How can we treat psychosis if we don’t know what we are treating? Over the years, attempts at defining psychosis subtypes have met with dead ends. However, recent research supports a new approach that offers a rational classification model organized according to 5 specific comorbid anxiety and depressive disorder diagnoses.

Anxiety and depressive symptoms are not just the result of psychotic despair. They are specific diagnoses, they precede psychosis onset, they help define psychotic syndromes, and they can point to much more effective treatment approaches. Most of the psychotic diagnoses in this schema are already recognized or posited. And, just as patients who do not have psychotic illness can have more than 1 anxiety or depressive disorder, patients with psychosis can present with a mixed picture that reflects more than 1 contributing comorbidity. Research further suggests that each of the 5 psychosis comorbidity diagnoses may involve some similar underlying factors that facilitate the formation of psychosis.

This article describes the basics of 5 psychosis subtypes, and provides initial guidelines to diagnosis, symptomatology, and treatment. Though clinical experience and existing research support the clinical presence and treatment value of this classification model, further verification will require considerably more controlled studies. An eventual validation of this approach could largely supplant ill-defined diagnoses of “schizophrenia” and other functional psychoses.

Recognizing the comorbidities in the context of their corresponding psychoses entails learning new interviewing skills and devoting more time to both initial and subsequent diagnosis and treatment. In our
5 psychosis subtypes

Psychosis-proneness underlies functional psychoses

Functional (idiopathic) schizophrenia and psychotic disorders have long been difficult to separate, and many categorizations have been discarded. Despite clinical dissimilarities, today we too often casually lump psychoses together as schizophrenia.2,3 Eugen Bleuler first suggested the existence of a “group of schizophrenias.”4 It is possible that his group encompasses our 5 psychoses from 5 inbuilt emotional instincts, each corresponding to a specific anxiety or depressive subtype.

The 5 anxiety and depressive subtypes noted in this article are common, but psychosis is not. Considerable research suggests that certain global “psychotogenic” factors create susceptibility to all psychoses.5-7 While many genetic, neuroanatomical, experiential, and other factors have been reported, the most important may be “hypofrontality” (genetically reduced frontal lobe function, size, or neuronal activity) and dopaminergic hyperfunction (genetically increased dopamine activity).5-7

An evolutionary perspective

One evolutionary theory of psychopathology starts with the subtypes of depression and anxiety. For example, major depressive disorder and generalized anxiety disorder may encompass 5 commonplace and more specific anxiety and depressive subtypes. Consideration of the emotional, cognitive, and functional aspects of those subtypes suggests that they may have once been advantageous for primeval human herds. Those primeval altruistic instincts may have helped survival, reproduction, and preservation of kin group DNA.5

More than any other species, humans can draw upon consciousness and culture to rationally overcome the influences of unconscious instincts. But those instincts can then emerge from the deep, and painfully encourage obedience to their guidance. In nonpsychotic anxiety and depressive disorders, the specific messages are experienced as specific anxiety and depressive symptoms.5 In psychotic disorders, the messages can emerge as unreasoned and frightful fears, perceptions, beliefs, and behaviors. With newer research, clinical observation, and an evolutionary perspective, a novel and counterintuitive approach may improve our ability to help patients.8

Five affective comorbidities evolved from primeval altruistic instincts

Melancholic depression

Melancholic depression is often triggered by serious illness, group exclusion, pronounced loss, or purposelessness. We hear patients talk painfully about illness, guilt, and death. Indeed, some increased risk of death, especially from infectious disease, may result from hypercortisolemia (documented by the dexamethasone suppression test). Hypercortisolemic death also occurs in salmon after spawning, and in male marsupial mice after mating. The tragic passing of an individual saves scarce resources for the remainder of the herd.

Obsessive-compulsive disorder

Factor-analytic studies suggest 4 main obsessive-compulsive disorder (OCD) subtypes: cleanliness, hoarding, intrusive thoughts, and organizing. Obsessive-compulsive traits can help maintain a safe and efficient environment in humans and other species, but OCD is dysfunctional.

Panic anxiety

Panic anxiety is triggered by real, symbolic, or emotional separation from home and family. In toddlers, separation anxiety can reduce the odds of getting lost and hurt.

