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Supplementary Material

Cognitive performance in Idiopathic Intracranial Hypertension and relevance of intracranial pressure

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Supplementary methods

Participants

IIH:WT was a five year randomized, controlled, parallel-group, multicenter trial. IIH:WT recruited participants at three UK National Health Service (NHS) hospitals between July 25, 2014 and May 25, 2017. Participants were identified from neurology and ophthalmology clinics from seven NHS hospitals. The National Research Ethics Committee West Midlands approved the trial (14/WM/0011).

Exclusion criteria were: pregnancy, significant comorbidity (including Cushing's, Addison's or use of steroids), undergone optic nerve sheath fenestration (due to their effects on long term OCT outcomes), specific medical or psychiatric contraindication for bariatric surgery and inability to provide informed consent. Those with central visual field defects that would impair ability to perform the screen based cognitive assessments were excluded. Those with a previous CSF shunt were included, but only if the shunt had failed and they had recurrence of active papilloedema (and ICP >25cmCSF), thus everyone in the cohort had active disease. For controls inclusion criteria were: female, BMI >35kg/m², able to give informed consent and aged between 18 and 55 years. Exclusion criteria were: pregnancy, inability to give informed consent and diagnosis of IIH.

Clinical measurements

Steroid hormone profiling

After an internal standard mixture was added to 400 μl of serum, steroids were extracted via liquid/liquid extraction with 2 ml of tert-butyl methyl ether (MTBE). The MTBE layer was removed, evaporated to dryness, and reconstituted in methanol/water prior to LC-MS/MS analysis. The extracts were analyzed on a Xevo TQ-XS triple quadrupole mass spectrometer (Waters) coupled to an Acquity ultra-high performance liquid chromatography system (UPLC) (Waters). Steroids were separated on a HSS T3, (1.8 μm) column (Waters) using a methanol/water gradient (both with 0.1 % formic acid). Starting conditions was 45% methanol, which was held for 1 minute, followed by a linear gradient to 75 % methanol at 5 minutes. Subsequently, the column was washed at 98 % methanol and reconditioned at starting condition prior to the next injection. Steroid hormones were identified and quantified

via comparison to reference standards; positive identification was confirmed via matching retention time and two identical mass transitions. Steroid hormones quantified were cortisol and cortisone. The calibration series ranged from 0.01 to 250 ng/ml (including a blank and a 0 ng/ml calibrator).

Cognitive tests

Raven's Standard Progressive Matrices (fluid intelligence)

The Raven's Standard Progressive Matrices fluid intelligence test was given in accordance with the standard instructions. The test consists of five sections, with 12 problems in each section. The problems get progressively more difficult within each section. Participants complete the task using a scoring sheet to mark their responses. The measurements for analysis are the total percentage correct and the time taken to complete the test.

Attention network test

The attention network test is used to measure the alerting, orienting, and executive components of attention, and the interactions among the three attention networks.¹⁻³ Each trial was 3.5 seconds long, consisting of a fixation cross in the middle of the screen throughout the trial. After 1 second, on half of trials, an orienting cue appeared above or below the central fixation cross for 0.4 seconds. If the orienting cue had not appeared at 1 second, it would appear above or below fixation for 0.4 seconds at 1.4 seconds. Also at 1.4 seconds, on half of trials, an auditory alerting tone was presented for 0.1 seconds. At 1.8 seconds, five arrows appeared either above or below the central fixation cross for 1.7 seconds or until response. The target display was composed of a target central arrow pointing towards the left or right, and two flanking arrows on either side of the central arrow, all of which were pointing in the same direction (left or right). The task was to press the left or right arrow key on the keyboard to indicate whether the central arrow was pointing to the left or the right, as quickly and accurately as possible. If the target display appeared in the same location as the orienting cue, the orienting cue was valid; otherwise the orienting cue was invalid. If the flanking arrows were pointing in the same direction as the central target arrow, they were congruent with the target, if they were pointing in the opposite direction they were incongruent.

