Emotional disorders: Cluster 4 of the proposed meta-structure for DSM-V and ICD-11

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Emotional disorders: Cluster 4 of the proposed meta-structure for DSM-V and ICD-11

Paper 5 of 7 of the thematic section: ‘A proposal for a meta-structure for DSM-V and ICD-11’

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Background. The extant major psychiatric classifications DSM-IV, and ICD-10, are atheoretical and largely descriptive. Although this achieves good reliability, the validity of a medical diagnosis would be greatly enhanced by an understanding of risk factors and clinical manifestations. In an effort to group mental disorders on the basis of aetiology, five clusters have been proposed. This paper considers the validity of the fourth cluster, emotional disorders, within that proposal.

Method. We reviewed the literature in relation to 11 validating criteria proposed by a Study Group of the DSM-V Task Force, as applied to the cluster of emotional disorders.

Results. An emotional cluster of disorders identified using the 11 validators is feasible. Negative affectivity is the defining feature of the emotional cluster. Although there are differences between disorders in the remaining validating criteria, there are similarities that support the feasibility of an emotional cluster. Strong intra-cluster co-morbidity may reflect the action of common risk factors and also shared higher-order symptom dimensions in these emotional disorders.

Conclusion. Emotional disorders meet many of the salient criteria proposed by the Study Group of the DSM-V Task Force to suggest a classification cluster.

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Key words: Co-morbidity, DSM-V, emotional disorders, neuroticism.

Introduction

Emotional (or internalizing) disorders form the largest group of mental disorders, consisting of states with increased levels of anxiety, depression, fear and somatic symptoms. They include generalized anxiety disorder (GAD), unipolar depression, panic disorder, phobic disorders, obsessional states, dysthymic disorders, post-traumatic stress disorder (PTSD) and somatoform disorders. We have also included neurasthenia, as this diagnosis is commonly made in many parts of the world, and is in the ICD-10. We have preferred the term ‘emotional’ because we include somatoform disorders in the group. Depressive, anxious and somatoform symptoms occur together in general medical settings, and share many common features. Within this class there are differences in the genetic factors, the early environments and the biological measures, but there are also important similarities that justify bringing the disorders into a single group.

The current classifications are based upon similarities between the clinical manifestations of these disorders. The purpose of the proposed meta-structure, of which this paper is a part, is to determine whether it is feasible to identify disorder groupings based on aetiology (Andrews et al. 2009a, b; Carpenter et al. 2009; Krueger & South, 2009; Sachdev et al. 2009). The focus of this paper is whether the existing anxiety, mood and somatoform disorders could be grouped as aetologically similar disorders. ‘Similar’ is used here in the sense that the pattern of risk factors implicated in the development of emotional disorders is consistent across the disorders rather than suggesting that these disorders have a single, common cause.

Method

A Study Group of the DSM-V Task Force of the American Psychiatric Association (APA) has recently recommended 11 ‘validating criteria’ that could be used to identify groups of aetiologically related
disorders without altering the current diagnostic criteria (Hyman et al., personal communication, 3 December 2007). These are:

1. genetic factors;
2. familiality;
3. early environmental adversity;
4. temperament antecedents;
5. neural substrates;
6. biomarkers;
7. cognitive and emotional processing;
8. differences and similarities in symptomatology;
9. co-morbidity;
10. course;

Scopus, EMBASE, PsychINFO and Medline searches were conducted to identify English-language literature that considered the risk associated with each validating criterion and the emotional disorders. Large epidemiological data samples using the current classificatory criteria were preferred over small clinical samples. The literature that would support or negate the present thesis is selectively reviewed. If the Committees responsible for the revision of DSM-IV and ICD-10 agree that an aetiologically driven classification is feasible, the final cluster-membership of disorders can be determined by systematic reviews.

**Results**

**Genetic factors**

Genetic-epidemiological twin data have been used to identify two broad genetic risks of the common mental disorders: internalizing (emotional) and externalizing liabilities (e.g. Kendler et al. 2003). The Kendler group report that the internalizing genetic risk factor is composed of discrete but intercorrelated factors: one for anxious-misery with major depressive disorder (MDD) and GAD, and the other a fear risk factor with phobic disorders. Panic loaded moderately on the first genetic risk factor, however, was the only investigated internalizing syndrome to have a significant disorder-specific genetic risk (Kendler et al. 2003). This internalizing tripartite model is somewhat similar to that advocated initially by Clark & Watson (1991). Kendler et al. (2003) did not include PTSD, neurasthenia, obsessive-compulsive disorder (OCD), somatoform disorders or dysthymia. Nevertheless, co-morbidity studies that have implemented similar statistical analyses to those used by Kendler and colleagues have reported that PTSD (Cox et al. 2002; Slade & Watson, 2006), neurasthenia and dysthymia (Slade & Watson, 2006) load on the anxious-misery/distress factor and OCD is more related to the fear risk factor (Slade & Watson, 2006). Twin data have also been used to show that the genetic risk associated with MDD is shared substantially with GAD (Kendler et al. 1992), and both are strongly associated with neuroticism (Hettema et al. 2004; Kendler et al. 2006). Neuroticism also explains some, but not all, of the common genetic risk of several of the internalizing/emotional disorders (e.g. Hettema et al. 2006: MDD, GAD, Panic, Agoraphobia, Social Phobia, Animal and Situational Phobias; Kendler et al. 2007: GAD and MDD).

Studies of the genetic factors of the emotional disorders have often compared probands with a particular disorder with normal controls, or have tended to stay within a particular chapter of the major classifications; so that unipolar depression has been compared with other mood disorders, and various anxiety disorders have been compared with one another. However, there have been notable exceptions to this, and these are shown as Table 1.

