Neuropsychiatric Disease and Treatment downloaded from https://www.dovepress.com/

Functioning outcomes with adjunctive treatments for major depressive disorder: a systematic review of randomized placebo-controlled studies

Emmanuelle Weiller¹ Catherine Weiss² Christopher P Watling³ Christopher Edge³ Mary Hobart² Hans Eriksson¹ Maurizio Fava⁴⁻⁶

¹H. Lundbeck A/S, Valby, Copenhagen, Denmark; ²Otsuka Pharmaceutical Development & Commercialization Inc., Princeton, NJ, USA; 3Cambridge Medical Communication Ltd, Cambridge, UK; 4Division of Clinical Research of the MGH Research Institute, ⁵Department of Psychiatry, ⁶Clinical Trials Network & Institute (CTNI), Massachusetts General Hospital, Boston, MA, USA

Objective: Patients with major depressive disorder (MDD) with inadequate response to antidepressant treatment (ADT) may suffer a prolonged loss of functioning. This review aimed to determine if self-rated functional measures are informative in randomized placebo-controlled studies of adjunctive therapy in patients with MDD and inadequate response to ADT.

Methods: This was a systematic literature review of articles in any language from the MEDLINE database published between January 1990 and March 2017. Eligible studies met the following criteria: patients with MDD; inadequate response to at least one ADT; adjunctive therapy (pharmacological or otherwise) to ADT; placebo control group; randomized controlled trial or a post hoc analysis of a randomized controlled trial; reported a self-rated functioning scale. Study characteristics and functioning efficacy data were extracted.

Results: A total of 2,090 discrete records were screened, 293 full-text articles were assessed for eligibility, and 26 studies were included. All studies were acute (6-12 weeks) except for one 52-week study. The only self-rated functioning scale used in the included studies was the Sheehan Disability Scale (SDS). Of the 13 adjunctive agents identified, aripiprazole, brexpiprazole, edivoxetine, and risperidone improved functioning versus placebo (p < 0.05), as measured by the SDS total or mean score. On the SDS "work/studies" item, only aripiprazole had a statistically significant benefit, in one study out of four. Thus, where a benefit was observed on the SDS total or mean, this was generally driven by improvement on the "social life" and "family life" items. A limitation of the review is that it only considered published literature from one database.

Conclusion: The SDS, a self-rated functional measure, is informative in acute randomized placebo-controlled studies of adjunctive therapy in patients with MDD and inadequate response to ADT. However, the item that measures work performance may be less relevant to this population than the items that measure social and family life.

Keywords: depression, antidepressant, adjunct, Sheehan Disability Scale, functional, work

Introduction

Major depressive disorder (MDD) is characterized by symptoms including depressed mood and a loss of interest or pleasure in activities. 1 As a consequence of depressive symptoms, patients with MDD typically have impaired functioning across multiple domains, including work, social, and family functioning.^{2,3} For example, depressive symptoms are associated with reduced marital quality, reduced work performance, and lower earnings.2

Key goals for patients with MDD experiencing a depressive episode are remission and full recovery.^{4,5} Recovery should be considered in broad terms, encompassing work, social, and family functioning as well as improvement of depressive symptoms.^{6,7}

Correspondence: Emmanuelle Weiller H. Lundbeck A/S, Ottiliavej 9, 2500 Valby, Denmark Tel +33 6 8042 1831 Email emmanuelle.weiller@gmail.com

Indeed, patients with MDD consider a return to normal levels of functioning as one of the most important factors in defining remission from depression.⁸ Although numerous standardized assessments are available to monitor functioning outcomes in the clinic and in research, functioning scales are used less frequently and less consistently than symptom severity scales.⁹ Furthermore, functioning may be less responsive to treatment than symptoms, meaning that functional improvement can lag behind symptomatic outcomes.^{10,11}

Despite being the mainstay of pharmacological treatment for MDD, more than half of patients do not respond to antidepressant treatment (ADT), as shown by the Sequenced Treatment Alternatives to Relieve Depression (STAR*D) study.¹² Patients with inadequate response to ADT have a prolonged loss of functioning and a lower likelihood of employment than those patients who do respond. 13,14 Treatment strategies for patients with inadequate response to an optimized dose of ADT include switching to another antidepressant, combining the initial antidepressant with a second antidepressant that has a different mode of action, or augmenting the antidepressant with a non-antidepressant drug. 4,15-18 Of the various options for augmentation, second-generation antipsychotics are best supported by the evidence. ¹⁹ However, the effects of different adjunctive therapies on patient functioning have not been consistently studied, and it is not clear whether existing measures of functioning are useful among patients with MDD and inadequate response to ADT.

A recent systematic review investigated the effect of ADT (selective serotonin reuptake inhibitors, serotonin—norepinephrine reuptake inhibitors, other antidepressants, and psychotherapies such as cognitive behavior therapy [CBT]) on functional outcomes in MDD.²⁰ The review, which excluded clinical studies of adjunctive pharmacotherapies, concluded that functioning improves together with depressive symptoms, but that functional deficits often remain, even among patients who achieve symptomatic remission.²⁰ Given the importance of functioning to the overall well-being of patients, the aim of the present systematic literature review was to determine if self-rated functional measures are informative in randomized placebo-controlled studies of adjunctive therapy in patients with MDD and inadequate response to ADT.

Methods

This systematic review adheres to PRISMA.²¹

Eligibility criteria

Studies were included if they were published, randomized, placebo-controlled studies of adjunctive therapy to ADT in

patients with MDD and inadequate response to at least one ADT, and reported a self-rated scale of functioning. The literature search was performed on 8 March 2017. Reports were limited to those published on or after 1 January 1990. No language exclusions were applied.

Search strategy

The aim of the initial top-level search strategy was to identify studies that satisfied three criteria: 1) the study included patients with MDD; 2) the study was of antidepressant augmentation (with any pharmacological or non-pharmacological approach, such as CBT or deep brain stimulation); and 3) the study included a placebo or sham control group. The US National Library of Medicine's MEDLINE database was searched, via PubMed, using the terms: (depress* OR MDD) AND (adjunct* OR "add-on" OR augment* OR resist* OR refractory OR inadequate OR incomplete OR suboptimal) AND (placebo).

Following the initial search it became apparent that these terms were likely to miss some publications of olanzapine–fluoxetine combination (OFC) studies. OFC is indicated for the treatment of treatment-resistant depression in the US,²² and, due to its availability as a single tablet, has been tested against fluoxetine with no need for placebo. Consequently, a second search was performed on 21 March 2017 to capture OFC studies, using the terms: (depress* OR MDD) AND ((olanzapine AND fluoxetine) OR OFC).

Study selection

Following the top-level database searches, duplicates were excluded and records were screened to exclude unsuitable articles based on titles and abstracts (Figure 1). At this stage, studies were not excluded based on a lack of functioning outcomes, because these are often secondary outcomes not mentioned in abstracts.

After screening, full-text articles for the remaining records were assessed for eligibility, defined as meeting all of the following criteria: 1) the study included patients with MDD; 2) the patients had inadequate response (by any definition) to at least one ADT; 3) the study investigated an adjunctive therapy to ADT; 4) the study included a placebo or sham control group (or a fluoxetine control group, for OFC); 5) the study was a randomized controlled trial or a post hoc analysis of a randomized controlled trial; and 6) the study reported self-rated functioning scale (or subscale) outcomes. Any self-rated functioning scale was eligible, defined as a scale that reflects the user's actual behavior in the world and is assessed in ways that emphasize doing, performing,

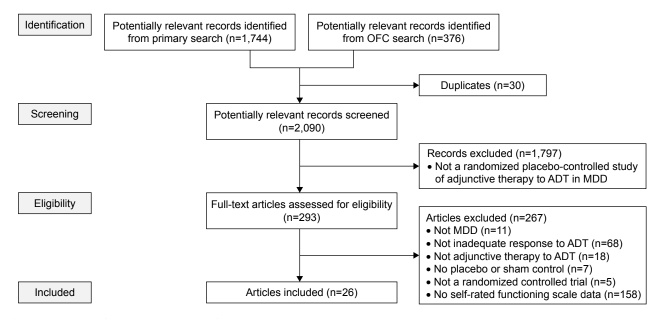


Figure 1 Flow diagram of published articles examined for inclusion in a systematic review.

Abbreviations: ADT, antidepressant treatment; MDD, major depressive disorder; OFC, olanzapine–fluoxetine combination.

maintaining, etc (this is distinct from quality of life – a measure based on self-perception and context with an emphasis on satisfaction, contentment, or enjoyment in various aspects of life). Eligibility assessment was performed in duplicate by two reviewers (split between CPW, CE and JARM), and disagreements were resolved by consensus.

