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This is the author's manuscript
Original Citation:
Availability:
This version is available http://hdl.handle.net/2318/1520886 since 2017-01-16T17:02:46Z
Published version:
DOI:10.1002/eat.22399
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This is the accepted version of the following article:

Does depression matter in neuropsychological performances in anorexia nervosa? A descriptive review

International Journal of Eating Disorders 48 (6) 2015 : 736–745

DOI: 10.1002/eat.22399

which has been published in final form at

http://onlinelibrary.wiley.com/doi/10.1002/eat.22399/full

Does depression matter in neuropsychological performances in anorexia nervosa? A descriptive review

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ABSTRACT

Objective. This review aims to examine the impact of depressive symptoms on the assessment of cognitive flexibility, central coherence, and decision-making in individuals with anorexia nervosa (AN).

Method. An online search was carried out using PubMed and PsycInfo. Articles were selected for review if they were published in English between 1990 and 2014 and used the Wisconsin Card Sorting Test, the Trail Making Task parts A and B, the Brixton Test, the Rey-Osterrieth Complex Figure Test, and/or the Iowa Gambling Task.

Results. Sixty-two studies were included. Thirty (48%) of the studies statistically assessed the association between depression and neurocognition in AN versus healthy controls. Where significant correlations were found, it became clear that the more serious the depression, the greater the neuropsychological impairment. Only six (10%) studies examined whether increased depressive symptoms were able to eliminate the differences between individuals with AN and healthy controls, and one study found that depressive symptoms did eliminate group differences in cognitive flexibility and decision-making.

Discussion. Only a subgroup of articles on neuropsychology in AN adjusted for depression. However, given the role of depression that some articles suggest, future studies should pay closer attention to the evaluation of this potential confounder.

Keywords: anorexia nervosa; neuropsychology; depression; cognitive flexibility; central coherence; decision making.

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EMPIRICAL ARTICLE

Introduction

Major depressive disorder is a frequent comorbid condition of anorexia nervosa (AN),[1, 2] and the impact of depressive symptoms on neurocognition has been acknowledged.[3] In fact, literature has reported depressed patients to have altered set-shifting[4-6] and decision-making[7] abilities. Several mechanisms may underlie such alterations, including attention, memory, emotional information, motivation, rumination, and response to failure[8].

However, the available body of evidence on the potential role of depression on neuropsychological performances of patients with eating disorders is still debated.[9, 10] Regarding AN, results are even more mixed, with studies proposing a depression-related impairment in cognitive flexibility, mainly with respect to attention[11] and serotonin dysregulation.[12] A few reviews on neuropsychological domains in AN exist,[13-18] but none of them specifically address depression-related aspects.

The main aim of this work is twofold: (1) to examine whether depressive symptoms were assessed in patients with AN in studies evaluating cognitive flexibility, central coherence, and decision-making. The latter domains were chosen given their well-established alterations[17] in AN; (2) to outline the body of evidence currently available on the effects of depression on neuropsychology in AN. In fact, where depression is present, the different performances on neuropsychology found between healthy controls (HC) and those with AN might be due to depression rather than the AN pathology.

Methods

Two independent researchers (S.B. and M.A.) carried out an online search on PubMed and PsycInfo databases. A hand search of the reference lists of all articles meeting the inclusion criteria was also performed.

The following inclusion criteria were adopted: (1) articles published between 1990 and 2014; (2) studies focusing on currently ill adults with AN; (3) works on cognitive flexibility, decision-making, and central coherence; (4) original research articles; (5) English language; (6) HC as a comparison group; (7) use of the following tests: Wisconsin Card Sorting Test (WCST), Trail Making Task parts A and B (TMT-A/-B), Brixton Test, Rey-Osterrieth Complex Figure Test (ROFT), and Iowa Gambling Task (IGT). Reviews and case reports were excluded.

To ascertain the second aim of this review, we focused on two main statistical methods: (1) correlational analyses (Pearson's or Spearman's linear correlations), and (2) statistical adjustments for depression (e.g., univariate general linear model [UGLM], multivariate analysis of variance [MANOVA], multivariate analysis of covariance [MANCOVA]). The former analyses identify a bidirectional evaluation between depression and neuropsychology. The latter instead aim to verify whether depression can explain the difference on neuropsychological performance between AN and HC.