Genomic screens have not yet identified specific genes for the majority of the emotional disorders, notwithstanding recognition of the broad genetic liabilities to mental illness. The only gene identified for an emotional disorder is the 5-hydroxytryptamine (5-HT) transporter gene, first shown to have an important gene–environment interaction in the aetiology of depression by Caspi et al. (2003), and since confirmed by several different studies (e.g. Eley et al. 2004; Kendler et al. 2005; Wilhelm et al. 2006). This gene has also been implicated in anxious traits including neuroticism and harm avoidance by Lesch et al. (1996), who found that individuals with one or two short forms of the allele had higher rates of neuroticism and its anxious, depressive and angry hostility subfacets. Nevertheless, the role of the 5-HT gene remains inconclusive, as other studies have failed to replicate it (Gillespie et al. 2005; Willis-Owen et al. 2005).

We seem to have two overlapping groups of genes dealing with anxious-misery and fear, but there are also other genes not shared with neuroticism. For example, Kendler et al. (2006) show that although the association between neuroticism and MDD results from shared genetic risk factors, a substantial proportion of the genetic vulnerability to MDD is not reflected in neuroticism. There is indirect evidence for common genes for PTSD, somatoform disorders, neurasthenia, obsessive disorder and dysthymia but the degree of overlap remains to be clarified.

**Familiality**

The present contention is not that there are no differences between probands with different emotional disorders, it is that there is important common ground between them. This is confirmed by the higher rates of other emotional disorders in the first-degree relatives.
(FDRs) of affected probands than is expected in the FDRs of normal controls (see Table 2). Eley et al (2002) show that although the rate for GAD (with or without other disorders such as panic or MDD) varies from 9% to 20% in FDRs of GAD probands, the risk for HC ranges from 2% to 4%. There is considerable variation in rates of GAD in FDRs of patients with fear disorders, although they are always raised relative to normal controls. There is also some support for the specific transmission of individual emotional disorders. For example, Mendlewicz et al. (1993) showed that the morbid risk of panic disorder is significantly higher in panic-affected probands than FDRs of probands with GAD, MDD and HC. It is also noteworthy that, though not reaching significance the risk of GAD, MDD and panic disorder in FDRs of affected probands were greater than in FDRs of HC. Consistent with Mendlewicz et al. and the later findings of Kendler et al. (2003), as noted above, Goldstein et al. (1994) report that panic disorder has a specific genetic component with a possible increased risk of social phobia. Fyer et al. (1995) also argue that ‘pure’ panic and fear disorders have some disorder-specific genetic risk. In terms of OCD, Hanna et al. (2005) report that there are higher rates of anxiety disorders in FDRs of probands with familial OCD in comparison to sporadic forms of OCD.

Complementary to the findings of studies where offspring are the affected proband, investigations of depressed parents and grandparents have also found increased risk of anxiety (Lieb et al. 2002a; Weissman et al. 2005), depression and substance disorders in offspring (e.g. Lieb et al. 2002a). The inter-cluster familiarity of some of the emotional disorders and some of the externalizing disorders may be explained by common genetically determined vulnerability or by social learning within families. Family studies are a weaker test of the hypothesis being put forward than

### Table 1. Summary of studies that have linked genetic factors across the main symptom domains

<table>
<thead>
<tr>
<th>Diagnoses</th>
<th>Reference</th>
<th>Main relevant findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anxious-misery prominent</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Major depression and GAD</td>
<td>Kendler et al. (2003)</td>
<td>Anxious-misery and fear disorders regarded as ‘affective spectrum disorder’</td>
</tr>
<tr>
<td></td>
<td>Kendler et al. (2006)</td>
<td>Swedish twins: shared risk N and MDD genetic correlation of +0.46</td>
</tr>
<tr>
<td></td>
<td>Kendler et al. (2007)</td>
<td>N accounts for 25% of GAD and MDD genetic overlap</td>
</tr>
<tr>
<td></td>
<td>Hettema et al. (2006)</td>
<td>Similar genetic risk factors for lifetime MDD and GAD; most covariance not shared with N</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Genetic factor independent of N increases the risk for MDD, GAD and Panic</td>
</tr>
<tr>
<td>Dysthmic disorder</td>
<td>No data</td>
<td></td>
</tr>
<tr>
<td>Neurasthenia</td>
<td>Angst et al. (2006)</td>
<td>Neurasthenia co-morbid with atypical depression</td>
</tr>
<tr>
<td>Post-traumatic stress</td>
<td>Koenen et al. (2003)</td>
<td>Shared familial vulnerability contributes to the association between PTSD and MDD, and PTSD and DD. This vulnerability is mediated by genetic factors</td>
</tr>
<tr>
<td></td>
<td>Skre et al. (1993)</td>
<td>PTSD more prevalent in co-twins of anxiety probands</td>
</tr>
<tr>
<td></td>
<td>Chantarujakapong et al. (2001)</td>
<td>PTSD has common additive genetic liability with GAD and Panic</td>
</tr>
<tr>
<td>Fear and avoidance prominent</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Phobias and panic disorder</td>
<td>Kendler et al. (2003)</td>
<td>Two genetic factors one for anxious misery (MDD, GAD and Panic) the other for fear disorders (Animal and situational phobias), with common path coefficient of +0.34 between them</td>
</tr>
<tr>
<td></td>
<td>Hettema et al. (2006)</td>
<td>Each disorder correlates with N</td>
</tr>
<tr>
<td>Obsessional disorder</td>
<td>Clifford et al. (1984)</td>
<td>Heritability 44% shared with N</td>
</tr>
<tr>
<td>Somatic symptoms prominent</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Somatoform disorder</td>
<td>Gillespie et al. (2000)</td>
<td>Partial support for common genetic factors between somatic distress and anxious-misery</td>
</tr>
<tr>
<td></td>
<td>Torgersen (1986)</td>
<td>Link between anxiety and somatic disorders</td>
</tr>
</tbody>
</table>

DD, Dysthymic disorder; GAD, generalized anxiety disorder; MDD, major depressive disorder; N, neuroticism; PTSD, post-traumatic stress disorder.
genetic-epidemiological twin data because familial aggregation of different emotional disorders reflect not only common genetics but also social learning within families. The various studies of familiality reported in Table 2 do not report rates for neurasthena or somatoform disorders, so it cannot be claimed that there is at present complete evidence for the proposed cluster from familiality data. To the extent that familiality data support the present hypothesis, it is in the higher rates of anxiety disorders in FDRs of the diagnoses included in the cluster.