Data extraction

The following data were extracted for each study, where available: 1) the definition of inadequate response to ADT, both historical (prior to study enrollment) and prospective/ ongoing (during the study); 2) the adjunctive treatment (type, dose, and duration); 3) the number of randomized patients; 4) the primary efficacy endpoint and whether or not it was met (in order that failed or negative studies might be identified); 5) the name of the self-rated functioning scale/subscale/items used; 6) self-rated functioning scale scores at baseline (ie, randomization to adjunctive therapy) and the mean change from baseline to the study endpoint (including error measurements, p-values versus placebo, and patient numbers); 7) the setting; and 8) the source of funding. Published data were supplemented with data from ClinicalTrials.gov and study protocols/reports, where available. One reviewer extracted the data from included studies (CPW) and a second reviewer checked the extracted data (CE). Disagreements were resolved by checking the original data source. To assess the risk of bias in individual studies, one reviewer (CPW) judged the adequacy of randomization, blinding, and outcome reporting for each study.

Results

Study characteristics

A total of 2,090 discrete records were identified and screened (Figure 1).

Of the 293 full-text articles assessed for eligibility, 26 articles were included, of which 20 reported one or more primary study, and six reported post hoc analyses of already identified studies. In total, these articles described 26 different studies, the characteristics of which are shown in Table 1. Five of the post hoc analyses were pooled analyses of aripiprazole, and are not discussed further. ^{23–27} The sixth post hoc analysis reported self-rated functioning data from a risperidone study in more detail than in the primary manuscript. ²⁸ No sources of bias were identified in individual studies.

The only self-rated functioning scale used in the included studies was the Sheehan Disability Scale (SDS),^{49–51} which was always a secondary or exploratory outcome. The SDS comprises three visual analog scales on which patients self-rate the extent to which symptoms have disrupted their: 1) work/studies (including paid and unpaid volunteer work and training); 2) social life or leisure activities; and 3) family life or home responsibilities. Each of these items is scored from 0 (not at all) to 10 (extremely). Patients can skip the work/studies item if they have not worked/studied in the last week for reasons unrelated to their disorder (eg, retirement); the instructions are unclear for patients who have stopped working because of their depression. The majority of studies calculated the SDS total score, obtained by summing the scores for the three items (range 0 to 30). ^{37–43,45,46,48}

	201	3
•	0	, י
	01100110	5
٠	Ċ	5
	LTION I	ב ב
•	Toriot	
	haracteri	3
(_)
•	٥)
	2	3

Reference	Adjunctive	Adjunctive	Randomized	Inadequate	Inadequate response definition	nse definition	Primary	Setting	Source of
(ClinicalTrials. gov identifier)	treatment	treatment duration	patients	response to (current episode)	Historical ADT ^a	Prospective/ongoing ADT ^b	efficacy measure)	funding
Berman et al, 2007 ²⁹ (NCT00095823)	Aripiprazole 2–20 mg	6 weeks	362	1–3 historical ADTs + 1 prospective SSRI/SNRI	<50% improved on ATRQ	<50% reduction in HAM-D ₁ , total score from start of prospective treatment to week 8; HAM-D ₁ , total score \geq 14 and CC1 Long > 2 a mod 8.	MADRS	24 outpatient sites in the US	Bristol-Myers Squibb and Otsuka
Marcus et al, 2008³º (NCT00095758)	Aripiprazole 2–20 mg	6 weeks	381	l–3 historical ADTs + l prospective SSRI/SNRI	<50% improved on ATRQ	$<$ 50% reduction in HAM-D ₁₇ total score from start of prospective treatment to week 8; HAM-D ₁₇ total score \ge 14 and CGLI score \ge 3 at week 8.	MADRS	36 outpatient sites in the US	Bristol-Myers Squibb and Otsuka
Berman et al, 2009 ³¹ (NCT00105196)	Aripiprazole 2–20 mg	6 weeks	349	I–3 historical ADTs + I prospective SSRI/SNRI	<50% improved on ATRQ	<50% reduction in HAM-D ₁₇ total score from start of prospective treatment to week 8; HAM-D ₁₇ total score ≥ 14 at week 8; CGI-I score ≥ 3 at weeks 6 and 8	MADRS	36 outpatient sites in the US	Bristol-Myers Squibb and Otsuka
Kamijima et al, 2013 ³² (NCT00876343)	Aripiprazole 3 mg/3–15 mg	6 weeks	586	I–3 historical ADTs + I prospective ADT	Not specified	<50% reduction in HAM-D ₁₇ total score from start of prospective treatment to week 8; HAM-D ₁₇ total score \geq 14 and CGI-I score \geq 3 at week 8°	MADRS	169 sites in Japan	Otsuka
Quiroz et al, 2016 ³³ (NCT01437657)	Basimglurant MR 0.5 mg/1.5 mg	6 weeks	333	I–3 historical ADTs (I ongoing SSRI/ SNRI)	Investigator judgment	MADRS total score \geq 25 and CGI-S score \geq 4 at screening ^d	MADRS	59 outpatient sites in Chile, Europe, Japan, Mexico, and the US	Hoffmann-La Roche
Thase et al, 2015 ³⁴ (NCT01360645)	Brexpiprazole 2 mg	6 weeks	379	I–3 historical ADTs + I prospective SSRI/SNRI	<50% improved on ATRQ	<50% reduction in HAM-D ₁₇ total score from start of prospective treatment to week 8; HAM-D ₁₇ total score \geq I4 at week 8; <50% reduction in MADRS total score from start of prospective treatment to weeks 2, 4, 6, and 8; CGI-I score \geq 3 at weeks 2, 4, 6, and 8°	MADRS	59 outpatient sites in Canada, Europe, and the US	Otsuka and Lundbeck
Thase et al, 2015 ³⁵ (NCT01360632)	Brexpiprazole I mg/3 mg	6 weeks	677	I–3 historical ADTs + I prospective SSRI/SNRI	<50% improved on ATRQ	<59% reduction in HAM-D ₁₇ total score from start of prospective treatment to week 8; HAM-D ₁₇ total score \geq I4 at week 8; <50% reduction in MADRS total score from start of prospective treatment to weeks 2, 4, 6, and 8; CGI-I score \geq 3 at weeks 2, 4, 6, and 8°	MADRS	92 outpatient sites in Canada, Europe, Russia, and the US	Otsuka and Lundbeck
Fava et al, 2016³6 (NCT01500200)	Buprenorphine + samidorphan 2 mg + 2 mg/8 mg + 8 mg	Stage 1: 5 weeks; stage 2: 5 weeks ^f	142	I–2 historical ADTs (I ongoing)	<50% improved on ATRQ	For entry into stage 2!: <50% reduction in HAM-D ₁ , total score from start of stage I to week 5; HAM-D ₁ , total score \geq I4 at week 5	HAM-D ₁₇	31 sites in the US	Alkermes

(Continued)