The search keywords included the following: "eating disorders" OR "anorexia nervosa", AND "neuropsychology", OR "cognitive flexibility", OR "decision-making", OR "rigidity", OR "setshifting", OR "central coherence", OR "Wisconsin Card Sorting Test", OR "Iowa Gambling Task", OR "Trail Making Task", OR "Brixton Test", OR "Rey-Osterrieth Complex Figure Test".

Results

In total, the initial search yielded 77 studies (72 using the online search and five with the hand search); however, 15 articles were excluded because six were reviews, [13-18] two were case reports, [19, 20] four had no HC group, [21-24] and three showed recovered individuals [25-27]. Thus, 62 studies were finally included.

Of all studies considered, 30 (48%)[1, 9, 28-55] statistically addressed the influence of depression on cognitive flexibility, central coherence, and decision-making. However, only six studies[1, 30, 36, 37, 41, 51] statistically controlled for depression, and one article[1] concluded that depression was able to erase the difference between AN and HC, concerning both cognitive flexibility and decision-making. Of 16 studies[9, 31, 32, 35, 38, 41-46, 48, 49, 53-55] on cognitive flexibility performing correlations, 14 studies[31, 32, 35, 38, 41, 42, 44-46, 48, 49, 53-55] found nonsignificant findings and two studies[9, 43] reported significant positive correlations between depression and suboptimal neurocognition. Moreover, of those papers that did not take depression as a limitation. In addition, 29 studies[9, 29, 31, 39, 43, 46-50, 52, 55-57, 62, 63, 66-73, 79-83] were conducted on a sample of less than 30 affected participants.

Assessment of Neuropsychological Domains

Cognitive Flexibility

Both not computerized and adapted computerized versions of WCST[84], TMT-A/-B[85], and Brixton Test[86] have been included.

Forty-three studies[1, 9, 31, 32, 35-38, 41-46, 48, 49, 51, 53-56, 58, 60-62, 64, 67-73, 75-77, 79, 81-83, 87-89] investigated cognitive flexibility and 29 articles[1, 31, 32, 35-38, 41, 43, 44, 46, 48, 49, 51, 53, 61, 62, 67, 68, 71, 72, 75-77, 81-83, 88, 89] found significant differences between patients with AN and HC on this neuropsychological domain (Table 1).

Thirty-five studies[1, 9, 31, 32, 35-38, 41-46, 48, 49, 51, 53-55, 60,61, 67-73, 75, 79, 82, 83, 88, 89] investigated depression and 20 articles[1, 9, 31, 32, 35-38, 41-46, 48, 49, 51, 53-55] included this assessment in subsequent statistical analyses yielding significant findings in three cases[1, 9, 43] (Fig. 1a). With more detail, of 16 studies[9, 31, 32, 35, 38, 41-46, 48, 49, 53-55] performing correlations, 14 studies[31, 32, 35, 38, 41, 42, 44-46, 48, 49, 53-55] found nonsignificant findings, whereas two studies[9, 43] reported instead significant positive correlations demonstrating that the greater the depression score, the more impaired the neuropsychological performance. Five studies[1, 36, 37, 41, 51] controlled the difference between AN and HC for depression, and in one case[1], such a difference did not hold significant after statistical control.

