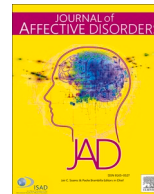




Contents lists available at ScienceDirect

## Journal of Affective Disorders

journal homepage: [www.elsevier.com/locate/jad](http://www.elsevier.com/locate/jad)

## Real-world evidence from a European cohort study of patients with treatment resistant depression: Treatment patterns and clinical outcomes

K. Heerlein<sup>a,\*</sup>, G. Perugi<sup>b</sup>, C. Otte<sup>c</sup>, T. Frodl<sup>d</sup>, G. Degraeve<sup>e,f</sup>, W. Hagedoorn<sup>g</sup>, A. J. Oliveira-Maia<sup>h,i</sup>, V. Perez Sola<sup>j</sup>, S. Rathod<sup>k</sup>, G. Rosso<sup>l</sup>, P. Sierra<sup>m</sup>, S. Malynn<sup>n</sup>, J. Morrens<sup>o</sup>, C. Verrijcken<sup>p</sup>, B. Gonzalez<sup>q</sup>, A.H. Young<sup>r,s</sup>

<sup>a</sup> Janssen EMEA, Neuss, Germany<sup>b</sup> University of Pisa, Department of Clinical and Experimental Medicine, Pisa, Italy<sup>c</sup> Charité Universitätsmedizin, Berlin, Germany<sup>d</sup> Universitätsklinikum Magdeburg, Otto von Guericke Universität Magdeburg, Magdeburg, Germany<sup>e</sup> AZ Alma General Hospital, Eeklo, Belgium<sup>f</sup> PC Dr Guislain Hospital, Ghent, Belgium<sup>g</sup> Practice for Psychiatry and Psychotherapy, Heerde, Netherlands<sup>h</sup> Champalimaud Research and Clinical Centre, Champalimaud Centre for the Unknown, Lisbon, Portugal<sup>i</sup> NOVA Medical School, NMS, Universidade Nova de Lisboa, Lisbon, Portugal<sup>j</sup> Institut de Neuropsiquiatria i Addiccions, Hospital del Mar, Barcelona IMIM Hospital del Mar Medical Research Institute, Univ Autònoma de Barcelona, CIBERSAM, Department of Psychiatry, Barcelona, Spain<sup>k</sup> Southern Health NHS Foundation Trust, Research Department, Tom Rudd Unit, Southampton, United Kingdom<sup>l</sup> San Luigi Gonzaga Hospital, Department of Neurosciences, University of Turin, Turin, Italy<sup>m</sup> University and Polytechnic Hospital La Fe, Valencia, University of Valencia, Spain<sup>n</sup> Janssen EMEA, Dublin, Ireland<sup>o</sup> Janssen EMEA, Beerse, Belgium<sup>p</sup> Janssen EMEA, Paris, France<sup>q</sup> Janssen EMEA, Madrid, Spain<sup>r</sup> Institute of Psychiatry, Psychology and Neuroscience, King's College London, Department of Psychological Medicine, London, United Kingdom<sup>s</sup> South London and Maudsley NHS Foundation Trust, Bethlem Royal Hospital, Beckenham, United Kingdom

### ABSTRACT

**Background:** Treatment resistant depression (TRD) characterizes a subgroup of 10–30% of patients with major depressive disorder, and is associated with considerable morbidity and mortality. A consensus treatment for TRD does not exist, which often leads to wide variations in treatment strategies. Real-world studies on treatment patterns and outcomes in TRD patients in Europe are lacking and could help elucidate current treatment strategies and their efficacy.

**Methods:** This non-interventional cohort study of patients with TRD (defined as treatment failure on  $\geq 2$  oral antidepressants given at adequate dose and duration) with moderate to severe depression collected real-world data on treatment patterns and outcomes in several European countries. Patients were started on a new treatment for depression according to routine clinical practice.

**Results:** Among 411 patients enrolled, after 6 months, only 16.7% achieved remission and 73.5% showed no response. At Month 12, while 19.2% achieved remission and 69.2% showed no response, 33.3% of those in remission at Month 6 were no longer in remission. Pharmacological treatments employed were heterogenous; 54 different drugs were recorded at baseline, and the top 5 treatment types according to drug classes accounted for 40.0% of patients. Even though remission rates were very low, at Month 12, 60.0% of patients had not changed treatment since enrolment.

**Conclusions:** The heterogeneity of treatments highlights a lack of consensus. Moreover, despite low response rates, patients often remained on treatments for substantial periods of time. These data further support existence of an unmet treatment need for TRD patients in Europe.

\* Corresponding author.

E-mail address: [kheerlei@its.jnj.com](mailto:kheerlei@its.jnj.com) (K. Heerlein).

<https://doi.org/10.1016/j.jad.2021.03.073>

Received 10 February 2021; Received in revised form 22 March 2021; Accepted 24 March 2021

Available online 1 April 2021

0165-0327/© 2021 The Author(s).

Published by Elsevier B.V. This is an open access article under the CC BY-NC-ND license

(<http://creativecommons.org/licenses/by-nc-nd/4.0/>).

## Author Interview, Animated Summary and Plain Language Summary

To view an interview with one of the authors, Professor Giulio Perugi, summarizing this publication, please see the video below, or visit the manuscript on line at doi:[10.1016/j.jad.2021.03.073](https://doi.org/10.1016/j.jad.2021.03.073).

### 1. Introduction

Major depressive disorder (MDD) is a debilitating condition that results in considerable morbidity and mortality, in part due to high suicide risk (Cavanagh et al., 2003). MDD is thought to affect 4–10% of the general population across their lifetime (National Institute for Health and Care Excellence, 2020) and depressive disorders are a leading cause of disability (James et al., 2018). Approximately 10–30% of patients with MDD have treatment resistant depression (TRD) (Rush et al., 2006b; Jaffe et al., 2019; Al-Harbi et al., 2012; Voineskos et al., 2020), defined as a major depressive episode (MDE) that fails to respond to two or more different antidepressants, given at adequate dose and duration (European Medicines Agency, 2013; Souery et al., 1999). Other definitions of TRD have also been used, precluding consensus on the prevalence of the condition (Wiles et al., 2014; Conway et al., 2017; Nemeroff et al., 2007).

Current treatments for MDD include medication, psychological and neurostimulation therapies (National Institute for Health and Care Excellence, 2020; European Medicines Agency, 2013; Cleare et al., 2015). Clinical management of TRD can include different combinations of these treatments, including antidepressant and non-antidepressant drugs, as well as non-pharmacological therapies (European Medicines Agency, 2013; Voineskos et al., 2020; Bennabi et al., 2019; Ionescu et al., 2015). All antidepressant drugs can be used in the treatment of TRD including selective serotonin reuptake inhibitors (SSRIs), dual serotonin and norepinephrine reuptake inhibitors (SNRIs), tricyclic antidepressants (TCA), monoamine oxidase inhibitors (MAOIs), and ‘other’ antidepressants such as tianeptine, agomelatine and  $\alpha$ 2-antagonists (Ionescu et al., 2015; Kim et al., 2019; Tobe and Rybakowski, 2013). Medications without primary antidepressant properties can also be used in the treatment of TRD to potentiate the effects of an antidepressant drug. These include lithium, thyroid hormone and antipsychotics such as quetiapine (Bauer et al., 2013). Pharmacological treatment strategies for TRD can be categorized as switching (from one antidepressant to another); combination therapy (adding another antidepressant to the current one); and augmentation/add-on therapy (use of a non-antidepressant medication in addition to a current antidepressant) (Barowsky and Schwartz, 2006; Ionescu et al., 2015; Voineskos et al., 2020). Non-pharmacological (psychotherapeutic and neurostimulatory) therapies, have also been developed for TRD (Voineskos et al., 2020; van Bronswijk et al., 2019; Lefaucheur et al., 2020; Lisanby, 2007) but will not be discussed in detail here.

While treatment patterns and outcomes in patients with TRD in clinical practice have been studied in the US (Kubitz et al., 2013; Amos et al., 2018), recent research in Europe is limited. The US Sequenced Treatment Alternatives to Relieve Depression (STAR\*D) trial is the largest multistep treatment study of patients with depression to date and provided key insights into treatment failure in the clinical setting (Rush et al., 2006b). STAR\*D showed that most patients with TRD fail to achieve remission with current treatments. Remission rates of approximately 33%, 25–33%, 14% and 13% were achieved during the first, second, third and fourth treatment steps in the study, respectively, leading the authors to suggest that the likelihood of remission is reduced in patients with TRD (equating to those reaching steps three and four). In patients that achieved remission, loss of response was common; 60% of TRD patients that responded to treatment became unresponsive to treatment within 6 months. A prospective, 2-year observational study of patients with TRD being treated in routine clinical practice, reported a 12-month remission rate of 3.6%. Loss of remission between Month 12

and 24 occurred in 75% of patients, highlighting that maintenance of therapeutic effect is poor (Dunner et al., 2006). Data from European patients, although limited, suggest similar outcomes. In a study in the UK, the 18-month remission rate for patients with TRD receiving treatment as usual, as directed by their primary care practitioner, was 6.5% and dropped to 4.4% at 42-month follow up (Fonagy et al., 2015).

Regarding European-wide market approval, a single pharmacological treatment, esketamine nasal spray (in combination with an SSRI or SNRI) was the first treatment approved for TRD specifically (European Medicines Agency, 2013; Mahase, 2019; European Medicines Agency, 2019). Importantly, however, it was not approved until after this study ended (European Medicines Agency, 2013; Mahase, 2019; European Medicines Agency, 2019). Additionally, there is no consensus on treatment pathways for TRD, and evidence suggests wide variation between and within European countries (MacQueen et al., 2017). The current cohort study was established to collect data from TRD patients with moderate to severe depression being treated in routine clinical practice in a sample of European countries. The aim of the study was to describe real-world clinical features and, as such, did not include a specific hypothesis. Treatment patterns and outcomes were followed for up to 21 months in patients starting a new therapy, having already experienced at least two treatment failures in the current MDE. The objectives of the study were to describe the disease-related and socio-demographic characteristics and disease burden of TRD patients in Europe; treatment patterns used to manage TRD in European clinical practice, and the resulting clinical outcomes; and healthcare resource utilisation associated with TRD. The baseline characteristics of the patient cohort have been published recently, demonstrating that patients had low health-related quality of life (HRQoL) and reduced function (Heerlein et al., 2021). This paper reports data supporting the second objective, focusing on treatment patterns and clinical outcomes among patients from this cohort.