Durgam et al, 2016 ³⁷ (NCT01469377)	Cariprazine 1–2 mg/ 2–4.5 mg	8 weeks	618	I–2 historical ADTs (1 ongoing)	Not specified	ATHF resistance rating ≥ 3 with global confidence ≥ 3 at screening $^\circ$	MADRS	76 outpatient sites in Europe and the US	Forest Laboratories (Allergan) and Gedeon Richter
Fava et al, 2015 ³⁸ (NCT01098240)	CP-601,927 2–4 mg	6 weeks	162	1–3 historical ADTs + 1 prospective SSRI	<50% improved on ATRQ	<50% reduction in HAM-D ₁ total score from start of prospective treatment to week 8; HAM-D ₁ total score \geq 16 at week 8	MADRS	25 outpatient sites in the US	Pfizer
Vieta et al, 2014³9 (NCT01157078, NCT01180400)	Dexmecamylamine 2–8 mg (2 studies)	8 weeks	319, 295	0–I historical ADTs + I prospective SSRI/SNRI	Not specified	<50% reduction in HAM-D ₁ , total score from start of prospective treatment to week 8; HAM-D ₁ , total score \ge 16 and CGI-S score \ge 4 at week 8	MADRS	70 outpatient sites in India and the US; 80 outpatient sites in Europe	AstraZeneca and Targacept
Möller et al, 2015 ⁴⁰ (NCT01153347, NCT01197508)	Dexmecamylamine 0.2 mg/1 mg/2 mg/ 4 mg/8 mg (across 2 studies)	8 weeks	641, 696	0–1 historical ADTs + 1 prospective SSRI/SNRI	Not specified	<50% reduction in HAM-D ₁₇ total score from start of prospective treatment to week 8; HAM-D ₁₇ total score \geq 16 and CGI-S score \geq 4 at week 8	MADRS	114 outpatient sites in India and the US; 132 outpatient sites in Europe, South Africa, and South	AstraZeneca and Targacept
Tummala et al, 2015 ⁴¹ (NCT01152554)	Dexmecamylamine 2–8 mg	52 weeks	813 (769 de novo ⁸)	0–I historical ADTs + I prospective SSRI/SNRI	Not specified	De novo patients%. HAM-D $_{17}$ total score \geq 10 and CGI-S score \geq 3 at week 6 of prospective treatment	Not applicable (safety)	America I I 5 outpatient sites in the US	AstraZeneca
Ball et al, 2014 ⁴² (NCT00840034)	Edivoxetine 6–18 mg	8 weeks	227	I ongoing SSRI	Not applicable	Investigator judgment at screening; <25% reduction in QIDS-SR ₁₆ score from start of adjunctive placebo phase to week 2	MADRS	25 outpatient sites in the US	Eli Lilly and Company
Ball et al, 2016 ⁴³ (NCT01173601, NCT01187407, NCT01185340)	Edivoxetine 6 mg/12 mg/12–18 mg/18 mg (across 3 studies)	8 weeks	701, 689, 449	l ongoing SSRI	Not applicable	Investigator judgment at screening and start of adjunctive placebo phase; $<25\%$ reduction in MADRS total score from start of adjunctive placebo phase to week 3; MADRS total score ≥ 14 at week 3	MADRS	Multiple outpatient sites in Australia, Europe, Japan, Russia, South Africa, and the US	Ei Lilly and Company
Barbee et al, 2011 ⁴⁴ (NCT00901407)	Lamotrigine 100–400 mg	10 weeks	96	≥1 historical ADT + 1 prospective paroxetine/ paroxetine CR	Not specified	$HAM-D_{17}$ total score ≥ 1.5 at week 8 of prospective treatment	MADRS	19 sites in the US	GlaxoSmithKline
Sanacora et al, 2017 ⁴⁵ (NCT01482221)	Lanicemine 50 mg/100 mg (IV regimen)	12 weeks (primary analysis at week 6)	302	≥ I historical ADT (I ongoing)	Investigator judgment	Investigator judgment	MADRS	49 outpatient sites in Chile, Europe, South Africa, and the US	AstraZeneca
Richards et al, 2016 ⁴⁶ (NCT01436149, NCT01436162)	Lisdexamfetamine dimesylate 20–70 mg (2 studies)	8 weeks	404, 426	0–2 historical ADTs + 1 prospective SSRI/SNRI	Not specified	<50% reduction in MADRS total score from start of prospective treatment to week 8; MADRS total score \geq 18 at week 8	MADRS	76 sites in Canada, Europe, Mexico and the US; 94 sites in Europe, South Africa, and the US	Shire

ColinicalTrials. treatment durationtreatment durationresponse to durationHistorical DDTa durationHistorical DDTa durationHistorical durationHistorical durationHistorical durationHistorical durationHistorical durationHistorical durationHistorical durationHistorical durationHinvestigator duration in HAM-D ₁ total score ≥ 18 at duration in HAM-D ₁ total score ≥ 18 at duration	Reference	Adjunctive	Adjunctive	Adjunctive Randomized Inadequate	Inadequate	Inadequate response definition	onse definition	Primary Setting	Setting	Source of
Olanzapine 8 weeks 605 (pooled) I historical Investigator <25% reduction in HAM-D ₁₇ total score MADRS Multiple outpatient ADT+I prospective idgment from start of prospective treatment to total stess in Canada and the US week 8; HAM-D ₁₇ total score ≥18 at score from start of prospective treatment to total stess in Canada and the US week 8; ≤15% reduction in HAM-D ₁₇ total score ≥18 at score from week 7 to week 8; ≤15% reduction in HAM-D ₁₇ total score ≥20 at stess in Canada and the US score in HAM-D ₁₇ total score ≥20 at score ≥20 at stess in the US stess in Canada and the US score ≥20 at score ≥20 at stess in the US score ≥20 at score ≥20 at stess in the US stess in Canada and the US score ≥20 at score ≥20 at stess in the US score ≥20 at score ≥	(ClinicalTrials. gov identifier)	treatment	treatment duration	patients	response to (current episode)	Historical ADT ^a	Prospective/ongoing ADT⁵	efficacy measure		funding
6–18 mg (as OFC; ADT+1 prospective indgment from start of prospective treatment to total sites in Canada sites in Canada fluoxetine fluoxetine fluoxetine heek 8; HAM-D ₁₇ total score \geq 18 at and the US score from week 8; \leq 15% reduction in HAM-D ₁₇ total score \geq 10 mg oing ADT Not applicable CGI-S score \geq 20 at HAM-D ₁₇ 75 outpatient week 4 of prospective phase total sites in the US sites in the US	Thase et al,	Olanzapine	8 weeks	605 (pooled)	historical	Investigator	<25% reduction in HAM-D ₁₇ total score		Multiple outpatient	Eli Lilly and
12 Studies pooled) fluoxetine week 8; HAM-D $_{17}$ total score \geq 18 at and the US week 8; \leq 15% reduction in HAM-D $_{17}$ total score \geq 15 m and the US score from week 7 to week 8 sore from week 7 to week 8 sore \geq 15 outpatient length on 15 m longoing ADT	2007 ⁴⁷	6–18 mg (as OFC;			ADT + I prospective	judgment	from start of prospective treatment to		sites in Canada	Company
Signature A seeks 274 I ongoing ADT Not applicable CGI-S score \geq 4 and CDS score \geq 20 at HAM-D ₁₇ 75 outpatient legal week 4 of prospective phase total sites in the US	(NCT00035321)	2 studies pooled)			fluoxetine		week 8; HAM-D $_{17}$ total score \ge 18 at week 8; \le 15% reduction in HAM-D $_{17}$ total	_	and the US	
Risperidone 6 weeks 274 I ongoing ADT Not applicable CGI-S score ≥ 4 and CDS score ≥ 20 at HAM-D ₁₇ 75 outpatient I-2 mg week 4 of prospective phase total sites in the US							score from week 7 to week 8			
I–2 mg veek 4 of prospective phase total sites in the US	Mahmoud	Risperidone	6 weeks	274	I ongoing ADT	Not applicable	CGI-S score ≥4 and CDS score ≥20 at	HAM-D ₁₇	75 outpatient	Ortho-McNeil-
	et al, 2007 ⁴⁸	I–2 mg					week 4 of prospective phase	total	sites in the US	Janssen

Notes: "Historical" describes an ADT that was discontinued prior to study enrollment." "Prospective" describes an ADT that was initiated during the study, whereas "ongoing" describes an ADT that was initiated prior to study enrollment comparison with the other aripiprazole studies, we assume this is an error. "In addition, where available, a patient-rated MADRS Carroll Depression Scale; CGI-I/S, Clinical MR, modified release; OFC olarzapine-fluoxetine combination; QIDS-SR₁₆, 16-item Quick Inventory of Depressive Symptomatology (Self-Report); SNR1, serotonin–norepinephrine reuptake inhibitor; SSR1, selective serotonin reuptake inhibitor and was continued during the study. The manuscript states that any one of these criteria is sufficient; however, by Global Impressions - Improvement/Severity total score ≥23

If the work/studies item was unrated, as was the case for 10%–35% of patients (in studies where patient numbers by item were available), it was generally unclear whether these patients were excluded from the SDS total score, or whether the SDS total score was calculated by imputing the mean of the other two items in place of the work/studies item. One study calculated the SDS sum of items 2 and 3 score (range 0 to 20), since a large proportion of patients (25%–35%) had no work/studies rating.³³ Six studies calculated the SDS mean score, obtained by taking an average of the item scores for all items that were rated (range 0 to 10).^{29–32,34,35} Finally, four studies did not specify how SDS scores were calculated.^{36,44,47}

As can be seen in Table 1, 13 different adjunctive agents were used across the studies, of which five were second-generation antipsychotics. All were short-term studies (6–12 weeks), except for one 52-week study. The minimum number of ADTs (historical plus prospective) to which patients were required to show inadequate response prior to randomization ranged from one to two. Definitions of inadequate response varied, from "investigator judgment", to strict definitions with qualifying scores on multiple rating scales at multiple time points.

Efficacy

Each of the second-generation antipsychotics (aripiprazole, brexpiprazole, cariprazine, olanzapine as OFC [pooled data], and risperidone) had at least one dose that met the primary efficacy endpoint of their respective studies (improvement of depressive symptoms, measured by either Montgomery–Åsberg Depression Rating Scale total score or 17-item Hamilton Depression Rating Scale total score; Table 1).^{29–32,34,35,37,47,48} The combination of buprenorphine and samidorphan also met its primary efficacy endpoint at one dose.³⁶ All other agents failed to meet the primary efficacy endpoint of their respective studies, and were therefore failed or negative studies.