Anticle	Sample	Mean Age (years)	Gender	Neuropsychological Bomains Investigated and Main Results Companing AN versus HC	Instrument Used to Assess Depression	Correlations Between Neuropsychological Performance and Depression in AN	Control for Depression
Abbate-Daga et al., 2011 ¹	30 AN; 30 HC	24.1 AN; 24.7 HC	All females	Cognitive flexibility: ≠ Decision making: ≠	801	ï	UGLM Depression Sig. (F and p values not available)
Adoue et al., 2014 ³⁸	63 AN; 49 HC	24.8 AN; 30.3 HC	All females	Decision making: ≠	BDI	NS (rho and p values not available)	1
Arbel et al., 2013 ⁵⁶	15 AN; 10 EDNOSAN; 25 HC	23.8 AN, 23.1 HC	All females	Cognitive flexibility: =	I	1	1
Bodell et al., 2014 ²⁹	15 AN-R: 7 AN-BP. 20 HC	25.6 AN	All females	Decision making: ≠	HDRS	NS (r and p values not available)	Ĩ
Brogan et al., 2010 ⁸⁰	22 AN; 17 BN; 18 08; 20 HC	29.1 AN; 29.9 BN; 52.1 08: 27.7 HC	All females	Decision making: ≠	I	1000 July 1	1
Castro-Fornieles et al., 200957	12 AN, 9 HC	14.5 AN; 14.6 HC	AN: 91.67% females HC 88.89% females	Central coherence: =	CDI	Ē	Ē
Cavedini et al., 2004 ⁹⁸	26 AN-R; 33 AN-BP, 82 HC	21.7 AN-R; 23.4 AN- BP, 30.9 HC	AN-R: 96.1% females AN-BP: 96.6% females HC: 52.4% females	Cognitive flexibility: = Decision making: ≠	I	1	I
Cavedini et al., 2006 ¹⁹	18 AN-R; 20 AN-BP; 30 HC	23.8 AN-R; 21.5 AN- BP: 22.6 HC	All females	Decision making: ≠	I	Ē	Ē
Chan et al., 2014 ³⁰	94 ANI 63 BNI 67 HC	25.6 AN; 26.9 BN; 25.5 HC	97% females	Decision making: =	801	NS (r and p values not available)	Test statistics not available Depres- sion NS (F and p val- ues not available)
Danner et al., 2012 ³¹	16 AN; 15 AN-REC 15 HC	25.6 AN; 24.3 AN-rec; HC 25.8	All females	Cognitive flexibility: ≠ Decision making = Central coherence: =	11/108	Cognitive flexibility: WCST perseveration r = 0.13, $p = NS$ Deci- sion making: 1GT r = 0.24, $p = NS$ Cen- tral coherence: ROFT copyr = -0.12, p = NS ROFT recall r = 0.10, $p = NS$	1
Dmitrzak-Weglarz et al., 2013 ⁴⁰	46 AN-R; 14 AN-BP; 45 HC	15.7 AN-R; 16.5 AN- BP: 37.7 HC	All females	Cognitive flexibility: =	801	1	t
Fagundo et al., 201261	35 AN; 52 08; 137 HC	28.1 AN; 40.5 0B; 24.8 HC	All females	Cognitive flexibility: ≠ Decision making: ≠	SCL-90-R	Ĕ	Ē
Fassing et al., 2002 ⁶²	20 AN-R; 20 HC	23.8 AN; 23.1 HC	All females	Cognitive flexibility: ≠	1	1	
Favaro et al., 2012 ¹³	29 AN; 16 AN-Rec; 26 HC	25,8 AN; 23,8 AN-Rec; 26,7 HC	All females	Central coherence: ≠	1)KH	Î	Ē
Favaro et al., 2013 ³²	73 AN-R; 93 AN-Rec 140 HC	25.0 AN, 27.2 HC	All females	Gognitive flexibility: ≠	105H	Cognitive Flexibility. WCST global score r = -0.03, $p = NS$	E)
Fitzpatrick et al., 2012/07	32 AN; 22 HC	14.9 AN; 15.4 HC	All females	Cognitive flexibility: =	I	T	Ē
Galimberti et al., 2013 ⁸¹	29 AN; 29 UR-AN; 29 HC: 29 UR-HC	24.1 AN; 43.8 UR-AN; 28.6 HC: 43.3 UR-HC	All females	Cognitive flexibility: ≠ Decision makine: ≠	Î)	Ē	Ē
Garrido et al., 201331	27 AN-R; 24 AN-BP; 20 BN; 38 HC	25.9 AN-R; 28.2 AN- BP + BN; 23.3 HC	All females	Decision making: #	801	NS (r and p-values not available)	T

TABLE 1. Studies on cognitive flexibility, decision-making, central coherence, and their evaluation of depression (N = 62)