**Early environmental adversity**

Most adult psychiatric disorders including the ‘stress-related and fear circuitry’ disorders, and a range of other syndromes such as behaviour disorders and substance use disorders, have their roots in early life (Kim-Cohen et al. 2003). However, evidence for specificity between adult diagnoses is less impressive than the similarities (see Table 3).

For most members of this cluster, early environmental factors that predispose to the disorder have much in common, but co-morbidity may account for some of the reported findings; so that the factors may be related only to anxious and depressive syndromes. However, for obsessional and somatoform disorders, other specific factors may be operating. Findings of early adversity in obsessional disorder have been inconsistent and apparent support for early adversity may be due to accompanying anxious symptoms such as cognitive/perceptual biases (Alonso et al. 2004). In somatoform disorders, several studies report higher

### Table 2. Summary of studies that have linked familial factors across the main symptom domains

<table>
<thead>
<tr>
<th>Diagnoses</th>
<th>Reference</th>
<th>Main relevant findings</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Anxious-misery prominent</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Major depression and</td>
<td>Reich (1995)</td>
<td>Both GAD and MDD co-aggregate, but frequency of GAD in FDRs of MDD, and of MDD in FDRs of GAD, is always higher than either in HCs</td>
</tr>
<tr>
<td>generalized anxiety disorder</td>
<td>Kendler et al. (1997)</td>
<td>Risk for anxiety disorders increased in FDRs of both GAD and MDD, but rates for MDD only higher in FDRs of MDD probands. Underlying vulnerabilities to internalizing and to externalizing disorders transmitted across generations with moderate fidelity</td>
</tr>
<tr>
<td></td>
<td>Lieb et al. (2002a)</td>
<td>One parent with MDD: rates in offspring raised for MDD, GAD, DD, Phob, AG</td>
</tr>
<tr>
<td><strong>Dysthmic disorder</strong></td>
<td>Klein et al. (1995)</td>
<td>Two parents with MDD: rates raised for PTSD and Panic</td>
</tr>
<tr>
<td></td>
<td>Donaldson et al. (1997)</td>
<td>Strong familial relationship between DD and MDD</td>
</tr>
<tr>
<td></td>
<td>Rashed et al. (2001)</td>
<td>Sixty-six per cent of FDRs of dysthymics had dysthymia compared to 36% of major depressives and 22% of normals</td>
</tr>
<tr>
<td>Neurasthena</td>
<td>No data</td>
<td>Four groups of probands: PTSD, MDD, GAD and HC. Rates for each higher in FDRs, but always higher than rates in the FDRs of HC</td>
</tr>
<tr>
<td>Post-traumatic stress</td>
<td>Davidson et al. (1998)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Fear and avoidance prominent</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Phobias, panic disorder</td>
<td>Hudson et al. (2003)</td>
<td>Affective spectrum disorder (includes all disorders in anxious-misery and fear) aggregates strongly in families, and MDD displays a significant familial co-aggregation with other forms of affective spectrum disorders taken collectively</td>
</tr>
<tr>
<td></td>
<td>(evidence above, in anxious misery, also applies to fear disorders)</td>
<td></td>
</tr>
<tr>
<td>Obsessional disorder</td>
<td>Nestadt et al. (2001)</td>
<td>Higher lifetime rates of GAD, Panic, Separation anxiety disorder and MDD in FDRs, after adjustment for independent transmission, MDD drops out</td>
</tr>
<tr>
<td></td>
<td>Ettelt et al. (2008)</td>
<td>FDRs of OCD cases have higher harm-avoidance than HC</td>
</tr>
<tr>
<td></td>
<td>Pauls et al. (1995)</td>
<td>Early onset cases more familial than late onset</td>
</tr>
<tr>
<td><strong>Somatic symptoms prominent</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Somatoform disorders</td>
<td>No data</td>
<td></td>
</tr>
</tbody>
</table>

AG, Agoraphobia; DD, dysthmic disorder; FDR, first-degree relative; GAD, generalized anxiety disorder; HC, healthy controls; MDD, major depressive disorder; OCD, obsessive-compulsive disorder; Phob, phobic disorder; PTSD, post-traumatic stress disorder.
rates of childhood illness in addition to measures of childhood adversity. Sexual abuse in childhood has been found to be associated with a wide range of functional symptoms, such as somatization disorders (Walker et al., 1995), somatoform symptoms (Hexel & Sonneck, 2002), irritable bowel disorder (Reilly et al., 1999), conversion disorder (Roelofs et al., 2002) and functional pelvic pain (Reiter et al., 1991).

Heim and colleagues (Heim et al., 2000, Heim & Nemeroff, 2001) argue that children exposed to traumatic life experiences develop an increased sensitization of those parts of the nervous system related to stress and emotion, and in consequence develop an increased vulnerability to later stress due to hyper-reactivity of corticotrophin-releasing factor, as well as to other neurotransmitter systems. In summary, early adversity seems to increase the probability of most later disorders, but there is little reason to suggest effects on specific disorders.

