Personalizing depression treatment:
Approximately one-half of patients with major depressive disorder (MDD) will have partial or nonresponse to first-line antidepressant monotherapy, despite receiving an adequate dosage and sufficient duration of treatment. This has led to the definition of treatment-resistant depression (TRD) as a depressive episode that has shown insufficient response to ≥1 trial of an antidepressant that has demonstrated efficacy in clinical trials. Depressed patients should be treated to full remission because absence of complete remission is associated with:

- a more recurrent and chronic illness course
- increased medical and psychiatric comorbidities
- greater functional burden
- increased social and economic costs linked with impaired social functioning.

Clinicians need to properly identify MDD and treatment resistance to guide optimal treatment choices. Additional tools are necessary to accurately identify, document, and communicate about symptoms commonly experienced by depressed patients but not fully characterized by DSM-IV-TR MDD criteria. Finally, in many cases, trait or situational factors might obfuscate accurate diagnosis and the natural course of illness, and tools that can be implemented practically will help identify patients with MDD.

Our group has created and implemented 2 clinician-administered tools—the SAFER Interview and the Antidepressant Treatment Response Questionnaire (ATRQ)—to enrich the qualitative assessment of MDD and treatment resistance.

SAFER: Assessing the diagnosis, symptom severity
The SAFER interview refines the diagnosis of depressed patients by assessing the state vs trait nature of the symptoms, their assessability, their face and ecological validity, and if they pass the rule of the 3...
Ps: pervasiveness, persistence, and pathological nature of the current MDD episode (Table 1 and Table 2). This reliable assessment of the patient’s diagnosis and symptom severity is made in a way that reflects the illness in a real-world setting.

Clinical application of SAFER. Implementing SAFER in clinical settings promotes a personalized, dimensional approach by taking into account a varying degree of symptom severity in depressed patients, in contrast to relying on symptom lists as found in the DSM-IV-TR. Using the SAFER interview deepens the typical psychiatric diagnostic process, allows for a more precise understanding of the patient’s situation, and may help clinicians select effective treatments that target specific symptoms, thus resulting in more rapid alleviation of MDD.

Using SAFER to evaluate Ms. Y, a clinician would ask: Does she have symptoms that are present primarily during episodes of acute illness? Does the episode constitute a measurable exacerbation of preexisting symptoms? This clinical vignette illustrates the importance of the first SAFER criterion, state vs trait nature of the symptoms. Ms. Y is a SAFER “pass”—meaning consistent with a major depressive episode—because exacerbation of preexisting symptoms is measurable. However, if her symptoms represented a chronic, long-standing trait, she would be a SAFER “fail” based on this criterion, and her symptoms likely would not improve during a brief pharmacologic intervention. For such patients, SAFER would have oriented the clinician toward alternative therapies such as psychotherapy or a combination of longer, more complex pharmacologic treatment and psychotherapy.

By helping clinicians refine MDD diagnoses, SAFER draws attention to specific MDD symptoms as assessed by the Hamilton Depression Rating Scale, eg, ongoing depressive mood, feelings of guilt, major sleep disturbances, and work impairments.
depression entities (eg, MDD or dysthymia) and directs them toward appropriate guidelines, treatment algorithms, and precautions (eg, when treated with pharmacotherapy, dysthymic patients are particularly sensitive to unwanted effects and adherence is a serious issue).7 The SAFER interview also can help identify when what appears to be a depressive episode clearly is precipitated by situational factors, and is more likely to be influenced by changes in the patient’s life than by treatment. For such patients, first consider nonpharmacologic interventions to avoid unnecessary medication exposure.

**ATRQ: Efficacy and adequacy**

The ATRQ examines the efficacy (improvement from 0% [not improved at all] to 100% [completely improved]), and adequacy (adequate duration and dose) of any antidepressant treatment in a step-by-step procedure.1,8,9 For a copy of the ATRQ, visit this article at CurrentPsychiatry.com.