Article	Sample	Mean Age (years)	Gender	Neuropsychological Domains Investigated and Main Results Comparing AN versus HC	Instrument Used to Assess Depression	Correlations Between Neuropsychological Performance and Depression in AN	Control for Depression
Giel et al., 2013 ⁹	15 AN; 20 UD; 35 HC	23.9 AN; 36.3 UD; 30.2 HC	AN: 100% females UD: 60% females HC 77% females	Cognitive flexibility. =	3550IQ	Cognitive Hexibility: WCST: $hho = 0.33$, p = 0.006 TMT: tho = 0.31, $p = 0.041$	ĩ
Giltberg et al., 200764 Goddard et al., 201482	51 AN; 51 HC 29 AN; 42 HC	24.5 AN; 24.2 HC 26.2 AN; 26.4 HC	Not available All males	Cognitive flexibility: = Cognitive flexibility: ≠	DAS5-21	11	11
Guillaume et al., Sono ⁹⁰	49 AN; 38 BN; 83 HC	233 AN; 23 BN, 28 HC	All females	Decision making: =	HDRS	I	ĵ
Harrison et al., 2011 ³⁴	35 AN-R; 15 AN-BP; 48 BN; 35 AN-Rec; 89 HC	27.1 ED group; 29 AN- Rec, 28.5 HC	All females	Central coherence: ≠	DASS-21	Overall ED group: NS (rho and p values not available)	ŀ
Heled et al., 2014 ⁶⁵	30 AN; 30 AN-WR; 44 HC	23.2 AN; 24.6 AN-WR; 24.5 HC	All females	Central coherence: #	E	I	Ē
Holliday et al., 2005 ⁵⁵	47 AN; 47 US-AN; 47 HC	26.3 AN. 27.6 USAN. 26.5 HC	All females	Cognitive flexibility: #	SOLH	Overall ED group: NS (r and p values not available)	1
Jones et al., 1991 ¹⁶	30 AN; 20 AN-WR, 38 BN; 39 HC	24.4 ANI, 26 AN WR, 24.1 BN, 24.9 HC	All females	Cognitive flexibility: # Central coherence #	MMPI Sale 2	1	MANCOVA Depression NS /(15, 326.16) = 1.48 p = 0.11
Karrakam et al., 2013 ⁸⁸	41 MZ-ED; 11 MZ-H; 12 DZ-ED; 8 DZ-H; 42 Control twins	31 MZ-ED; 54 MZ-H; 35 DZ-ED; 52 DZ-H; 45 Control twins	All females	Cognitive flexibility: # Central coherence: =	DAS5-21	ľ.	l
Kim et al., 2010 ⁵⁷	40 AN; 28 BN; 34 HC	22.8 AN; 23 BN; 22.6 HC	All females	Cognitive flexibility: #	BDI	t	MANOVA Depression NS (F and p values not available)
Kim et al., 2011 th	22 AN; 28 BN; 26 HC	22 AN: 23 BN: 23.5	All females	Central coherence: #	BDI	1	1
Kingston et al., 1996 ¹⁸	46 AN, 41 HC	22.1 AN; 22.HC	All females	Cognitive flexibility: \neq Central coherence: \neq	BDI	NS Central coherence: ROFT copy r = -0.253, $p = NS (r)and p values notavailable?$	ł
Konstantakopoulus et al. 2011 ⁶⁷	25 AN; 15 BN; 35 HC	28.6 AN; 27.2 BN; 24.9 HC	All females	Cognitive flexibility: ≠	SOLH	1	1
Liao et al. 2009 ¹⁸	29 AN; 26 BN; 51 HC	28.5 AN, 27.8 BN, 29.4 HC	All females	Decision making: #	BDI	NS (r and p values not available)	1
Lopez et al., 2008 ⁴⁰	42 AN; 42 HC	28:4 AN; 26.3 HC	All females	Central coherence: ≠	HADS	NS (r and p values not available)	Ē
Lounes et al., 2011 ⁴¹	45 AN; 49 HC	27.6 AN; 24.1 HC	All fernales	Cognitive flexibility: \neq	SOLH	Cognitive flexibility: r = 0.130, $p = NS$	MANCOVA Depression NS F(1,80)=0.007; p>0.05
Mathias and Kent, 1998 ⁴²	34 AN; 31 HC	22 AN: 20.8 HC	All females	Cognitive flexibility: = Central coherence: ≠	109	Cognitive flexibility: TMT-A $r = 0.001$, p = NS TMT-B r = 0.157, $p = NS Cen- tral coherence: ROFT result r = -0.156, p = NSp = NS ROFT copyr = -0.354, p = NS$	1

