Adaptive Affect: The Nature of Anxiety and Depression

This article offers another way of thinking about anxiety and depression. Both are viewed as extensions of normal biological drives that work in parallel to keep us safe, healthy, and functioning well. But when pushed to the extremes these same adaptive drives are responsible for anxiety and depression. We are constantly faced with stressors great and small, ranging from requirements to meet basic physical, social, and emotional needs such as acquiring food or finding a partner, to making less imperative decisions such as selecting a pair of socks to wear in the morning. Normally we are able to effectively and efficiently manage the stressors that are encountered. However, if these stressors, when combined, are too great and beyond the individual’s ability to manage, impairment in functioning begins to occur. Initially, the stressors cause anxiety, but if they remain unresolved, over time symptoms of depression may begin to appear in increasing number and intensity.

Adaptive Affect

Adaptive characteristics, meaning those that are helpful to us, are retained in the general population. As such, the continued presence of anxiety and depression suggests the processes reflect something beneficial. While the adaptive role anxiety plays in preserving and protecting us is not difficult to see (eg, avoiding potential harm), at the extremes of severity and duration, anxiety can become problematic (eg, clinically significant impairment). The adaptive role of depression is less clear. On the surface, it does not seem plausible that depression would be adaptive, as a hallmark feature is impairment in functioning. Yet, depression is present across cultures and has been preserved over time. This manuscript describes a slightly different perspective on the origin and relationship between anxious and depressive mood states.

Abstract: An approach viewing anxiety and depression as extensions of normal adaptive biologic drives is discussed. Anxiety is viewed as the result of an underlying biological drive to preserve and maintain our wellbeing. At the extremes, if unresolved, this drive can be maladaptive, particularly if activated over prolonged periods of time. This paper proposes that depression is the result of a biological drive that mediates the effects of maladaptive levels of anxiety. These two processes are thought to be acting simultaneously. When operating in the normal range, these drives are helpful; in the extremes, they are associated with impairment. Over time, if unresolved, symptoms of anxiety will begin to become associated with increasing levels of depression.

Keywords: depression, anxiety, adaptive affect, stress

Plain Language Summary

This article offers another way of thinking about anxiety and depression. Both are viewed as extensions of normal biological drives that work in parallel to keep us safe, healthy, and functioning well. But when pushed to the extremes these same adaptive drives are responsible for anxiety and depression. We are constantly faced with stressors great and small, ranging from requirements to meet basic physical, social, and emotional needs such as acquiring food or finding a partner, to making less imperative decisions such as selecting a pair of socks to wear in the morning. Normally we are able to effectively and efficiently manage the stressors that are encountered. However, if these stressors, when combined, are too great and beyond the individual’s ability to manage, impairment in functioning begins to occur. Initially, the stressors cause anxiety, but if they remain unresolved, over time symptoms of depression may begin to appear in increasing number and intensity.

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Adaptive characteristics, meaning those that are helpful to us, are retained in the general population. As such, the continued presence of anxiety and depression suggests the processes reflect something beneficial. While the adaptive role anxiety plays in preserving and protecting us is not difficult to see (eg, avoiding potential harm), at the extremes of severity and duration, anxiety can become problematic (eg, clinically significant impairment). The adaptive role of depression is less clear. On the surface, it does not seem plausible that depression would be adaptive, as a hallmark feature is impairment in functioning. Yet, depression is present across cultures and has been preserved over time.
Affective symptoms caused by stressors both internal and external have been the subject of a tremendous amount of study over the last century, and documented for several millennia. During this time, any number of models explaining the development of affective symptoms have been presented, as these models have increased in accuracy and relevance, they have also become more complex. The goal of this manuscript is to provide a simple model explaining the natural development of affective symptoms over time. The approach described is straightforward. Individuals are constantly under stress to adapt to the environment (internal and external), required to maintain an adaptive balance between action and non-action. The thought being stress produces anxiety that is adaptive when the individual is able to respond and alter themselves or the environment, but in situations that a satisfactory response is not possible it is maladaptive to remain in a highly activated state for prolonged periods of time. In these situations it would be beneficial to reduce the call to action, to decrease the anxiety response. It is possible that some of the symptom presentations we associate with clinical depression are the result of the extension of an adaptive biological mechanism that mediates the effects of prolonged anxiety. If we associate anxiety with activation and depression with deactivation at the extremes, the idea can be displayed in a diagram (see Figure 1).

Figure 1 is a “temporal” phase diagram that allows for activation and deactivation to occur simultaneously over time. The axes “y” (Activation/Arousal) and “x” (Deactivation/Suppression) display the interaction between underlying biological drives that are responsible for symptom presentation at the extremes. Clinically significant impairment and/or distress define the boundary for normal mood states and those requiring treatment (expressed as dashed line “Imp” in Figure 1). The lines displaying boundaries between normal, anxiety, depression, and mixed are dashed, indicated they are imprecise and individualized. The lines between anxiety, depression, and mixed presentations were included to illustrate the progression of symptoms, with ΔT indicating the direction of change over time. Holding severity constant, the natural progression over time is indicated as (A→D)ΔT (anxiety yields depression over time).

If these underlying biological drives are developed, the question of why would activation and deactivation occur simultaneously remains, as it would appear to be more efficient simply to reduce or stop the activation process rather than initiate a mediating process. There is an advantage associated with being able to reduce the effects of anxiety-producing stimuli while maintaining a level of arousal that does not impair functioning. In the individual that is functioning normally, this may achieve this balance and create a space to process (conscious and nonconscious) as needed, increasing positive outcomes. In the person who is experiencing clinically significant distress and/or impairment, this space may appear symptomatic, replete with maladaptive levels of anxiety and depression.

While, in general, if stressors remain constant, the relationship (A→D)ΔT would be expressed. Key prognostic features, such as the rate at which the symptom presentation moves from being predominantly anxious to including symptoms of depressed mood, and the magnitude of stressors required for impairment to occur are unique to the individual (genetic vulnerability and resiliency factors, as well as characteristics that moderate disease burden and coarse of illness have been identified). A person’s present state (position on the diagram) is the result of a summation of multiple inputs (internal and external) calling for action and non-action simultaneously. The specific symptom presentation would depend upon the individual’s location on the time line (anxiety-mixed-depression).

Disorders associated with primary physiologic hyperarousal (panic and phobic disorders) would be located close to the y-axis of Figure 1, and those associated with neurovegetative symptoms would be close to the x-axis (atypical depression). Disorders showing symptoms of both anxiety and depression, such as melancholic depression,
would be located in the middle portion of the diagram in the mixed section. This approach addresses normal mood and affective symptoms of anxiety and depression; mania and psychosis are thought to involve additional underlying systems and are not included in this model. Normal mood such as being excited or relaxed would be below the impairment line, close to the y- and x-axis, respectively. In most patients, exposure to sustained levels of unresolvable anxiety-producing stressors leads to depression. However, patients with disorders hallmark by brief episodic hyperarousal experience clinical anxiety as discrete events, and may lack the sustained exposure to stressors required to develop clinical depression.

An adaptive affect approach offers a slightly different perspective on what we currently understand about anxiety and depression. By looking at the summation of concurrent activating and deactivating drives, the model explains much of the broad spectrum of symptom presentations associated with affective disorders, and based upon trajectory, provides a framework for understanding the progression of an individual’s affective states over time. The approach is simple, and the information required is generally gathered as part of a standard clinical interview within or outside of a research protocol. Whether in clinical practice or research, this perspective provides a simple way to interpret a person’s history and current functioning, as well as a “common sense” model that is easy to understand.

Disclosure
The author reports no conflicts of interest in this work.

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