A summary of the SDS results from the included studies is given in Table 2. Baseline SDS scores were similar between treatment groups in each study. All groups (active and control) improved numerically from baseline to endpoint, as measured by the SDS total or SDS mean. Most active treatments showed a numerically greater improvement than placebo (except for dexmecamylamine, which had inconsistent results). However, the majority of agents, and studies, failed to show a statistically significant benefit versus placebo on the SDS total or SDS mean. Only four agents demonstrated efficacy (p<0.05 versus placebo) on the SDS total or SDS mean in at least one study: aripiprazole, brexpiprazole,

 Table 2
 Change from baseline to endpoint in self-rated functioning scale scores (for the population who received ≥ I dose of randomized treatment and had ≥ I post-baseline efficacy
 evaluation)

(H	0										
Kererence	I reatment	SDS summary score	ary score				SDS Work/studies	ndies	SDS social life	e	SUS ramily life	_ -
	arm	Baseline		Chang	Change to endpoint		Change to endpoint	ndpoint	Change to endpoint	dpoint	Change to endpoint	Idpoint
		u	Mean (SD)	٦	Mean (SE)	p-value	Mean (SE)	p-value	Mean (SE)	p-value	Mean (SE)	p-value
						versus		versus		versus		versus
Aripiprazole		SDS mean										
Berman	Placebo	164	$5.4 (0.2)^a$	164	-0.7 (0.2)		-0.7 (0.3) ^b		-0.8 (0.2)		-0.5 (0.2)	
et al, 2007 ²⁹	2-20 mg	167	5.7 (0.2) ^a	167	-1.1 (0.2)	0.055	-0.7 (0.2)°	0.98	-I.4 (0.2)	0.030	-1.1 (0.2)	0.017
Marcus et al,	Placebo	168	5.4 (0.2) ^a	891	-0.7 (0.2)		−0.4 (0.2) ^d		-0.6 (0.2) ^e		-1.0 (0.2) ^e	
200830	2-20 mg	180	5.1 (0.2) ^a	180	-I.3 (0.2)	0.012	$-0.5 (0.2)^{f}$	0.62	-I.4 (0.2) ^g	0.002	-I.8 (0.2) ^g	0.002
Berman	Placebo	091	5.9 (0.2) ^a	091	-0.8 (0.2)		-0.7 (0.3) ^h		-0.7 (0.2)		-0.8 (0.2)	
et al, 2009 ³¹	2-20 mg	091	5.7 (0.2) ^a	091	-1.2 (0.2)	0.075	-0.8 (0.3)	0.79	-I.2 (0.2)	0.052	-I.4 (0.2)	0.037
Kamijima	Placebo	195	5.3 (0.1) ^a	193	-0.5 (0.1)		-0.4 (0.1)		-0.6 (0.1)		-0.3 (0.1)	
et al, 2013^{32}	3 mg	197	5.0 (0.1) ^a	197	-1.0 (0.1)	0.001	(1.0) 6.0-	0.003	-1.1 (0.1)	0.003	(0.1)	0.003
	3–15 mg	194	5.0 (0.1) ^a	193	-1.0 (0.1)	<0.001	-1.0 (0.1)	<0.001	-1.2 (0.1)	<0.001	-0.9 (0.1)	0.003
Basimglurant MR	MR	SDS sum o	SDS sum of items 2 and 3									
Quiroz et al,	Placebo	i601	13.2 (3.7)	601	-5.1 (0.6)							
201633	0.5 mg	112	13.8 (4.0)	112	-5.1 (0.6)	0.94						
	I.5 mg	<u>-</u>	13.5 (3.6)	Ξ	-6.4 (0.6)	60.0						
Brexpiprazole	40	SDS mean										
Thase et al,	Placebo	161	6.3 (2.1)	178	-0.9 (0.2)		-1.0 (0.2)		-I.0 (0.2)		-0.7 (0.2)	
201534	2 mg	188	6.0 (2.0)	175	-1.4 (0.2)	0.035	-1.1 (0.2)	19.0	-I.6 (0.2)	0.022	-I.3 (0.2)	0.011
Thase et al,	Placebo	221 ^k	5.6 (1.9)	203	-0.8 (0.2)		-0.7 (0.2)		-0.9 (0.2)		-0.8 (0.2)	
201535	l mg	226 ^k	5.9 (2.0)	211	-1.3 (0.2)	910.0	-1.1 (0.2)	0.082	-I.3 (0.2)	0.035	-I.3 (0.2)	610.0
	3 mg	230 ^k	5.7 (2.2)	213	-1.3 (0.2)	0.019	-0.9 (0.2)	0.30	-I.4 (0.2)	0.028	-I.4 (0.2)	0.0077
Buprenorphine +	ie +											
samidorphan												
Fava et al,	Placebo											
201636	2 mg + 2 mg					SZ						
	8 mg + 8 mg					SZ						
Cariprazine		SDS total										
Durgam	Placebo	264	18.5 (4.7)	264	-6.6 (0.5)						-2.3	
et al, 2016 ³⁷	I–2 mg	273	18.7 (4.7)	273	-7.7 (0.5)	0.24		SZ		SN		NS
	2–4.5 mg	271	18.8 (4.8)	271	-8.0 (0.5)	0.11		NS		NS	-2.9	0.014
CP-601,927		SDS total										
Fava et al,	Placebo	85	18.5 (5.3)	82	-5.7 (8.2)							
201538	2–4 mg	77	19.0 (5.9)	77	-6.6 (8.0)	SN		NS		NS		NS
												(Continued)