TABLE 1. Continued

Article	Sample	Mean Age (years)	Gender	Neuropsychological Domains Investigated and Main Results Comparing AN versus HC	Instrument Used to Assess Depression	Correlations Between Neuropsychological Performance and Depression in AN	Control for Depression
McAnarney et al.,	24 AN; 37 HC	16.3 AN; 15.9 HC	All females	Cognitive flexibility: \neq	BDI-II	I	I
Murphy et al., 2002 ⁶⁹	16 AN; 16 BN; 16 HC	22.3 AN; 22 BN; 25.3 HC	All females	Cognitive flexibility: =	BDI	I	I
Murphy et al., 2004 ⁷⁰	16 AN; 16 BN; 16 OCD; 16 HC	22.3 AN; 22.0 BN; 25.1 OCD: 25.3 HC	All females	Cognitive flexibility: = Cognitive flexibility: =	BDI	I	I
Nakazato et al., 2000 ⁷¹	29 AN; 18 AN Rec; 28	28.3 AN; 32.2 AN-Rec;	All females	Cognitive flexibility: ≠	HADS	I	I
2009 Nakazato et al., 2010 ⁷²	27 AN; 18 AN-Rec; 28 HC	27.7 AN; 32.2 AN-Rec; 26 9 HC	All females	Cognitive flexibility: \neq	HADS	I	I
Ohrmann et al., 2004 ⁴³	11 AN; 12 HC	22.7 AN; 27.5 HC	All females	Cognitive flexibility: ≠	BDI; MADRS	Cognitive flexibility: WCST number of cards rho = 0.888 , $p < 0.001$ WCST false category rho = 0.672 , $p < 0.05$ wCST perservations	I
Oltra-Cucarella et al.,	12 AN; 12 AN-WR; 16	21.7 AN; 22.2 AN-WR; 18.6 HC	All females	Cognitive flexibility: =	BDI-II	100.0 × 4 .000.0 - 0111	I
Pignatti and Bernas- coni. 2013 ⁸³	23 AN; 17 BN; 20 HC	29.1 AN; 29.9 BN; 27.8 HC	All females	Cognitive flexibility: ≠	SCL-90-R	I	I
Roberts et al., 2010 ⁴⁴	35 AN-R; 33 AN-BP; 30 BN; 30 AN-Rec; 30 US- AN; 20 US-BN; 88 HC	23.7 AN-R; 25.6 AN- BP; 26.4 BN; 32.1 AN- Rec; 24.2 US-AN; 27.6 US-RN: 28.4 HC	All females	Cognitive flexibility: \neq	HADS	NS (r and <i>p</i> values not available)	I
Roberts et al., 2013^{74}	35 AN-R; 33 AN-BP; 30 BN; 30 AN-Rec; 30 US- AN; 20 US-BN; 88 HC	23.7 AN PK 25.6 AN- 23.7 AN PK 25.6 AN- BP; 26.4 BN; 32.1 AN- Rec; 24.2 US-AN; 27.6 US-AN; 28.4 HC	All females	Central coherence: ≠	HADS	I	I
Sarrar et al., 2011 ⁴⁵	30 AN; 28 HC	16.2 AN; 16.7 HC	All females	Cognitive flexibility: =	DIKJ	NS (<i>r</i> and <i>p</i> values not	Ι
Sato et al., 2013 ⁴⁶	15 AN; 15 HC	23 AN; 22 HC	All females	Cognitive flexibility: \neq	MMPI Scale 2	NS (<i>r</i> and <i>p</i> values not available)	Ι
Sherman et al., 2006^{47}	18 AN; 19 HC	25.6 AN; 25.7 HC	All females	Central coherence: ≠	BDI	NS (r and p values not	I
Stedal et al., 2012 ⁸⁹	155 AN; 66 HC	17.1 AN; 15.4 HC	AN: 95.5% females	Cognitive flexibility: ≠ Central coherence: ≠	BDI		Ι
Steinglass et al., 2006 ⁴⁸	15 AN; 11 HC	24 HC; 25.6 AN	All females	Cognitive flexibility: ≠	BDI	NS (<i>r</i> and <i>p</i> values not available)	I
Szmukler et al., 1992 ⁴⁹	18 AN; 18 HC	Mean age not available	All females	Cognitive flexibility: \neq	BDI	NS (<i>r</i> and <i>p</i> values not available)	I
Tapajóz P de Sampaio et al., 2013 ⁵⁰	8 AN-R; 1 AN-BP; 15 EDNOS-AN; 15 BN-P; 3 BN-NP; 6 EDNOS-BN; 24 HC	24.5 AN; 24.4 BN; 25.2 HC	All females	Central coherence: ≠	BDI	NS (rho and p values not available)	I

