Risperidone for the Management of Treatment-Resistant Anxiety in a Patient with Ehlers-Danlos Syndrome: A Case Report

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Abstract

Patients with Ehlers-Danlos syndrome (EDS) have a higher prevalence of comorbid anxiety disorders. Due to the complex nature of these patients, there is often a delay in the diagnosis of these disorders as well as insufficient management of their anxiety symptoms. Current treatment options are often insufficient or poorly tolerated by patients, creating barriers to proper treatment. We hypothesized that patients with EDS and comorbid anxiety, who have failed multiple medication trials, may benefit from a trial of risperidone. In this case report, we discuss the successful management of treatment-resistant anxiety in a patient with EDS with the use of risperidone. Management of comorbid anxiety in these patients is essential, as untreated anxiety can result in increased somatic sensation sensitivity and poor social functioning. Once underlying anxiety disorders are addressed, patients with EDS can better cope with chronic pain symptoms and are more likely to build a therapeutic alliance with their treating physicians. This results in an improved prognosis, social functioning, and overall quality of life.

Categories: Psychiatry, Rheumatology
Keywords: clinical anxiety, ehlers danlos syndrome, risperidone

Introduction

Ehlers-Danlos syndrome (EDS) refers to a hereditary group of disorders that affect connective tissue matrix proteins. Characteristics of EDS include joint hypermobility, skin hyperextensibility, and tissue fragility [1]. EDS is a multisystemic disease, affecting the musculoskeletal system, cardiovascular, gastrointestinal, mucocutaneous, urogynecological, ocular, and neuropsychiatric systems. Neuropsychiatric manifestations include proprioception dysfunction, dysautonomia, and anxiety [1].

Literature has demonstrated that there is a strong association between EDS and comorbid psychiatric disorders. Some examples of these disorders include anxiety, depression, eating disorders, and neuro-developmental disorders [2]. In a systematic, retrospective chart review of 391 patients, 49.4% were reported to have a psychiatric disorder, with 28.9% having two or more disorders, and 26.6% having an anxiety disorder [3]. These patients often also suffer from chronic pain, which is associated with an increased incidence of comorbid psychiatric disorders [3]. For example, one study found that 29% of patients with chronic pain also suffer from depression [4]. Furthermore, in rheumatologic patients with joint hypermobility syndrome, also known as Ehlers-Danlos III, 70% of patients were found to have some kind of anxiety disorder earlier in life [4]. Overall, these patients were ten times more likely to suffer from anxiety than patients without joint hypermobility [4].

Anxiety disorders in patients with EDS and joint hypermobility syndrome often go unrecognized, and despite the high prevalence of anxiety disorders in patients with EDS, less than 25% of patients receive treatment [4]. Current treatments for anxiety disorders in these patients include cognitive-behavioral therapy (CBT) for the management of comorbid chronic pain, as well as pharmacologic treatments [4]. Pharmacological therapies include selective serotonin reuptake inhibitors (SSRIs), which are first line, tricyclic antidepressants (TCAs), such as imipramine and clomipramine, monoamine oxidase inhibitors (MAOIs), benzodiazepines, and beta-blockers [5]. Each of these medications has the potential for drug-drug interactions.

A prior case report discussed the management of anxiety, depression, and borderline personality disorder in a 25-year-old female with EDS who was treated with sertraline 150 mg qd (once a day), gabapentin 600 mg BID (twice a day), and lithium 300 mg BID (to help augment SSRI and attempt to decrease suicidality). However, this patient’s anxiety symptoms persisted despite the use of medications [6]. A large number of psychiatric medication regimens have been employed for the management of anxiety in patients with EDS, however they are often unsuccessful or poorly tolerated [6]. This requires providers to better tailor therapies to each individual in order to sufficiently treat patients with treatment-resistant anxiety disorders. Our hypothesis is that patients with EDS and comorbid anxiety disorders, who have had failed multiple...
Delays in diagnosis and treatment of anxiety can lead to poorer social functioning, as well as pain.

A cross-sectional study that assessed 80 patients with hypermobile EDS in France divided patients into two groups based on self-report: low and high anxiety levels. 51.2% of participants were categorized as having a high level of anxiety. The delay in diagnosis and uncertainty of the source of underlying pain, which are due in part to both society and physician’s unfamiliarity with EDS, contributes to an increased level of anxiety. The high level of anxiety can lead to hopelessness and fear that her condition would not improve, and concern that she would continue to require lengthy inpatient hospitalizations.

Upon initial interview, the patient denied suicidal ideation, intent, and plan, but complained of ongoing anxiety symptoms and appeared acutely anxious. Her level of anxiety was interfering with her social functioning, as well as her ability to comply with medical treatment. She had been unable to work and was no longer able to travel because of worsening physical and mental health. The high level of anxiety led to hopelessness and fear that her condition would not improve, and concern that she would continue to require lengthy inpatient hospitalizations.

