active cognitive training tool (Lumosity web-based cognitive games; Lumos Labs, Inc). Chang et al⁴⁰ used an adaptive working memory training platform (Cogmed; Neural Assembly), Ezeamama et al⁴² used computerized cognitive rehabilitation therapy software (Captain's Log MindPower Builder; Brain Train, Inc), and Livelli et al⁴⁵ combined paper and pencil and computer-based exercises. Vance et al^{44,55,56} used a computer program (InSight; Posit Science) in their 2012 study⁴⁴ and an individualized targeted cognitive training framework in their 2021 studies.^{55,56}

The control conditions were divided into 3 types: placebo (6 studies \$^{41,42,45,48,49,54}\$ used sham tDCS or standard of care), no contact (3 studies \$^{44,55,56}\$), and other (2 studies \$^{46,47}\$ used nonactive cognitive training and 1 study \$^{40}\$ used nonadaptive working memory training [Cogmed; Neural Assembly]). Six studies \$^{44,45,48,49,54,56}\$ reported outcomes for the abstraction/executive function domain, 9 studies \$^{40,41,44-47,49,54,56}\$ for the attention/working memory domain, 5 studies \$^{41,42,49,54,56}\$ for the memory domain, 5 studies \$^{42,45,49,54,56}\$ for the motor skills domain, 5 studies \$^{41,42,44,47,54}\$ for the speed of information processing domain, 3 studies \$^{44,45,55}\$ for the daily function domain, 1 study \$^{56}\$ for the sensory/perceptual skills domain, and 4 studies \$^{42,45,47,49}\$ for the verbal/language skills domain.

Meta-analysis of Cognitive and Daily Function Domains

Computerized cognitive training was significantly associated with improvements in 6 of the 8 domains: abstraction/executive function, attention/working memory, memory, motor skills, speed of information processing, and daily function. The detailed results of the meta-analysis for each domain are shown in **Figure 3** and **Figure 4**.

Abstraction/Executive Function

Among 6 studies 44,45,48,49,54,56 including 240 participants, the SMD for abstraction/executive function was 0.58 (95% CI, 0.26-0.91; P < .001). No significant heterogeneity (Q = 7.11; $I^2 = 30\%$; P = .21) or publication bias (intercept, -0.70; 95% CI, -5.92 to 4.52; P = .73) was found.

Attention/Working Memory

Among 9 studies $^{40,41,44\cdot47,49,54,56}$ involving 376 participants, the SMD for attention/working memory was 0.62 (95% CI, 0.33-0.91; P < .001). Moderate heterogeneity was detected (Q = 14.04; $I^2 = 43\%$; P = .08). No significant publication bias was found (intercept, 1.98; 95% CI, -1.69 to 5.65; P = .24).

Memory

Among 5 studies^{41,42,49,54,56} comprising 245 participants, the SMD for memory was 0.59 (95% CI, 0.20-0.97; P < .001). Moderate heterogeneity was detected (Q = 7.64; $I^2 = 48\%$; P = .11). No significant publication bias was found (intercept, 1.13; 95% CI, -5.29 to 7.55; P = .61).

Motor Skills

Among 5 studies 42,45,49,54,56 involving 229 participants, the SMD for motor skills was 0.50 (95% CI, 0.24-0.77; P < .001). No significant heterogeneity (Q = 3.82; $I^2 = 0\%$; P = .43) or publication bias (intercept, 1.29; 95% CI, -3.71 to 6.30; P = .47) was found.