### 2. Methods

#### 2.1. Patients

Patients aged 18 to 74 years fulfilling the criteria for TRD were eligible. A diagnosis of MDD (fulfilling the Diagnostic and Statistical Manual of Mental Disorders Fifth Edition [DSM-5] or the International Statistical Classification of Diseases and Related Health Problems, Tenth Revision [ICD-10] criteria for MDD or Depressive disorder) was required for inclusion (European Medicines Agency, 2013), as well as a Montgomery-Åsberg Depression Rating Scale (MADRS; Montgomery and Åsberg, 1979) score  $\geq 20$ , indicating moderate to severe depression. TRD was defined by treatment failure ( $\leq 25\%$  improvement on best day) on  $\geq 2$  different oral antidepressants, taken for  $\geq 6$  weeks, at adequate dose in the same MDE, as determined using the Massachusetts General Hospital-Antidepressant Treatment Response Questionnaire (MGH-ATRQ; Chandler et al., 2010).

For inclusion, patients needed to be starting a new antidepressant treatment, as an inpatient or outpatient. Treatment was according to routine clinical practice in that setting; choice of treatment, dose and method of administration was at the discretion of the treating physician. In the context of the study, any pharmacological or non-pharmacological treatment, including neurostimulation and psychotherapeutic interventions, prescribed to replace, or in addition to, the previous treatment, was considered a new antidepressant treatment. Switches to a generic drug or changes in dose did not count as a new treatment.

The following were exclusion criteria: current or prior diagnosis of psychotic disorders, MDD with psychotic features, bipolar disorders or intellectual disability according to DSM-5 or ICD-10; history of suicidal behavior within one year prior to enrolment; homicidal ideation or intent, or suicidal ideation or intent, within one month prior to entering the study; moderate to severe substance misuse (including alcohol)

within six months before enrolment. Written informed consent was provided by all participants, and their capability for doing so was judged by the treating physician. Local ethics review boards provided approval for the study, which adhered to the Declaration of Helsinki.

## 2.2. Study design

A prospective, multicenter, observational cohort study of patients with TRD in Europe was conducted. The study has been described in detail elsewhere (Heerlein et al., 2021). Briefly, the study was comprised of baseline data collection, a 12-month observational period with a minimum follow-up of approximately 6 months for each enrolled patient, and an extended observation period up to 6 months from recruitment of the last patient, resulting in a maximum of 21 months of follow-up. Analysis was conducted only on data collected at 6 and 12 months after baseline (**Supplementary Fig. 1**). Patients were enrolled from Belgium, Germany, Italy, the Netherlands, Portugal, Spain and the United Kingdom.

## 2.3. Treatments

Antidepressant treatments (pharmacological, psychotherapeutic and neurostimulatory) prescribed on enrolment and during the observational period were documented in medical records. Pharmacological treatment strategies were categorized as monotherapy, combination therapy or augmentation therapy. Treatment could be changed at any time during the study, at the discretion of the treating physician.

## 2.4. Study procedures and evaluations

Data were collected at baseline, on scheduled visits every 6 months and at the end of the study, which was planned to run until 6 months after enrolment of the last patient. The data collection planned for the end of the study was also performed in the event of a patient leaving the study before the end of the observational period and, for patients who withdrew consent, were lost to follow-up or died before the end of the study, as many data were collected as possible. Additional data collection was encouraged in case of clinically relevant events, namely: admission to, or discharge from, inpatient settings; relapse of symptoms (including suicidal ideation/intent/behavior); remission of the current MDE (as confirmed by MADRS  $\leq 10$ ); visit/consultation with treating physician due to worsening or improvement of the current MDE (confirmed by Clinical Global Impression of Change [CGI-C; Busner and Targum, 2007]  $\leq 3$  or  $\geq 5$ ); change in pharmacological treatment (medication switch; dose changes except those relating to titration during switching; start of augmentation therapy); change in non-pharmacological treatment.

Baseline data were collected on patient socio-demographics, disease history and current clinical characteristics as well as details of all antidepressant treatments used in the current MDE prior to study entry. Assessment at any visit could include documentation of: spectrum and severity of depression symptoms (MADRS; Clinical Global Impression of Severity [CGI-S] score); change in depression severity since initiating last treatment (CGI-C score); suicidality (Columbia Suicide Severity Rating Scale; Posner et al., 2011); HRQoL (EuroQoL 5-dimension 5-level [EQ-5D-5L] patient-reported questionnaire, including the EuroQoL Visual Analog Scale [EQ-VAS]; Herdman et al., 2011); functional impairment or disability in and outside of work (Work Productivity and Activity Impairment [WPAI; Reilly et al., 1993] and the Sheehan Disability Scale [SDS; Sheehan et al., 1996] patient-reported questionnaires).

## 2.5. Data processing and statistical analysis

The remission or response status of each patient (irrespective of treatment strategy) was assessed at 6 and 12 months after baseline, according to MADRS scores. Remission was defined as a score  $\leq 10$ , while scores  $>10$  but with improvement from baseline  $\geq 50\%$  defined response

without remission. Visits were considered as occurring in Month 6 if they occurred 150–216 days after enrolment, while Month 12 visits were defined as those occurring 330–402 days after enrolment. Data obtained outside these windows were excluded from the relevant time point.

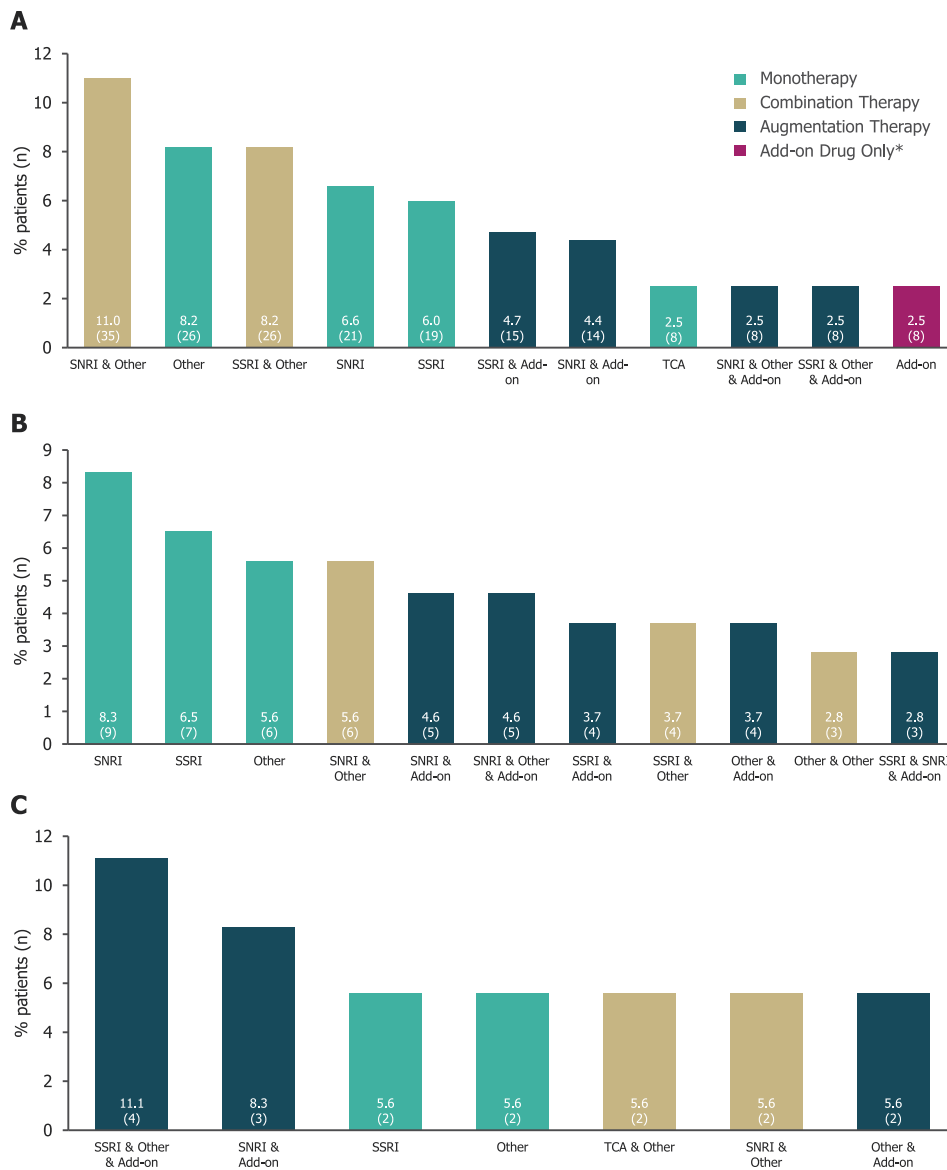
Treatment steps for each patient were reconstructed by matching MADRS scores to the start and end dates of each treatment strategy (dates and treatments obtained from medical records), with the end representing a change to a different treatment strategy. Only pharmacological treatment strategy changes were considered in the treatment strategy analysis, excluding any  $<30$  days duration. The MADRS values at the start and end of each treatment strategy were used to determine whether the patient had experienced remission, response without remission, or no response. When the MADRS score was not available, a response/remission proxy was inferred based on CGI scores, to increase the number of treatment strategies for which a corresponding treatment outcome could be reported. If the end CGI-S score was 1 or 2 (out of 7, with higher scores indicating greater severity), the treatment strategy was considered to have resulted in remission (Turrina et al., 2015). When this was not the case, yet the CGI-C score was 1 or 2 (out of 7, where the lowest scores indicate substantial improvement and scores  $>4$  indicate a worsening condition) then the treatment strategy was considered to have resulted in response without remission (Fava et al., 2017).