### Temperament antecedents
Negative affect (neuroticism) makes a strong contribution to all the emotional disorders. All those who

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Table 3. Summary of studies that have disadvantages in shared early life across the main symptom domains

<table>
<thead>
<tr>
<th>Diagnoses</th>
<th>Reference</th>
<th>Main relevant findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anxious-misery prominent</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Major depression</td>
<td>Rapee &amp; Bryant (2009)</td>
<td>Parents more likely to show low care and high overprotection for MDD, GAD and fear disorders. All are associated with parental neglect and both sexual and physical abuse</td>
</tr>
<tr>
<td>Generalized anxiety disorder</td>
<td>Moffitt et al. (2007 a)</td>
<td>All three forms of abuse and parental divorce are more common in MDD, GAD and fear</td>
</tr>
<tr>
<td></td>
<td>Kessler et al. (in press)</td>
<td>Childhood risk factors are almost identical, with all three forms of abuse and parental divorce being more common in the early lives of people with GAD</td>
</tr>
<tr>
<td></td>
<td>Kendler et al. (2004)</td>
<td>Children who have been sexually abused are at higher risk for both depression and GAD</td>
</tr>
<tr>
<td></td>
<td>Jaffee et al. (2002)</td>
<td>Sexual abuse common in future depressives</td>
</tr>
<tr>
<td></td>
<td>Hawker &amp; Boulton (2000)</td>
<td>Depression: onset &lt;15 years report a wide range of early traumas; onset &gt;15 years report sexual abuse only. Teasing and bullying are associated with increases in anxiety and depression</td>
</tr>
<tr>
<td>Dysthmic disorder</td>
<td>No data</td>
<td></td>
</tr>
<tr>
<td>Neurasthenia</td>
<td>No data</td>
<td></td>
</tr>
<tr>
<td>Post-traumatic stress</td>
<td>Koenen et al. (2007)</td>
<td>Age ≥5 years childhood risks: 50% develop PTSD after trauma; 8% after only 1 year</td>
</tr>
<tr>
<td></td>
<td>Koenen et al. (2003)</td>
<td>Neglect, sexual and physical abuse are at increased risk for developing PTSD</td>
</tr>
<tr>
<td></td>
<td>Widom (1999)</td>
<td>Family, individual and lifestyle variables also place individuals at risk</td>
</tr>
<tr>
<td></td>
<td>Breslau (2002)</td>
<td>Traumas sufficient to cause PTSD very common, those who do not develop PTSD after trauma are not at increased risk of MDD</td>
</tr>
<tr>
<td>Fear and avoidance prominent</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Phobias, panic disorder</td>
<td>Rapee &amp; Bryant (2009), Moffitt et al. (2007 a)</td>
<td>Parents more likely to show low care and high overprotection, associated with all forms of childhood abuse</td>
</tr>
<tr>
<td>Obsessional disorder</td>
<td>Alonso et al. (2004)</td>
<td>OCD reported higher levels of paternal rejection, hoarding predicted by low parental emotional warmth</td>
</tr>
<tr>
<td>Somatic symptoms prominent</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Somatoform disorder (see text for sexual abuse)</td>
<td>Craig et al. (2002)</td>
<td>Somatizers report childhood neglect and to physical illness in a parent (OR 2.9)</td>
</tr>
<tr>
<td></td>
<td>Craig et al. (2004)</td>
<td>Mothers reported frequent headaches and stomachache in childhood; were emotionally flatter</td>
</tr>
<tr>
<td></td>
<td>Egle &amp; Nickel (1998)</td>
<td>Parents more frequently chronically ill or disabled, more sexual and physical abuse and family disharmony</td>
</tr>
</tbody>
</table>

GAD, Generalized anxiety disorder; MDD, major depressive disorder; OCD, obsessive–compulsive disorder; PTSD, post-traumatic stress disorder; OR, odds ratio.
have reported multivariate analyses of self-report instruments of these disorders find a large general distress factor running across items, and this has been variously named ‘neuroticism’ (Eysenck, 1964; and many other authors), ‘negative emotionality’ (Harkness et al. 1995) and ‘negative affectivity’ (Watson et al. 1984). Jardine et al. (1984) studied the co-variation between neuroticism and current depressive and anxious symptoms and reported that, in both sexes, there is a genetic correlation between symptom scores and neuroticism of approximately 0.8. Khan et al. (2005) studied emotional disorders using the Virginia Twin Register, and concluded that ‘neuroticism not only contributed to individual diagnoses but also accounted for a significant part of the life-time comorbidity of common psychiatric disorders. The most striking finding was that neuroticism, on average, accounted for 26% of the comorbidity among the[se] disorders.’ Hettema et al. (2006) concluded that more than half of the genetic correlations between the internalizing disorders can be explained by the genetic risk of neuroticism (see Table 4). In the case of emotional disorders and, to a much lesser extent, externalizing disorders, high scores for negative emotionality precede the development of these disorders (Kendler et al. 1993; Krueger, 1999b). We are not aware of data showing the same for the other proposed clusters.

In summary, there is strong support for high neuroticism scores in all the proposed disorders in the cluster. However, on its own this is not a sufficient cause of emotional disorders because neuroticism scores are raised relative to healthy controls in other mental disorders; for example, neuroticism also contributes to the externalizing disorders to a smaller extent (10–12%) and novelty seeking (7–14%) also contributes to the co-morbidity between these disorders (Khan et al. 2005).

Neural substrates

Neuroimaging studies

The problem with making definitive statements about this topic is that, although there is a great deal of information about the neural substrate of both depression and the fear disorders, there have been remarkably few imaging studies that have examined functional and structural central nervous system disruptions in GAD (Martin & Nemeroff, in press). Nonetheless, and despite differences between anxious and depressed subjects, there are substantial similarities between the majority of the emotional disorders.