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			Control condition	Nonadaptive Cogmed working memory training	Sham tDCS	Standard of care	Sham tDCS	Standard of care	Sham tDCS	Sham tDCS	:
				Attention and working memory	Speed of information processing; attention and working memory	Attention and working memory; memory	Speed of information processing	Attention and working memory; abstraction and executive function; weehal and language skills	Speed of information processing; abstraction and executive function	Speed of information processing	
		Follow-up	duration, wk	4	2	2	4	9	m	NA A	
		Session	frequency per wk	3.5	5	7	A A	₹ Z	m	4	
			Sessions, No.	25	10	7	10	36	9	10	
	CCT intervention		Training	Adaptive Cogmed working memory training	Posit Science (BrainHQ.com); Target Tracker; tic's	Captain's Log MindPower Builder	Posit Science (BrainHQ.com); tDCS	Combination of paper and pencil with computer-based exercises	GT Racing 2 (Gameloft); tDCS	Posit Science (BrainHQ.com); tDCS	
		Educational level, mean, y	Control arm	14.9	12.6	9.2	12.6	0.0	10.2	12.3	
		Educatio mean, y	CCT arm	14.2	12.5	8.3	12.5	10.0	12.0	12.5	
		sex, %	Control n arm	06	31	50	31	19	40	40	
		Female sex, %	CCT arm	94	35	59	35	31	0	33	
		an, y	Control arm	57.0	55.6	0.09	55.6	50.0	52.8	53.7	
		Age, mean, y		50.3	56.0	59.7	56.0	47.5	50.3	55.3	
			Control arm	20	16	40	16	16	r.	15	
		Sample size, No.	CCT arm	34	17	41	17	16	9	15	
acteristics		Sample	Total	54	33	81	33	35	Ξ	30	
Table. Study and Participant Characteristics	Sample		Description of participants	Adults with documented HIV seropositivity receiving ART for 26 mo	Older adults with or without HIV and without other physical or mental health disorders, hearing or vision problems, or history of brain trauma	Older adults (aged >50 y) with HIV and without other physical or mental health disorders	Older adults (aged 250 y) with HIV and without other physical or mental health disorders, hearing or vision problems, or history of brain trauma	Adults receiving care in division A infectious disease unit of Amedeo di Savoia Hospital	Adults with HIV, self- reported cognitive difficulties, and cognitive impairment in 2 neuropsychological domains, no history of seizures or bipolar disorder; receiving psychotropic medications	Older adults (aged 250 y) with HIV and without other physical or mental health disorders, hearing or vision problems, or history of brain trauma	
Table. Study			Source (location)	Chang et al, ⁴⁰ 2017 (US)	Cody et al, ⁴¹ 2020 (US)	Ezeamama et al, ⁴² 2020 (Uganda)	Fazeli et al, ⁴⁹ 2019 (US)	Livelli et al, ⁴⁵ 2015 (Italy)	Ownby et al, ⁵⁴ 2017 (US)	Pope et al, ⁴⁸ 2018 (US)	

Study	Table. Study and Participant Characteristics (continued)	cteristics	(continued,													
	Sample										CCT intervention					
		Sample	Sample size, No.		Age, mean, y	۱, y	Female sex, %	%,%	Educational level, mean, y	al level,			Session	Follow-up	Cognitive	
	Description of participants	Total	CCT arm	Control arm	CCT arm	Control arm	CCT arm	Control arm	CCT arm	Control arm	Training	Sessions, No.	frequency per wk	duration, wk	targeted by training	Control condition
	Adults (aged 18-65 y) with HIV receiving ART for >3 mo	21	11	10	51.3	44.2	27	20	14.1	12.6	Active cognitive training	12	2	10	Attention and working memory	Nonactive cognitive training
	Adults (aged 18-64 y) with HIV infection, self-reported history of cocaine use lasting 21 y, receiving ART for >3 mo, English speaking, educational level >8 y, no other substantial neuromedical comorbidities or mental impairment, and not pregnant	28	29	29	49.0	48.3	71	38	12.1	11.9	Active cognitive training (Luminosity web-based cognitive games)	48	NA	10	Attention and working memory	Nonactive cognitive training
Vance et al, ⁴⁴ 2012 (US)	Adults (aged ≥40 y) with HIV for ≥1 y without other significant neuromedical comorbidities or mental impairment	46	22	24	50.1	52.9	23	29	13.3	13.1	Posit ScienceInSight computer program	٧ ٧	۷ ع	Ю	Speed of information processing	No contact
	Adults (aged ≥40 y) with HIV for ≥1 y with HIV for ≥1 y living within 100 miles of research center and without other substantial neuromedical comorbidities or mental impairment	88	48	40	54.2	54.2	27	38	12.4	12.5	Individualized targeted cognitive training framework	٧ ٧	٧ ع	12	Speed of information processing	No contact
Vance et al, ⁵⁵ 2021 (US)	Adults (aged ≥40 y) with HIV for ≥1 y living within 100 miles of research center and without other substantial neuromedical comorbidities or mental impairment	109	64	45	53.4	53.8	28	33	12.1	12.4	Individualized targeted cognitive training framework	NA	NA	12	Speed of information processing	No contact

Abbreviations: ART, antiretroviral therapy; CCT, computerized cognitive training; NA, not applicable; tDCS, transcranial direct current stimulation.

