Characteristics, correlates, and outcomes of childhood and adolescent depressive disorders.
Developmental psychopathology has identified the defining clinical and contextual features of depression in youngsters. In particular, empirical studies have characterized the longitudinal course of depressive illness and common patterns of co-occurring psychiatric conditions. The functional consequences of early-onset illness have also been documented. A growing body of research is identifying the neurobiological and psychological correlates. In addition, studies are beginning to identify specific genetic and experiential risk factors. In general, the core patterns of depressive disorders across the lifespan are emerging. This paper details the phenomenology, correlates, clinical course, and consequences of pediatric depression, highlighting the similarities and differences in the characteristics of depression among children, adolescents, and adults. A few caveats are warranted before...
proceeding to the following sections. The term “depression” refers to unipolar mood disorders only. Up to now, most of the research on pediatric depression was conducted in major depressive disorder, and therefore, the reported findings are primarily for this condition.

**Historical context of depression in children and adolescents**

Case reports of youngsters exhibiting symptoms resembling depressive disorders in adults were described as early as the 17th century. Nevertheless, early theories of depression discounted the validity of the disorder in youth, suggesting that the necessary psychological mechanisms were not yet present for the experience of depression, or that depression was “masked” in the form of other disorders. In particular, it was considered that children did not have a well-developed superego. In 1975, the National Institute of Mental Health convened a meeting to discuss the incidence and diagnosis of depression in children. When the existence of depression in children became acceptable and the basic diagnostic criteria were established, research on childhood depression burgeoned, resulting in the growth of theoretical models as well as empirical databases, and depression is no longer considered “an adult disease.” Despite this burgeoning research, some obstacles remained with regard to the pursuit of knowledge on adolescent depression. The early “storm and stress” theories of development suggested that many of the problems associated with depression, such as sadness, irritable mood, self-doubt, and social withdrawal, were normative expressions of adolescent angst. It is now established, however, that many youngsters do not go through such emotional distress, and that adolescent depression is a serious disorder, often heralding chronic or recurrent problems into adulthood.

**A developmental framework in understanding childhood and adolescent depression**

In the past three decades, depression research in children and adolescents has progressed from applying simple extensions of clinical descriptions and theories developed in adults to generating an increasingly sophisticated understanding of these disorders informed by the emerging field of developmental psychopathology. Research adopting this framework has taken into account the normative developmental processes influencing differences in the etiology, phenomenology, correlates, and outcomes of depression in children, adolescents, and adults. It is important to note, however, that this new field of research often does not differentiate among particular stages of development through childhood and adolescence. Although some continuity is likely across childhood and adolescence in the experience and expression of depression, the underlying risk mechanisms and the consequences of depression, some differences are also plausible.

When applying a developmental perspective to psychopathology, one important issue to consider is the conceptualization of different life stages. For example, the transition from childhood to adolescence involves changes in multiple domains, including physical, sexual, cognitive, and social development, with a considerable range of individual differences in the age at which each of these changes occur. At present, there is no consensus on the clear boundaries in defining child and adolescent populations. Since gathering information on these multiple domains is complex, for pragmatic reasons, the majority of the studies have used chronological age to define these boundaries; children ≤12 years and adolescents between 13 and 18 years. In some cases, however, studies are reviewed in which these ages overlap (eg, some studies included 13-year-olds in the child samples, whereas others included 12-year-olds among adolescent samples, and still others reported findings according to grade level or physical pubertal status).

**Epidemiology of unipolar depression in children and adolescents**

**Prevalence and incidence**

Prevalence estimates of unipolar depression vary with the time period of reference and method of assessment. The reported point prevalence rates (30-day or 1-year) of major depressive disorder in nonreferred samples range between 0.4% and 2.5% in children, and between 0.7% and 9.8% in adolescents. Elevated risk for the disorder begins in the early teens, and continues to rise in a linear fashion throughout adolescence, with lifetime rates estimated to range from 15% to 25% by late adolescence. These prevalence estimates of adolescent depression are comparable to the lifetime rates reported in adults, suggesting that the rates of depression begin to plateau by early adult life. These data also indicate
that, for a substantial proportion of adult cases, the onset occurred during adolescence.\textsuperscript{14} The prevalence of depression in youngsters is even greater when minor depression and subsyndromal depressive symptoms are considered. In the National Comorbidity Study, the only nationally representative community study in the United States that included adolescents, the lifetime prevalence of minor depression in 15- to 18-year-olds was 11\%.\textsuperscript{15} In a large sample of high-school students, up to 40\% of adolescents exceeded the cutoff point for high symptom levels on self-reported depressive symptoms.\textsuperscript{16} Subsyndromal depression is associated with high levels of distress and impaired functioning,\textsuperscript{17} and prospective studies indicated that it is a strong predictor of major depressive disorder.\textsuperscript{18,19}