The medial prefrontal cortex has a general role in emotional processing, being activated in multiple individual emotions (activated in four of five specific emotions in at least 40% of studies meta-analysed by Phan et al. 2002). The medial prefrontal cortex seems to be involved in any situation where there is a relative focus on internal state or self-reference rather than attention to the outside world; thus, inappropriate attention is paid to internal stimuli at the expense of the external world. The insula is a region of limbic sensory cortex responsible for the generation of one’s mental image of one’s physical state (Rauch & Drevets, 2009). The anterior insular cortex of the non-dominant (right) hemisphere is thought to more specifically subserve evaluation of self-awareness, the ‘feeling self’. It is reciprocally connected to the right orbitofrontal cortex, a region further implicated in reward evaluation and decision making (Craig, 2002). This may account for the close association between the somatoform disorders and the anxious-misery disorders. There is considerable overlap between the clinical syndromes and the neural circuits involved, and it is clear that different circuits are involved across different diagnostic subgroups.

The emotional disorders as currently defined share certain common features, namely activation of visceral brain centres (amygdala, hypothalamus, locus coeruleus, dorsal raphe), and involve interpretation of novel stimuli that may have survival consequences (loss, threat, fear). The amygdala organizes the emotional response to stress; it is thought to be overactive in MDD and fear-disorder patients and may underlie the rumination on aversive or guilt-provoking memories that is common to both the mood and anxiety disorders (for mood disorders: Drevets, 2001, 2003; for anxiety disorders: Charney, 2003; Rauch et al. 2003; Etkin & Wagner, 2007). In addition to exaggerated amygdala responses, Rauch et al. (2006) argue that PTSD is characterized by deficient frontal cortical and hippocampal functioning. Etkin & Wager (2007) reported that PTSD, social anxiety disorder and specific phobia all showed greater activity than matched comparison subjects in the amygdala and insula, but PTSD showed hypo-activation in the dorsal and rostral anterior cingulate cortices and the ventromedial prefrontal cortex, structures linked to the experience and regulation of emotion. Rapoport & Shaw (2008) implicate the ‘cortico-striato-thalamocortical loop’ in OCD; limbic structures may account for the strong anxiety component. The role of the amygdala in GAD patients is not well established. Nonetheless, a recent functional magnetic resonance imaging (fMRI) study of youth with GAD found that right amygdala activation associated with emotional stimuli was positively correlated with anxiety severity. The right amygdala and the right ventrolateral prefrontal cortex also had strong negative coupling in reaction
to the emotional stimuli (Monk et al. 2008). Although
more investigation is required into the neural substrates of GAD, this finding and its similarities to other mood and anxiety neuroimaging studies support the notion of a similar ‘emotional’ neural substrate.

**Neurotransmitters**

Abnormalities of the 5-HT transporter gene are associated with trait neuroticism as measured by the Neuroticism–Extroversion–Openness (NEO) Personality Inventory (Schinka et al. 2004; Sen et al. 2004). Individuals with this abnormality may also be at risk for unipolar depression (Lotich & Pollock, 2004; also see above in ‘Genetic factors’ section), and have increased amygdala activity in response to fearful stimuli, when compared to those with a normal homozygous long gene (Hariri et al. 2002). There are abnormalities of the norepinephrine and serotonin systems in most of the emotional disorders, although there are also specific differences between anxiety and depression (Charney, 2003; Martin & Nemeroff, in press).

### Table 4. Summary of studies that have linked temperament across the main symptom domains

<table>
<thead>
<tr>
<th>Diagnoses</th>
<th>Reference</th>
<th>Main relevant findings</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Anxious-misery prominent</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Major depression and</td>
<td>Hettema et al. (2006)</td>
<td>N accounts for between one-third and one-half of the genetic contribution to distress and fear disorders (somatoform and OCD not included). A second genetic factor, independent of N, makes a smaller contribution for depression, GAD and Panic</td>
</tr>
<tr>
<td>generalized anxiety</td>
<td>Kendler et al. (1993)</td>
<td>N robustly predicts MDD in a prospective study</td>
</tr>
<tr>
<td>disorder</td>
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<tr>
<td><strong>Dysthmic disorder</strong></td>
<td>Bienvenu et al. (2004)</td>
<td>All distress and fear disorders, including DD, associated with high N</td>
</tr>
<tr>
<td></td>
<td>Williamson et al. (2005)</td>
<td>All distress and fear disorders, including DD, associated with high N</td>
</tr>
<tr>
<td></td>
<td>Angst (1998)</td>
<td>High levels of N, low quality of life in DD</td>
</tr>
<tr>
<td></td>
<td>Rhebergen et al. (2009)</td>
<td>N scores of DD higher than MDD</td>
</tr>
<tr>
<td><strong>Neurasthaenia</strong></td>
<td>Cao et al. (2005)</td>
<td>N and positive rate of serum Epstein–Barr virus were both pathogenic factors for neurasthaenic patients</td>
</tr>
<tr>
<td></td>
<td>De Gucht et al. (2003)</td>
<td>N was a significant predictor of both current somatization and functional somatic syndromes in patients with chronic fatigue syndrome</td>
</tr>
<tr>
<td><strong>Post-traumatic stress disorder</strong></td>
<td>Cox et al. (2004)</td>
<td>Association between PTSD and N</td>
</tr>
<tr>
<td></td>
<td>van den Hout &amp; Engelhard (2004)</td>
<td>N (measured before trauma) makes independent contribution to PTSD</td>
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<td></td>
<td>Gilbertson et al. (2006)</td>
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<td></td>
<td>Engelhard &amp; van den Hout (2007)</td>
<td></td>
</tr>
<tr>
<td><strong>Fear and avoidance prominent</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Phobias, panic disorder</td>
<td>Hettema et al. (2006), Bienvenu et al. (2004), Williamson et al. (2005)</td>
<td>Associated with high N</td>
</tr>
<tr>
<td><strong>Obsessional disorder</strong></td>
<td>Ettelt et al. (2008)</td>
<td>Harm avoidance high in OCD, also in FDRs</td>
</tr>
<tr>
<td><strong>Somatic symptoms prominent</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unexplained somatic symptoms</td>
<td>Kvaal &amp; Patodia (2000)</td>
<td>Demonstrate association between N and unexplained somatic symptoms</td>
</tr>
<tr>
<td></td>
<td>Ono et al. (2000)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Houtveen &amp; van Doornen (2007)</td>
<td></td>
</tr>
<tr>
<td>Somatoform disorder</td>
<td>McGrady et al. (1999)</td>
<td>Demonstrate association between N and somatoform disorders</td>
</tr>
</tbody>
</table>