The patient’s past psychiatric history included diagnoses of unspecified depression and anxiety. She had multiple prior suicide attempts, cutting behaviors, and three prior psychiatric hospitalizations. Patient was not following with any outpatient psychiatrist prior to our evaluation. She reported multiple prior drug trials ( durations and doses unknown) with adverse side effects, including paroxetine (suicidality), sertraline (suicidality), duloxetine (shakiness and tremors, poor impulse control, decreased sleep; discontinued due to concern for hypomania), amitriptyline (no improvement in symptoms),quetiapine (worsened anxiety), lorazepam, diazepam (both caused worsening paranoia), and gabapentin (vomiting, shakiness, and did not help improve pain symptoms). The only medications that the patient reported had been helpful in reducing anxiety were haloperidol and midazolam. Initially, venlafaxine 37.5 mg (crushed and administered through jejunostomy tube) was trialed, however, it caused worsening anxiety and gastrointestinal distress. Patient was then trialed on monotherapy of risperidone 0.5 mg oral at bedtime, with subsequent improvement in anxiety symptoms. She reported lessened anxiety and overall improved mood, and she was satisfied with the improvement in her symptoms. She became less preoccupied with somatic complaints and her lessened anxiety led to improvement in the quality of sleep. She was discharged home with this prescription and arranged with outpatient psychiatric follow up.

The chief complaint of patients with EDS frequently involves high levels of pain, which can lead to difficulties with diagnosis due to its subjective quality. Years of ongoing, and often untreated, suffering prior to a formal diagnosis can lead to patterns of negative thinking including catastrophizing, filtering, personalizing, generalizing, and polarizing [4]. These cognitive distortions can be precipitants for anxiety-related symptoms. The delay in diagnosis and uncertainty of the source of underlying pain, which are due in part to both society and physician’s unfamiliarity with EDS, contributes to an increased level of anxiety. The anxiety-related symptoms are often unrecognized or not prioritized because of the medical complexity of these patients. These patients often report feeling marginalized, alone, strange, or misunderstood [4]. These feelings can lead to psychological distress and these patients would benefit from earlier referral to psychologist and psychiatrists for support.

A cross-sectional study that assessed 80 patients with hypermobile EDS in France divided patients into two groups based on self-report: low and high anxiety levels. 51.2% of participants were categorized as experiencing a high level of anxiety [1]. In 2008, a postal survey was used to analyze 250 patients with EDS in Sweden, using the Hospital Anxiety and Depression Scale (HADS) and Short Form Health Survey (SF-36) scales. On the HADS, 74.8% scored high on anxiety, and 22.4% scored high on depression. The EDS group scored significantly lower on the SF-36 scale, indicating that patients with EDS perceived their physical and mental quality of health to be lower than the general population [7]. The high prevalence of anxiety in patients with EDS, as well as their perceived low quality of mental health, indicate the need for a multidisciplinary approach to these patients including psychiatric support and management.

Delays in diagnosis and treatment of anxiety can lead to poorer social functioning, as well as pain-related symptoms. The only medications that the patient reported had been helpful in reducing anxiety were haloperidol and midazolam. Initially, venlafaxine 37.5 mg (crushed and administered through jejunostomy tube) was trialed, however, it caused worsening anxiety and gastrointestinal distress. Patient was then trialed on monotherapy of risperidone 0.5 mg oral at bedtime, with subsequent improvement in anxiety symptoms. She reported lessened anxiety and overall improved mood, and she was satisfied with the improvement in her symptoms. She became less preoccupied with somatic complaints and her lessened anxiety led to improvement in the quality of sleep. She was discharged home with this prescription and arranged with outpatient psychiatric follow up.
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Introduction

It is well known that comorbid psychiatric conditions are frequent in patients with EDS. The EDS population is unique in that these patients already face a wide range of medical disorders however the more prevalent diseases end up being studied and spoken of in publications more often. This article emphasizes the importance of treating medical disorders that have a high association with anxiety disorders such as EDS. If we can have a means of treating debilitating anxiety in a patient with EDS, then this should be studied and spoken about because of the great physical burden these patients already face. If we can broaden the realm of psychiatric treatment to very specific disorders in patients with EDS, then this should be studied and spoken about because of the great physical burden these patients already face. If we can broaden the realm of psychiatric treatment to very specific disorders in patients with EDS, then this should be studied and spoken about because of the great physical burden these patients already face.

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Conclusions

Both literature and clinical experience indicate that psychiatric disorders, including anxiety disorders, are comorbid with EDS. However, there is minimal literature published regarding the management of treatment-resistant anxiety in this patient population. The above case describes the successful use of risperidone to reduce anxiety in a patient with EDS, indicating that risperidone may be helpful in managing anxiety disorders in these patients. The mechanism behind the improvement in anxiety symptoms may be due to risperidone’s dual antagonism of both serotonergic and dopaminergic receptors. We believe that further research into risperidone, as well as other atypical antipsychotics as a treatment option in this patient population is warranted. This can help attain the goal of enhanced quality of life for patients with anxiety disorders in these patients. The mechanism behind the improvement in anxiety symptoms may be due to risperidone’s dual antagonism of both serotonergic and dopaminergic receptors. We believe that further research into risperidone, as well as other atypical antipsychotics as a treatment option in this patient population is warranted. This can help attain the goal of enhanced quality of life for patients with anxiety disorders in these patients.
EDS who already suffer a great deal of physical burden from their illness.

Additional Information

Disclosures

**Human subjects:** Consent was obtained by all participants in this study. **Conflicts of interest:** In compliance with the ICMJE uniform disclosure form, all authors declare the following: **Payment/services info:** All authors have declared that no financial support was received from any organization for the submitted work. **Financial relationships:** All authors have declared that they have no financial relationships at present or within the previous three years with any organizations that might have an interest in the submitted work. **Other relationships:** All authors have declared that there are no other relationships or activities that could appear to have influenced the submitted work.

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