Continuous variables were summarized using descriptive statistics (N, mean, standard deviation [SD], median, minimum and maximum). Categorical variables were summarized by frequency distribution (number and percentage of patients in each category). Significance testing of data stratified by outcome (remission, response without remission, and non-response) was carried out first by using a Kruskal Wallis (KW) test to evaluate the null hypothesis that data from all three outcome groups were equal. If the KW test indicated a significant difference, it was followed by post-hoc Dunn's tests comparing the data between each of the three outcome groups. A p value  $<0.05$  was considered significant. Treatment strategy duration by response status was compared using a repeated-measures model using SAS proc mixed procedure. The model statement included categorical variables for response status (two categories: response, no response) along with treatment strategy number at baseline (three categories: first step, second step, third step or higher). To account for within-subject correlation of treatment strategy duration, a repeated-measures statement with unstructured covariance matrix was included. The p value corresponding to the response status factor in the model statement was evaluated for significance. Time-to-treatment strategy change analysis was performed by Kaplan-Meier time-to-event analysis.

## 3. Results

### 3.1. Baseline characteristics and treatments

The final analysis set included 411 eligible patients (**Supplementary Table 1**). The mean (SD) age of patients was 51.0 (10.8) years and 62.3% were female. The mean (SD) MADRS score was 31.8 (6.0) and 67.4% of patients had moderate depression (MADRS 20–34); the remainder (32.6%) had severe depression (MADRS  $>34$ ). In the current MDE, 53.8%, 31.1% and 14.6% of patients had experienced two, three or four or more treatment failures, respectively. At baseline, 343/411 (83.4%) patients were taking at least one pharmacological treatment. Psychotherapeutic treatment and neurostimulation therapy were being prescribed to 19.2% and 6.6% of patients, respectively. A full description of the baseline characteristics of this patient population has been published previously (Heerlein et al., 2021).



**Fig. 1.** MADRS responses at Month 6 and Month 12. **A.** Patient outcomes at Month 6 and 12 (total population). **B.** Patient outcomes at Month 12, by remission status at Month 6. **C.** Patient outcomes at Month 12, by response status at Month 6. Dataset includes all patients, irrespective of treatment strategy. Only patients with MADRS scores for both Month 6 and 12 were included in B and C. Remission: MADRS score  $\leq 10$ ; response: MADRS improvement from baseline  $\geq 50\%$ ; response without remission: MADRS improvement from baseline  $\geq 50\%$  and MADRS score  $> 10$ ; no response: MADRS improvement from baseline  $< 50\%$  and MADRS score  $> 10$ . MADRS: Montgomery-Åsberg Depression Rating Scale.

### 3.2. Overall treatment outcomes at Month 6

At Month 6, 58 patients were reported as discontinued: 1 patient ended the study after 99 days<sup>1</sup>; 37 patients were lost to follow up; 7 withdrew consent; 3 died; 9 discontinued due to ‘other’ reasons; 1 was of unknown status. Data were excluded from a further 47 patients who were still in the study at Month 6, but whose visits did not meet the defined cut-off dates for a Month 6 visit. All other patients were included in this analysis, irrespective of treatment strategy. Of these patients, 73.5% showed no response, 9.8% showed response without remission and 16.7% were in remission (Fig. 1A). As per the definitions used to stratify patients by outcome status, the mean (SD) MADRS score for patients achieving remission was 6.0 (2.6) compared to 14.1 (2.7) in patients with response without remission and 26.2 (7.5) in patients with

no response (Table 1). Using KW tests followed by post-hoc Dunn’s tests to analyse mean change from baseline, greater change was seen across EQ-5D-5L, EQ-VAS and SDS in patients achieving remission compared to those with no response ( $p < 0.001$  for all comparisons; Table 1). Greater mean change was also observed for patients achieving response without remission in the same comparison ( $p < 0.05$  or  $p < 0.01$ ; Table 1).

### 3.3. Overall treatment outcomes at Month 12

At Month 12, 244 patients in total were not included in the analysis set: for 164 patients the study ended prior to their 12-month assessment; 45 were lost to follow up; 9 withdrew consent; 4 died; 22 discontinued due to ‘other’ reasons; 1 was of unknown status. Data were excluded from a further 21 patients who were still in the study at Month 12, but whose visits did not meet the defined cut-off dates for a Month 12 visit. All other patients were included in this analysis, irrespective of treatment strategy. Of these patients, 69.2% had not responded, 11.6% showed response without remission and 19.2% were in remission (Fig. 1A). Using KW tests followed by post-hoc Dunn’s tests to analyse mean change from baseline, greater changes in EQ-5D-5L, EQ-VAS and

<sup>1</sup> Whilst the protocol recommended a minimum of 6 months follow-up, this patient was reported to have ended the study after 99 days, even though the study was on-going. No data from this patient were recorded after 99 days, so their data are not included in 6 or 12-month analyses.

**Table 1**  
Patient outcomes at Month 6.

N Mean (SD)	All (N=306)	No Response (n=225)	Response without Remission (n=30)	Remission (n=51)
Total MADRS score	306 21.6 (10.2)	225 26.2 (7.5)	30 14.1 (2.7)	51 6.0 (2.6)
CGI-C score, n (%)				
Very much improved	28 (9.2)	2 (0.9)	2 (6.7)	24 (47.1)
Much improved	73 (23.9)	31 (13.8)	16 (53.3)	26 (51.0)
Minimally improved	92 (30.1)	82 (36.4)	9 (30.0)	1 (2.0)
No change	73 (23.9)	71 (31.6)	2 (6.7)	0
Minimally worse	25 (8.2)	24 (10.7)	1 (3.3)	0
Much worse	15 (4.9)	15 (6.7)	0	0
EQ-5D-5L (change from baseline)		206 0.11 (0.25)	26 0.26 (0.21)*	46 0.34 (0.28)***
EQ-VAS (change from baseline)		210 6.32 (21.20)	26 21.00 (19.21)**	48 32.35 (20.02)***
Total SDS (change from baseline)		153 -2.67 (5.85)	19 -7.58 (5.65)**	38 -12.53 (6.88)***

Dataset includes all patients, irrespective of treatment strategy. \* $p < 0.05$ ; \*\* $p < 0.01$ ; \*\*\* $p < 0.001$  when compared to no response (Kruskal Wallis test followed by post-hoc Dunn's test). Remission: MADRS score  $\leq 10$ ; response without remission: MADRS improvement from baseline  $\geq 50\%$  and MADRS score  $> 10$ . CGI-C: Clinical Global Impression of Change scale; EQ-5D-5L: European Quality of Life (EuroQol)-5-dimension 5-level; EQ-VAS: EuroQol-visual analog scale; MADRS: Montgomery-Åsberg Depression Rating Scale; SD: standard deviation; SDS: Sheehan Disability Scale.

**Table 2**  
Patient outcomes at Month 12.

N Mean (SD)	All (N=146)	No Response (n=101)	Response without Remission (n=17)	Remission (n=28)
Total MADRS score	146 20.0 (9.7)	101 25.1 (6.6)	17 13.6 (2.2)	28 5.7 (3.3)
CGI-C score, n (%)				
Very much improved	20 (13.7)	1 (1.0)	4 (23.5)	15 (53.6)
Much improved	29 (19.9)	13 (12.9)	7 (41.2)	9 (32.1)
Minimally improved	43 (29.5)	38 (37.6)	3 (17.6)	2 (7.1)
No change	34 (23.3)	31 (30.7)	2 (11.8)	1 (3.6)
Minimally worse	13 (8.9)	11 (10.9)	1 (5.9)	1 (3.6)
Much worse	6 (4.1)	6 (5.9)	0	0
EQ-5D-5L (change from baseline)		94 0.11 (0.26)	16 0.31 (0.26)*	26 0.35 (0.28)***
EQ-VAS (change from baseline)		97 9.73 (20.11)	16 23.13 (13.35)*	27 35.19 (24.40)***
Total SDS (change from baseline)		67 -2.91 (6.73)	9 -7.00 (6.34)	18 -14.44 (7.76)***

Dataset includes all patients, irrespective of treatment strategy. \* $p < 0.05$ ; \*\* $p < 0.01$ ; \*\*\* $p < 0.001$  when compared to no response (Kruskal Wallis test followed by post-hoc Dunn's test). Remission: MADRS score  $\leq 10$ ; response without remission: MADRS improvement from baseline  $\geq 50\%$  and MADRS score  $> 10$ . CGI-C: Clinical Global Impression-Change scale; EQ-5D-5L: European Quality of Life (EuroQol)-5-dimension 5-level; EQ-VAS: EuroQol-visual analog scale; MADRS: Montgomery-Åsberg Depression Rating Scale; SD: standard deviation; SDS: Sheehan Disability Scale.

SDS were observed in patients who had achieved remission at Month 12 than for those who had not responded ( $p < 0.001$ ; Table 2), and changes in EQ-5D-5L and EQ-VAS were greater in patients achieving response without remission when compared to those with no response ( $p < 0.05$ ; Table 2).