Change to enclopint Change to encloped Change to enclop	Reference	Treatment	SDS sumn	SDS summary score				SDS work/studies	udies	SDS social life	ife	SDS family life	ife
Parameter Para		arm	Baseline		Chang	ge to endpoint		Change to e	ndpoint	Change to e	ndpoint	Change to e	ndpoint
wreying versus versus versus - Placebo 151 -5.5 (0.7) -1.7 (0.2)* -1.9 (0.2) 0.33 - Placebo 143 -6.1 (0.7) -0.2 (0.2)* -1.9 (0.2) 0.33 - Lebrago 143 -6.1 (0.7) -0.2 (0.2)* 0.2 -1.9 (0.2) 0.33 - Lebrago 143 -6.1 (0.7) 0.42 -2.0 (0.2)* 0.7 -1.9 (0.2) 0.33 - Lebrago 1 1.3 -6.8 (0.5) 0.96 -2.0 (0.2) 0.7 -1.9 (0.2) 0.7 - Lebrago 1 1.0 -2.8 (0.5) 0.42 -1.7 (0.2) 0.7 -1.9 (0.2) 0.7 - Long 1 1.0 -2.5 (0.6) 0.42 -1.7 (0.2) 0.7 -1.9 (0.2) 0.7 - Long 1 1.0 -2.1 (0.2) 0.42 -1.9 (0.2) 0.7 -1.9 (0.2) 0.7 - Long 1 1.0 -2.2 (0.2) 0.42 -1.7 (0.2) 0.7 -1.9 (0.2) 0.7 - Long - 1.0 -2.0 (0.2)			_	Mean (SD)	_	Mean (SE)	p-value	Mean (SE)	p-value	Mean (SE)	p-value	Mean (SE)	p-value
Precedo SDS total							versus		versus		versus		versus
Pincebo 151 -5.5 (0.7) -1.7 (0.2)* -1.9 (0.2)	Dexmecamy	amine	SDS total										
2-8 mg 1-3 mg	Vieta et al,	Placebo			151	-5.5 (0.7)		-1.7 (0.2) ^m		-1.9 (0.2)		-2.0 (0.3)	
Placebo Placebo 142 -58 (6.5) -18 (0.2) -19 (0.2) -1	201439	2–8 mg			143	-6.1 (0.7)	0.42	$-2.0 (0.2)^{n}$	0.21	-I.9 (0.2)	0.93	-2.1 (0.3)	97.0
1.8 2.8 mg 1.3 2.8 (0.5) 0.96 -2.2 (0.2) 0.76 -1.9 (0.2) 0.76 -1.9 (0.2) 0.76 -1.9 (0.2) 0.76 -1.9 (0.2) 0.76 -1.9 (0.2) 0.76 -1.9 (0.2) 0.76 -1.9 (0.2) 0.76 -1.9 (0.2) 0.75		Placebo			142	-5.8 (0.5)		−1.8 (0.2)°		-I.9 (0.2)		-2.0 (0.2)	
Hacebo 153		2–8 mg			138	-5.8 (0.5)	96.0	-2.2 (0.3) ^p	0.20	-2.0 (0.2)	92'0	-1.9 (0.2)	0.62
Hing Hing High High	Möller et al,	Placebo			153	-4.9 (0.6)		−1.7 (0.2) ^q		-I.9 (0.2)		-1.6 (0.2)	
Hag 4 mg Hag 55 (6.6) 0.49 -18 (0.2) 0.97 Placebo 172 -1.1 (0.2) 0.79 -1.9 (0.2) 0.24 Placebo 172 -1.1 (0.6) 0.18 -2.0 (0.2) 0.79 -1.1 (0.2) 0.24 Comman 172 -1.1 (0.6) 0.18 -2.0 (0.2) 0.24 Placebo 171 -6.1 (0.6) 0.18 -2.0 (0.2) 0.24 Placebo 171 -6.1 (0.6) 0.24 -2.0 (0.2) 0.24 Placebo 171 -1.1 (0.6) 0.27 -1.9 (0.2) 0.44 Placebo 171 17.9 (6.9) 6.8 -2.6 0.039 -1.9 (0.1) 0.10 -2.1 (0.2) 0.44 Placebo 171 17.9 (6.9) 6.8 -2.6 0.039 -1.9 (0.10 -2.1 0.046 Placebo 171 17.9 (6.9) 6.3 -6.0 0.039 -1.9 (0.10 -2.1 0.046 Placebo 171 2.40 4.5 (0.4) NS -1.9 (0.10 -2.1 0.046 Placebo 171 2.40 4.5 (0.4) NS -1.9 (0.10 -2.1 0.046 Placebo 171 2.40 4.5 (0.4) NS -1.9 (0.10 -2.1 0.046 Placebo 2.50 2.31 -2.4 (0.4) NS -2.5 (0.4) NS Placebo 2.50 2.51 -2.5 (0.4) NS -2.5 (0.4	201540	l mg			152	-5.5 (0.6)	0.42	-1.7 (0.2) ^r	0.95	-I.9 (0.2)	0.92	-I.9 (0.2)	0.21
Range Rang		4 mg			148	-5.5 (0.6)	0.49	-I.8 (0.2) ^s	0.87	-I.9 (0.2)	0.97	-I.8 (0.2)	0.34
Placebo Placebo 172 -7.1 (0.6) -2.1 (0.2) -2.4 (0.2) 0.25 2 mg		8 mg			149	-4.5 (0.7)	09:0	-1.7 (0.2) ^t	0.79	-I.6 (0.2)	0.24	-I.6 (0.2)	0.97
171 -61 (0.6) 0.18 -2.0 (0.2) 0.25		Placebo			172	-7.1 (0.6)		-2.1 (0.2) ^u		-2.4 (0.2)		-2.3 (0.2)	
Sumg 1.88 0.63 0.63 0.18 0.27 0.45 0.20 0.50 0.50 Sumg 1.68 -6.3 (0.6) 0.27 -1.9 (0.2) 0.66 -2.2 (0.2) 0.44 0.54 Sumg 1.63 -6.2 (0.6) 0.27 -1.9 (0.2) 0.66 -2.2 (0.2) 0.44 Interplace SDS total 1.9 (6.8) 6.8 -2.6 0.039 -1.9 0.10 -2.1 0.046 0.44 Interplace P. 6.9 1.7 (6.8) 6.8 -2.6 0.039 -1.9 0.10 -2.1 0.046 0.44 Interplace P. 6.9 1.149 E. 1.149		0.2 mg			171	-6.1 (0.6)	0.18	-2.0 (0.2)	16.0	-2.1 (0.2)	0.25	-2.0 (0.2)	0.17
Shing Shin		2 mg			891	-6.3 (0.6)	0.33	-I.8 (0.2) ^w	0.45	-2.2 (0.2)	0.50	-2.1 (0.2)	0.40
Placebo 2–8 mg 185 -7.0 (7.3)		8 mg			163	-6.2 (0.6)	0.27	-I.9 (0.2)×	99.0	-2.2 (0.2)	0.44	-2.0 (0.2)	0.20
Flacebo	Tummala	Placebo			70	-7.4 (7.5)							
ine SDS total ¹ = -26 = -0.4 = -1.0 = -1	et al, 2015 ⁴¹	2–8 mg			185	-7.0 (7.9)							
Placebo 71k 192 (6.8) 68 -2.6 -0.4 -1.0 -1.0 6-18 mg 67k 17.9 (6.9) 63 -6.0 0.039 -1.9 0.10 -2.1 0.046 12 mg 67k 17.9 (6.9) 63 -6.0 0.039 -1.9 0.10 -2.1 0.046 12 mg E: 1,149k E: 18.1 (5.8) 231 -5.4 (0.4) NS NS 12 mg (pooled) (pooled) 230 -5.3 (0.4) NS NS 12-18 mg 226 -6.3 (0.4) NS NS 12-18 mg 232 -5.3 (0.4) NS NS 12-18 mg 230 -4.5 (0.5) NS NS 10-400 mg 230 -4.5 (0.5) NS NS 10-400 mg 101 -7.1 (1.0) 0.89 100 mg 100	Edivoxetine		SDS totaly										
Hander	Ball et al,	Placebo	71k	19.2 (6.8)	89	-2.6		-0.4		0.1-		0.1-	
al, Placebo P: 690* P: 18.1 (6.1) 240 -4.5 (0.4) 12 mg E: 1,149* E: 18.2 (5.8) 231 -5.4 (0.4) 18 mg (pooled) (pooled) 230 -5.3 (0.4) Placebo 226 -6.3 (0.4) 12-18 mg 226 -6.3 (0.4) 12-18 mg 219 -4.4 (0.5) 12-18 mg 230 -4.5 (0.5) rigine et al, Placebo Indo -400 mg SDS total a Placebo Indo mg Indo mg Indo mg	2014 ⁴²	6–18 mg	4∠9	17.9 (6.9)	63	-6.0	0.039	6. I-	0.10	-2.1	0.046	6.I-	0.11
12 mg E: 1,149 ^k E: 18.2 (5.8) 231 -5.4 (0.4) 18 mg (pooled) (pooled) 230 -5.3 (0.4) 18 ng (pooled) 231 -4.3 (0.4) 12 - 18 mg 226 -6.3 (0.4) 12 - 18 mg 232 -5.3 (0.4) 12 - 18 mg 232 -5.3 (0.4) 12 - 18 mg 232 -4.5 (0.5) 12 - 18 mg 230 -4.5 (0.5) 13 Placebo 230 -4.5 (0.5) 100 - 400 mg 250 total 27 -6.9 (1.0) 100 mg 100 mg 100 -6.9 (1.0) 100 mg 100 -6.9 (1.0)	Ball et al,	Placebo	P: 690 ^k	P: 18.1 (6.1)	240	-4.5 (0.4)							
18 mg (pooled) (pooled) 230 -5.3 (0.4) Placebo 226 -6.3 (0.4) 12-18 mg 226 -6.3 (0.4) 12-18 mg 232 -5.3 (0.4) Placebo 219 -4.4 (0.5) 12-18 mg 232 -5.3 (0.4) 12-18 mg 232 -4.5 (0.5) 12-18 mg 230 -4.5 (0.5) 12-18 mg 230 -4.5 (0.5) 100-400 mg 20 20 20 100-400 mg 20 20 20 100 mg 20 20 100 mg 20 20 20 100 mg 20 20 100 mg 20 20 20 100 mg 20 20 20 100 mg 20 20	201643	I2 mg	E: 1,149 ^k	E: 18.2 (5.8)	231	-5.4 (0.4)	SZ						
Placebo 231 -4.3 (0.4) 6 mg 226 -6.3 (0.4) 12–18 mg 232 -5.3 (0.4) Placebo 219 -4.4 (0.5) 12–18 mg 230 -4.5 (0.5) 12–18 mg 230 -4.5 (0.5) 12–18 mg 230 -4.5 (0.5) 13–18 mg 230 -4.5 (0.5) 13		I8 mg	(polood)	(booled)	230	-5.3 (0.4)	SZ						
6 mg 226 -6.3 (0.4) 12-18 mg 232 -5.3 (0.4) Placebo 219 -4.4 (0.5) 12-18 mg 230 -4.5 (0.5) 219 -4.4 (0.5) 210 -4.0 (0.5) 230 -4.5 (0.5) 230 -4.5 (0.5) 230 -4.5 (0.5) 230 -4.5 (0.5) 240 -4.5 (0.5) 250 -		Placebo			231	-4.3 (0.4)							
12–18 mg		9 mg			226	-6.3 (0.4)	≥0.05						
Placebo 219 -4.4 (0.5) 12–18 mg 230 -4.5 (0.5) ringine et al, Placebo mine (IV regimen) SDS total a Placebo 97 -6.9 (1.0) 100 mg 100 mg 100 mg 100 mg -6.9 (1.0)		12–18 mg			232	-5.3 (0.4)	SZ						
12–18 mg		Placebo			219	-4.4 (0.5)							
et al, Placebo et al, 100–400 mg mine (IV regimen) SDS total 97 –6.9 (1.0) 1745 50 mg 100 mg 100 mg 100 –6.9 (1.0)		12–18 mg			230	-4.5 (0.5)	SZ						
et al, Placebo 100–400 mg mine (IV regimen) SDS total 97 –6.9 (1.0) 1745 50 mg 100 mg 100 –6.9 (1.0)	Lamotrigine												
100–400 mg mine (IV regimen) SDS total a Placebo 97 –6.9 (1.0) 100 mg 100 mg	Barbee et al,	Placebo											
E (IV regimen) SDS total 97 –6.9 (1.0) Placebo 100 mg 100 –6.9 (1.0)	201144	100-400 mg					NS						
Placebo 97 –6.9 (1.0) 50 mg 101 –7.1 (1.0) 100 mg –6.9 (1.0)	Lanicemine (IV regimen)	SDS total										
50 mg 100 mg 100 — 6.9 (1.0)	Sanacora	Placebo			4	-6.9 (1.0)							
100 –6.9 (1.0)	et al, 2017 ⁴⁵	50 mg			101	-7.1 (1.0)	0.89						
		100 mg			00	-6.9 (1.0)	0.99						