**Secular trends**

Retrospective data from successive cohorts born since World War II suggest that the diagnosis of unipolar depression may be becoming increasingly common and beginning earlier in life.\textsuperscript{20,21} Interpreting secular trends is complicated because of increased clinical awareness of early-onset depression and changing diagnostic practices. However, the recent replication results from the National Comorbidity Survey and some studies of pediatric clinical cohorts are very compelling, because adjusted lifetime hazard rates of depression are based on the same interview methods with participants across different age groups ascertained at the same time.\textsuperscript{13,22,23}

**Gender differences**

Epidemiological studies have consistently demonstrated that females are two to three times more likely than males to develop depression.\textsuperscript{24} The female predominance in depression has been observed across many countries and cultures, as well as in cohorts across multiple generations.\textsuperscript{25,26} A developmental trend has been observed for sex differences in the prevalence of depression. Prior to adolescence, the rate of depressive disorders is about equal in boys and girls, or even higher among boys.\textsuperscript{5,27} During early to middle adolescence, the rate of depressive symptoms and disorders in girls rises by two to three times that of boys, a trend that continues through adult life.\textsuperscript{11-13,28} Explanations for this gender difference have included hormonal changes, increased stress, tendencies toward rumination and other maladaptive responses to stress, and differences in interpersonal orientation and socialization experiences.\textsuperscript{7,29,30}

**Effect of social status**

The effect of social class on depression has been well-documented in adults.\textsuperscript{31} Studies in children and adolescents, using both symptom levels and diagnostic criteria, have linked depression to lower income and socioeconomic status.\textsuperscript{32,34} Lower socioeconomic status may be a marker of specific risk factors associated with depression, rather than exerting a direct influence. For example, low socioeconomic status is associated with high levels of chronic stress due to economic difficulties, adverse environmental conditions, and family disruption.

**Ethnic and cultural differences**

Among adults, ethnic/racial differences have been reported with regard to prevalence rates of depression, with ethnic minority groups having lower rates compared with non-Hispanic whites.\textsuperscript{35,36} Among child and adolescent samples, either no ethnic differences were detected in depression, or ethnic minority groups, Hispanics, in particular, had higher symptom levels.\textsuperscript{33,37-39} However, analysis of data from more nationally representative samples in the United States revealed complex relationships between ethnicity and depression. For example, one study found that African-American girls did not manifest the puberty-related increase in depressive symptoms that is commonly observed in non-Hispanic white girls.\textsuperscript{40} It appears that both individual- and context-level characteristics exert effects on depressive symptomatology. For instance, African-American teens living within predominantly non-Hispanic white neighborhoods were at especially high risk for depressive symptoms.\textsuperscript{41} Similarly, adolescents of Mexican descent living in the United States, but not Mexican adolescents, had higher rates of depression compared with non-Hispanic white youth.\textsuperscript{37} Future investigations should attempt to disentangle the effects of adverse conditions that might be associated with ethnic status as well as cultural expressions of depressive symptoms among different ethnic groups.\textsuperscript{41,42}

**Functional consequences and socioeconomic burden**

Depressive disorders in children and adolescents are associated with significant economic and social burden
on individuals, families, and societies.\textsuperscript{43-46} The functional consequences that characterize depressed youngsters suggest that the disorder can interfere with developmental milestones. For example, depression is frequently associated with problems in interpersonal relationships and school performance, as well as delays in social, emotional, and cognitive development.\textsuperscript{47-52} It is not clear, however, whether these psychosocial disturbances are precursors or consequences of depression. Moreover, other factors frequently associated with depression, such as comorbid psychiatric disorders, poor family functioning, low socioeconomic status, and exposure to stressful life events, impact psychosocial functioning.\textsuperscript{47-55} Depression in children and adolescents is also associated with an increased frequency of suicidal behaviors, delinquency, and alcohol and drug use.\textsuperscript{50,53} Prospective studies found that after recovery, children and adolescents continue to manifest impaired psychosocial functioning in multiple domains.\textsuperscript{56-58} Moreover, children and adolescents with depression have persistent psychosocial problems in adult life, including criminal behavior, dysfunctional interpersonal relationships, early pregnancy, low educational attainment, poor occupational functioning, unemployment, and suicidal behavior.\textsuperscript{60} Some studies also reported high rates of psychiatric hospitalization and mental health services compared with their counterparts without depression.\textsuperscript{61} Data in adults suggest that depressed patients with early-onset illness have more impaired social and occupational functioning and poorer quality of life compared with patients whose episode(s) first started in adult life.\textsuperscript{59}