### 3.4. Changes in remission or response status between Month 6 and 12

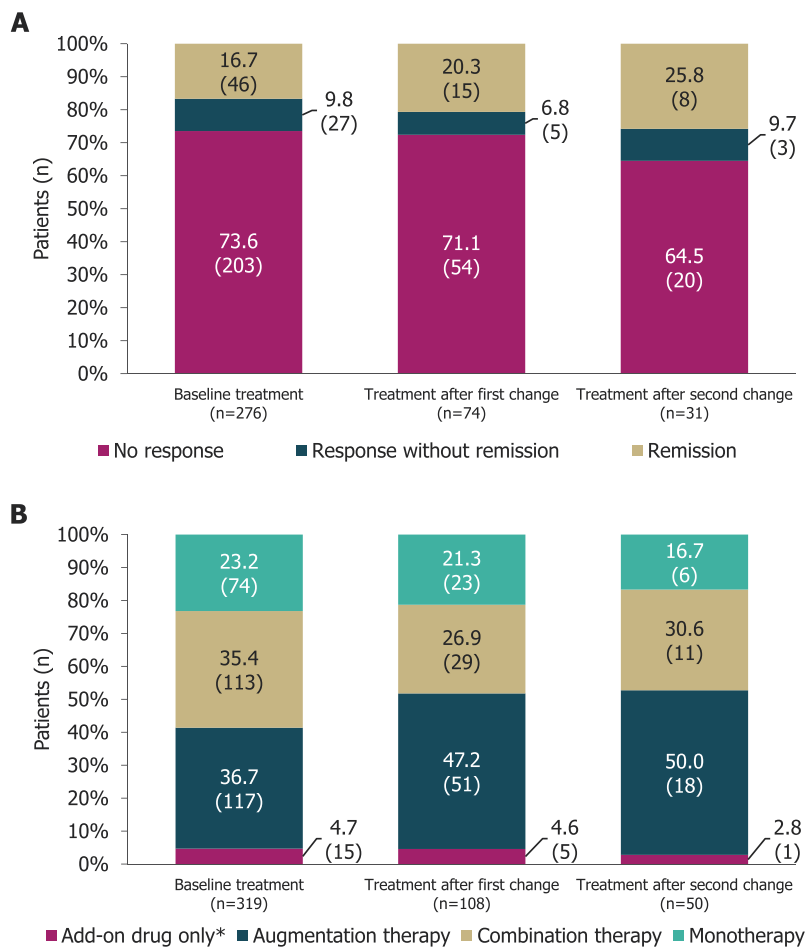
Maintenance of remission or response was analysed in patients for whom data at both timepoints were available. Of patients who had not achieved remission at Month 6, 86.6% of those that were assessed again at Month 12 still had not achieved remission (Fig. 1B). Among patients who achieved remission at Month 6, 33.3% of those that were assessed again at Month 12 were no longer in remission (Fig. 1B). When these patients were stratified by response vs non-response, 51.7% of Month 6 responders had lost response by Month 12, and among non-responders at Month 6, 72.9% remained unresponsive at Month 12 (Fig. 1C).

### 3.5. Treatment patterns and response by number of treatments since enrolment

Data from patients taking  $\geq 1$  pharmacological treatment with a duration  $\geq 30$  days were included in this analysis. The rate of response to treatment increased with the number of treatment changes after baseline (Fig. 2A). With treatment implemented at baseline, 26.4% of patients responded, while response was achieved in 35.5% of patients completing a third treatment since entering the study (Fig. 2A). For baseline treatment, the proportions of patients on combination or augmentation strategies were 35.4% and 36.7%, respectively, 23.2% were on monotherapy (Fig. 2B). The proportion of patients on monotherapy decreased as treatment was changed, with 16.7% on monotherapy in the third treatment since starting the study (Fig. 2B). There was a concurrent increase in the percentage of patients on augmentation therapy that accounted for 50.0% of patients completing a third treatment since starting the study (Fig. 2B).

Together, the top five treatments, taken by patients at baseline





**Fig. 2.** Pharmacological strategies and treatment outcomes per treatment step during the study. A. Treatment outcomes by treatment step during the study from baseline. B. Pharmacological strategies per treatment step during the study from baseline; data include only patients that were taking  $\geq 1$  pharmacological treatment and who had been taking it for  $\geq 30$  days. All patients have already experienced at least two treatment line failures prior to study entry, as per study inclusion criteria; since the number of treatment lines prior to enrollment varies across the cohort, the number of treatment changes does not necessarily relate to the total number of treatment lines a patient has been prescribed. Figure shows results from the total population. \*Add-on drug only therapy: prescription of an add-on medication in the absence of regular oral antidepressant(s). Augmentation therapy: prescription of an add-on medication in addition to regular oral antidepressant (s). Combination therapy: prescription of  $\geq 2$  antidepressant medications. Monotherapy: prescription of 1 antidepressant medication. Remission: MADRS score  $\leq 10$ ; response without remission: MADRS improvement from baseline  $\geq 50\%$  and MADRS score  $> 10$ ; no remission: MADRS improvement from baseline  $< 50\%$  and MADRS score  $> 10$ . MADRS: Montgomery-Åsberg Depression Rating Scale.

defined according to drug classes, accounted for 40.0% of all treatments used at that timepoint (Fig. 3A). The most common treatment at baseline was the combination of an SNRI plus “other” antidepressant, used in 11.0% of patients. “Other” antidepressants as monotherapy and SSRI plus “other” antidepressant were both taken by 8.2% of patients (Fig. 3A). In patients whose treatment was changed following failure of baseline treatment, the top two individual treatment classes taken on changing were SNRIs (8.3%) and SSRIs (6.5%) both used as a monotherapy (Fig. 3B). The top two treatment classes and combinations used for patients changing treatment a second time in the study were a combination of SSRI plus “other” antidepressant plus an add-on drug (11.1%) and an SNRI plus add-on medication (8.3%; Fig. 3C).

### 3.6. Treatment strategy duration, per treatment strategy used since baseline

Data from 392 treatment strategies were available for analysis, after excluding non-pharmacological strategies. Of these, outcomes (remission, response, non-response) were assigned based on CGI-S and/or CGI-C in 3.6% (14/392) of treatment strategies where a corresponding MADRS score was not available. Pharmacological strategies  $\geq 30$  days duration were analysed. Where a treatment strategy resulted in response (including remission), the mean (SD) duration of treatment was 250.6 (136.0) days. When patients did not achieve a response, the mean (SD) treatment strategy duration before changing was shorter (196.2 [128.3] days;  $p < 0.001$ ; repeated measures model; Fig. 4A). Kaplan-Meier analysis was used to analyse time-to-treatment change endpoints. After approximately 6 months of treatment, 68.4% of patients had not changed treatment strategy (including prescription of an additional

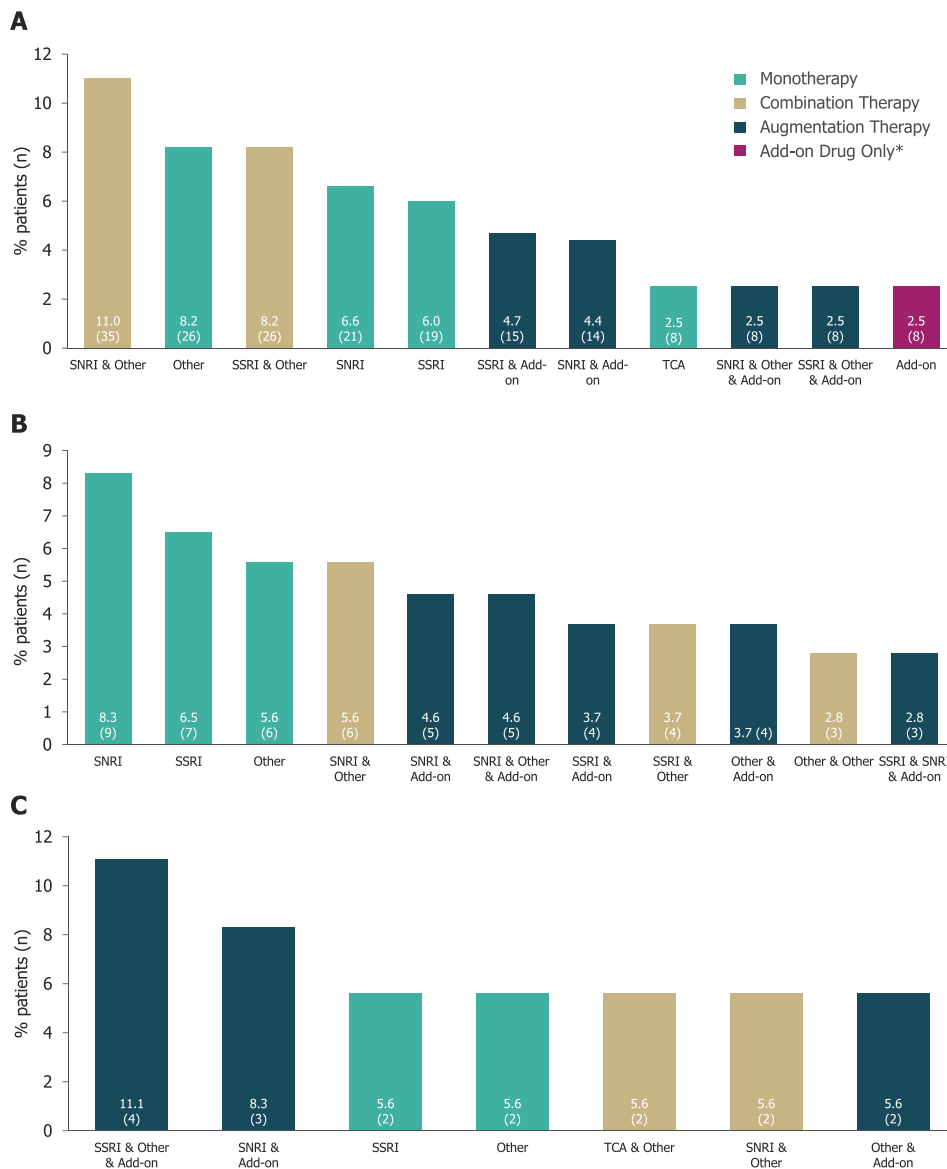
drug) since enrolment. At Month 12, approximately 60.0% of patients had not changed their treatment strategy since entry into the study (Fig. 4B).