Lisdexamfetamine	tamine	SDS total										
dimesylate												
Richards	Placebo	198	15.6 (5.7)	164	$-4.3 (-5.3, -3.3)^{z}$							
et al, 2016 ⁴⁶	20-70 mg	861	15.9 (6.0)	162	$-4.7 (-5.6, -3.7)^{z}$	0.58		SZ		NS		NS
-nic	Placebo	213	16.8 (5.8)	189	$-4.3 (-5.2, -3.4)^{z}$							
Disc	20–70 mg	209	16.8 (5.3)	180	$-4.9 (-5.8, -4.0)^{z}$	0.35		SZ		NS		NS
Olanzapine (as OFC;	(as OFC;											
2 studies pooled)	(paloc											
Thase et al,	Fluoxetine			203					-1.1 (2.6)		-I.2 (2.6) ¹	
2007 ⁴⁷	6–18 mg			198				0.46	-I.6 (2.8)	0.027	-1.7 (2.7)	0.047
Risperidone	6	SDS total										
Mahmoud	Placebo	113	20.2 (0.6) ^a	101	-3.9		-0.8		-1.5		4. -	
et al, 2007 ⁴⁸	I-2 mg	011	19.9 (0.6) ^a	001	-7.6	<0.00	7.1-	0.12	-2.9	<0.001	-2.5	<0.001
Pandina												
et al, 2009 ²⁸												

"n=170. fn=142. 8n=181. hn=126. hn=115. For the population who received \geq 1 dose of randomized treatment. For the randomized population. SD. "n=134. Abbreviations: E, edivoxetine; IV, intravenous; MR, modified release; NS, not statistically significant (and p-value not reported); OFC, olanzapine-fluoxetine combination; P, placebo; SDS, Sheehan Disability Scale. n=115.9=137.7=132.1=134.1=134.1=137.7=141.1=140.7=126.1f the world/studies item was missing, the mean of the other two items was imputed to calculate the total score. 195% CI. bn=130. cn=127. dn=136. **Notes:** Highlighted cells indicate $\rho \le 0.05$ versus placebo. ^aSE. n=124. °n=120.

edivoxetine, and risperidone. A large variation in placebo response was seen between studies.

Considering the individual SDS items, only one agent (aripiprazole) in one study had a statistically significant benefit on the SDS work/studies item. The study in question (Kamijima et al)³² investigated two doses (3 mg and 3–15 mg) of adjunctive aripiprazole in a 6-week randomized treatment phase, and both doses showed a benefit on the work/studies item. Three other studies of aripiprazole were included in the review; none of these showed efficacy on the SDS work/studies item, despite having almost identical designs to Kamijima et al.

On the SDS social life item, aripiprazole, brexpiprazole, edivoxetine, olanzapine (as OFC), and risperidone showed a benefit over placebo (p<0.05) in at least one study. On the SDS family life item, a benefit (p<0.05) was observed for aripiprazole, brexpiprazole, cariprazine, olanzapine (as OFC), and risperidone in at least one study.

Discussion

This review of 26 randomized placebo-controlled studies showed that, of the 13 adjunctive agents identified, only aripiprazole, brexpiprazole, edivoxetine, and risperidone statistically significantly improved functioning versus placebo, as measured by the SDS total or mean, in patients with MDD and inadequate response to at least one ADT. In a previous meta-analysis of adjunctive second-generation antipsychotics in MDD, which was conducted prior to the availability of brexpiprazole data, only aripiprazole and risperidone were found to provide a benefit based on a composite endpoint of patient-reported functioning and quality of life. 52 Furthermore, a systematic review in patients with MDD who received ADT (but no adjunctive pharmacotherapy) found that many patients, particularly partial responders, continued to experience functional impairments after treatment, highlighting an unmet need in MDD.20

Other than edivoxetine, a selective norepinephrine reuptake inhibitor, the only agents to show statistically significant benefits on any SDS items in the present review were second-generation antipsychotics. Of these agents, aripiprazole and brexpiprazole have an indication for the adjunctive treatment of MDD, and OFC has an indication for the treatment of treatment-resistant depression (all in the US). ^{22,53,54} The development of edivoxetine as an adjunctive treatment for MDD was halted in 2013 because it failed to meet the primary endpoint in three Phase 3 studies. ⁵⁵ Indeed, more than half of the adjunctive agents identified in this review failed to meet the primary efficacy endpoint of their respective studies. Nonetheless, the findings of this review suggest that the SDS

as a whole is a useful scale to track changes in functioning among patients with inadequate response to ADTs, and that adjunctive second-generation antipsychotics or edivoxetine may improve functioning in such patients.

The SDS was the only self-rated measure of functional impairment that was used in the retrieved records. This observation is in line with a meta-analysis of second-generation antipsychotics for the adjunctive treatment of MDD.⁵² Indeed, the SDS appears to be the most widely used functioning measure in studies of MDD.20 The SDS is considered to be a reliable and valid measure of functioning impairment, originally developed in 1981 for use in treatment outcome studies in psychiatry. 49-51 To the authors' knowledge, no minimal clinically important difference has been established for the SDS. Sheehan selected the three items of work/studies. social life, and family life after reviewing other impairment instruments and consulting with patients and colleagues.⁵¹ In general, all three items, including the work/studies item, are sensitive to treatment effects across a variety of psychological disorders.⁵¹ In the present review, however, in the population of patients with MDD and inadequate response to ADTs, only aripiprazole had a statistically significant benefit on the SDS work/studies item, and this was only in one study out of four. Thus, where a benefit was observed on the SDS total or mean, this was generally driven by improvement on the social life and family life items.

There are several possible reasons for a lack of effect on the work/studies item in this population. Since the work/studies item is not rated for patients who are not working, one possibility is that the studies were underpowered to measure this item. In general, as shown in a US nationwide survey of patients with MDD, inadequate responders to ADT (based on self-reports) are less likely to be employed than responders. ¹⁴ Unfortunately, the majority of studies in the present review did not separate out the number of patients who rated each item of the SDS. Where such data were available, 10%–35% of patients across the studies with a rating on the social life and family life items did not rate the work/studies item. Thus, the power to show a difference between treatment groups was reduced for the work/studies item compared with the other items.

Nevertheless, 65%–90% of patients did rate the work/studies item, and therefore were in employment or studying. In general, studies have shown that people with depression who are in employment are less severely ill than those who are unemployed. Thus, on average, the subset of patients who rated the work/studies item may be less severely ill than the total population in each study. Meta-analyses have

investigated the question of whether or not antidepressant efficacy increases with baseline illness severity, with varying results. 58,59 If, as some have suggested, antidepressants are more efficacious in more severely ill patients, then the drugplacebo difference may be expected to be greater in the total population (ie, on the social life and family life items) than in the subset of patients in employment (ie, on the work/studies item).

Finally, it is possible that the studies were too short to show a benefit on the work/studies item. With a few exceptions, the included studies assessed functioning after 6 or 8 weeks of adjunctive treatment. In general, job performance deficits can still remain after 18 months among patients whose depressive symptoms have improved, 60 and patients with inadequate response to ADT are particularly at risk of persisting impairment.⁶¹ Thus, acute studies may not be able to detect a benefit in occupational functioning in this population of inadequate responders. Only one of the included studies was a long-term study, and, over 52 weeks, adjunctive dexmecamylamine did not show a notable difference to adjunctive placebo on the SDS total.⁴¹ However, dexmecamylamine failed as an adjunctive agent in MDD, having shown no differences to placebo on the primary efficacy outcome in four acute studies, 39,40 and thus no benefit on the SDS might be expected.

Recent literature has acknowledged the difficulty in using the work/studies item among populations with a high proportion of non-workers, and attempts have been made to modify the SDS accordingly. Sonne et al proposed a rewording of the first item from "work/studies" to "work/daily tasks", so that patients without a job could still rate the item. Similarly, Bech reported a modified version of the SDS in which the work/studies item was replaced by an overall rating, "Your daily activities all things considered".

The present review is limited because it only considered published literature, leading to a risk of publication bias (as positive studies are more likely to be published than failed or negative studies). However, even with the risk of publication bias, fewer than a third of the included studies reported a treatment benefit on the SDS. In addition, the review is limited since MEDLINE (via PubMed) was the only database searched. Nonetheless, the 26 studies identified had a consistent message, that the work/studies item is less informative in this population than the other two items.