TABLE 1. Continued

Correlations Between Neuropsychological Performance and Depression in AN Coi			Decision making: IGT performance r = -0.283, $p = NS$	I	1	I	NS (r and p values not available)	Π	NS (r, rho and p values not available)	
0. Instrument Used to Assess Depression	HADS	HADS	BDI D	I	1	1	HSCL NC	DASS-21		
Domains Investigated and Main Results Comparing AN versus HC	Cognitive flexibility: \neq	Cognitive flexibility: \neq	Decision making: ≠	Cognitive flexibility: ≠	Cognitive flexibility: ≠	Decision making: ≠	Cognitive flexibility: \neq Central coherence: \neq	Cognitive flexibility: =	Cognitive flexibility: =	
Gender	All females	All females	All females	All females	All females	AN or sub-threshold AN: 60.4%	All females	All females	All females	
Mean Age (years)	26.7 AN; 26.5 BN; 24.8 HC	27.2 AN; 28.4 AN-Rec; 25.9 HC	28.5 AN; 28.9 AN-Rec; 26.3 HC	26.9 AN; 30.2 AN-Rec; 27.7 BN; 26.5 EDNOS; 27 HC	25.4 AN-R; 30.7 AN- Rec; 27.3 BN; 27.7 HC	27.2 AN male; 27.5 AN female; 25.4 HC male; 22.2 HC female	25.7 AN; 24.5 AN-WR; 30.8 AN-Rec; 27.5 US- AN; 27.4 HC	21.8 AN; 22 HC	26 AN-R; 20 AN-BP; 19 HC	A 1 4 6 6 111 4 80
Sample	34 AN; 19 BN; 35 HC	20 AN-R; 14 AN-BP; 18 AN-R Rec; 36 HC	29 AN; 14 AN-Rec, 29 HC	215 AN; 72 AN-Rec, 69 BN; 29 EDNOS; 216 HC	171 AN-R; 90 AN-Rec; 82 BN; 199 HC	19 AN or sub- threshold AN male; 29 AN or sub-threshold AN female; 20 HC male; 41 HC female	60 AN; 63 AN-WR; 29 AN-Rec; 28 US-AN; 120 HC	24 AN; 24 HC	31 AN-R; 20 AN-BP; 26 HC	AL CUAL A ALLER
Article	Tchanturia et al., 2004 ⁵¹	Tchanturia et al., 2004 ⁷⁵	Tchanturia et al., 2007 ⁵²	Tchanturia et al., 2011 ⁷⁶	Tchanturia et al., 2012 ⁷⁷	Tchanturia et al., 2012 ⁷⁸	Tenconi et al., 2010 ⁵³	Tokley and Kemps, 2007 ⁷⁹	3°th	