DD, Dysthymic disorder; FDR, first-degree relative; GAD, generalized anxiety disorder; MDD, major depressive disorder; N, Neuroticism; OCD, obsessive–compulsive disorder; PTSD, post-traumatic stress disorder.
Biomarkers

We are not aware of a biomarker that features in all the emotional disorders. Individuals reporting elevated levels of negative affect consistently show augmented base startle reactivity (Cook et al. 1991; Lang et al. 1993). Tomarken & Keener (1998) report that depressed patients are characterized by increased activity in the right frontal cortex; however, not all depressed patients show this, so it cannot be thought of as a universal biomarker. Biomarkers for depression include abnormalities of the P300 response (e.g. Kemp et al. 2009). The problem is that these studies do not include patients with the range of disorders that would be necessary to give a firm view about this validator.

Cognitive and emotional processing

It is not the present contention that there are no differences between the various diagnoses contained in the emotional disorders cluster, and there are indeed major differences between them in terms of cognitive and emotional processes. Nonetheless, there are some common themes. Beck (1976) related anxiety to helplessness and depression to hopelessness. Those who become uncertain about their ability to control outcomes (entrapment, in terms of their life situation) and who are relatively certain about negative outcomes (i.e. hopeless) may be expected to be both depressed and anxious. Thus, those who are both helpless and hopeless may be expected to be ‘co-morbid’.

Anxious and depressed subjects differ in several important respects, but share biased judgements on the likelihood that negative events will occur (Krantz & Hamm, 1979; Butler & Mathews, 1983). MacLeod & Byrne (1996) reported that those who are both depressed and anxious showed both greater anticipation of negative experiences and reduced anticipation of positive experiences, whereas those who are only anxious showed only the former.

Gilboa-Schechtman et al. (2002) found that the well-documented memory biases for linguistic material in depressed subjects also apply to visual material. They reported that co-morbid anxious depressives, relative to normal controls, exhibited an enhanced memory for sad and angry versus happy expressions, whereas those who were only anxious did not display this bias.

Mineka et al. (1998) conclude that most evidence suggests that anxiety is associated with automatic attentional biases for emotion-relevant (threatening) material, that depression is associated with memory biases for emotion-relevant (negative) information, and that both anxiety and depression are associated with judgemental or interpretive biases. They conclude that the view that anxious and depressive disorders are two different disorders is replaced by ‘a more nuanced view in which anxiety and depression are posited to have both shared, common components and specific, unique components’.

Differences and similarities in symptomatology

As both the DSM and ICD systems require very different sets of symptoms to achieve diagnostic status within the class of emotional disorders, it is paradoxical that psychiatric screening questionnaires (e.g. the 12-item General Health Questionnaire) are effective in detecting a wide range of emotional disorders. The reason for this is that all the emotional disorders share a common core of symptoms that are of fairly low severity (Grayson et al. 1987; Goldberg et al. 1997). Examples of these milder symptoms are sleeping badly, lacking energy, feeling tired, feeling irritable, worrying and feeling gloomy. It is possible for individuals with an externalizing disorder such as drug dependence, a neurocognitive disorder such as mild dementia or a psychosis such as early schizophrenia to have these symptoms as well; but most of them do not. However, by the time a person is diagnosable with an emotional disorder, they will possess these non-specific symptoms.

Latent structure of symptoms

The underlying relationships between individual symptoms can also be established by latent trait analysis and latent class analysis (Goldberg et al. 1987). When latent trait analysis is applied to data from research interviews such as the Present State Examination, both from community samples and primary care attenders, this produces three correlated traits: anxious symptoms, depressive symptoms and fear symptoms (Ormel et al. 1995). This high correlation is of course due to the large general factor in an unrotated factor analysis of such symptoms.

Factor analysis

Jacob (in press) analysed data from confirmatory factor analysis of psychiatric research interviews given to subjects in seven different countries, and reported that a single factor solution provided only a marginally less good solution than one with two, highly correlated factors of anxiety and depression. Krueger et al. (1998) compared mental disorders at aged 18 and 21 of a prospectively assessed birth cohort, and argued that an internalizing/externalizing model provided a more optimal representation of an individual’s course than more complex models. They argued that co-morbidity may result from common mental disorders being
reliable, covariant indicators of stable, underlying processes, such as ‘internalization’. Krueger et al. (2003) reported similar findings using exploratory factor analysis using the CIDI (Primary Care Version) followed by confirmatory factor analysis on a large data set collected in general health-care settings in 14 different countries. Here, a two-factor model with depression, anxious symptoms (worry and arousal), neurasthenia, somatization and hypochondriasis on the first, and alcohol use disorders on the second provided the best fit for the data. With the US and German data, a three-factor model with depression and anxiety symptoms on an ‘anxious misery’ factor, neurasthenia, somatization and hypochondriasis loading on a ‘somatization factor’, and alcohol use disorders on a third also provided a reasonably good fit, although correlations between the various factors were substantial, at around +0.70.