\textbf{Clinical presentation of depression in children and adolescents}

\textbf{Developmental influences}

The diagnosis of dysthymic disorder and major depressive disorder are based on similar criteria for children, adolescents, and adults, with two exceptions. First, the \textit{Diagnostic and Statistical Manual of Mental Disorders} (4th edition with text revisions; \textit{DSM-IV-TR}) has allowed the substitution of irritability for depressed mood in children and adolescents.\textsuperscript{62} Second, the duration criterion for dysthymic disorder in children and adolescents is 1 year instead of 2.\textsuperscript{63} Empirical data also suggest that the clinical syndrome of depression is remarkably similar among children, adolescents, and adults.\textsuperscript{64,65} There are some developmental differences, however.\textsuperscript{66} Specifically, hypersomnia shows a developmental trend, with a higher prevalence in depressed adolescents than in children.\textsuperscript{65-66} Suicide attempts, particularly those involving high lethality, also increase with age.\textsuperscript{66,67} Melancholic and psychotic symptoms may occur less frequently in children, whereas somatic complaints and behavior problems are more common during this developmental period.\textsuperscript{64,67-69} Psychotic depression in children appears to be manifested by auditory hallucinations instead of delusions, as seen in adolescents and adults. Although the reasons for the developmental variations in depressive symptoms are not known, maturational effects on emotional and behavioral regulation and cognitive function might contribute to these differences.

\textbf{Gender differences}

Gender differences have also been documented with respect to the severity and symptom profiles of unipolar depression among children, adolescents, and adults although no compelling gender effects were found on the salient features.\textsuperscript{62,67-70} Among both adolescents and adults, females typically report higher levels of symptoms.\textsuperscript{71-74} With regard to symptom patterns, somatic symptoms, such as changes in appetite and weight, sleep problems, and psychomotor retardation are more common in females.\textsuperscript{75,76} Increased crying, feelings of failure, guilt, poor self-esteem, and other cognitive symptoms may also be more frequent in females.\textsuperscript{75-77,79-81} In contrast, depressed males more frequently report anhedonia, diurnal variation in mood and energy, social withdrawal, and work impairment.\textsuperscript{75,76,80} The reasons for gender differences in unipolar depression are not well understood. One model suggests that females are more prone to exhibit a cognitive style characterized by negative self-evaluation and rumination.\textsuperscript{82,83} Gender differences in brain function have also been postulated as one potential reason for the symptom variability.\textsuperscript{84,85}

\textbf{Comorbidity}

Both clinical and epidemiological studies have shown that up to 40\% to 70\% of children and adolescents with depression also suffer from another psychiatric disorder, and many youngsters have two or more comorbid diagnoses.\textsuperscript{86-88} Approximately 70\% of children and adolescents with dysthymic disorder will eventually develop an episode of major depressive disorder, resulting in “double depression.”\textsuperscript{89,90} Other frequent comorbid diagnoses include anxiety disorders, disruptive disorders, and sub-
stance use disorders. Although it is not clear whether these comorbid conditions represent a developmental sequence, shared genetic or environmental risk factors or a separate subtype of the disorder, it is likely that one or more of these factors contribute to comorbidity. Age and gender can influence the patterns of comorbidity. Specifically, separation anxiety disorder and attention-deficit hyperactivity disorder are more common in children, whereas conduct disorder, panic disorder, and substance abuse are more common in adolescents. Similarly, disruptive and substance use disorders are less likely, and eating disorders are more likely, in girls than boys. The presence of comorbidity has important clinical and functional implications. In particular, youth with comorbid dysthymia and major depression had more severe and longer depressive episodes, and higher frequency of suicidality and social impairment than those who had a single mood disorder. Similarly, comorbid anxiety disorder was associated with increased severity and duration of depressive symptoms, increased suicidality, poor response to psychotherapy, and elevated risk for addictive disorders. In contrast, depressed youth with co-occurring disruptive disorders tended to have fewer melancholic symptoms, fewer recurrent episodes of depression, a lower frequency of familial mood disorders, a higher incidence of criminal behavior, and a higher response to placebo than patients with pure depressive illness. Comorbid substance abuse is associated with earlier onset and more severe substance-related problems, increased frequency of behavioral problems, more prolonged and recurrent depressive episodes, more severe impairment in family, school, and legal domains, higher risk for suicidal behavior, and increased utilization of health services and substantially higher treatment costs. Examination of data in adults suggest that, compared with depressed patients whose first depressive episode occurs in adult life, patients with early-onset illness have increased rates of anxiety disorders and substance use disorders, as well as personality disorders, resulting in more chronicity and disability.