Figure 3. Pooled Effects of Abstraction and Executive Function, Attention and Working Memory, Memory, and Motor Skills

Abstraction and executive function

	Particip	ants, No.		SMD			Fave	ors : Fa	ivors			Weight,
Source	CCT	Control	SE	(95% CI)			C	CT co	ntrol			%
Fazeli et al, ⁴⁹ 2019	17	16	0.35	0.24 (-0.45 to 0.92	2)				-			16.2
Livelli et al, ⁴⁵ 2015	16	16	0.38	1.14 (0.39 to 1.88))							14.3
Ownby et al, ⁵⁴ 2017	6	5	0.61	0.02 (-1.17 to 1.21	L)			-				6.6
Pope et al, ⁴⁸ 2018	15	15	0.38	0.93 (0.18 to 1.68))					_	→	14.1
Vance et al, ⁴⁴ 2012	22	24	0.30	0.18 (-0.40 to 0.76	5)			-		-		20.3
Vance et al, ⁵⁶ 2021	48	40	0.22	0.74 (0.31 to 1.18))							28.4
Total	124	116	NA	0.58 (0.26 to 0.91))					>		100.0
Heterogeneity: $\tau^2 = 0.05$;	$\chi^2 = 7.11$; df	$=5; P=.21; I^2=3$	30%									
Test for overall effect: z =	= 3.50; <i>P</i> < .00	1			-1.5	-1.0	-0.5	0	0.5	1.0	1.5	
							SN	ID (95%	CI)			

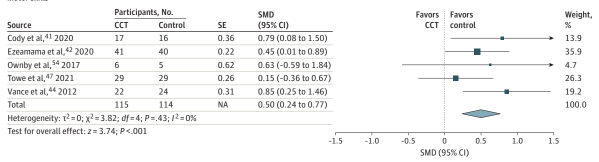
Attention and working memory

	Participa	ants, No.		SMD			Fav	ors : F	avors			Weight,
Source	CCT	Control	SE	(95% CI)			C	CT c	ontrol			%
Chang et al, 40 2017	34	20	0.30	1.01 (0.43 to 1.60)						_		12.9
Cody et al, ⁴¹ 2020	17	16	0.35	0.03 (-0.66 to 0.71)				-				10.9
Fazeli et al, ⁴⁹ 2019	17	16	0.35	0.38 (-0.31 to 1.07)			_	-	_			10.8
Livelli et al, ⁴⁵ 2015	16	16	0.40	1.54 (0.75 to 2.32)							→	9.1
Ownby et al, ⁵⁴ 2017	6	5	0.63	0.82 (-0.41 to 2.06)				- 1		-		4.6
Towe et al, ⁴⁶ 2017	29	29	0.48	1.22 (0.28 to 2.15)						_		7.1
Towe et al, ⁴⁷ 2021	11	10	0.27	0.50 (-0.02 to 1.02)				-	_			14.4
Vance et al, ⁴⁴ 2012	22	24	0.30	0.32 (-0.27 to 0.90)			-		_	_		13.0
Vance et al, ⁵⁶ 2021	48	40	0.22	0.41 (-0.01 to 0.83)				-	_	_		17.2
Total	200	176	NA	0.62 (0.33 to 0.91)					$\langle \rangle$	>		100.0
Heterogeneity: $\tau^2 = 0.08$;	$\chi^2 = 14.04$; d	f=8; P=.08; I ² =	43%									
Test for overall effect: z =	= 4.21; <i>P</i> < .00	1		1	_	-1.0	-0.5	-	0.5	1.0	1.5	
				-1	.5	-1.0		u 1D (95%		1.0	1.5	

Memory

	Particip	ants, No.		SMD			Favo	's : Fa	ors/			Weight,
Source	ССТ	Control	SE	(95% CI)			CC	Т со	ntrol			%
Ezeamama et al, ⁴² 2020	41	40	0.23	0.64 (0.20 to 1.08)	_							28.1
Fazeli et al, ⁴⁹ 2019	17	16	0.35	0.11 (-0.57 to 0.79)			-		_		18.6
Livelli et al, ⁴⁵ 2015	16	16	0.40	1.45 (0.67 to 2.22)					-			15.9
Ownby et al, ⁵⁴ 2017	6	5	0.62	0.61 (-0.60 to 1.82)						→	8.3
Vance et al, ⁵⁶ 2021	48	40	0.22	0.36 (-0.06 to 0.78)			-		_		29.1
Total	128	117	NA	0.59 (0.20 to 0.97)						>		100.0
Heterogeneity: $\tau^2 = 0.09$; χ	² =7.64; df	= 4; P=.11; I ² =4	48%									
Test for overall effect: $z = 2$	2.97; P=.00	3										
					-1.5	-1.0	-0.5	0	0.5	1.0	1.5	
							SMI	(95%	CI)			

Motor skills



CCT indicates computerized cognitive training; NA, not applicable; and SMD, standardized mean difference.