Participants completed a practice block of 24 trials before completing two blocks of 128 trials. The task has a two (auditory signal: yes or no) x two (orienting cue: valid or invalid) x two (flanker congruency: congruent or incongruent) design, such that there were 16 trials per condition in each block (32 trials per condition in total). The trials were presented in a random order in each block. Measurements for analysis included reaction times for correct responses and percentage correct. Condition differences (the alerting effect, the orienting effect, and the flanker effect) were computed as the difference in average reaction times and percentage of correct responses in each condition comparison (no alerting - alerting; invalid orienting cue - valid orienting cue; incongruent flankers - congruent flankers).

Operation span task

The operation span verbal working memory task (⁴ modified from ⁵) required that participants try to remember a series of words in the correct serial order, while trying to solve mathematical problems. Each trial started with the presentation of a math problem in the general form of "is (a x b) +/- c = d?". On each trial, participants read the equation out loud followed by "yes" or "no", if the equation was correct or incorrect. Half of presented equations were correct. Immediately following the response to the equation, a word appeared on the screen for one second, which participants read aloud. Following a 0.5 second delay, another trial was presented, or the recall instruction was presented. When presented with the recall instruction, participants recalled, out loud, the words that were presented in that block of trials, in the serial order in which they were presented. If a word could not be recalled, participants were instructed to replace the word with "can't remember" to preserve the serial order of words. Blocks consisted of two, three, four or five trials before the recall instruction, with three blocks of each size presented in a pseudorandom order. The measurement used for analysis was the overall percentage correct of 36 trials (blocks of trials with a size of two were excluded from the analysis due to performance ceiling effects).

Sustained attention task

In the sustained attention task, 225 digits (25 of each of the digits from 1-9) were presented for 0.25 seconds each. They were immediately followed by a mask (comprised of the overlaid capital letters O and X) for 0.9 seconds, to ensure speeded evaluation. The digits and mask on a given trial were of varying size on the screen (five sizes for each digit: 2, 2.5, 3, 3.5, and 4

normalised units, which were presented five times each). Participants were given the target number six, and only pressed the response button when the target number appeared (25 times total) and were told to respond as quickly and as accurately as possible. The task began after a practice block of 18 trials. Measurements for analysis included reaction time for correct responses, total percentage correct, and percentage correct to the target (errors of omission: when the target was presented but the response was incorrectly withheld).

Word span task

The word span task is the verbal short-term memory counterpart to the operation span task. On each trial, the participant is presented with a word on the computer screen for 1 second, followed by a 0.5 second delay, and the presentation of another word or the recall instruction. Participants read each word out loud when it appeared. When presented with the recall instruction, participants recalled aloud, the words presented during that block of trials in the correct serial order. As with the operation span task, participants were instructed to replace forgotten words with 'can't remember' to preserve the serial order of the recalled words. Blocks consisted of 2, 3, 4, 5, 6, or 7 words before the recall instruction, with each size presented three times in a pseudorandom order. The measurement used for analysis was the overall percentage correct of 66 trials (blocks of trials with sizes 2 and 3 were excluded from the analysis due to ceiling effects).