**Developmental course and outcomes of childhood and adolescent depression**

**Episode duration**

Considerable variations have been found in the duration of depressive episodes in nonreferred and clinical samples of youth. For example, in a large sample of high-school students, the duration of major depressive episode ranged from 2 weeks to 250 weeks, with a mean duration of 26 weeks. The probability of remission was 3 weeks in 25% of the sample, 8 weeks in 50%, and 24 weeks in 75% of the sample. Longer durations were reported in clinical samples, with a mean length of 6 to 9 months. Up to 30% to 40% of patients can be expected to recover by 6 months and 70% to 80% by 12 months, and 5% to 10% of patients have a protracted episode, lasting longer than 2 years. Dysthymic episodes tend to be more protracted, with an average duration of 2.5 to 3.5 years. In a prospective study of a clinical sample, only 7% of youth with dysthymia showed evidence of recovery 2 years after the onset of a first episode. Overall, children and adolescents have similar recovery patterns, and these patterns also are comparable to the data in adults. Among the baseline demographic and clinical variables that were examined, none has yet been shown to consistently predict recovery from a depressive episode in youngsters. Age at onset of illness, greater severity, presence of comorbid disorders, and parental history of depression potentially influence the time to recovery. Among adults, greater severity, longer duration of episode at the time of recruitment, pre-existing dysthymic disorder, and co-occurring anxiety disorders and personality problems were associated with longer time to recovery.

**Recurrence and continuity into adulthood**

Longitudinal studies of both epidemiological and clinical samples consistently demonstrated that children and adolescents with depression tend to have recurrent episodes. The probability of recurrence following the recovery of a major depressive episode is approximately 40% by 2 years and 70% by 5 years. These rates are comparable to the rates of recurrence in adult samples. In addition to recurrent episodes during childhood and adolescence, longitudinal studies of depressed youngsters documented recurrent episodes in adult life. There also appears to be some specificity in the continuation of psychopathology in adult life, particularly with respect to adolescent-onset depression. Several studies of depressed adolescents documented increased risk for recurrent depressive episodes, but not other psychiatric disorders, when compared with their counterparts without depression. In contrast, there is some evidence that child-

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hood-onset depression is not necessarily predictive of depression in adulthood, except for a subsample with symptoms characteristic of the adult disorder.128,130,131 Among children, adolescents, and adults, few baseline demographic or clinical characteristics predict who will or will not experience a recurrent depressive episode. Some potential predictors of recurrence in adults include early onset, number of prior episodes, stressful experiences, cognitive vulnerability, negative family interaction patterns, comorbid personality disorders, and persistent biological dysregulation during recovery.59,132 Among youth, co-occurring personality problems, specifically borderline personality disorder symptoms, were associated with recurrence.125,133 There is disagreement regarding whether girls are at increased risk for recurrent depressive episodes than boys.60,111,112,133 although no gender differences were found in recurrence rates among adults.134

Other psychiatric outcomes

Although recurrent unipolar depression is the primary outcome for depressed youth, development of other psychiatric disorders has also been documented. Longitudinal studies reported that 20% to 40% of children and adolescents with major depressive disorder developed bipolar disorder within a period of 5 years.48,129,135,136 The clinical characteristics associated with increased risk for bipolar disorder in youngsters and adults with depression include early-onset illness, mood lability, depressive episode accompanied by psychomotor retardation or psychotic features, atypical depression, protracted depressive episodes, family history of bipolar disorder or heavy familial loading for mood disorders, and pharmacologically induced hypomania.94,136-139 Depressed youngsters are also at risk for developing substance use disorders in adolescence and adulthood.12,88,101,140-142 Protracted depressive episodes, comorbid anxiety or conduct disorder, and hypothalamic-pituitary-adrenal (HPA) dysregulation may be associated with increased risk for substance abuse in depressed youth.101,103,142