Speed of Information Processing

Among studies 41,42,44,47,54 including 246 participants, the SMD for speed of information processing was 0.65 (95% CI, 0.37-0.94; P < .001). No significant heterogeneity (Q = 4.60; $I^2 = 13\%$; P = .33) or publication bias (intercept, -0.22; 95% CI, -5.72 to 5.29; P = .91) was detected.

Daily Function

Among 3 studies 44,45,55 comprising 187 participants, the SMD for daily function was 0.44 (95% CI, 0.02-0.86; P = .04). No significant heterogeneity (Q = 3.60; $I^2 = 45\%$; P = .16) or publication bias (intercept, 3.87; 95% CI, -12.58 to 20.31; P = .21) was detected.

Figure 4. Pooled Effects of Sensory and Perceptual Skills, Information Processing Speed, Verbal and Language Skills, and Daily Function

Sensory and perceptual skills Participants, No. SMD Favors Favors Weight, SE (95% CI) Source Control control Vance et al,⁵⁶ 2021 0.06 (-0.36 to 0.48) 100 48 40 0.21 Total 48 40 NA 0.06 (-0.36 to 0.48) 100 Heterogeneity: NA Test for overall effect: z = 0.29; P = .78-1.5 -1.0 1.0 1.5 SMD (95% CI)

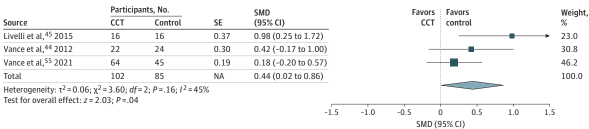
Speed of information processing

	Particip	ants, No.		SMD			Favo	ors : Fav	ors .			Weight,
Source	ССТ	Control	SE	(95% CI)			C	CT co	ntrol			%
Cody et al, ⁴¹ 2020	17	16	0.36	0.85 (0.14 to 1.57)				-		_	→	14.4
Ezeamama et al, ⁴² 2020	41	40	0.22	0.35 (-0.09 to 0.79)				+		_		32.8
Fazeli et al, ⁴⁹ 2019	17	16	0.35	0.49 (-0.20 to 1.18)				_	_			15.2
Ownby et al, ⁵⁴ 2017	6	5	0.61	0.49 (-0.71 to 1.70)					_			5.4
Vance et al, ⁵⁶ 2021	48	40	0.23	0.98 (0.54 to 1.43)								32.1
Total	129	117	NA	0.65 (0.37 to 0.94)						>		100.0
Heterogeneity: $\tau^2 = 0.01$; χ	2 = 4.60; df	= 4; P = .33; I ² = 1	13%									
Test for overall effect: z = 4	1.50; P<.00	1										
				-1	5	-1.0	-0.5	0	0.5	1.0	1.5	
							SN	D (95%	CI)			

Verbal and language skills

	Particip	ants, No.		SMD			Favor	: Fav	ors			Weight,
Source	ССТ	Control	SE	(95% CI)			CC.	con	trol			%
Ezeamama et al, ⁴² 2020	41	40	0.22	0.22 (-0.21 to 0.66	<u> </u>		-	-				29.5
Fazeli et al, ⁴⁹ 2019	17	16	0.35	0.22 (-0.47 to 0.90)			-		_		22.8
Livelli et al, ⁴⁵ 2015	16	16	0.40	1.53 (0.74 to 2.32)							→	20.3
Towe et al, ⁴⁷ 2021	29	29	0.26	0.14 (-0.38 to 0.65)			-				27.4
Total	103	101	NA	0.46 (-0.07 to 0.99)							100.0
Heterogeneity: $\tau^2 = 0.20$; χ^2	² =9.72; df	= 3; P=.02; I ² =6	59%									
Test for overall effect: $z = 1$.72; P=.09											
					-1.5	-1.0	-0.5	0	0.5	1.0	1.5	
							SMD	(95% C	1)			

Daily function



CCT indicates computerized cognitive training; NA, not applicable; and SMD, standardized mean difference.