ontrol for Depression

IANCOVA Depression NS [F(4,82) = 1.40,

I 1

p = 0.24

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osa; BN-NP: bulimia nervosa-nonpurging; BN-P: bulimia nervosa-purging; EDNOS-BN: eating disorder not otherwise specified; EDNOS-AN: eating disorder not otherwise specified-nonewise specified-bulimia nervosa type; DD: unipolar depression; OCD: obsessive-compulsive disorder; UR-AN: unaffected relatives of AN; UR-HC: unaffected relatives of HC; US-AN: unaffected siters of Notes: AN: anorexia nervosa; AN-BP: anorexia nervosa-binge-purging subtype; AN-R: anorexia nervosa-restricting subtype; AN-Rec: anorexia nervosa-recovered; AN-WP: anorexia nervosa-binge-purging subtype; AN-Rec: anorexia nervos AN; USBN: unaffected sisters of BN; MZ-ED, monozygotic eating disorder probands; MZ-H: monozygotic non-eating-disorder cotwin; DZ-ED: dizygotic eating disorder probands; DZ-H: dizygotic non-eating-disorder cotwin; LO-HC: health control with low obsessionality; HO-HC: health control with high obsessionality; HC: healthy controls.

als; r = -0.02, p = NSp = NS WCST total tri-

Cognitive flexibility: WCST perseverative p = NS WCST totalerrors: r = -0.03, errors: r = 0.06.

DASS-21

Cognitive flexibility: =

All females

27.6 AN-R; 22.6 LO-HC: 21.1 HO-HC

22 AN-R; 21 LO-HC; 20

Wilsdon and Wade, 2006⁵⁵

HO-HC HC

Neuropsychological domain investigated and main results comparing AN versus HC; \neq : the performance of patients with anorexia nervosa were significantly worse than healthy controls; =: no significant differences were found between patients and controls.

Instrument used to assess depression: BDI: Beck Depression Inventory; CDI: Children"s Depression Inventory; BDI-II: Beck Depression Inventory-III; SCL-90-R: Symptom Check List-90-Revised; HSCL: Hopkins Symptoms Check Ust; QID5-SR: Quick Inventory of Depressive Symptomatology; DAS5-21: Depression, Anxiety, and Stress Scale-21; HDRS: Hamilton Depression Rating Scale; MMPI: Minnesota Multiphasic Personality Inventory; HADS: Hospital Anxiety and Depression Scale; MADRS: Montgomery Asberg Depression Rating Scale; DIKJ: Depressionsinventar für Kinder und Jugendliche; No: depression not evaluated.

Correlations between neuropsychological performance and depression in AN: --: no correlations were performed; NS (not significant): no significant correlations were found.

Depression NS: depression was found not to reach significance; Depression Sig: depression was found to reach significance.

Continued TABLE 1.

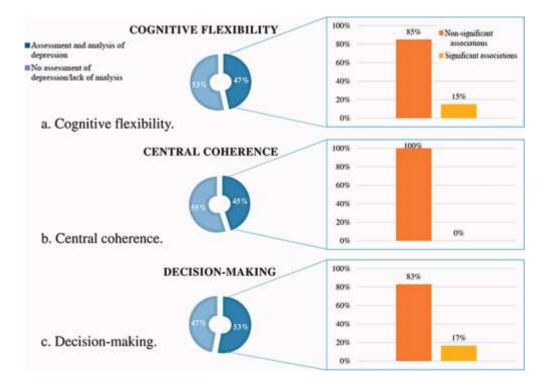


FIGURE 1. Proportion of studies on neuropsychological impairments in anorexia nervosa that took depression statistically into account and their reported presence of significant versus nonsignificant associations. [Color figure can be viewed in the online issue, which is available at wileyonlinelibrary.com.]

Central Coherence

Studies assessing central coherence using the ROCF[90], a test used to assess visuospatial abilities, were included. A Central Coherence Index can be computed resulting from the order of construction and style indices. The drawing style can be assessed according to the scoring systems of Savage and colleagues[91] and of Booth[92]. Of 20 articles[1, 34, 36, 38, 40, 42, 47, 50, 53, 57, 63, 65, 66, 69, 70, 73, 74, 82, 88, 89] on central coherence, 15 studies[34, 36, 38, 40, 42, 47, 50, 53, 63, 65, 66, 73, 74, 82, 89] found significant differences in global score between AN and HC on this measure (Table 1).

The vast majority of studies investigated depression (19[31, 34, 36, 38, 40, 42, 47, 50, 53, 57, 63, 66, 69, 70, 73, 74, 82, 88, 89]) with studies[31, 34, 36, 38, 40, 42, 47, 50, 53] including such data in subsequent statistical analyses. No articles reported significant findings on the role of depression using either correlations[31, 34, 38, 40, 42, 47, 50, 53] or MANCOVA[36] (Fig. 1b).