## 4. Discussion

This study assessed the naturalistic treatment patterns and clinical outcomes of 411 patients with TRD in Europe. Results at 6 months after initiating a new treatment strategy, as per routine clinical practice, showed most patients did not respond to treatment or achieve remission, remained very unwell and had poor HRQoL and reduced function. The number of different treatment strategies reported was high; the top five treatment types accounted for only 40.0% of treatments, suggesting that current treatment strategies employed for patients with TRD are very heterogeneous. Importantly, and considering that there is no consensus strategy across Europe for treating TRD, the wide variation in treatment strategies being used in clinical practice makes it difficult to build a robust evidence base of which strategies are most efficacious to inform clinical guidelines. This, in turn, makes it more challenging for clinicians to make informed treatment decisions.

The results reported here largely align with other studies, in terms of the variety of treatments used and treatment outcomes (Rush et al., 2006b, Ionescu et al., 2015).

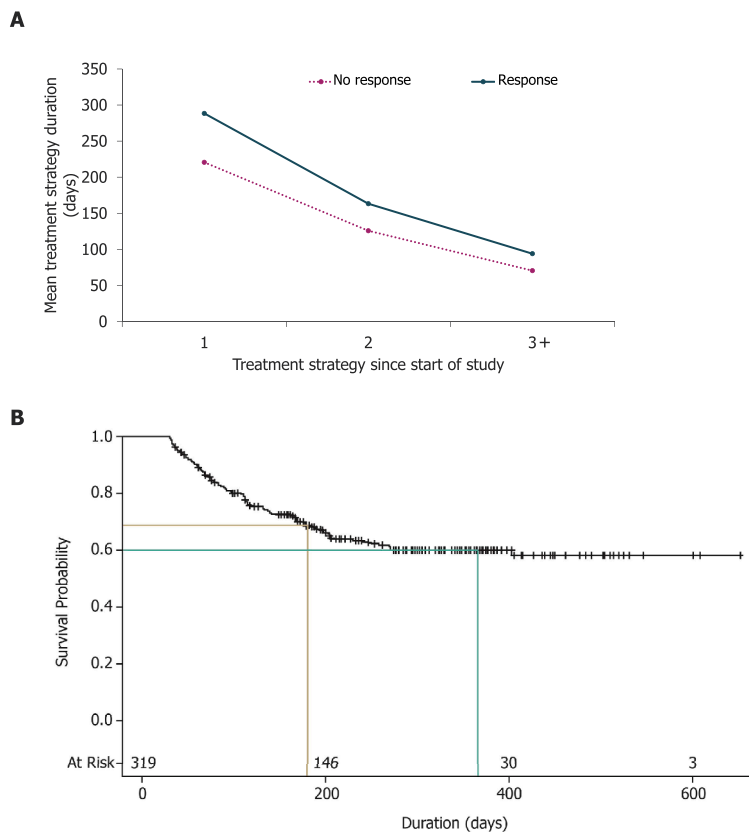
In this study, most patients failed to achieve or sustain a clinical response, with less than 20% of patients in remission at 12 months. In a 2-year observational study of patients with TRD in which patients received a variety of treatments (“treatment as usual”), that could be changed at any time during the study, a 12-month remission rate of 3.6% was reported (Dunner et al., 2006). This is considerably lower than the



**Fig. 3.** Treatment strategies used at each treatment step, by treatment classes. \*Add-on drug only therapy: the use of an add-on medication in the absence of regular oral antidepressant(s). **A.** Baseline treatment. **B.** Treatment after first treatment change in the study. **C.** Treatment after second treatment change in the study. All patients have already experienced  $\geq 2$  treatment line failures prior to study entry, as per study inclusion criteria. Augmentation therapy: the prescription of an add-on medication in addition to regular oral antidepressant(s). Combination therapy: the prescription of  $\geq 2$  antidepressant medications. Figure shows most common treatment classes prescribed in  $\geq 2\%$ ,  $\geq 5\%$  and  $>2$  patients (**A**, **B** and **C**, respectively). SNRI: serotonin-norepinephrine reuptake inhibitor; SSRI: selective serotonin reuptake inhibitor; TCA: tricyclic antidepressant.

remission rate observed in the current study. Notable differences that may have contributed towards the lower remission rate include the inclusion of patients with bipolar disorder, higher number of mean treatment failures and longer mean duration of the current MDE, indicating a cohort with more difficult to treat depression. Furthermore, in that study the self-rated 30-item Inventory of Depressive Symptomatology-Self-Report was used to assess changes in the degree of depression. Other studies of long-term outcomes for TRD patients in routine clinical practice have also reported lower remission rates than the current study. As in this study, the US Treatment-Resistant Depression Registry study (Aaronson et al., 2017) used clinician rated assessments (MADRS), and the baseline depression severity was comparable. However, only 12% of patients achieved remission after 1 year, possibly because the cohort included patients with bipolar disorder and overall had a considerably worse clinical history with a mean of 7.3 treatment failures. Regarding treatment practices in Europe, a long-term study of UK TRD patients in routine clinical practice reported an 18-month remission rate of 6.5%, decreasing to 4.4% after 42 months (Fonagy et al., 2015). Compared with the current study, patients in this cohort had a greater mean number of prior treatment failures, as well as greater mean current MDE duration. Notwithstanding differences in the

baseline clinical characteristics of these other patient cohorts, the higher rates of remission reported in the current study may also be explained by low patient numbers at Month 12, possibly reflecting differential loss to follow up. In the current study, one third of Month 6 remitters were no longer in remission at Month 12, suggesting that treatment efficacy was lost. Remission was lost by an even higher proportion of patients (75%) between Month 12 and 24 in another observational study, possibly due to key differences in the patient baseline characteristics and study, as was highlighted above (Dunner et al., 2006). When analyzed in terms of response, rather than remission, more than 50.0% of Month 6 responders lost response by Month 12, suggesting response is a less stable outcome for patients than remission. Both analyses support the argument that current treatment strategies are inadequate to maintain long-term treatment success in many TRD patients, but that remission is a more robust target outcome than simply response. That remission should be the aim of treatment strategies is further supported by the greater improvement in functioning/disability in patients who achieved remission compared to those who only achieved a response without remission in this study. Others have also acknowledged that failure to achieve full remission is associated with an increased risk of relapse and recurrent episodes, as well as the personal and societal burden resulting



**Fig. 4.** Pharmacological treatment duration and time to first treatment change. **A.** Least square mean plot of treatment number and duration, by outcome. Treatment outcome based on MADRS score, or CGI-C/S where a MADRS score was unavailable (14/392 [3.6%]). Treatment durations were right censored at the time corresponding to the last study visit. Since some patients (most likely) continued treatment after the study had ended, their treatment durations would in fact be longer; this analysis therefore underestimates treatment duration. **B.** Time to first treatment change from study entry (excluded treatment lines <30 days in duration and non-pharmacological treatments). Censoring was applied to treatments not stopped at the moment corresponding to the last study visit. Response: MADRS improvement from baseline  $\geq 50\%$  or MADRS score  $> 10$ , or (if MADRS score unavailable) CGI-C score  $\leq 2$  or CGI-S score  $\leq 2$ ; no response: MADRS improvement from baseline  $< 50\%$  and MADRS score  $> 10$ , or (if MADRS score unavailable) CGI-C score  $> 2$  or CGI-S score  $> 2$ . CGI-C: Clinical Global Impression of Change; CGI-S: Clinical Global Impression of Severity; MADRS: Montgomery-Åsberg Depression Rating Scale.

from residual symptoms (Mendlewicz, 2008; Rush et al., 2006a).

Despite high levels of non-response, patients continued with pharmacological treatments for long time periods. The mean length of time on the baseline treatment strategy was 220.1 days and after one year, 60.0% of patients were still on their first treatment strategy since entering the study. Given all patients in this study had already experienced at least two treatment failures prior to enrolling, and their continued lack of response, this finding is unexpected. The factors that contributed to the continuation of treatments by patients for such long periods in this study are not clear. One possibility is that treating physicians have low expectations of added treatment responses. Following change of treatment strategy, low levels of patient response were observed, with only 20.3–25.8% of patients responding. This was despite a wide range of different treatment strategies employed, further indicating the inefficacy of treatment strategy alternatives in this population.

In the European region, there are few specific recommendations for medication strategies to treat TRD as market approval submissions for most MDD treatments have not included efficacy and safety studies on the TRD subpopulation. An extended-release formulation of quetiapine is indicated in the EU for use as an add-on treatment for patients with MDD in which a first antidepressant has failed (European Medicines Agency, 2020). In December 2019, esketamine nasal spray, an N-methyl-D-aspartate receptor (NMDAR) antagonist and new mechanism of action, was granted EU market approval for the treatment of TRD in adults, when used in combination with either an SSRI or a SNRI (European Medicines Agency, 2020). The same drug had already obtained FDA approval for use in combination with an oral antidepressant for TRD earlier that same year (FDA, 2019). In the US, an olanzapine/fluoxetine hydrochloride combination was granted FDA approval for the treatment of TRD in 2009 (FDA, 2020) and aripiprazole and brexpiprazole are also approved in this indication (FDA, 2020; FDA, 2018). However, none of these treatments have EMA approval for use in the

TRD subpopulation. Since the 1950s, pharmacological treatments for MDD have targeted the monoamine pathway (Hillhouse and Porter, 2015), but the results described here suggest that, for many patients with TRD, treatment strategies involving these drugs may be ineffective. The poor HRQoL experienced by TRD patients in whom treatment continues to be ineffective points to an urgent need for investigators to develop alternative treatments for TRD and investigate new treatment strategies. A pipeline of drugs to treat MDD are in development that target the glutamate pathway. These alternative mechanisms of action may open new possibilities for treating TRD, if there is sufficient evidence to support the need for efficacy and safety trials that include the TRD subpopulation as a separate entity.