Conclusion

The SDS, a self-rated functional measure, is informative in acute randomized placebo-controlled studies of adjunctive

therapy in patients with MDD and inadequate response to ADT. However, the item that measures work performance may be less relevant to this population than the items that measure social and family life.

Acknowledgments

This review was funded by Otsuka Pharmaceutical Development & Commercialization Inc. (Princeton, NJ, USA) and H. Lundbeck A/S (Valby, Denmark). Dr Jenny AR Muiry of Cambridge Medical Communication Ltd (Cambridge, UK) assisted with data collection. Cambridge Medical Communication Ltd also provided editorial support.

Author contributions

CPW and CE performed data collection and analysis. All authors contributed toward data analysis, drafting and revising the paper, and agree to be accountable for all aspects of the work.

Disclosure

EW and HE are employees of H. Lundbeck A/S. CW and MH are employees of Otsuka Pharmaceutical Development & Commercialization Inc. CPW and CE are employees of Cambridge Medical Communication Ltd, which received funding from Otsuka Pharmaceutical Development & Commercialization Inc. and H. Lundbeck A/S for this work. MF has received research support from, and has been a consultant or advisory board member for, all major pharmaceutical companies with drugs used in MDD. He owns the copyright for the Massachusetts General Hospital Antidepressant Treatment Response Questionnaire (ATRQ) and a number of other psychiatric rating scales. Full lifetime disclosures can be viewed online at: http://mghcme.org/faculty/faculty-detail/ maurizio fava. He received no remuneration for his involvement in this review. The authors report no other conflicts of interest in this work.

References

- American Psychiatric Association. Diagnostic and Statistical Manual of Mental Disorders. 5th ed. Arlington, VA: American Psychiatric Association; 2013.
- Kessler RC. The costs of depression. Psychiatr Clin North Am. 2012; 35(1):1–14.
- Druss BG, Hwang I, Petukhova M, Sampson NA, Wang PS, Kessler RC. Impairment in role functioning in mental and chronic medical disorders in the United States: results from the National Comorbidity Survey Replication. *Mol Psychiatry*. 2009;14(7):728–737.
- National Institute for Health and Clinical Excellence. The NICE Guideline on the Treatment and Management of Depression in Adults (Updated Edition). NICE; 2010. Available from: https://www.nice.org.uk/guidance/cg90/evidence/full-guidance-243833293. Accessed November 16, 2017.

- Nierenberg AA, DeCecco LM. Definitions of antidepressant treatment response, remission, nonresponse, partial response, and other relevant outcomes: a focus on treatment-resistant depression. *J Clin Psychiatry*. 2001;62 Suppl 16:5–9.
- Angst J, Kupfer DJ, Rosenbaum JF. Recovery from depression: risk or reality? Acta Psychiatr Scand. 1996;93(6):413

 –419.
- Fava GA, Bech P. The concept of euthymia. Psychother Psychosom. 2016;85(1):1–5.
- Zimmerman M, McGlinchey JB, Posternak MA, Friedman M, Attiullah N, Boerescu D. How should remission from depression be defined? The depressed patient's perspective. *Am J Psychiatry*. 2006;163(1):148–150.
- Evans VC, Lam RW. Assessments of functional improvement: selfversus clinician-ratings. *Medicographia*. 2014;36(4):512–520.
- Saltiel PF, Silvershein DI. Major depressive disorder: mechanism-based prescribing for personalized medicine. Neuropsychiatr Dis Treat. 2015; 11:875–888.
- McKnight PE, Kashdan TB. The importance of functional impairment to mental health outcomes: a case for reassessing our goals in depression treatment research. *Clin Psychol Rev.* 2009;29(3):243–259.
- Rush AJ, Trivedi MH, Wisniewski SR, et al. Acute and longer-term outcomes in depressed outpatients requiring one or several treatment steps: a STAR*D report. Am J Psychiatry. 2006;163(11):1905–1917.
- Mauskopf JA, Simon GE, Kalsekar A, Nimsch C, Dunayevich E, Cameron A. Nonresponse, partial response, and failure to achieve remission: humanistic and cost burden in major depressive disorder. *Depress Anxiety*. 2009;26(1):83–97.
- Knoth RL, Bolge SC, Kim E, Tran QV. Effect of inadequate response to treatment in patients with depression. Am J Manag Care. 2010;16(8): e188–e196
- American Psychiatric Association. Practice Guideline for the Treatment of Patients with Major Depressive Disorder. 3rd ed. American Psychiatric Association; 2010. Available from: http://psychiatryonline.org/pb/assets/raw/sitewide/practice_guidelines/guidelines/mdd.pdf. Accessed November 16, 2017.
- Kennedy SH, Lam RW, McIntyre RS, et al. Canadian Network for Mood and Anxiety Treatments (CANMAT) 2016 clinical guidelines for the management of adults with major depressive disorder: Section 3. Pharmacological treatments. Can J Psychiatry. 2016;61(9):540–560.
- Fava M. Pharmacological approaches to the treatment of residual symptoms. J Psychopharmacol. 2006;20(3 Suppl):29–34.
- Fava M, Rush AJ. Current status of augmentation and combination treatments for major depressive disorder: a literature review and a proposal for a novel approach to improve practice. *Psychother Psychosom*. 2006;75(3):139–153.
- Connolly KR, Thase ME. If at first you don't succeed: a review of the evidence for antidepressant augmentation, combination and switching strategies. *Drugs*. 2011;71(1):43–64.
- Sheehan DV, Nakagome K, Asami Y, Pappadopulos EA, Boucher M. Restoring function in major depressive disorder: a systematic review. *J Affect Disord*. 2017;215:299–313.
- Liberati A, Altman DG, Tetzlaff J, et al. The PRISMA statement for reporting systematic reviews and meta-analyses of studies that evaluate health care interventions: explanation and elaboration. *PLoS Med*. 2009;6(7):e1000100.
- Symbyax® (olanzapine and fluoxetine) capsules for oral use [prescribing information]. Indianapolis, IN: Lilly USA, LLC; 2017.
- Thase ME, Trivedi MH, Nelson JC, et al. Examining the efficacy of adjunctive aripiprazole in major depressive disorder: a pooled analysis of 2 studies. *Prim Care Companion J Clin Psychiatry*. 2008;10(6): 440–447
- Trivedi MH, Corey-Lisle PK, Guo Z, Lennox RD, Pikalov A, Kim E. Remission, response without remission, and nonresponse in major depressive disorder: impact on functioning. *Int Clin Psychopharmacol*. 2009;24(3):133–138.
- Steffens DC, Nelson JC, Eudicone JM, et al. Efficacy and safety of adjunctive aripiprazole in major depressive disorder in older patients: a pooled subpopulation analysis. *Int J Geriatr Psychiatry*. 2011;26(6):564–572.

Dovepress

 Fabian TJ, Cain ZJ, Ammerman D, et al. Improvement in functional outcomes with adjunctive aripiprazole versus placebo in major depressive disorder: a pooled post hoc analysis of 3 short-term studies. *Prim Care Companion CNS Disord*. 2012;14(6).