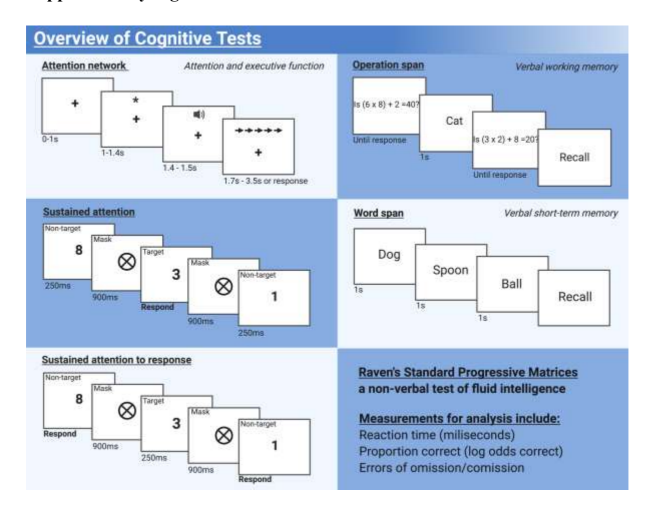
Sustained attention to response task

The sustained attention to response task was presented in exactly the same fashion as the sustained attention task, except that the instructions were for the participant to press the response key for every digit that appeared except if the target (the digit three) appeared.^{6,7} The task began after a practice block of 18 trials. Measurements for analysis included reaction time for correct responses, total correct, total correct to the target (errors of commission: when the target was presented but a response was incorrectly made). The sustained attention to response task was carried out twice in a sub-group of IIH and control participants before and after a lumbar puncture. This was to allow the assessment of the effect of acute reduction in intracranial pressure.

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Supplementary Figure



Supplementary Fig. 1 Overview of cognitive tests completed by control and IIH participants.

All participants completed the tasks in order: the attention network test, operation span, sustained attention, word span, and the sustained attention to response task. Attention network: A fixation cross would remain in the middle of the screen throughout the 3.5s trial. In half of the trials; an orientating cue appeared above or below, after that an alerting tone was presented and if the cue had not appeared previously it did at this time. Then five arrows would appear below or above the fixation cross. The task was to press the left or right arrow key to indicate if the central arrow was pointing in the left or right direction. Operation span: Participants were required to remember a series of words in the correct order while trying to solve mathematical problems. The participants were required to read the equations aloud followed by yes or no and asked to recall words at the end of the trial. Sustained attention: Participants were given the target number 6 and presented with 225 digits (25 of each of the digits 1-9) which were masked between digits. Participants were required to

respond when the target number appeared. **Word span**: Participants are presented with a series of words and asked to recall the list in order. **Sustained attention to response**: Presented in the same fashion as SA task however participants were required to press the response key for every digit that appeared except the target number (3).

Test results were measured in reaction time (milliseconds), proportion correct (log odds correct) and errors of omission/commission (percentage correct when the target was presented but the response was incorrectly withheld).

Supplementary Table Guide

Supplementary Table 1. Summary of current medication use in IIH participants at baseline.

Supplementary Table 2. Descriptive statistics for attention network test conditions in IIH and controls.

Supplementary Table 3. Descriptive characteristics of IIH participants at baseline and 12 months follow up

Supplementary Table 4. Baseline and follow up descriptive statistics and within-group comparisons

Supplementary Table 5. Comparison of change in cognitive performance between community weight management intervention and surgery groups over 12 month period

Supplementary Table 1. Summary of current medication use in IIH participants at baseline.

Medication	IIH	Control	
	(n =66)	(n = 25)	
Acetazolamide, n (%)	19 (29%)	0	
Daily dose (mg); mean (SD)	855 (509)	-	
Topiramate, n (%)	6 (9%)	1 (4%)	
Daily dose (mg); mean (SD)	75 (67)	200	
Diuretics	3 (5%)	0	
Bendroflumethiazide	1 (33%)	-	
Furosemide	1 (33%)	-	
Co-amilofruse	1 (33%)	-	
Antihypertensive	5 (8%)	2 (8%)	
B-blockers	1 (20%)	0	
ACE inhibitors	1 (20%)	2 (100%)	
A-II antagonist (ARB)	1 (20%)	0	
Other	2 (40%)	0	
Other headache preventatives	18 (27%)	2 (8%)	
Beta-blocker	1 (6%)	2 (100%)	
Tricyclic	7 (39%)	0	
Anticonvulsant	8 (44%)	0	
Other	2 (11%)	0	

ACE = Angiotensin-converting-enzyme; ARB = angiotensin II receptor blockers

Supplementary Table 2. Descriptive statistics for attention network test conditions in IIH and controls.