The data presented here, together with those previously described on HRQoL and functionality in TRD in the first paper published from this study (Heerlein et al., 2021) demonstrate the substantial impact of TRD on patients and society. These data add to the body of real-world evidence demonstrating that treatment strategies currently employed in routine clinical practice in Europe, lack efficacy in most patients with TRD.

To allow more in-depth analysis of the relationship between different treatment strategies used in routine clinical practice and clinical outcomes for patients with TRD, it would be valuable to conduct larger-scale observational studies with a similar design to the current study. As more treatments gain market approval in Europe for TRD specifically, it will be important to assess how these impact patient outcomes in a real-world setting, beyond the controlled environment of clinical trials. Newer treatments require safety and efficacy studies in well-defined TRD cohorts to support market approval and improve access to potentially beneficial treatments for these patients.

#### 4.1. Limitations

The limitations of the current study include its relatively small size



compared with other studies, and the absence of a control group to allow comparison of the cohort with patients not starting on a new treatment strategy. For analysis of treatment strategies, a small proportion of outcomes were assigned based on CGI rather than MADRS scores. Furthermore, since the number of treatment lines failed prior to enrolment varies across the cohort, the number of treatment changes in the study does not relate directly to the total number of treatments a patient had been prescribed during the current MDE. As expected in a real-world study, patients for whom data were available for analysis decreased over time, but this was for many reasons and only a small number were true patient drop out. Of note, the study design meant that the later a patient enrolled in the study, the less likely they were to reach 12 months before the study end date; this was the case for approximately 70% of patients. A small proportion of Month 6 or Month 12 visits did not happen within the post-study defined cut-off dates, and in those that did, some did not have a MADRS score (or other outcome measures) recorded at one or more visits. Furthermore, some assessments and/or questionnaires were not completed fully, so numbers of responses for some outcome measures vary. A number of patients were lost to follow up after attending only some visits, possibly due to reduced functioning caused by TRD. Thus, overall, patient numbers are much lower for analysis of Month 12 data than for data collected at Month 6, and results for Month 12 should therefore be interpreted with caution. Finally, comparisons of patient subgroups did not take confounding variables, such as country of origin, into account. Importantly, since patient numbers in the study were approximately proportional to the population of each country, this led to substantial variations in absolute numbers across the countries studied.

## 5. Conclusions

In this study, patients with TRD were treated with many different treatment strategies suggesting that there is no consensus on the standard of care to be used in this population. Despite the wide range of treatments used in these patients, treatment response rates were low, regardless of the number of treatment steps tried, indicating that overall treatment outcomes for patients with TRD are poor. Additionally, data presented here suggest that patients may spend a substantial amount of time on each treatment without a response.

## Authors' contributions

Substantial contributions to study conception and design: KH, GP, WH, AJOM, VPS, SR, GR, SM, JM, CV, BG, AHY; substantial contributions to acquisition, analysis or interpretation of the data: KH, GP, CO, TF, GD, WH, AJOM, VPS, SR, GR, SP, SM, JM, CV, BG, AHY; drafting the article or revising it critically for important intellectual content: KH, GP, CO, TF, GD, WH, AJOM, VPS, SR, GR, SP, SM, JM, CV, BG, AHY; final approval of the version of the article to be published: KH, GP, CO, TF, GD, WH, AJOM, VPS, SR, GR, SP, SM, JM, CV, BG, AHY.

## Data sharing statement

Janssen EMEA's Data Sharing Policy does not include non-interventional studies.

## Funding

This study was sponsored by Janssen EMEA. This article was based on the original study 54135419DEP4001 sponsored by Janssen EMEA. Support for third-party writing assistance for this article, provided by Julia Stevens, Ph.D., and Emma Phillips, Ph.D., Costello Medical, UK, was funded by Janssen EMEA in accordance with Good Publication Practice (GPP3) guidelines (<http://www.ismpp.org/gpp3>).

## Disclosures

KH: Employee of Janssen EMEA

GP: Grant/research support from Angelini; speaker/consultant for Angelini, Janssen, Lundbeck, Neuraxpharm, Sanofi Aventis

CO: Speaker/consultant for Allergan, Ferring, Fortbildungskolleg, Limes Kliniken, MedOnline, Medical Tribune, Neuraxpharm, Sage Therapeutics, and Stillachhaus; Research funding from Deutsche Forschungsgemeinschaft, German Federal Ministry of Education and Research and European Union

TF: Speaker for Janssen-Cilag and Recordati

GD: Consultant/speaker for AstraZeneca, BMS, Eli Lilly, EuroGenerics, GSK, Janssen, Lundbeck, Pfizer and Sanofi

WH: Consultant for Janssen

AJOM: Grants from Schuhfried GmbH, Janssen and Compass Pathways, Ltd; investigator-driven research funded by Fundação para Ciência e Tecnologia (PTDC/MED-NEU/31331/2017), Fundação para Ciência e Tecnologia and FEDER (FCT-PTDC/MEC-PSQ/30302/2017-IC&DTLISBOA-01-0145-FEDER), the European Commission Horizon 2020 program (H2020-SC1-2017-CNECT-2-777167-BOUNCE; H2020-SC1-DTH-2019-875358-FAITH) and the European Research Council (grant agreement 950357)

VPS: Consultancy fees, honoraria or grants from AB-Biotics, AstraZeneca, Bristol-Myers-Squibb, CIBERSAM, Esteve, FIS-ISCiii, Janssen, Lundbeck, Otsuka, Pfizer and Servier

SR: Consultancy fees from Janssen, Lundbeck and Otsuka; grants from Janssen

GR: Speaker/consultant for Angelini, Innova Pharma, Janssen, Lundbeck and Otsuka

SP: Speaker/consultant for Angelini, Janssen, Lundbeck and Sanofi

SM: Employee of Janssen EMEA

JM: Employee of Janssen EMEA

CV: Employee of Janssen EMEA

BG: Employee of Janssen EMEA

AHY: Grants from Janssen; speaker/consultant for Allergan, AstraZeneca, Bionomics, Eli Lilly, Janssen, Johnson & Johnson, Livanova, Lundbeck, Servier, Sumitomo Dainippon Pharma and Sunovion; independent research is funded by the National Institute for Health Research (NIHR) Biomedical Research Centre at South London and Maudsley NHS Foundation Trust and King's College London. The views expressed are those of the authors and not necessarily those of the NHS, the NIHR, or the Department of Health

## Principal investigators and associated study sites

Belgium: Degraeve, Gunther: Private Practice, G. Degraeve; Domken, Marc-André: ISO SL - Site du Petit Bourgogne; Geerts, Stefaan: A. Z. St-Lucas; Gillain, Benoit: Clinique Saint-Pierre; Hauwaert, An: Hauwaert An; Kornreich, Charles: C.H.U. Brugmann - Site Victor Horta; Scantamburlo, Gabrielle: CHU de Liege; Souery, Daniel: Psy pluriel (Uccle); Vandewalle, Ward: St-Andries Ziekenhuis; Germany: Adli, Mazda: Fliedner Klinik Berlin; Barth, Thomas: Klinikum Chemnitz gGmbH; Benes, Heike: Somni Bene GmbH; Bodenschatz, Ralf: Pharmakologisches Studienzentrum Chemnitz GmbH; Cindik-Herbrueggen, Elif: NPZR - Neuropsychiatrisches Zentrum Riem; Englisch, Susanne: Universitaetsmedizin der Johannes Gutenberg-Universitaet Mainz; Frodl, Thomas: Universitaetsklinikum Magdeburg A.o.e.R; Hädrich, Florian: Privat-nerven-klinik Dr. Med. Kurt Fontheim; Hahn, Kirsten: Praxis Dr. med. Kirsten Hahn; Hajak, Goeran: Sozialstiftung Bamberg; Kuehn, Frank: Praxis Kuehn; Kuhn, Jens: Johanniter Krankenhaus Oberhausen; Kusserow, Stefan: Praxis Dr. Stefan Kusserow; Otte, Christian: Charite - Campus Benjamin Franklin; Reif, Andreas: Klinikum der Johann Wolfgang Goethe-Universitaet; Sallach, Klaus: Gemeinschaftspraxis f. Neurologie, Psychiatrie und Psychotherapie Dres. Leonhardt u. Sallach; Schaefer, Martin: Kliniken Essen-Mitte; Schlegel, Eugen: Zentrum f. Neurologisch-Psychiatrische Studien und Begutachtung; Schulze,