Risk factors for depression in children and adolescents

Developmental influences on vulnerability to depression

As described above, the risk for depression increases markedly during the transition from childhood to adolescence. Adolescence is a crucial developmental stage, marked by a confluence of biological, psychological, and social challenges.143,146 There are significant physical maturational changes (eg, the onset of puberty), social-cognitive advances (eg, ability for more abstract thinking and generalizations across situations and time), interpersonal transitions (eg, changes in social roles in family and peer relationships), and social-contextual changes (eg, school transitions). Although these maturational transitions offer tremendous opportunities for youth, because the developing brain and behavioral and cognitive systems mature at different rates, and because these systems are under the control of both common and independent biological processes, this developmental period also is marked by heightened vulnerability. The normative developmental transitions associated with adolescence might serve as sensitive periods for the activation of specific processes involved in the onset, persistence, and recurrence of depressive episodes.147,148

Family-genetic factors

There is clear evidence of familial transmission of depression.149-151 These data, however, cannot distinguish environmental from genetic causes of transmission. Family, twin, and adoption studies indicated effects of both genetic and environmental factors for unipolar depression.152,153 Based on epidemiological data, the proportion of variance attributed to genetic factors is between 24% and 58% for depressive illness.154 Genetic influences have been found to vary with age and sex. Shared environmental influences may be more important in younger children, and these influences may be replaced by new genetic and unique environmental influences as children grow older.150,155 In one study, the increased heritability effect in adolescents was found only for girls, and not boys.136 Research on behavioral genetics initially partitioned population variance into two components, one due to genetic factors and the second due to environmental influences. The implication was that the two were separate, and it was assumed that gene-environment interactions were usually of so little importance that they could be ignored. Theoretical considerations suggested that this was not likely to be true, and empirical findings are now accumulating on the interactions between identified common single genetic variants and environmentally-mediated risks.157 Indeed, the important role of environmental factors in modulating vulnerability and their
interactions with genetic variants has been specifically demonstrated for depression.\textsuperscript{152,158,159}

Recent research on genetic liability for depression has begun to address the mode of inheritance, such as temperament characteristics associated with emotionality and emotional regulation, a tendency toward stress exposure and reactivity, and alterations in neurobiological regulation.\textsuperscript{160-162} Molecular genetic effects related to early-onset depression have been observed on chromosome 11p14 that has been associated with brain-derived neurotrophic factor (BDNF).\textsuperscript{163,164} BDNF is a nerve growth factor involved in the regulation of cellular development, neuronal survival, synaptic plasticity, and resistance to stress.\textsuperscript{165} A growing body of evidence has implicated BDNF-dependent processes in the pathophysiology of depressive disorders and the therapeutic action of antidepressant agents.\textsuperscript{166} Developmental and gender-related differences have been documented with respect to BDNF expression.\textsuperscript{167,168} These findings highlight the dynamic changes in neurobiological processes underlying depressive disorders that may be shaped by environmental inputs.

**Neurobiology**

In contrast to the wealth of information on the neurobiology of adult depression, there are relatively few studies in pediatric samples, although this is a burgeoning area of investigation. Most studies of childhood and adolescent depression have followed up the observations and methods used in adult studies, and they focused primarily on electrophysiological, neuroendocrine, and neuroimaging techniques.\textsuperscript{5,169,170} Aside from cross-sectional designs during the acute depressive episode, some studies applied these measures to at-risk youth, or employed longitudinal designs to examine their relation to the clinical course of depression. It is important to note, however, that the sample sizes in many of these studies are modest. Nevertheless, convergent patterns across studies are informative in determining developmental continuities and discontinuities with adult depression.