Social anxiety

Social anxiety includes fear of self-embarrassment, exposure as a pretender to higher social rank, and thus often a reluctant avoidance of increased social rank. While consciousness and cultural...
encouragement can overcome that hesitation and thus lead to greater success, social anxiety activation can still cause painful anxiety. The social hierarchies of many species include comparable biological influences, and help preserve group DNA by reducing hierarchical infighting.

**Atypical depression and bipolar I mania**

Atypical depression includes increased rejection sensitivity, resulting in inoffensive behavior to avoid social rejection. This reduces risk of isolation from the group, and improves group harmony. Unlike the 4 other syndromes, atypical depression and bipolar I mania may reflect 2 separate seasonal mood phases. Atypical depression (including seasonal affective disorder) often worsens with shortened winter daylight hours, akin to hibernation. Initial bipolar I mania is more common with springtime daylight, with symptoms not unlike exaggerated hibernation awakening.

Primeval biological altruism has great evolutionary value in many species, and even somewhat in modern humans. But it is quite different from modern rational altruism. Although we sometimes override our instincts, they respond with messages experienced as emotional pain—they still tell us to follow instructions for primeval herd survival. In an earlier book, I (JPK) provide a lengthier description of the evidence for this evolutionary psychopathology theory, including interplay of the 5 instincts with psychotogenic factors.

**Five comorbidity psychoses from 5 primeval instincts**

The 5 affective comorbidities described above contribute to the presence, subtype, and treatment approaches of 5 corresponding psychoses. Ordinary panic attacks might occur when feeling trapped or separated from home, so people want to flee to safety. Nonhuman species with limited consciousness and language are unlikely to think “time to head for safety.” Instead, instincts encourage flight from danger through internally generated perceptions of threat. Likewise, people with psychosis and panic, without sufficient conscious modulation, may experience sensory perceptions of actual danger when feeling symbolically trapped.

One pilot study carefully examined the prevalence of these 5 comorbidities in an unselected group of psychotic patients. At least 85% met criteria for ≥1 of the 5 subtypes. Moreover, organic psychoses related to physical illness, substances, and iatrogenesis may also predict future episodes of functional psychoses.

Using statistical analysis of psychosis rating scales, 2 studies took a “transdiagnostic” look at psychoses, and each found 5 psychosis subtypes and a generalized psychosis susceptibility factor. Replication of that transdiagnostic approach, newly including psychosis symptoms and our 5 specific comorbidities, might well find that the 5 subtype models resemble each other.

Our proposed 5 comorbidity subtypes are:

**Delusional depression (melancholic depression).** Most common in geriatric patients, this psychosis can also occur at younger ages. Prodromal melancholic depression can include guilt and hopelessness, and is acute, rather than the chronic course of our other 4 syndromes. Subsequent delusional depression includes delusions of bodily decay, illness, or death, as well as overwhelming guilt, shame, and remorse. The classic vegetative symptoms of depression continue. In addition to infectious disease issues, high suicide risk makes hospitalization imperative.

**Obsessive-compulsive schizophrenia.** Just as OCD has an early age of onset, obsessive-compulsive schizophrenia begins earlier than other psychoses. Despite preserved cognition, some nonpsychotic patients with OCD have diminished symptom insight. OCD may be comorbid with schizophrenia in 12% of cases, typically preceding psychosis onset. Obsessive-compulsive schizophrenia symptoms may include highly exaggerated doubt or ambivalence; contamination concerns; eccentric, ritualistic, motor stereotypy, checking, disorganized, and other behaviors; and paranoia.

**Schizophrenia with voices (panic anxiety).** Classic paranoid schizophrenia with voices...
appears to be the most similar to a “panic psychosis.” Patients with nonpsychotic panic anxiety have increased paranoid ideation and ideas of reference as measured on the Symptom Checklist-90. Schizophrenia is highly comorbid with panic anxiety, estimated at 45% in the Epidemiologic Catchment Area study. These are likely underestimates: cognitive impairment hinders reporting, and psychotic panic is masked as auditory hallucinations. A pilot study of schizophrenia with voices using a carbon dioxide panic induction challenge found that 100% had panic anxiety. That study and another found that virtually all participants reported voices concurrent with panic using our Panic and Schizophrenia Interview (PaSI) (Box 1, page 28). Panic onset precedes schizophrenia onset, and panic may reappear if antipsychotic medications sufficiently control voices: “voices without the voices,” say some.