Weiller et al

- Nelson JC, Rahman Z, Laubmeier KK, et al. Efficacy of adjunctive aripiprazole in patients with major depressive disorder whose symptoms worsened with antidepressant monotherapy. CNS Spectr. 2014;19(6): 528–534.
- Pandina GJ, Revicki DA, Kleinman L, et al. Patient-rated troubling symptoms of depression instrument results correlate with traditional clinicianand patient-rated measures: a secondary analysis of a randomized, double-blind, placebo-controlled trial. *J Affect Disord*. 2009;118(1–3): 139–146.
- Berman RM, Marcus RN, Swanink R, McQuade RD, Carson WH, Corey-Lisle PK, Khan A. The efficacy and safety of aripiprazole as adjunctive therapy in major depressive disorder: a multicenter, randomized, double-blind, placebo-controlled study. *J Clin Psychiatry*. 2007; 68(6):843–853.
- Marcus RN, McQuade RD, Carson WH, et al. The efficacy and safety of aripiprazole as adjunctive therapy in major depressive disorder: a second multicenter, randomized, double-blind, placebo-controlled study. J Clin Psychopharmacol. 2008;28(2):156–165.
- Berman RM, Fava M, Thase ME, et al. Aripiprazole augmentation in major depressive disorder: a double-blind, placebo-controlled study in patients with inadequate response to antidepressants. CNS Spectr. 2009; 14(4):197–206.
- Kamijima K, Higuchi T, Ishigooka J, et al. Aripiprazole augmentation to antidepressant therapy in Japanese patients with major depressive disorder: a randomized, double-blind, placebo-controlled study (ADMIRE study). *J Affect Disord*. 2013;151(3):899–905.
- Quiroz JA, Tamburri P, Deptula D, et al. Efficacy and safety of basimglurant as adjunctive therapy for major depression: a randomized clinical trial. *JAMA Psychiatry*. 2016;73(7):675–684.
- 34. Thase ME, Youakim JM, Skuban A, et al. Efficacy and safety of adjunctive brexpiprazole 2 mg in major depressive disorder: a Phase 3, randomized, placebo-controlled study in patients with inadequate response to antidepressants. *J Clin Psychiatry*. 2015;76(9):1224–1231.
- 35. Thase ME, Youakim JM, Skuban A, et al. Adjunctive brexpiprazole 1 and 3 mg for patients with major depressive disorder following inadequate response to antidepressants: a Phase 3, randomized, double-blind study. J Clin Psychiatry. 2015;76(9):1232–1240.
- Fava M, Memisoglu A, Thase ME, et al. Opioid modulation with buprenorphine/samidorphan as adjunctive treatment for inadequate response to antidepressants: a randomized double-blind placebocontrolled trial. *Am J Psychiatry*. 2016;173(5):499–508.
- Durgam S, Earley W, Guo H, et al. Efficacy and safety of adjunctive cariprazine in inadequate responders to antidepressants: a randomized, double-blind, placebo-controlled study in adult patients with major depressive disorder. *J Clin Psychiatry*. 2016;77(3):371–378.
- Fava M, Ramey T, Pickering E, Kinrys G, Boyer S, Altstiel L. A randomized, double-blind, placebo-controlled Phase 2 study of the augmentation of a nicotinic acetylcholine receptor partial agonist in depression: is there a relationship to leptin levels? *J Clin Psychopharmacol*. 2015; 35(1):51–56.
- Vieta E, Thase ME, Naber D, et al. Efficacy and tolerability of flexiblydosed adjunct TC-5214 (dexmecamylamine) in patients with major depressive disorder and inadequate response to prior antidepressant. *Eur Neuropsychopharmacol*. 2014;24(4):564–574.
- 40. Möller HJ, Demyttenaere K, Olausson B, et al. Two Phase III randomised double-blind studies of fixed-dose TC-5214 (dexmecamylamine) adjunct to ongoing antidepressant therapy in patients with major depressive disorder and an inadequate response to prior antidepressant therapy. World J Biol Psychiatry. 2015;16(7):483–501.
- 41. Tummala R, Desai D, Szamosi J, Wilson E, Hosford D, Dunbar G, Eriksson H. Safety and tolerability of dexmecamylamine (TC-5214) adjunct to ongoing antidepressant therapy in patients with major depressive disorder and an inadequate response to antidepressant therapy: results of a long-term study. *J Clin Psychopharmacol*. 2015;35(1):77–81.

42. Ball S, Dellva MA, D'Souza DN, Marangell LB, Russell JM, Goldberger C. A double-blind, placebo-controlled study of edivoxetine as an adjunctive treatment for patients with major depressive disorder who are partial responders to selective serotonin reuptake inhibitor treatment. *J Affect Disord*. 2014;167:215–223.

- 43. Ball SG, Ferguson MB, Martinez JM, et al. Efficacy outcomes from 3 clinical trials of edivoxetine as adjunctive treatment for patients with major depressive disorder who are partial responders to selective serotonin reuptake inhibitor treatment. J Clin Psychiatry. 2016;77(5):635–642.
- Barbee JG, Thompson TR, Jamhour NJ, et al. A double-blind placebocontrolled trial of lamotrigine as an antidepressant augmentation agent in treatment-refractory unipolar depression. *J Clin Psychiatry*. 2011; 72(10):1405–1412.
- 45. Sanacora G, Johnson MR, Khan A, et al. Adjunctive lanicemine (AZD6765) in patients with major depressive disorder and history of inadequate response to antidepressants: a randomized, placebocontrolled study. *Neuropsychopharmacology*. 2017;42(4):844–853.
- 46. Richards C, McIntyre RS, Weisler R, et al. Lisdexamfetamine dimesylate augmentation for adults with major depressive disorder and inadequate response to antidepressant monotherapy: results from 2 Phase 3, multicenter, randomized, double-blind, placebo-controlled studies. *J Affect Disord*. 2016;206:151–160.
- 47. Thase ME, Corya SA, Osuntokun O, et al. A randomized, double-blind comparison of olanzapine/fluoxetine combination, olanzapine, and fluoxetine in treatment-resistant major depressive disorder. *J Clin Psychiatry*. 2007;68(2):224–236.
- Mahmoud RA, Pandina GJ, Turkoz I, Kosik-Gonzalez C, Canuso CM, Kujawa MJ, Gharabawi-Garibaldi GM. Risperidone for treatmentrefractory major depressive disorder: a randomized trial. *Ann Intern Med.* 2007;147(9):593–602.
- Sheehan DV. The Anxiety Disease. New York: Charles Scribner & Sons: 1983.
- Sheehan DV, Harnett-Sheehan K, Raj BA. The measurement of disability. Int Clin Psychopharmacol. 1996;11 Suppl 3:89–95.
- Sheehan KH, Sheehan DV. Assessing treatment effects in clinical trials with the Discan metric of the Sheehan Disability Scale. *Int Clin Psychopharmacol*. 2008;23(2):70–83.
- Spielmans GI, Berman MI, Linardatos E, Rosenlicht NZ, Perry A, Tsai AC. Adjunctive atypical antipsychotic treatment for major depressive disorder: a meta-analysis of depression, quality of life, and safety outcomes. *PLoS Med.* 2013;10(3):e1001403.
- 53. Abilify® (aripiprazole) tablets; Abilify Discmelt® (aripiprazole) orally disintegrating tablets; Abilify® (aripiprazole) oral solution; Abilify® (aripiprazole) injection for intramuscular use only [prescribing information]. Tokyo, Japan: Otsuka Pharmaceutical Co., Ltd.; 2017.
- Rexulti® (brexpiprazole) tablets, for oral use [prescribing information].
 Tokyo, Japan: Otsuka Pharmaceutical Co., Ltd.; 2017.
- 55. Eli Lilly and Company. Lilly announces edivoxetine did not meet primary endpoint of Phase III clinical studies as add-on therapy for major depressive disorder [press release]. Indianapolis, IN: Eli Lilly and Company; 2013. Available from: https://investor.lilly.com/releasedetail. cfm?ReleaseID=811751. Accessed November 17, 2017.
- Beck A, Crain AL, Solberg LI, Unützer J, Glasgow RE, Maciosek MV, Whitebird R. Severity of depression and magnitude of productivity loss. *Ann Fam Med*. 2011;9(4):305–311.
- 57. Wiles NJ, Mulligan J, Peters TJ, et al. Severity of depression and response to antidepressants: GENPOD randomised controlled trial. *Br J Psychiatry*. 2012;200(2):130–136.
- Kirsch I, Deacon BJ, Huedo-Medina TB, Scoboria A, Moore TJ, Johnson BT. Initial severity and antidepressant benefits: a meta-analysis of data submitted to the Food and Drug Administration. *PLoS Med*. 2008;5(2):e45.
- Fountoulakis KN, Veroniki AA, Siamouli M, Möller HJ. No role for initial severity on the efficacy of antidepressants: results of a multimeta-analysis. *Ann Gen Psychiatry*. 2013;12(1):26.
- Adler DA, McLaughlin TJ, Rogers WH, Chang H, Lapitsky L, Lerner D. Job performance deficits due to depression. *Am J Psychiatry*. 2006; 163(9):1569–1576.

- 61. Trivedi MH, Morris DW, Wisniewski SR, et al. Increase in work productivity of depressed individuals with improvement in depressive symptom severity. *Am J Psychiatry*. 2013;170(6):633–641.
- Sonne C, Carlsson J, Bech P, Elklit A, Mortensen EL. Treatment of trauma-affected refugees with venlafaxine versus sertraline combined with psychotherapy – a randomised study. *BMC Psychiatry*. 2016; 16(1):383.
- Bech P. Measurement-Based Care in Mental Disorders. Cham, Switzerland: Springer International Publishing AG; 2016.

Neuropsychiatric Disease and Treatment

Publish your work in this journal

Neuropsychiatric Disease and Treatment is an international, peerreviewed journal of clinical therapeutics and pharmacology focusing on concise rapid reporting of clinical or pre-clinical studies on a range of neuropsychiatric and neurological disorders. This journal is indexed on PubMed Central, the 'PsycINFO' database and CAS, and is the official journal of The International Neuropsychiatric Association (INA). The manuscript management system is completely online and includes a very quick and fair peer-review system, which is all easy to use. Visit http://www.dovepress.com/testimonials.php to read real quotes from published authors.

 $\textbf{Submit your manuscript here:} \ \texttt{http://www.dovepress.com/neuropsychiatric-disease-and-treatment-journal} \\$