**Electrophysiological studies**

Baseline electroencephalographic (EEG) studies documented reduced left frontal electrical activity in infant and adolescent offspring of depressed mothers,\textsuperscript{171-173} and in adolescents with major depressive disorder.\textsuperscript{174,175} Decreased left frontal EEG activity presumably reflects an underactivation of the approach system and reduced positive emotional expression, which also may be a vulnerability marker for depression.\textsuperscript{176} In a study of young adults with a history of childhood depression, frontal EEG asymmetry differed between men and women and varied in relation to longitudinal clinical course.\textsuperscript{177} Men showed more decreased alpha power at all sites than women, and women with history of childhood depression had greater right frontal alpha suppression, whereas men with childhood depression had greater left frontal alpha suppression. Participants who developed bipolar disorder had the most extreme patterns of frontal EEG asymmetry. In the same sample, eye-blink responses to affective stimuli also were associated with variations in clinical outcome in adult life.\textsuperscript{177} These electrophysiological measures were acquired in adult life, and, therefore, the observed changes might be “scar” markers of repeated depressive and/or manic episodes rather than premorbid markers.

EEG sleep measures have shown considerable variability with regard to group differences between depressed youngsters and matched controls.\textsuperscript{5,178} Age and gender have a significant influence on these measures,\textsuperscript{179,180} and depressed adolescents seem to have relatively more frequent disturbances in circadian rest-activity rhythms, sleep architecture, and EEG rhythms during sleep compared with depressed children.\textsuperscript{5,180,181} Among adolescents, the EEG sleep measures were remarkably stable when examined both during the acute depressive episode and during sustained remission, suggesting that these measures are trait-like.\textsuperscript{182,183} Changes in sleep architecture and sleep-related EEG rhythms also were documented in healthy adolescents at high risk for depression, and these changes were associated with vulnerability for depression during prospective follow-up.\textsuperscript{184-186} Additionally, baseline EEG sleep patterns differed between depressed adolescents who subsequently had a recurrent unipolar course versus those who developed bipolar illness; adolescents with unipolar course had predominantly rapid eye movement (REM) sleep changes while adolescents with bipolar course had non-REM sleep changes.\textsuperscript{8} In the same study, adolescents who subsequently developed substance use disorders had relatively normal EEG sleep patterns.\textsuperscript{187} Although EEG sleep changes in pediatric depression, particularly the childhood-onset type, show discontinuities with findings in adult depression,\textsuperscript{180,189} it is important
to also emphasize the variability across studies of both children and adolescents. The observed variability in EEG sleep changes in depressed youngsters may reflect, at least in part, heterogeneity in the longitudinal clinical course of these disorders. For example, sleep data in adults suggest distinct biological substrates in unipolar and bipolar mood disorders. REM latency changes were observed less frequently in bipolar depression. Sleep loss can effectively trigger the onset of mania in patients with bipolar illness, but has minimal euphorigenic effect in unipolar depression. Therapeutic sleep deprivation also appears to have different clinical effects in unipolar and bipolar patients. As described above, a substantial minority of youngsters initially identified as having unipolar depression subsequently develop bipolar disorder, and those with early-onset illness in particular. Among children, studies that excluded depressed patients with family history of bipolar disorder were more likely to demonstrate EEG sleep changes compared with controls.

Neuroendocrine studies

There has been considerable interest in the HPA system, consistent with the possibility that depression is linked to altered responses to stress, and numerous studies have documented HPA dysregulation in adult depression. HPA findings in depressed children and adolescents were inconsistent. In particular, depressed children did not display changes in 24-hour cortisol patterns. Few differences in basal cortisol secretion have been observed between depressed adolescents and controls, and when group differences were detected, they tend to be subtle alterations in normal diurnal patterns. These subtle changes, however, were relatively robust in predicting the longitudinal clinical course; higher cortisol secretion in the evening or during sleep, a time when the HPA axis is relatively quiescent, was associated with a longer time to recovery from the depressive episode, a propensity for recurrence, and suicide attempts. Higher cortisol secretion also was detected in at-risk youth who subsequently developed depression. Another neuroendocrine marker possibly related to depression is growth hormone, which is secreted by the anterior pituitary and follows a circadian pattern with increased secretion during slow-wave sleep. Although the precise role of growth hormone secretion in depression is not known, it appears to be a marker of central noradrenergic and serotonergic (5-HT) systems. Reduced growth hormone secretion during sleep has been observed in adult depression, but findings in children and adolescents have been variable, with some studies showing no differences whereas others showing reduced or increased secretion. One study found that depressed children with stressful life events had increased growth hormone secretion compared with their counterparts who did not experience recent stress, suggesting that environmental factors have a moderating influence and also underscoring the need for integrative models in examining the pathophysiology of pediatric depression. In another study, depressed adolescents who subsequently made suicide attempts had increased growth hormone secretion during sleep, and when this group was separated, depressed adolescents manifested blunted growth hormone secretion compared with controls, again highlighting the value of neuroendocrine measures in predicting the longitudinal course in depressed youngsters. In contrast to the findings in basal secretion, pharmacological challenge studies documented blunted growth hormone response to a variety of pharmacological agents in depressed children, similar to those reported in depressed adults. In contrast, data in adolescents were predominantly negative. Although the sample sizes were modest in these adolescent studies, pubertal changes and gender might account for some variability among child, adolescent, and adult samples.