**Persecutory delusional disorder (social anxiety).** Some “schizophrenia” without voices may be misdiagnosis of persecutory (paranoid) delusional disorder (PDD). Therefore, the reported population prevalence (0.02%) may be underestimated. Social anxiety is highly comorbid with “schizophrenia” (15%). Case reports and clinical experience suggest that PDD is commonly preceded by social anxiety. Some nonpsychotic social anxiety symptoms closely resemble the PDD psychotic ideas of reference (a perception that low social rank attracts critical scrutiny by authorities). Patients with PDD may remain relatively functional, with few negative symptoms, despite pronounced paranoia. Outward manifestation of paranoia may be limited, unless quite intense. The typical age of onset (40 years) is later than that of schizophrenia, and symptoms can last a long time.

**Bipolar I mania with delusions (atypical depression).** Atypical depression is the most common depression in bipolar I disorder. Often more pronounced in winter, it may intensify at any time of year. Long ago, hypersomnia, lethargy, inactivity, inoffensiveness, and craving high-calorie food may have been conducive to hibernation.

Bipolar I mania includes delusions of special accomplishments or abilities, energetically focused on a grandiose mission to help everyone. These intense symptoms may be related to reduced frontal lobe modulation. In some milder form, bipolar I mania may once have encouraged hibernation awakening. Indeed, initial bipolar I mania episodes are more common in spring, as is the spring cleaning that helps us prepare for summer.

**Recognizing affective trees in a psychotic forest**

Though long observed, comorbid affective symptoms have generally been considered a hodgepodge of distress caused by painful psychotic illness. But the affective symptoms precede psychosis onset, can be masked during acute psychosis, and will revert to ordinary form if psychosis abates.

Rather than affective symptoms being a consequence of psychosis, it may well be the other way around. Affective disorders could be important causal and differentiating components of psychotic disorders. Research and clinical experience suggest that adjunctive treatment of the comorbidities with correct medication can greatly enhance outcome.

**Diagnostic approaches**

Because interviews of patients with psychosis are often complicated by confusion, irritability, paranoid evasiveness, cognitive impairment, and medication, nuanced diagnosis is difficult. Interviews should explore psychotic syndromes and subtypes that correlate with comorbidity psychoses, including pre-psychotic anxiety and depressive diagnoses that are chronic (though unlike our 4 other diagnoses, melancholic depression is not chronic).

Establishing pre-psychotic diagnosis of chronic syndromes suggests that they are still present, even if they are difficult to assess during psychosis. Re-interview after some improvement allows for a significantly better diagnosis. Just as in
nonpsychotic affective disorders, multiple comorbidities are common, and can lead to a mixed psychotic diagnosis and treatment plan.  

Structured interview tools can assist diagnosis. The PaSI (Box 1, page 28) elicits past, present, and detailed history of DSM panic, and has been validated in a small pilot randomized controlled trial. The PaSI focuses patient attention on paroxysmal onset voices, and then evaluates the presence of concurrent DSM panic symptoms. If voices are mostly psychotic panic, they may well be a proxy for panic. Ultimately, diagnosis of 5 comorbidities and associated psychotic symptoms may allow simpler categorization into 1 (or more) of the 5 psychosis subtypes.

**Treatment by comorbidity subtype**

Treatment of psychosis generally begins with antipsychotics. Nominal psychotherapy (presence of a professionally detached, compassionate clinician) improves compliance and leads to supportive therapy. Cognitive-behavioral therapy and dialectical behavior therapy may help later, with limited interpersonal approaches further on for some patients.

The suggested approaches to pharmacotherapy noted here draw on research and clinical experience. All medications used to treat comorbidities noted here are approved or generally accepted for that diagnosis. Estimated doses are similar to those for comorbidities when patients are nonpsychotic, and vary among patients. Doses, dosing schedules, and titration are extremely important for full benefit. Always consider compliance issues, suicidality, possible adverse effects, and potential drug/drug interactions. Although the medications we suggest using to treat the comorbidities may appear to also benefit psychosis, only antipsychotics are approved for psychosis per se.

**Delusional depression.** Antipsychotic + antidepressant. Tricyclic antidepressants are possibly most effective, but increase the risk of overdose and dangerous falls among fragile patients. Electroconvulsive therapy is sometimes used.