**Co-morbidity**

High rates of co-morbidity are encountered in both primary care (Ustun & Sartorius, 1995) and community settings (Kessler et al. 1996, 2005, in press). This is because emotional disorders reflect common dimensions of symptoms, and the various symptoms that characterize each emotional disorder represent different combinations of phobic, anxious, depressive or somatic components (Krueger et al. 2003). As severity of disorder increases, so does the likelihood of an individual satisfying more than one of these disorders. If ‘co-morbid’ cases are merely more severe examples of an underlying distress syndrome, it remains to ask how they differ from cases of uncomplicated disorder, such as depressive episode or GAD. In the Dunedin study, co-morbid cases had lower self-esteem and higher neuroticism in adolescence than either disorder on its own (Moffitt et al. 2007a). In a prospective study of children with OCD, Swedo et al. (1989) found these disorders co-morbid with MDD in 35%, overanxious in 18%, phobias in 17% and only occurring on its own in 26% of cases.

Factor analysis of the relationship between diagnoses in this cluster shows that a three-factor model produces the best fit, with the three factors being anxious-misery disorders, fear disorders and externalizing disorders. Somatic symptoms were not assessed in this study (Krueger, 1999a, see Fig. 1). These relationships have now been confirmed by community surveys in The Netherlands (Vollebergh et al. 2001) and Australia (Slade & Watson, 2006). A very similar pattern is reported in two other studies (Cox et al. 2002; Kendler et al. 2003). Other data sets have also identified the internalizing/emotional and externalizing disorders factors (Krueger et al. 1998, 2001, 2003; Kessler et al. 2005; Lahey et al. 2008). They have led to a call for a revised arrangement of diagnostic constructs in DSM-V (Clark & Watson, 2006).

**Course**

Anxiety tends to have an earlier onset than depression, often beginning in childhood and being followed by adolescent depression, and adult depression being preceded by adolescent anxiety. When cases are followed into adult life, the diagnostic overlap increases dramatically between them (Pine et al. 1998; Regier et al. 1998; Wittchen et al. 2000; Moffitt et al. 2007b). Anxiety and depression seem to be a risk factors for each other; in a subsample who had current co-morbid MDD and GAD in the 32-year follow-up of the Dunedin cohort (n = 117), anxiety disorders preceded depression in 42%, depression coming first in 32%, and both occurring simultaneously in 26% (Moffitt et al. 2007b). Bittner et al. (2004) show that any anxiety disorder at age 14 is a risk factor for later depression, with severe impairment being the best predictor. Beesdo et al. (2007) confirm this for social anxiety disorder, with parental anxiety or depression and behavioral inhibition being distal risk factors, and the severity and persistence of earlier symptoms being proximal risk factors. In the Zurich prospective cohort study, co-morbid anxiety and depression is more stable than either disorder on its own, with anxiety on its own being unstable over time. Once co-morbidity develops, the probability of recurrence of either

![Fig. 1. Best-fitting model for the entire National Comorbidity Survey, a three-factor variant of the two-factor internalizing/externalizing model. All parameter estimates are standardized and significant at p <0.05 (after Krueger, 1999a).](image-url)
disorder alone, and particularly anxiety, is far lower than that of co-morbidity (Merikangas et al. 2003).

The course of the emotional disorders is one of episodes of disorder followed by remission, but with high probability of relapse. The mean age of onset of anxiety disorders in the Epidemiologic Catchment Area Study was 16, with depression starting on average 5 years later (Regier et al. 1998), and with rates falling after age 55 (Narrow et al. 2002). In the UK the peak age for neurotic disorders was between 40 and 55 years, with rates falling in older age groups (Office of National Statistics, 2000). Thus, rates of recovery exceed new onsets in those above an approximate age of 50. Over a 3-year period, Lieb et al. (2002b) found that about half the cases of somatoform disorders remitted, but the incidence of new cases was about 26%. Female gender, lower social class, substance use, anxiety and affective disorders and also the experience of traumatic sexual and physical threat events predicted new onsets of somatoform conditions.

These disorders tend to have a relapsing course; in the large UK birth cohort that has been followed to the age of 53, 70% of adolescents who had emotional disorders at both ages 13 and 15 had mental disorders at age 36, 43 or 53, compared with about 25% of the mentally healthy adolescents (Colman et al. 2007). Studies of single disorders tend to have a shorter follow-up period, and depend on the severity of the first episode of disorder; thus, Eaton et al. (2008) followed first episodes of depression in a community sample for at least 13 years and showed that half had only a single episode, whereas 35% had recurrent episodes and 15% were unremitting. Brodaty et al. (2001) followed depressives who had been admitted to hospital for 25 years, and showed that only 12% had recovered and were well whereas 84% had had recurrences. Patients attending specialist clinics for anxiety disorders have been followed for shorter periods. In the 12-year follow-up in the Harvard/Brown Anxiety Disorders Research Program (HARP), only panic disorder had a favourable course, with 82% achieving recovery, compared with 58% for GAD, 48% recovery for panic with agoraphobia and only 37% for social phobia patients. The equivalent figures for recurrences during the follow-up period are 45%, 58% and 39% respectively (Bruce et al. 2005). Major depression is not included in the HARP data set. Fergusson et al. (2006) analysed a longitudinal study of 953 New Zealand children at ages 18, 21 and 25 years, considering the diagnoses of MDD, GAD, panic and phobias. Although a factor labelled by them as an ‘internalizing factor’ makes a substantial contribution to the longitudinal component of each disorder, there was only a disorder-specific factor in two of the disorders: MDD and phobias.

**Treatment**

Once more, our argument is not that there are no differences between various emotional disorders, it is that there are sufficient commonalities to conceptualize these disorders as part of a coherent spectrum. In the short term at least, all emotional disorders respond to some extent to a wide range of psychological interventions, including such simple measures as ‘case management’, with interest and concern, with regular visits and administration of a placebo tablet, as in the control arm of a randomized controlled trial. Cognitive behaviour therapy, with suitable adaptations, can be effective in all of them. Consistent with the data on the neural substrate, selective serotonin reuptake inhibitors (SSRIs) are often effective in all the emotional disorders. However, a thorough meta-analysis by Furukawa et al. (in press) showed that benzodiazepines and azopirones were also equally effective in both anxious and depressive symptoms, as measured by the Hamilton scales on anxiety and depression. Studies were included if they were on the Cochrane Collaboration Depression, Anxiety and Neurosis Group (CCDAN) Registers in December 2006. One possible explanation is that cases of either has some symptoms of both, alternatively it may be that any psychotropic can be shown to be superior to placebo. However, they could also be explained if the two diagnostic entities were very similar to one another.