While conducting the interview, clinicians ask about treatment adherence to each medication trial. A treatment-resistant patient may go through many types of trials, from monotherapy to combination to augmentation.10 For each trial, the ATRQ systematically reviews 4 strategies to enhance treatment response:

- increasing the initial antidepressant dosage11
- combining the initial antidepressant with another antidepressant, typically from another class12
- augmenting the initial antidepressant with a nonantidepressant12
- switching from the initial antidepressant to another antidepressant.13

These strategies also are applied in cases of lost sustained antidepressant efficacy or depressive relapse/recurrence, although empirical evidence supporting these strategies is lacking, with the possible exception of dose increase.14,15

In the convention our group adopted, an adequate antidepressant trial must be ≥6 weeks in total length, with a dose within an adequate range as specified in the medication’s package insert. In addition, for the purposes of conducting TRD trials, we have considered a patient treatment-resistant if response to adequate dose and duration is <50%. On the ATRQ, 50% improvement refers to 50% symptom reduction from baseline without achieving remission. In an initial clinical trial that lasts ≥6 weeks, any dose increase (for ≥4 weeks) represents optimization and is not considered a new or separate trial, whereas augmentation or combination therapy (for ≥3 weeks) or a switch to another antidepressant (for ≥6 weeks) are considered new trials/treatments.

**Decision tree for ATRQ.** Because antidepressant treatment always is constrained by personal (eg, adherence, insurance coverage, etc.) and clinical (eg, contraindications due to comorbid conditions, side effects, etc.) considerations, we propose a decision tree to help clinicians determine the number of failed antidepressant trials their patient experienced (Figure, page 30). Although this tree does not represent all treatment scenarios, we hope it could help clinicians implement TRD treatment strategies because it highlights proper assessment of treatment duration, dosage, and changes before applying a TRD diagnosis.

The ATRQ meticulously examines patients’ antidepressant history to identify:

- pseudo-resistance, to guide adequate dosing and/or duration, and

Pseudo-resistance refers to treatment failures that can be attributed to factors...
such as inadequate treatment dosing or duration, atypical pharmacokinetics that reduce agents’ effectiveness, patient nonadherence (eg, due to adverse effects), or misdiagnosis of the primary disorder (ie, other mood disorders or depressive subsets such as dysthymia or minor depression mistreated as unipolar depression).\textsuperscript{16,17} Studies show that many patients with MDD referred to specialty settings are undertreated and receive inadequate antidepressant doses,\textsuperscript{18} which suggests that many referrals for TRD are in fact pseudo-resistance.\textsuperscript{19} Despite the lack of consensus on criteria for TRD,\textsuperscript{1} standardization of what constitutes treatment adequacy during antidepressant trials (eg, adherence, dose, duration) is indispensable.

**Clinical application of the ATRQ.** Because TRD may require specific interventions, we first need to properly identify treatment resistance. Also, systematic use of a classification system enhances the ability of clinicians and patients to provide meaningful descriptions of antidepressant resistance.

In clinical practice, choice of treatment strategy is based on factors that include partial or nonresponse, tolerability, avoiding withdrawal symptoms, the need to target side effects of a current antidepressant by administering another drug, cost, avoiding drug-drug interactions, and patient preference. Because a treatment-resistant patient may go through many types of trials, it is essential to obtain information about all treatments to determine the number of failed clinical trials a patient may have had for the current MDD episode and lifetime episodes. The importance of asking about adherence to each trial cannot be overemphasized.

**CASE REPORT**

**Limited improvement**

Mr. T, age 45, reported that his current depressive episode started several years ago. The
first antidepressant trial he received, sertraline, 100 mg/d for 3 months, resulted in 0% improvement. Next he received citalopram, 20 mg/d, for 1 month, without any improvement. The next trial, venlafaxine, 75 mg/d, for 7 weeks, resulted in a 30% response. More recently Mr. B tried duloxetine, up to 90 mg/d, for 2 years with an 80% improvement during the first 3 months and then a decrease of response to <40%. He then received aripiprazole, 10 mg/d, in combination with duloxetine for approximately 4 weeks with no response.

Using the ATRQ to evaluate Mr. B, a clinician would consider the sertraline trial as a failed trial. The citalopram treatment would not be considered an adequate trial because it was too short. The venlafaxine course wouldn’t count as an adequate trial because the dosage was too low. The trial with duloxetine would count as a response (80% improvement) followed by a “poop out” or tachyphylaxis (40% improvement). The fifth trial with the combination of duloxetine and aripiprazole would count as a failed trial. The ATRQ highlights which drugs have been used for too short a duration or at too low a dosage. In Mr. B’s case, using the ATRQ revealed that of 5 trials, only 2 showed antidepressant resistance.