**Obsessive-compulsive schizophrenia.** Antipsychotic + selective serotonin reuptake inhibitor (SSRI). Consider aripiprazole (consider long-acting injectable formulation for increased compliance). Aripiprazole also may enhance the benefit of fluoxetine for comorbid OCD. Carefully titrate, as tolerated, to optimal dose of fluoxetine (40 to 80 mg/d; the long half-life of fluoxetine and its metabolite norfluoxetine also improves compliance), while watching for activation and other adverse effects. Limited clinical experience suggests that lower-dose clonazepam every 12 hours may reduce the adverse effects of fluoxetine.

**Schizophrenia with voices.** Antipsychotic + clonazepam. Concurrent usage may stabilize psychosis more rapidly, and with a lower antipsychotic dose. Titrate a fixed dose of clonazepam every 12 hours (avoid as-needed doses), starting low (ie, 0.5 mg) to limit initial drowsiness (which typically diminishes in 3 to 10 days). Titrate to full voice and panic cessation (1 to 2.5 mg every 12 hours). Exercise caution about excessive drowsiness, as well as outpatient compliance and abuse. Besides alprazolam, other antipanic medications have little incidental benefit for psychosis.

**Persecutory delusional disorder.** Antipsychotic + SSRI. Aripiprazole (consider long-acting injectable for compliance) also enhances the benefits of fluoxetine for social anxiety. Long half-life fluoxetine (20 mg/d) improves compliance and near-term outcomes.

**Bipolar I mania: mania with delusions.** Consider olanzapine for acute phase, then add other antimanic medication (commonly lithium or valproic acid), check blood level, and then taper olanzapine some weeks later. Importantly, lamotrigine is not effective for bipolar I mania. Consider suicide risk, medical conditions, and outpatient compliance. Comorbid panic anxiety is also common in bipolar I mania, often presenting as nontrengthening voices.

**Seasonality:** Following research that bipolar I mania is more common in...
**Box 1**

**Panic and Schizophrenia Interview**

Let’s talk for a minute about your voices.

**[IDENTIFYING PAROXYSMAL MOMENTS OF VOICE ONSET]**

Do you hear voices at every single moment, or are they sometimes silent? Think about those times when you are not actually hearing any voices.

Now, there may be reasons why the voices start talking when they do, but let’s leave that aside for now.

So, whenever the voices do begin speaking—and for whatever reason they do—is it all of a sudden, or do they start very softly and then very gradually get louder?

If your voices are nearly always there, then are there times when the voices suddenly come back, get louder, get more insistent, or just get more obvious to you?

**[Focus patient on sudden moment of voice onset, intensification, or awareness]**

Let’s talk about that sudden moment when the voices begin (or intensify, or become obvious), even if you know the reason why they start.

I’m going to ask you about some symptoms that you might have at that same sudden moment when the voices start (or intensify, or become obvious). If you have any of these symptoms at the other times, they do not count for now.

So, when I ask about each symptom, tell me whether it comes on at the same sudden moments as the voices, and also if it used to come on with the voices in the past.

For each sudden symptom, just say “YES” or “NO” or “SOMETIMES.”

**[Begin each query with: “At the same sudden moment that the voices come on”]**

1. Sudden anxiety, fear, or panic on the inside? Y N S
2. Sudden anger or rage on the inside? [ANGER QUERY] Y N S
3. Sudden heart racing? Heart pounding? Y N S
5. Sudden sweating? Y N S
6. Sudden trembling or shaking? Y N S
7. Sudden shortness of breath, or like you can’t catch your breath? Y N S
8. Sudden choking or a lump in your throat? Y N S
9. Sudden nausea or queasiness? Y N S
10. Sudden dizziness, lightheadedness, or faintness? Y N S
11. Sudden feeling of detachment, sort of like you are in a glass box? Y N S
12. Sudden fear of losing control? Fear of going crazy? Y N S
13. Sudden fear afraid of dying? Afraid of having a heart attack? Y N S
14. Sudden numbness or tingling, especially in your hands or face? Y N S
15. Sudden feeling of heat, or cold? Y N S
17. Sudden fear that people want to hurt you? [EXCESS FEAR QUERY] Y N S

**Clinical Point**

Case reports and clinical experience suggest that persecutory delusional disorder is commonly preceded by social anxiety.