Neuroimaging studies

Studies using various neuroimaging techniques provided converging lines of evidence supporting prefrontal cortical-striatal and medial temporolimbic dysfunction in adult depression. There is a striking paucity of neuroimaging studies in pediatric depression, and existing studies are marked by small sample sizes and inconsistent findings. Within this context, volumetric studies documented reduced left frontal lobe volume, particularly in those with familial depression. Alterations in amygdala and hippocampal volumes also were found, although the effects appear to be moderated by anxiety and manic symptoms. In neurochemical studies, reduced glutamate and creatinine/phosphocreatinine concentrations in the anterior cingulate, and increased choline concentrations in the left dorsolateral prefrontal cortex, were documented in pediatric depression.
Summary

Neurobiological research in pediatric depression suggests that neurobiological factors change during the course of development, and developmentally influenced neurobiological processes may become disrupted during depressive episodes. Longitudinal studies that account for familial and clinical variability allude to this possibility, whereas cross-sectional studies that fail to account for developmental changes, gender differences, and family history produced inconsistent findings. These data also indicate that early-onset depressive disorders may not necessarily result from the same etiological processes, and the specific subtype with a recurrent unipolar course is associated with neurobiological changes typically observed in adult unipolar depression.

Temperament and personality

Temperament is thought to have a genetic/biological basis, although experience and learning, particularly within the social context, also can influence its development and expression. The trait that is associated with most emotional disorders has been given various labels by different theorists, including behavioral inhibition, harm avoidance, negative affectivity, neurotism, and trait anxiety, although the conceptual and empirical overlap among these constructs far outweighs the differences. Negative affectivity is the propensity to experience negative emotions, and it reflects sensitivity to negative stimuli, increased wariness, vigilance, physiological arousal, and emotional distress. In contrast, positive affectivity is characterized by sensitivity to reward cues, sociability, and adventurousness. Depression is characterized by high levels of negative affectivity and low levels of positive affectivity, and these features have also been found in depressed children. Elevated levels of behavioral inhibition have been observed in laboratory tasks with young offspring of depressed parents. Longitudinal studies have shown that children with inhibited, socially reticent, and easily upset temperament at age 3 had elevated rates of depressive disorders at age 21 than those who did not demonstrate these characteristics. Similarly, physicians’ ratings of behavioral apathy (ie, lack of alertness) at ages 6, 7, and 11 predicted adolescent mood disorders and chronic depression in middle adulthood. Difficult temperament, characterized by inflexibility, low positive mood, withdrawal, and poor concentration correlated with depressive symptoms both concurrently and prospectively in adolescents. The relation between temperament and depression may vary somewhat by age. In one study, neurotic-like symptoms predicted the first episode of depression in 31- to 41-year-old individuals, but this was not the case for 17- to 30-year-olds. Similarly, adult participants who experienced a first episode of depression had exhibited elevated levels of dependent traits 2 to 3 years earlier. However, no differences were found with regard to dependent traits between adolescents who later developed depression and those who did not develop the disorder. Gender might also moderate the relationship between temperament and depression; while females with higher levels of chronic depression during young adulthood had been described as shy and withdrawn at 3 to 4 years of age, males with chronic depression exhibited higher levels of under-controlled behavior as young children.