**Limitations to the validity of the cluster**

Of the ‘validity criteria’ it is clear that the most important are the first four factors. In the emotional disorders, temperamental characteristics are definitely the strongest of these. However, most mental disorders also show higher negative affect than healthy controls, so even this difference is quantitative rather than qualitative. Nor should it be thought that all disorders within the cluster resemble one another closely; for example, a patient with long-standing anxiety (‘GAD’) will have a different pattern of neural activation and endocrine abnormalities than one with an acute depression (Martin & Nemeroff, in press). However, they are both likely to show high negative affect, a higher familial rate of anxiety disorders, and a relatively disadvantaged early life.

Another problem is that a disorder may manifest itself differently at different ages; for example, pre-pubertal anxiety may be followed by an episode of adolescent depression, as the adolescent confronts major problems in peer popularity, educational achievement or sexual choice. Nor is there always a linear relationship between childhood problems and adult disorder; conduct problems at ages 7–9 years...
may be associated with increased risk for antisocial personality disorder and crime in early adulthood (ages 21–25 years), but also with adverse sexual and partner relationships (including domestic violence), early parenthood, and increased risks of substance use, mood and anxiety disorders and suicidal acts (Fergusson et al. 2005). In the Dunedin study, for example, conduct problems at ages 11–15 were associated with increased risk for all psychiatric disorders at age 26, including internalizing problems, schizophreniform disorders and mania, in addition to broadly externalizing phenomena such as substance abuse (Kim-Cohen et al. 2003).

Nor, of course, do abnormalities of personality exist on single dimensions; it is quite possible to be high on both negative emotion and low constraint, and this will influence the type of disorder developed (Krueger et al. 1996; Krueger, 1999b).

It must also be conceded that the evidence presented is stronger in some areas than in others, and the differing strengths of the evidence is shown in Table 5.

Although fairly complete arguments have been advanced for depression, GAD, panic and phobias, the evidence is patchy for PTSD and somatoform disorders, and is extremely sketchy for obsessive states and neurasthenia. The latter two have been included because they have high scores for negative affect, and both contain a common set of non-specific emotional symptoms. There have been few objective studies of neurasthenia as it is not recognized by the DSM system, but when it is diagnosed using the ICD-10 it falls well within the symptom complexes contained in this cluster (Ustun & Sartorius, 1995). The existing classifications are partly responsible for some of the gaps; the necessary work has often not been done.

Clinical usefulness of the proposed classification

The disorders in this cluster are all closely associated with one another, and frequently occur in combination with one another, and all respond to at least some extent to SSRIs and cognitive behaviour therapy. The most frequent example of this is the association of anxious symptoms with depressive symptoms, and a diagnosis of ‘anxious depression’ would allow clinicians to make a single diagnosis rather than declaring the patient to be ‘co-morbid’. In the National Morbidity Survey of 8580 UK respondents, Das-Munshi et al. (2008) showed that the subsyndromal disorder allowed in the ICD-10 had a prevalence of 8.8% and accounted for 20% of all days off work in the country. Taken together with the full syndromes of anxiety and depression, differences in health-related quality of life measures between diagnostic groups were accounted for by overall symptom severity. The finding that half of the anxiety, depression and subsyndromal cases and a third of the co-morbid depression and anxiety cases grouped into a single latent class challenges the notion of these conditions as having distinct phenomenologies. Mixed presentations may be the norm in the population.

For all emotional disorders, an assessment of anxious and depressive symptoms will always be necessary. It should no longer be necessary to diagnose co-morbidity between two very different classes of disorder (e.g. depression and somatoform disorder) when both these disorders occur in the same cluster. Some treatments such as SSRIs may be generally appropriate, but other interventions, such as specific forms of counselling or cognitive behaviour therapy, may be directed at salient symptoms. The proposal

<table>
<thead>
<tr>
<th>Validator</th>
<th>MDD</th>
<th>DD</th>
<th>GAD</th>
<th>Phob</th>
<th>Pan</th>
<th>OCD</th>
<th>NA</th>
<th>PTSD</th>
<th>Somat</th>
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<tr>
<td>Genetics</td>
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DD, Dysthymic disorder; GAD, generalized anxiety disorder; MDD, major depressive disorder; OCD, obsessive–compulsive disorder; Pan, panic disorder; Phob, phobic disorders; PTSD, post-traumatic stress disorder; Somat, somatoform disorders.

Strength of evidence: ★, good; ●, fair; ♦, poor or indirect; ○, absent.

Table 5. Availability of evidence over the 11 validators to support the proposed clusters
would also to decrease the use of NOS (Not Otherwise Specified) categories, which are at present in frequent use.

For internists and general practitioners, the classification will simplify an otherwise confusing system, and encourage clinicians to assess anxious and depressive symptoms whenever they are faced with a patient with other psychological symptoms, or with unexplained somatic symptoms.

Conclusions
There are important differences between individual members of the emotional cluster including the fact that each disorder is defined by some symptoms that do not occur in other disorders, there are differences in cognitive and emotional processing, the neural substrate and the rates for individual disorders in FDRs. These differences should not obscure the similarities between all members of the cluster and should not necessitate putting these disorders into separate chapters of the DSM and ICD classifications.

Acknowledgement
Dr H. Mayberg kindly commented on the ‘Neural substrate’ section.

Declaration of Interest
None.

References