		Control	ШН	
Condition Measure		Mean (SD), n	Mean (SD), n	
Alerting	RT	0.693 (0.145), 19	0.696 (0.090), 53	
No Alerting	RT	0.711 (0.153), 19	0.713 (0.092), 53	
Alerting	Correct	0.983 (0.028), 19	0.960 (0.059), 53	
No Alerting	Correct	0.975 (0.048), 19	0.961 (0.061), 53	
Valid Orienting	RT	0.667 (0.156), 19	0.663 (0.092), 53	
Invalid Orienting	RT	0.737 (0.145), 19	0.746 (0.091), 53	
Valid Orienting	Correct	0.981 (0.039), 19	0.967 (0.055), 53	
Invalid Orienting	Correct	0.977 (0.039), 19	0.954 (0.066), 53	
Congruent Flankers	RT	0.640 (0.152), 19	0.642(0.088), 53	
Incongruent Flankers	RT	0.764 (0.150), 19	0.767(0.097), 53	
Congruent Flankers	Correct	0.997 (0.006), 19	0.985 (0.031), 53	
Incongruent Flankers	Correct	0.961 (0.068), 19	0.936 (0.1020, 53	

RT = Reaction time; Correct = proportion correct

Supplementary Table 3. Descriptive characteristics of IIH participants at baseline and 12 months follow up

	Baseline	12 months	
Clinical measurement	Mean (SD), n	Mean (SD), n	- p
Body Mass Index (BMI), kg/m ²	43.9 (7.0), 66	39.0 (8.8), 59	<0.001
Intracranial opening pressure (cmCSF)	34.7 (5.7), 66	29.0 (7.7), 54	<0.001 *
Intracranial closing pressure (cmCSF)	19.4 (3.8), 61	17.7 (4.6), 52	0.062
Headache severity day of test	3.5 (2.8), 61	2.1 (3.0),51	0.009 *
Monthly headache days	22.2 (8.0), 63	14.8 (11.6), 53	<0.001 *
Headache severity	5.0 (2.0), 63	3.6 (2.9), 53	<0.001 *
Headache disability (HIT-6)	64.7 (7.3), 65	58.5 (10.7), 55	<0.001 *
Serum IL-6	6.0 (2.5), 61	5.7 (2.4), 50	0.313
CSF IL-6	6.5 (15.8), 54	3.6 (1.9), 47	0.269
OCT RNFL thickness (µM)	139.6 (5.8), 59	106.3 (28.0), 57	<0.001 *
Hospital anxiety and depression scale – Anxiety score (HAD A)	10.3 (4.9), 65	9.9 (4.9), 57	0.466
Hospital anxiety and depression scale - Depression score (HAD D)	7.6 (4.5), 65	6.70 (4.7), 57	0.108
Quality of life (PCS)	28.7 (12.7), 60	37.7 (14.9), 53	<0.001 *
Quality of life (MCS)	37.7 (11.0), 60	38.9 (12.2), 53	0.633
Apnea-hypopnea index	14.1 (20.5), 40	12.4 (21.0), 20	0.025 *
Humphrey visual field mean deviation	-3.6 (3.7), 65	-2.4 (2.5), 58	<0.001 *

HIT-6 = headache impact test; IL-6 = interleukin-6; MCS = mental component score; OCT = optical coherence tomography; PCS = physical component score.

