Introduction

According to the center of disease control statistics, stroke is the fifth leading cause of death worldwide. Lloyd-Jones, et al estimated that stroke attacks every 40 sec and causes loss of life every 4 minute. Stroke can be defined as a complex neurological condition that occurs when there is an interruption of blood supply to a part of the brain causing brain cells to die [1]. It is a medical emergency and immediate medical care is warranted.

Discussion

Research data over the years has already established a significance of post-stroke psychiatric manifestations needs fixing, et al states in psychiatric times that first 6 months after stroke is a time in which a person is highly vulnerable to develop these psychiatric conditions. This necessitates an obligation of a clinician and a family member as well to further explore these symptoms. We will discuss each of the possible psychiatric outcomes on individual grounds.

Depression

Post-stroke depression is a depression that occurs after the stroke and cannot be attributed to any other medical condition. Diagnostic and statistical manual of mental disorders V defines depression as depressed mood or loss of pleasure along with 5 of the following for 2 weeks. These include markedly diminished interest in activities, significant weight loss when not dieting or weight gain, insomnia or hypersomnia, psychomotor retardation, loss of energy, feelings of worthlessness or excessive or guilt, diminished ability to concentrate, recurrent thoughts of death, recurrent suicidal ideation. It is highly advised that the clinician should screen patients for these symptoms using an integrated clinical approach to screen, diagnose, and to manage them accordingly. This can certainly impact positively on their prognosis and subsequently on their quality of life.

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References

Pseudobulbar Affect

Pseudobulbar affect can be defined as an emotional disturbance that causes uncontrollable crying or laughing secondary to stroke or any other neurological condition. Various hypothesis has been involved in the pathogenesis of PBA. The most influential theory states that neurophysiological pathways from cortex to pons to cerebellum are disinhibited leading to inappropriate affective expression consist of an involuntary burst of laughter or excessive tearfulness [7]. Other theories have also validated the fact that people affected with cerebellar stroke demonstrate affective lability and PBA is more common with strokes within this territory. Neurochemical pathways explain the role of various neurotransmitters and neuromodulators in PBA but most important of which are serotonin and glutamate.

This is validated by the response of PBA to SSRI and glutamate inhibitors. PBA is often misdiagnosed as depression; therefore, a clinician should keep an eye to differentiate PBA from depressive symptoms as former last few seconds to few minutes while depressive symptoms can last longer from weeks to months. Also, the other symptoms of guilt, worthlessness, fatigue, anhedonia are missing from PBA. This is also important to consider that in PBA, there is no mood component in between crying or laughing spells which is opposed to Bipolar rapid cycler which also present with laughing or crying spells but with continued mood changes [8,9].

Post-Stroke Psychosis

It is a rare manifestation after stroke, however, a recent meta-analysis by Hele Strangerland, et al. states that the prevalence of post-stroke psychosis is about 4.6 percent. Research has evidenced that people affected with post-stroke psychosis usually have lesions in the right hemisphere. In majority of cases observed in the meta-analytic study, the most common artery that was found to be involved was right middle cerebral artery [10]. A person can present with psychosis with variable presentation depending on the territory of stroke. Risk factors that can increase the likelihood of developing psychosis following a stroke are the preexistent psychiatric condition, substance use, mood disorders, intellectual disability. This risk is greater than the population with no prior history of any psychiatric condition.

Anxiety

The Diagnostic and Statistical Manual of Mental Disorders (DSM-IV) classifies anxiety disorders as a collection of syndromes including generalized anxiety disorder (GAD), panic disorder (with or without agoraphobia), agoraphobia (with or without panic), specific phobia, social phobia, obsessive-co compulsive disorder (OCD), posttraumatic stress disorder (PTSD), acute stress disorder, anxiety disorder due to a general medical condition, substance-induced anxiety disorder, and anxiety disorder not otherwise specified [11]. Anxiety is one the most common psychological disorder post stroke and is associated with an increase in dependency in the activities of daily living, reduced social networking, the severity of Post-traumatic stress disorder, worsening disability over time and mortality.