Alexander: Praxis Dr. sc. med. Alexander Schulze; Thakkar, Sarang: Asklepios Klinik Nord-Ochsenzoll; Thomsen, Jana: Praxis Dr. med. Jana Thomsen; Italy: Amore, Mario: Azienda Ospedaliero Universitaria San Martino; Bellomo, Antonello: Azienda Ospedaliero Universitaria Ospedali Riuniti di Foggia; Bertolino, Alessandro: Azienda Ospedaliero Universitaria Consorziata Policlinico di Bari; Biondi, Massimo: Azienda Ospedaliero Universitaria Policlinico Umberto I - Università di Roma La Sapienza; Bondi, Emi: Azienda Socio Sanitaria Territoriale Papa Giovanni XXIII (Presidio Papa Giovanni XXIII); Bosi, Monica: Azienda Socio Sanitaria Territoriale Fatebenefratelli (Presidio Ospedale Sacco); Brambilla, Paolo: Fondazione IRCCS CA' Granda Ospedale Maggiore Policlinico; Clerici, Massimo: Azienda Socio Sanitaria Territoriale di Monza (Presidio San Gerardo); De Fazio, Pasquale: Azienda Ospedaliero Universitaria Mater Domini; De Filippis, Sergio: Casa di Cura Villa Von Siebenthal; De Giorgi, Serafino: AUSL LE di Lecce; Fagiolini, Andrea: A. O. U. Senese Policlinico Santa Maria alle Scotte; Janiri, Luigi: Fondazione Policlinico Universitario Agostino Gemelli IRCCS; Marchesi, Carlo: Azienda Unità Sanitaria Locale di Parma - Ospedale Maggiore; Muscatello, Maria Rosaria Anna: Azienda Ospedaliero Universitaria Policlinico G. Martino; Perugi, Giulio: Azienda Ospedaliero Universitaria Pisana; Petralia, Antonino: Azienda Ospedaliero Universitaria "Policlinico - Vittorio Emanuele" (Presidio Gaspare Rodolico); Rosso, Gianluca: Azienda Ospedaliero-Universitaria S. Luigi Gonzaga; Sani, Gabriele: Azienda Ospedaliero Sant'Andrea-Università di Roma La Sapienza; Vaggi, Marco: Azienda Sanitaria 3 Genovese; Vita, Antonio: Azienda Socio Sanitaria Territoriale degli Spedali Civili di Brescia (Presidio Spedali Civili); Zeppegno, Patrizia: Azienda Ospedaliero - Universitaria Maggiore delle Carità; Netherlands: Hagedoorn, Wolter: Praktijk voor Psychiatrie en Psychotherapie; Schlösser, Rutger: Psychiatriepraktijk Helmind; Witte, Roel: MAPTA; Portugal: Alcaface, João: Centro Hospitalar do Baixo Vouga, E.P.E. - Unidade de Aveiro; Bessa, João: Hospital de Braga; Fonseca, Sofia: Centro Hospitalar de Leiria; Freitas, João: Hospital Magalhães Lemos, EPE; Macedo, António: Centro Hospitalar e Universitário de Coimbra E.P.E - Hospitais da Universidade de Coimbra; Lara, Elsa: CUF - Infante Santo; Oliveira-Maia, Albino J: Fundação Champalimaud; Matos Pires, Ana: Unidade Local de Saúde do Baixo Alentejo, EPE; Serra, Madalena: Hospital Espírito Santo, EPE; Von Doellinger, Orlando: Centro Hospitalar do Tâmega e Sousa, EPE - Hospital Padre Americo, Vale do Sousa; Spain: Baca Garcia, Enrique: Fundacion Jimenez Diaz; Bobes Garcia, Julio Belarmino: CS Mental La Corredoria; Caballero, Luis: Hospital Universitario HM Puerta del Sur; Cardoner Alvarez, Narcis: Corporacio Sanitaria Parc Tauli; Gomez Carreno, Carlos Rodriguez: Hospital General Universitario de Ciudad Real; Hernandez Fleta, Jose Luis: Complejo Hospitalario Universitario de Gran Canaria Dr. Negrin; Menchon Magriña, Jose Manuel: Hospital Universitario de Bellvitge; Mesones Peral, Jesus Enrique: Hospital de Torrevieja; Perez Sola, Victor: Hospital del Mar; Sarro Maluquer, Salvador: Consulta Dr Salvador Sarro; Sierra San Miguel, Pilar: Hospital Universitario i Politecnico La Fe; Vazquez Noguero Mendez, Raul: Hospital Nicolas Peña; Vieta, Eduard: Hospital Clinic de Barcelona; Villanueva, Rosa: CSM Fuencarral; United Kingdom: Ahmed, Rais: Royal Derby Hospital; Anjum, Rubina: Burntwood and Lichfield CMHT; Gupta, Sumeet: West Park Hospital; Laugharne, Richard: Cornwall Learning Disabilities Service; Lawrence, Robert: Barnes Hospital; Lawrence, Ward: Abraham Cowley Unit; Macintyre, Donald: Royal Edinburgh Hospital; O'Neill-Kerr, Alexander: Berrywood Hospital; Rathod, Shanaya: Royal South Hants Hospital; Robinson, Andrew: Royal Cornhill Hospital; Sivasanker, Vimal: Kingfisher Court; Tremblay, Micheline: Vale House; Walters, Paul: Westhaven Hospital; Young, Allan H: Institute of Psychiatry

## Acknowledgements

The authors thank the patients, the investigators and their teams who took part in this study. The authors also acknowledge Julia Stevens, Ph.D., and Emma Phillips, Ph.D., from Costello Medical, UK, for medical

writing and editorial assistance based on the authors' input and direction, Yerkebulan Kambarov for publication coordination, and GAMIAN-Europe for their review and input into the manuscript. This study was funded by Janssen EMEA.

## Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.jad.2021.03.073.

## References

- Aaronson, S.T., Sears, P., Ruvuna, F., Bunker, M., Conway, C.R., Dougherty, D.D., Reimherr, F.W., Schwartz, T.L., Zajecka, J.M., 2017. A 5-Year Observational Study of Patients With Treatment-Resistant Depression Treated With Vagus Nerve Stimulation or Treatment as Usual: Comparison of Response, Remission, and Suicidality. *Am J Psychiatry* 174, 640–648.
- Al-Harbi, K.S., 2012. Treatment-resistant depression: therapeutic trends, challenges, and future directions. *Patient preference and adherence* 6, 369–388.
- Amos, T.B., Tandon, N., Lefebvre, P., Pilon, D., Kamstra, R.L., Pivneva, I., Greenberg, P.E., 2018. Direct and Indirect Cost Burden and Change of Employment Status in Treatment-Resistant Depression: A Matched-Cohort Study Using a US Commercial Claims Database. *J Clin Psychiatry* 79.
- Barowsky, J., Schwartz, T.L., 2006. An Evidence-Based Approach to Augmentation and Combination Strategies for: Treatment-Resistant Depression. *Psychiatry (Edmont)* 3, 42–61.
- Bauer, M., Dell'osso, L., Kasper, S., Pitchot, W., Dencker-Ransvik, E., Köhler, J., Jørgensen, L., Montgomery, S.A., 2013. Extended-release quetiapine fumarate (quetiapine XR) monotherapy and quetiapine XR or lithium as add-on to antidepressants in patients with treatment-resistant major depressive disorder. *J Affect Disord* 151, 209–219.
- Bennabi, D., Charpeaud, T., Yroni, A., Genty, J.B., Destouches, S., Lancrenon, S., Alaïli, N., Bellivier, F., Bougerol, T., Camus, V., Dorey, J.M., Doumy, O., Haesebaert, F., Holtzmann, J., Lançon, C., Lefebvre, M., Moliere, F., Nieto, I., Rabu, C., Richieri, R., Schmitt, L., Stephan, F., Vaiva, G., Walter, M., Leboyer, M., El-Hage, W., Llorca, P.M., Courtet, P., Auquier, B., Haffen, E., 2019. Clinical guidelines for the management of treatment-resistant depression: French recommendations from experts, the French Association for Biological Psychiatry and Neuropsychopharmacology and the foundation FondaMental. *BMC Psychiatry* 19, 262.
- Busner, J., Targum, S.D., 2007. The clinical global impressions scale: applying a research tool in clinical practice. *Psychiatry (Edmont)* 4, 28–37.
- Cavanagh, J.T., Carson, A.J., Sharpe, M., Lawrie, S.M., 2003. Psychological autopsy studies of suicide: a systematic review. *Psychol Med* 33, 395–405.
- Chandler, G.M., Iosifescu, D.V., Pollack, M.H., Targum, S.D., Fava, M., 2010. Validation of the Massachusetts General Hospital Antidepressant Treatment History Questionnaire (ATRQ). *CNS Neurosci Ther* 16, 322–325.
- Cleare, A., Pariante, C.M., Young, A.H., Anderson, I.M., Christmas, D., Cowen, P.J., Dickens, C., Ferrier, I.N., Geddes, J., Gilbody, S., Haddad, P.M., Katona, C., Lewis, G., Malizia, A., McAllister-Williams, R.H., Ramchandani, P., Scott, J., Taylor, D., Uher, R., Members of the Consensus Meeting, 2015. Evidence-based guidelines for treating depressive disorders with antidepressants: A revision of the 2008 British Association for Psychopharmacology guidelines. *J Psychopharmacol*, 29, 459–525.
- Conway, C.R., George, M.S., Sackeim, H.A., 2017. Toward an Evidence-Based, Operational Definition of Treatment-Resistant Depression: When Enough Is Enough. *JAMA Psychiatry* 74, 9–10.
- Dunner, D.L., Rush, A.J., Russell, J.M., Burke, M., Woodard, S., Wingard, P., Allen, J., 2006. Prospective, long-term, multicenter study of the naturalistic outcomes of patients with treatment-resistant depression. *J Clin Psychiatry* 67, 688–695.
- European Medicines Agency. 2020. Esketamine Summary of Product Characteristics. <https://www.ema.europa.eu/en/medicines/human/EPAR/spravato>. (Accessed 28 August 2020).
- European Medicines Agency. 2013. Guideline on clinical investigation of medicinal products in the treatment of depression. EMA/CHMP/185423/2010 Rev 2.
- European Medicines Agency. 2020. Seroquel SmPC, labelling and package leaflet. <https://www.ema.europa.eu/en/documents/referral/seroquel-seroquel-xr-associate-d-names-article-30-referral-annex-iii-en.pdf> (Accessed 20 December 2020).
- European Medicines Agency. 2019. Spravato EPAR Product Characteristics. [https://www.ema.europa.eu/en/documents/product-information/spravato-epar-product-information\\_en.pdf](https://www.ema.europa.eu/en/documents/product-information/spravato-epar-product-information_en.pdf). (Accessed 31 March 2021).
- Fava, M., Okame, T., Matsushima, Y., Perry, P., Weiller, E., Baker, R.A., 2017. Switching from Inadequate Adjunctive or Combination Treatment Options to Brepiprazole Adjunctive to Antidepressant: An Open-Label Study on the Effects on Depressive Symptoms and Cognitive and Physical Functioning. *Int J Neuropsychopharmacol* 20, 22–30.
- FDA. 2019. Esketamine US Prescribing Information, March 2019. Available at: [https://www.accessdata.fda.gov/drugsatfda\\_docs/label/2019/211243lbl.pdf](https://www.accessdata.fda.gov/drugsatfda_docs/label/2019/211243lbl.pdf). (Accessed 28 August 2020).
- FDA. 2017. Symbyax Prescribing Information. [https://www.accessdata.fda.gov/drugsatfda\\_docs/label/2017/021520s042lbl.pdf](https://www.accessdata.fda.gov/drugsatfda_docs/label/2017/021520s042lbl.pdf). (Accessed 14 April 2021).