Sensory/Perceptual Skills

One study⁵⁶ comprising 88 participants examined sensory/perceptual skills. The SMD was 0.06 (95% CI, -0.36 to 0.48; P = .78).

Verbal/Language Skills

Among 4 studies 42,45,47,49 involving 204 participants the SMD for verbal/language skills was 0.46 (95% CI, -0.07 to 0.99; P = .09). Moderate heterogeneity was detected (Q = 9.72; $I^2 = 69\%$; P = .02). However, no significant publication bias was found (intercept, 5.04; 95% CI, -9.55 to 19.63; P = .28).

Sensitivity Analysis

We conducted sensitivity analyses among all of the factors extracted from RCTs for each domain that had statistically significant results in the meta-analysis. Only 5 factors produced substantial changes in outcomes: age (memory domain: Q = 0.75; P = .02), session hours (attention/working memory domain: Q = 9.41; P < .001), time since training (abstraction/executive function domain: Q = 5.33; P = .02), CD4⁺ T-cell count (attention/working memory domain: Q = 4.78; P = .03), and HIV inhibition ratio (attention/working memory domain: Q = 4.52; P = .03; memory domain: Q = 4.76; P = .03). Detailed results from sensitivity analyses are available in eTable 6 in the Supplement.

Study Quality and Risk of Bias

Eleven studies^{40-42,44-49,54,55} had moderate-quality evidence, and 1 study⁵⁶ had very low-quality evidence (eTable 5 in the Supplement). Six domains (abstraction/executive function, attention/working memory, memory, motor skills, speed of information processing, and daily function) had moderate-quality evidence, for which the main reasons were moderate to high risk of bias, small samples, inclusion of few studies, and large 95% CIs. Two domains (sensory/perceptual and verbal/language skills) had very low-quality evidence.

Most studies had a low risk of bias for random sequence generation (11 studies^{40,41,} 44-49,54-56), selective reporting (10 studies^{41,42,44-46,48,49,54-56}), incomplete outcome data (8 studies^{40,42,45-47,49,55,56}), and other types of bias (10 studies^{40,42,44,45,47-49,54-56}) (Figure 2; eFigure in the Supplement). Six studies^{41,47,48,54-56} (50.0%) reported allocation concealment as a risk of bias. Eight studies^{40,42,44-47,55,56} (66.7%) did not blind those who implemented or assessed the interventions and/or those who reported relevant information.

Discussion

To our knowledge, this meta-analysis of 12 RCTs^{40-42,44-49,54-56} is the first to assess the association of CCT programs with cognitive and daily function among people living with HIV. Significant improvement was found in daily function and most cognitive domains, with the exception of sensory/perceptual and verbal/language skills, after CCT. Our results revealed no publication bias. The findings of the sensitivity analyses revealed that all pooled effects, with the exception of the memory and verbal/language skills domains, were statistically significant.

Moderate effect sizes for the speed of information processing and the attention/working memory domains suggested the potential for future improvement. Consistent with findings of previous meta-analyses of CCT interventions, ^{23,37,51,57} these 2 domains were also more likely to improve after CCT among individuals with mild cognitive impairment ^{23,37} and older adults with healthy cognitive status. ^{51,57} Because improvements after cognitive training typically reflect training content, ^{57,58} this result may be a result of sufficient training on these 2 subdomains within studies. Previous studies have also found that functional connectivity in the frontal-parietal brain network, mainly involved in the speed of information processing and the attention/working memory domains, increases after training. ⁵⁹⁻⁶¹ With regard to the sensory/perceptual and verbal/language skills domains, the null effect may be a result of the small number of studies and the presence of

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measurement bias because associations between these 2 domains and CCT have been found in other populations. ^{51,57} Future studies may consider dedicating more time to targeting these 2 domains. Multiple studies involving cognitively healthy adults⁵⁷ and individuals with Alzheimer disease⁶² have reported that training benefits extended beyond the cognitive domains assessed after training to include other cognitive domains, and these benefits have transferred to daily function, psychological health, and other higher-order competencies. ^{63,64} We also found an association between CCT and daily function among people living with HIV. Despite the limitations of these training tools, ⁶³ preliminary data suggest that CCT can, in principle, improve a broader range of essential functions, including cognitive and daily function. ⁵⁷

With regard to the sensitivity analysis, fewer results were statistically significant, and analysis of the same factor applied to different domains yielded substantially different results. We did not find any statistical difference in the characteristics of study participants. Our results partially replicated those reported in a meta-analysis of CCT programs among patients with Alzheimer disease. ⁶⁵ Karssemeijer et al ⁶⁵ found only a slight difference in the benefits of CCT among older adults with mild cognitive impairment vs dementia. Hill et al ²³ reported that CCT was beneficial for global cognition in the memory and learning domains among people with mild cognitive impairment. In comparison, the evidence for benefit among people with dementia was weak. ²³ Possible reasons might include (1) the cognitive enhancement mechanism after CCT may have differed between the 2 groups, and (2) differences in the training methods and measurement tools used may have produced different results.