The ATRQ and decision tree are meant to provide clinicians with user-friendly tools to more precisely determine the number of failed antidepressant trials a patient experienced. By assessing if an antidepressant trial had an adequate dose and duration, the ATRQ can help suggest the next treatment options. For example, if a trial was inadequate in dose and/or duration but the patient tolerated the medication, then optimizing treatment with the current drug would be a logical next step. If a patient does not respond to an adequate trial, clinicians have several options, such as switching to another antidepressant, using a combination of medications or an augmentation strategy, or increasing the dose of the original antidepressant.

Limitations of the ATRQ. Historical rating of treatment is not as accurate as a prospective trial. Another limitation of the ATRQ is that the minimum effective dose is accepted as adequate; many clinicians would suggest that such a dose is inadequate. Also, the duration specified for augmentation, dose increase, and monotherapy are based on expert consensus rather than systematic research. Nonetheless, this method of documenting prior trials and treatment adequacy is an important advance.

The ATRQ lacks a place to indicate discontinuation due to intolerance. Knowing if adverse events caused treatment nonadherence or discontinuation is relevant to selecting treatment.

The ATRQ considers only pharmacotherapy and electroconvulsive therapy. Comprehensive assessment of treatment resistance requires asking about depression-specific, evidence-based psychotherapies, including cognitive-behavioral therapy and interpersonal therapy. Another precaution is that the ATRQ and SAFER should be used in conjunction with structured or semi-structured clinical interviews and other clinical tools to rule out other primary diagnoses (eg, bipolar disorder, posttraumatic stress disorder, obsessive-compulsive disorder, or surreptitious substance abuse).

Using SAFER and ATRQ allows clinicians to make a proper MDD diagnosis with an accurate, reliable assessment of symptom severity and a precise count of antidepressant trials of adequate duration and dose. SAFER helps differentiate MDD from other depressive disorders and can be used to separate MDD from a similar presentation of a reaction to external circumstances that may remit if these circumstances change.

These 2 clinical tools focus on MDD and TRD. Compared with treatment resistance in MDD, treatment resistance in other forms of depression, such as minor depressive disorder or dysthymia, has been inadequately researched and should be addressed in large studies. However, to help patients achieve remission of depressive disorders—especially MDD—clinicians should use patient information gathered via measurement-based care in combination with algorithm recommendations. Persistence of clinical care of MDD is essential because several treatment steps
might be necessary for some patients to achieve remission. Throughout treatment, patients’ symptoms, adverse events secondary to ongoing treatment, and medication adherence should be assessed with appropriate tools. ATRQ and SAFER offer a clinician support in assessing treatment resistance history and a patient’s likelihood to respond to pharmacologic treatment.

Related Resources


Drug Brand Names

- Aripiprazole - Abilify
- Citalopram - Celexa
- Duloxetine - Cymbalta
- Sertraline - Zoloft
- Venlafaxine - Effexor

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Bottom Line

The SAFER interview promotes a personalized, dimensional approach to depression diagnosis by taking into account varying degrees of symptom severity. The Antidepressant Treatment Response Questionnaire and decision tree allow clinicians to more precisely determine the number of failed antidepressant trials a patient experienced, which can help suggest subsequent treatment options.


Mr. K, age 36, is referred to you by his primary care physician for “treatment-resistant depression.” Despite successive courses of fluoxetine, desvenlafaxine, and bupropion, he continues to experience fatigue, feelings of worthlessness, and depressed mood.

**How would you proceed?**

- [ ] Switch Mr. K to a different selective serotonin reuptake inhibitor
- [ ] Restart desvenlafaxine, 50 mg/d, and augment with aripiprazole, 5 mg/d
- [ ] Determine if the pharmacotherapy he received used an adequate dosage for an adequate duration, and if not, restart one of the initial medications
- [ ] Switch Mr. K to olanzapine/fluoxetine, 6 mg/25 mg

**JANUARY POLL RESULTS**

Ms. W, age 42, presents at the local emergency department (ED) with heart palpitations. Upset because a long-term relationship recently ended, she says she has been depressed, drinking alcohol each night to help her sleep, and considering ending her life. ED physicians rule out an acute medical illness.

**What would you do?**

- [ ] Start her on fluoxetine, 20 mg/d, and refer her for psychotherapy (21%)
- [ ] Recommend that she be admitted to the psychiatric unit (69%)
- [ ] Refer her for alcohol abuse treatment (5%)
- [ ] Prescribe lithium, 300 mg twice a day (5%)

**SUGGESTED READING:**