Decision-making

Fifteen studies[1, 28-31, 33, 39, 52, 58, 59, 61, 78, 80, 81, 93] investigated decision-making using the IGT[94] and 12 studies[1, 28, 29, 39, 52, 58, 59, 61, 78, 80, 81, 93] reported differences between AN and HC (Table 1).

Ten studies[1, 28-31, 33, 39, 52, 61, 93] investigated depression and 8 studies[1, 28-31, 33, 39, 52] included this assessment in subsequent statistical analysis. In one case[1], the difference between AN and HC on neurocognition was no longer significant after adjusting for depression.

Discussion

Most studies on neuropsychology in AN performed an assessment of depression using either self-report or clinician-rated instruments. However, about half of the articles included in this review statistically addressed (e.g., correlations, UGLM, MANOVA, etc.) the role of depression, and as a result, the state-of-the-art on this topic is mixed. Significant positive correlations were reported by two studies[9, 43] demonstrating that the greater the depression score, the more impaired the neuropsychological performance. Regarding depression, five studies[30, 36, 37, 41, 51] of six[1, 30, 36, 37, 41, 51] did not find depression to explain the difference between AN and HC on neuropsychology.

From a statistical standpoint, correlations can effectively identify a bidirectional association between depression and neurocognition; however, only a statistical adjustment for depression could ascertain whether the difference in the performance between AN and HC goes away. Nevertheless, only a minority of studies (i.e., 10%) used such a statistical analysis.

The plethora of instruments that have been used to assess depression hampers the generalizability of the available findings. In addition, the small sample size considered in some studies makes the statistical power of the analysis questionable.

Such methodological flaws and the scarcity of studies on this topic represent a finding in itself of great interest, given the relevant influence of depression on neurocognition[3]. This is even more important due to the fact that depressive symptoms frequently plague individuals with AN[2]. Moreover, the rationale for controlling for depression in AN has also been recently acknowledged[95] by a study showing that the adjustment for depression evened out the difference between AN and HC regarding speed of information processing and verbal fluency and overall reduced the differences with respect to a variety of neuropsychological domains[95].

Bearing in mind that only preliminary data exist, studies on cognitive flexibility seem to support the possibility of a marginal effect of depressive symptoms on this neuropsychological domain. In contrast, central coherence was consistently found not to be influenced by depression. Although one study found depression to influence decision-making[1], only eight studies[1, 28-31, 33, 39, 52] are available on the latter domain, so conclusions cannot be drawn in this regard.

Speculating on the possible reasons for the association between depression and cognitive flexibility is beyond the scope of this review. However, these findings are in line with a recent meta-analysis on depression[3] and multiple mechanisms may be involved[8]. Instead, central coherence seemed to be unrelated to depression, although the ROFC could be influenced not only by depressive symptoms but also by obsessive traits[96]. Studies on major depressive disorder showed decision-making to be impaired to different degrees in affected individuals depending on cognitive flexibility[97]. Further research is needed on this topic in AN because no definitive statements can be made yet.

Some limitations should be acknowledged: studies on recovered individuals have been excluded and differences between AN subtypes have not been considered. Also, other neuropsychological domains have not been included, as well as starvation and other psychiatric comorbidities. Still, some clinical characteristics of the sample may vary (e.g., age) or could not be evaluated because that information was not available in all articles (e.g., duration of illness and medications).

In closing, the study of the relationship between depression and neurocognition in AN is only in its infancy. However, the data seem to suggest such an association, mostly in regard to cognitive flexibility. Therefore, future studies comparing individuals affected by AN with and without comorbid major depression versus HC may shed light on this matter. The influence of depression on neuropsychological impairments in AN may have research (i.e., debate on cognition as candidate endophenotype[98]) and clinical (e.g., Cognitive Remediation Therapy[99, 100]) implications. For example, Cognitive Remediation Therapy may be tailored according to patients' needs and depending on their depressive symptoms.

ACKNOWLEDGEMENTS

The authors would like to thank Dr. Thomas Fummo, who revised the English of this article.

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