Source: Reference 15 © Jeffrey P. Kahn, MD
[PAST & PRODROMAL PANIC HISTORY]
At what age did you first see a therapist or psychiatrist?
At what age were you first hospitalized for an emotional problem?
At what age did you first start hearing voices?
At what age did you first start having strong fears of other people?
Before you ever heard voices, did you ever have any of the other sudden symptoms like the ones we just talked about? Y N S
Did those episodes back then feel sort of like your voices or sudden fears do now, except that there were no voices or sudden fears of people back then? Y N S
At what age did those sudden anxiety (or panic or rage) episodes begin?
Back then, was there MORE (M) sudden anxiety, or the SAME (S) sudden anxiety, or LESS (L) sudden anxiety than with your sudden voices now? M S L

[PAST & PRODROMAL PANIC SYMPTOMS]
Now let’s talk about some symptoms that you might have had at those same sudden anxiety moments, in the time before you ever heard any voices. So, for each sudden symptom just say “YES” or “NO” or “SOMETIMES.”

[Begin each query with: “At the same moment the sudden anxiety came on—but only during the time before you ever heard sudden voices”]
[Ask about the same 18 panic-related symptoms listed above]

[PHOBIA-RELATED PANIC AND VOICES]
Have you ever been afraid to go into a (car, bus, plane, train, subway, elevator, mall, tunnel, bridge, heights, small place, CAT scan or MRI, being alone, crowds)? Y N M

[If yes or maybe: Ask about panic symptoms in phobic situations]
Now let’s talk about some symptoms that you might have had at some of those times you were afraid. So, for each symptom just say “YES” or “NO” or “MAYBE.”

[Ask about the same 18 panic-related symptoms listed above]
At what age did you last have sudden anxiety without voices? Y N M
Has medication ever completely stopped your voices? Somewhat? Y N M
If so, did those other sudden symptoms still happen sometimes? Y N M

Thank you for your help, and for answering all of these questions!

Clinical Point
Bipolar I mania includes delusions of special accomplishments or abilities focused on a grandiose mission.
spring and summer, studies have shown beneficial clinical augmentation from dark therapy as provided by reduced light exposure, blue-blocking glasses, and exogenous melatonin (a darkness-signaling hormone).²⁴

**Bipolar I mania atypical depression (significant current or historical symptoms).** SSRI + booster medication. An SSRI (i.e., escitalopram, 10 mg/d) is best started several weeks after full bipolar I mania resolution, while also continuing long-term antimanic medication. Booster medications (i.e., buspirone 15 mg every 12 hours; lithium 300 mg/d; or trazodone 50 mg every 12 hours) can enhance SSRI benefits. Meta-analysis suggests SSRIs may have limited risk of inducing bipolar I mania.²⁵ Although not yet specifically tested for atypical depression, lamotrigine may be effective, and may be safer still.²³ However, lamotrigine requires very gradual dose titration to prevent a potentially dangerous rash, including after periods of outpatient noncompliance.

**Seasonality:** Atypical depression is often worse in winter (seasonal affective disorder). Light therapy can produce some clinically helpful benefits year-round.

To illustrate this new approach to psychosis diagnosis and treatment, our book includes detailed case studies on each of the 5 psychosis subtypes. The brief fictional case we present in **Box 2** describes a patient who had both premorbid social anxiety and panic anxiety, and then developed a mixed psychosis that reflected both of those contributing anxiety disorders.

**Larger studies are needed**

Current research supports the concept of a 5-diagnosis classification of psychoses, which may correlate with our comorbid anxiety and depression model. Larger diagnostic and treatment studies would invaluably examine existing research and clinical experience, and potentially encourage more
clinically useful diagnoses, specific treatments, and improved outcomes.

Acknowledgement
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References

Related Resources
• Veras AB, Kahn JP, eds. Psychotic Disorders: Comorbidity Detection Promotes Improved Diagnosis and Treatment. Elsevier; 2021.

Drug Brand Names
- Alprazolam - Xanax
- Aripiprazole - Abilify
- Buspirone - BuSpaR
- Clonazepam - Klonopin
- Escitalopram - Lexapro
- Fluoxetine - Prozac
- Lamotrigine - Lamictal
- Lithium - Eskalith, Lithobid
- Olanzapine - Zyprexa
- Trazodone - Oleptro
- Valproic acid - Depakote

Clinical Point
Larger studies would potentially encourage more clinically useful diagnoses and specific treatments

Bottom Line
New insights from evolutionary psychopathology, clinical research and observation, psychotogenesis, genetics, and epidemiology suggest that most functional psychoses may fall into 1 of 5 comorbidity-defined subtypes, for which specific treatments can lead to much improved outcomes.