Supplementary Table 4. Baseline and follow up descriptive statistics and within-group comparisons

				Baseline	Follow up		
Cognitive Test	Measure	Group	n	Score (SD)	Score (SD)	Change	p
Attention network (averaged)	RT	CWI	18	690 (89)	653 (83)	-37	0.079
		Surgery	21	699 (78)	661 (99)	-39	0.032
	Correct	CWI	18	0.971 (0.041)	0.985 (0.021)	0.013	< 0.001
		Surgery	21	0.957 (0.076)	0.950 (0.108)	-0.007	0.036
-	RT	CWI	18	464 (49)	436 (45)	-28	0.001
		Surgery	16	479 (55)	459 (40)	-0.020	0.087
Sustained attention	Correct	CWI	18	0.996 (0.005)	0.995 (0.007)	-0.001	0.516
		Surgery	16	0.996 (0.004)	0.994 (0.009)	-0.002	0.206
	Target Correct	CWI	18	0.987 (0.023)	0.977 (0.039)	-0.010	0.239
		Surgery	16	0.976 (0.027)	0.962 (0.059)	-0.014	0.190
Sustained attention to response	RT	CWI	20	400 (79)	357 (61)	-0.043	0.012
		Surgery	16	380 (66)	350 (59)	-0.029	0.097
	Correct	CWI	20	0.910 (0.082)	0.892 (0.113)	-0.017	0.007
		Surgery	16	0.894 (0.041)	0.901 (0.067)	0.008	0.294
	Target Correct	CWI	20	0.716 (0.296)	0.679 (0.323)	-0.037	0.007
		Surgery	16	0.562 (0.247)	0.628 (0.310)	0.066	0.300
Word span	Correct	CWI	21	0.647 (0.114)	0.630 (0.116)	-0.017	0.337
		Surgery	21	0.594 (0.182)	0.660 (0.148)	0.066	< 0.001
Operation span	n Correct	CWI	20	0.547 (0.201)	0.639 (0.193)	0.092	0.181
		Surgery	19	0.631 (0.180)	0.662 (0.143)	0.030	0.378

CWI = Community weight management intervention; RT = Reaction time; Correct = proportion correct; Target correct = proportion correct on target-present trials; Scores expressed as mean (SD) and compared using paired t-tests or z-tests as appropriate.

Supplementary Table 5. Comparison of change in cognitive performance between community weight management intervention and surgery groups over 12 month period

Interaction	Measure	p
Attention network task alerting (yes – no) x Group	RT	0.701
x Session	Correct	0.124
Attention network task alerting (valid – invalid) x	RT	0.581
Group x Session	Correct	0.217
Attention network task flanker (incongruent –	RT	0.037
congruent) x Group x Session	Correct	0.800
	RT	0.541
Sustained attention x Group x Session	Correct	0.741
	Target Correct	0.911
	RT	0.569
Sustained attention to response x Group x Session	Correct	0.009
	Target Correct	0.011
Word span x Group x Session	Correct	0.001
Operation span x Group x Session	Correct	0.420

RT = Reaction time (seconds); Correct = proportion correct; Target correct = proportion correct on target-present trials; comparisons made using mixed (within- and between-groups) analysis of variance.

Cogntive dysfunction in Idiopathic Intracranial Hypertension

n = 66

Idiopathic intracranial hypertension

n = 25

Gender, age and BMI matched controls



VS



Attention



Working memory



Executive function



Verbal short term memory



Attention and executive function (sustained attention and sustained attention to response tasks) is significantly impaired in Idiopathic Intracranial Hypertension

Cognitive function is driven by intracranial pressure

Acute
Post lumbar
puncture

Improves attention and executive function

Chronic Over 12 months Improves verbal short term memory

Factors impacting cognition



Headache severity



Depression



Obstructive Sleep Apnea



Disease duration



Serum cortisol

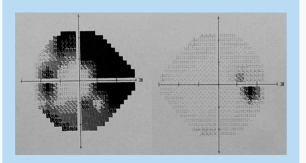
Potentially reversible with treatment

Cognitive impairment impacts visual field testing

Impaired attention and executive function



Impacts the ability to perform visual field testing as measured by false positive and negative indices, visual field index and test duration



Poor performance

Inacurate mean deviation evaluation