White JH et al. [12] conducted a prospective cohort study and concluded that post-stroke anxiety is positively associated with baseline PSD, baseline anxiety and less disability. Earlier it was thought that the prevalence of anxiety is low post stroke [13]. However, later, we got to know that anxiety occurs in 20-25% of the patients at any time after stroke, with Generalized anxiety disorder (GAD) and Phobic disorders being the most common types. The incidence of Post-stroke anxiety is 20% one month after stroke, increasing to 23% within five months and to 24% in 6 or more months [11].

De Wit L et. al. [14] assessed 532 patients at 2, 4 and 6 months at four rehabilitation centers using the Hospital Anxiety and Depression Scale. They find out that frequency of anxiety was 22% and 25% post-stroke at 2 and 4 months and 40% of the patients remained anxious at 6 months after stroke. GED Aiello et al. [15] in 2014 concluded that anxiety was more commonly reported than depression in the chronic stages of stroke i.e. 1 year post-stroke, no association of anxiety with lesion localization was found.

As far as the treatment is concerned, medications such as buspirone and paroxetine can be effective in reducing anxiety in patients with co-morbid anxiety and depression [16]. Manzoni et al. [17] because of their meta-analysis of 27 studies found relaxation training to be effective for reducing anxiety. Relaxation classes in patients post stroke led to increase motivation for participation in other activities, improve confidence, psychological functioning and quality of life [18].

Dementia

According to the American Psychiatric Association's Diagnostic and Statistical Manual of Mental Disorder post-stroke cognitive impairment or dementia is generally defined as dementia occurring after Stroke with the final diagnosis should be delayed up to 6 months [19]. However, many stroke survivors develop dementia 3 months after a stroke or after recurrent stroke. Recognizing cognitive disorder in the early phase help in doing early rehabilitation and thus improving fatality by improving management [20].

Dementia in stroke involve all types of dementia whether its vascular, degenerative or mix process. In a recent study, incorporating pathological examination, vascular dementia was found in 75% of the demented stroke survivors, 25% has...
mixed pathology with either Alzheimer’s type lesion or Lewy bodies [21]. Recent studies suggested that stroke survivors may trigger pathologies attributed to subcortical Vascular dementia, multi-infarct dementia, and strategic infarct dementia. Thus, the term vascular cognitive impairment was introduced, which incorporate all cognitive changes related to all causes of vascular diseases [20].

A meta-analysis done in 2009 by Pendlebury et al found that 10% of the patients were already suffering from dementia before the first stroke, 10% develop after first stroke and 30% developed after recurrent stroke. The prevalence of post-stroke dementia ranges from 6%-32% with the highest prevalence at 3 months which is about 28% [22]. Risk factors for PSD include increasing age, severe neurological deficits, hypertension, atrial fibrillation, diabetes mellitus, myocardial infarction [19]. To prevent post-stroke dementia management of the pre-stroke risk factors, play an important role, while early treatment and rehabilitation of stroke play an important role post stroke [23].

Mania

According to DSM V, Mania can be defined as a state of an abnormally and persistently elevated, expansive or irritable mood and persistently heightened energy for at least one week, and present for most of the day. Krauthammer C, Klerman G described mania as a secondary mania if it occurs in context of neurological conditions such as stroke or due to toxic and metabolic derangements [24]. It is a rare consequence after cerebrovascular insult with very few cases reported. It has a frequency of < 1 percent in comparison to depressive outcome in post stroke patients [25]. Post stroke mania presents with a similar presentation as primary mania, which is due to psychiatric illness. Lesions responsible for poststroke mania seems to be more related with right-sided lesions but there are also reports of mania following left-sided lesions [26].

Conclusion

Given the above review considering research articles the psychiatric consequences following a cerebrovascular accident are well understood. Despite abundant literature available, unfortunately these emotional and behavioral unsetslements are often undiscovered clinically. It is therefore imperative for a clinician to screen, recognize and to monitor these psychiatric symptoms for any progression on every follow up visit especially within first 3 years as this is time with extreme vulnerability to develop psychotic and mood symptoms. Also, neurology and psychiatry should work in collaboration to address these neuropsychiatric phenomena. There is also a need of extensive physical therapy and occupational therapy to improve clinical outcome.

References


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