Regarding CCT dose, our analyses suggested that better results occurred after longer CCT sessions (ie, >60 minutes), possibly because synaptic plasticity is more likely to occur after 30 to 60 minutes of stimulation. ^{57,66} A meta-analysis involving studies of patients with Alzheimer disease reached similar conclusions. ⁵⁷ Therefore, longer training sessions might be the recommended approach for people living with HIV. In contrast, many commercial products designed for at-home training use protocols consisting of shorter sessions, which may not be adequate. The findings of the current meta-analysis also provided a better understanding of viral load and current CD4⁺ T-cell counts among people living with HIV. Those with undetectable viral loads and current CD4⁺ T-cell counts of more than 500 cells/µL performed better than those with detectable viral loads and lower CD4⁺ T-cell counts, particularly in the attention/working memory domain. A previous study also found that people living with HIV who had current CD4⁺ T-cell counts lower than 500 cells/µL were more likely to have HAND. ⁴ The pathogenesis of HAND may explain this phenomenon. After entering the central nervous system, HIV can stimulate chronic neuroinflammation, which interacts with viral proteins and produces cognitive impairment. ⁶⁷ Therefore, people with lower viral loads and higher CD4⁺ T-cell counts, which indicate better immune status, may experience more benefit from CCT.

Only 50% of the included RCTs ^{41,47,48,54-56} reported allocation concealment as a risk of bias, producing inflated effect sizes and potential selection bias. ⁶⁸ Eight of 12 RCTs ^{40,42,44-47,55,56} (66.7%) did not blind those who implemented or assessed the interventions and/or those who reported relevant information. These limitations may have created implementation and assessor biases, producing increases in false-positive results. With regard to the quality of the evidence, 2 domains (sensory/perceptual and verbal/language skills) were considered to have very low-quality evidence, and the remaining 6 domains (abstraction/executive function, attention/working memory, memory, motor skills, speed of information processing, and daily function) had moderate-quality evidence, for which the main reasons were moderate to high risk of bias, small samples, inclusion of few RCTs, and wide 95% CIs. This result suggested that our results need to be further explored. We found no significant publication bias across outcomes, suggesting our outcomes were statistically significant.

Limitations

This study has limitations. First, the number of studies included in the meta-analysis was small, producing heterogeneity and low evidence quality. Therefore, the optimal intervention design for eliciting beneficial outcomes remains unclear. Second, the included RCTs also lacked measurement of

blood, cerebrospinal fluid, and brain imaging biomarkers after CCT. Including such measurements could have helped to clarify the potential mechanisms of the benefits observed after CCT. Third, the RCTs did not measure concurrent treatment (eg, ART, tDCS, physical exercise, or mindfulness) during the CCT intervention. Analysis of synergistic factors and outcomes can help physicians develop more beneficial plans for patients.

Conclusions

The findings of this meta-analysis of pooled data from RCTs suggested that CCT programs were associated with significant improvements in 6 cognitive and daily function domains (including abstraction/executive function, attention/working memory, memory, motor skills, speed of information processing, and daily function) among people living with HIV. Future studies are needed to clarify whether there is a difference in training benefits between CCT programs and to examine the synergistic factors and outcomes of different auxiliary interventions (eg. tDCS or exercise). In addition, more studies are needed to confirm the impact of potential factors and to assess training protocols among a large population of individuals living with HIV who are at risk of developing HAND. Studies in the field of implementation science are especially needed to address the challenge of removing barriers and bringing CCT from scientific research into clinical practice and implementing CCT programs in the real world.

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SUPPLEMENT.

eTable 1. Full Search Strategy

eTable 2. Supplementary Search Strategy

eTable 3. Reasons for Exclusion in Full-text Assessment

eTable 4. Categorization of Outcome Measures

eTable 5. Study Quality of Evidence According to GRADE Guidelines

eTable 6. Full Results of Sensitivity Analyses

eFigure. Risk of Bias Summary