- FDA. 2018. Rexulti Prescribing Information. Available at [https://www.accessdata.fda.gov/drugsatfda\\_docs/label/2018/205422s003lbl.pdf](https://www.accessdata.fda.gov/drugsatfda_docs/label/2018/205422s003lbl.pdf). (Accessed 20th December 2020).
- FDA. 2020. Ablify Prescribing Information. [https://www.accessdata.fda.gov/drugsatfda\\_docs/label/2020/021436s044s045,021713s035s036,021729s027s028,021866s029s030lbl.pdf](https://www.accessdata.fda.gov/drugsatfda_docs/label/2020/021436s044s045,021713s035s036,021729s027s028,021866s029s030lbl.pdf). (Accessed 20 December 2020).
- Fonagy, P., Rost, F., Carlyle, J.A., Mcpherson, S., Thomas, R., Pasco Fearon, R.M., Goldberg, D., Taylor, D., 2015. Pragmatic randomized controlled trial of long-term psychoanalytic psychotherapy for treatment-resistant depression: the Tavistock Adult Depression Study (TADS). *World Psychiatry* 14, 312–321.
- Heerlein, K., Young, A.H., Otte, C., Frodl, T., Degraeve, G., Hagedoorn, W., Oliveira-Maia, A.J., Perez Sola, V., Rathod, S., Rosso, G., Sierra, P., Morrens, J., Van Dooren, G., Gali, Y., Perugi, G., 2021. Real-world evidence from a European cohort study of patients with treatment resistant depression: Baseline patient characteristics. *J Affect Disorders* 283, 115–122.
- Herdman, M., Gudex, C., Lloyd, A., Janssen, M., Kind, P., Parkin, D., Bonnel, G., Badia, X., 2011. Development and preliminary testing of the new five-level version of EQ-5D (EQ-5D-5L). *Qual Life Res* 20, 1727–1736.
- Hillhouse, T.M., Porter, J.H., 2015. A brief history of the development of antidepressant drugs: from monoamines to glutamate. *Exp Clin Psychopharmacol* 23, 1–21.
- Ionescu, D.F., Rosenbaum, J.F., Alpert, J.E., 2015. Pharmacological approaches to the challenge of treatment-resistant depression. *Dialogues Clin Neurosci* 17, 111–126.
- Jaffe, D.H., Rive, B., Denece, T.R., 2019. The humanistic and economic burden of treatment-resistant depression in Europe: a cross-sectional study. *BMC Psychiatry* 19, 247.
- James, S.L., Abate, D., Abate, K.H., Abay, S.M., Abbafati, C., Abbasi, N., Abbastabar, H., Abd-Allah, F., et al., 2018. Global, regional, and national incidence, prevalence, and years lived with disability for 354 diseases and injuries for 195 countries and territories, 1990–2017: a systematic analysis for the Global Burden of Disease Study 2017. *The Lancet* 392, 1789–1858.
- Kim, T., Xu, C., Amsterdam, J.D., 2019. Relative effectiveness of tricyclic antidepressant versus monoamine oxidase inhibitor monotherapy for treatment-resistant depression. *Journal of Affective Disorders* 250, 199–203.
- Kubitz, N., Mehra, M., Potluri, R.C., Garg, N., Cossrow, N., 2013. Characterization of treatment resistant depression episodes in a cohort of patients from a US commercial claims database. *PLoS One* 8, e76882.
- Lefaucheur, J.P., Aleman, A., Baeken, C., Benninger, D.H., Brunelin, J., Di Lazzaro, V., Filipović, S.R., Grefkes, C., Hasan, A., Hummel, F.C., Jääskeläinen, S.K., Langguth, B., Leocani, L., Londero, A., Nardone, R., Nguyen, J.P., Nyffeler, T., Oliveira-Maia, A.J., Oliviero, A., Padberg, F., Palm, U., Paulus, W., Poulet, E., Quartarone, A., Rachid, F., Rektorová, I., Rossi, S., Sahlsten, H., Schecklmann, M., Szekely, D., Ziemann, U., 2020. Evidence-based guidelines on the therapeutic use of repetitive transcranial magnetic stimulation (rTMS): An update (2014–2018). *Clin Neurophysiol* 131, 474–528.
- Lisanby, S.H., 2007. Electroconvulsive therapy for depression. *N Engl J Med* 357, 1939–1945.
- Macqueen, G., Santaguida, P., Keshavarz, H., Jaworska, N., Levine, M., Beyene, J., Raina, P., 2017. Systematic Review of Clinical Practice Guidelines for Failed Antidepressant Treatment Response in Major Depressive Disorder, Dysthymia, and Subthreshold Depression in Adults. *Can J Psychiatry* 62, 11–23.
- Mahase, E., 2019. Esketamine is approved in Europe for treating resistant major depressive disorder. *BMJ* 367, 17069.
- Mendlewicz, J., 2008. Towards achieving remission in the treatment of depression. *Dialogues Clin Neurosci* 10, 371–375.
- Montgomery, S.A., Asberg, M., 1979. A new depression scale designed to be sensitive to change. *Br J Psychiatry* 134, 382–389.
- National Institute for Health and Care Excellence 2020. Clinical guideline [CG90]: The NICE Guideline on the Treatment and Management of Depression in Adults. Available from <https://www.nice.org.uk/guidance/cg90/evidence/full-guideline-pdf-4840934509>. (Accessed 6th December 2020).
- Nemeroff, C.B., 2007. Prevalence and management of treatment-resistant depression. *J Clin Psychiatry* 68 (Suppl 8), 17–25.
- Posner, K., Brown, G.K., Stanley, B., Brent, D.A., Yershova, K.V., Oquendo, M.A., Currier, G.W., Melvin, G.A., Greenhill, L., Shen, S., Mann, J.J., 2011. The Columbia-Suicide Severity Rating Scale: initial validity and internal consistency findings from three multisite studies with adolescents and adults. *Am J Psychiatry* 168, 1266–1277.
- Reilly, M.C., Zbrozek, A.S., Dukes, E.M., 1993. The validity and reproducibility of a work productivity and activity impairment instrument. *Pharmacoeconomics* 4, 353–365.
- Rush, A.J., Kraemer, H.C., Sackeim, H.A., Fava, M., Trivedi, M.H., Frank, E., Ninan, P.T., Thase, M.E., Gelenberg, A.J., Kupfer, D.J., Regier, D.A., Rosenbaum, J.F., Ray, O., Schatzberg, A.F., 2006a. Report by the ACNP Task Force on response and remission in major depressive disorder. *Neuropsychopharmacology* 31, 1841–1853.
- Rush, A.J., Trivedi, M.H., Wisniewski, S.R., Nierenberg, A.A., Stewart, J.W., Warden, D., Niederehe, G., Thase, M.E., Lavori, P.W., Lebowitz, B.D., Mcgrath, P.J., Rosenbaum, J.F., Sackeim, H.A., Kupfer, D.J., Luther, J., Fava, M., 2006b. Acute and longer-term outcomes in depressed outpatients requiring one or several treatment steps: a STAR\*D report. *Am J Psychiatry* 163, 1905–1917.
- Sheehan, D.V., Harnett-Sheehan, K., Raj, B.A., 1996. The measurement of disability. *Int Clin Psychopharmacol* 11 (Suppl 3), 89–95.
- Souery, D., Amsterdam, J., De Montigny, C., Lecrubier, Y., Montgomery, S., Lipp, O., Racagni, G., Zohar, J., Mendlewicz, J., 1999. Treatment resistant depression: methodological overview and operational criteria. *Eur Neuropsychopharmacol* 9, 83–91.
- Tobe, E.H., Rybakowski, J.K., 2013. Possible usefulness of tianeptine in treatment-resistant depression. *Int J Psychiatry Clin Pract* 17, 313–316.
- Turrina, C., Vita, A., Sacchetti, E. 2015. Remission in depression after treatment: Too obvious to clinicians, so why so difficult to measure? Available from [https://www.evidence-based-psychiatric-care.org/wp-content/uploads/2015/11/06\\_remission\\_EVIDENCE\\_BASED.pdf](https://www.evidence-based-psychiatric-care.org/wp-content/uploads/2015/11/06_remission_EVIDENCE_BASED.pdf). (Accessed 17th March 2021).
- Van Bronswijk, S., Moopen, N., Beijers, L., Ruhe, H.G., Peeters, F., 2019. Effectiveness of psychotherapy for treatment-resistant depression: a meta-analysis and meta-regression. *Psychol Med* 49, 366–379.
- Voineskos, D., Daskalakis, Z.J., Blumberger, D.M., 2020. Management of Treatment-Resistant Depression: Challenges and Strategies. *Neuropsychiatric disease and treatment* 16, 221–234.
- Wiles, N., Thomas, L., Abel, A., Barnes, M., Carroll, F., Ridgway, N., Sherlock, S., Turner, N., Button, K., Odoni, L., Metcalfe, C., Owen-Smith, A., Campbell, J., Garland, A., Hollinghurst, S., Jerrom, B., Kessler, D., Kuyken, W., Morrison, J., Turner, K., Williams, C., Peters, T., Lewis, G., 2014. Clinical effectiveness and cost-effectiveness of cognitive behavioural therapy as an adjunct to pharmacotherapy for treatment-resistant depression in primary care: the CoBaIT randomised controlled trial. *Health Technol Assess* 18, 1–167.