Cognitive vulnerability

Cognitive theories of depression assert that, when confronted with stressful experiences, individuals who have negative beliefs about the self, world, and future, and those who make global, stable, and internal attributions for negative events will appraise stressors and their consequences negatively, and therefore are more likely to become depressed than those who do not have such cognitive styles. Several types of cognitions have been proposed to be related to depression, including low self-esteem, negative automatic thoughts, dysfunctional attitudes, and cognitive distortions; self-control; control-related beliefs and self-efficacy; negative attributional style; and a ruminative response style. Cross-sectional studies with clinic and community samples of children and adolescents have consistently shown a strong correlation between a range of negative cognitions and depression. In prospective studies, negative cognitions predicted depression, often in interaction with negative life experiences. Developmental theorists have suggested that negative cognitions emerge over time, and that their relationship with depression becomes stronger with development. Indeed, the association between negative cognitions and depression is less robust in younger children than in older children and adolescents. If negative cognitions contribute to the development of depression, then offspring of depressed individuals should be more...
likely to exhibit cognitive vulnerability than children whose parents have not experienced depression. Indeed, children of depressed mothers reported lower perceived self-worth and greater negative attributional style than children of nondepressed mothers. Even though there is a concurrent and predictive relationship between negative cognitions and depression in youngsters, some have questioned whether negative cognitions are a concomitant or consequence of depression rather than part of a longitudinal chain. Future studies should examine the development of cognitive vulnerability over time, and whether it needs to be primed in children.

**Interpersonal relationships**

Interpersonal theories of depression emphasize the importance of social environment and the development of secure attachment. Vulnerability to depression presumably arises in early family environments in which the children’s needs for security, comfort, and acceptance are not met. Literature on the relationship between family environment and depression indicates that families of depressed individuals are characterized by problems with attachment, communication, conflict, cohesion, and support, as well as poor child-rearing practices. Additionally, perceived rejection by peers, family, and teachers predicts increases in depressive symptoms in children and adolescents. Interpersonal theories of depression propose that depressed individuals both react and contribute to interpersonal problems. Depressive symptoms and associated behaviors are presumed to elicit negative reactions from others; these aversive interpersonal experiences then foster the persistence or exacerbation of depression. Consistent with interpersonal models, depressed youngsters demonstrate difficulties in many aspects of relationships with peers and family members.

**Stress and coping**

**Stress**

Common to all definitions of stress is a focus on environmental conditions that threaten to harm the biological or psychological well-being. Stress may occur either as an acute event or as chronic adversity; and as a major life event or as minor accumulated events. Stressful events may be normative (eg, school transitions) or pathological (eg, abuse), and may be independent of, or dependent on an individual’s actions. Stress plays a prominent role in most theories of depression, and a clear empirical link exists between stress and depression in children and adolescents. Although no single or specific type of stressful event leads to depression, certain types of negative events consistently have been found to be associated with depression: child abuse/neglect, especially for women; socioeconomic disadvantage; personal disappointments, failures, and losses; and interpersonal problems. Early adversity may be a marker of continuing exposure to negative stressors, such that those with exposure to negative events and circumstances in childhood are more likely to continue to be exposed to stressful situations.

The relationship between stress and depression appears to be stronger in adolescents than in children, particularly in girls. The reasons for this are not entirely clear; hormonal effects, consolidation of cognitive styles, cumulative stress burden, and stress reactivity might have a potential role. One theory proposes that childhood adversity alters neurobiological and psychosocial processes, whereby individuals may be sensitized to the effects of recent stressful events, leading to depression at lower levels of stress, or with greater reactivity to the effects of stress. Another approach suggests that childhood stressors add to lifetime stress burden and independently predict depression along with recent stress.

Developmental models of psychopathology also suggest a transactional perspective in which stress exposure contributes to depressive symptoms and, in turn, depressed individuals contribute to negative events through their own behavior. Longitudinal studies have shown support for the stress-generation model, particularly with regard to interpersonal relationships. Factors that might contribute to the generation of stress include personality, lack of interpersonal competence, and comorbid psychopathology. The reciprocal model highlights the “vicious cycle” that can occur between stress and depression, and support for this reciprocal model has been found in a few studies of youngsters.

**Response to stress**

Although stress clearly plays a role in depression, individuals vary in their response to stress, and how they
respond to stress can affect their future adjustment and emotional well-being. Diathesis-stress models propose that depression results from the interaction between personal vulnerability and stressful events or circumstances. The majority of research testing diathesis-stress models of depression has construed vulnerability in terms of maladaptive appraisals of events. Several studies documented interactions between cognitive styles, such as negative attributional style and low perceived self-efficacy, and life stress in the prediction of depression in youngsters.278-280 Even further refining these theories, it has been speculated that a key determinant of depression may be the match between a particular cognitive vulnerability (eg, a tendency to base one’s self-worth on success in interpersonal relationships) and the nature of the stress (eg, interpersonal conflict). Supporting this theory, diathesis-stress interactions seem to be most powerful when there is a match between the type of cognitive vulnerability and the type of stressful experience.234 Consistent with the theory that cognitive styles may not yet be consolidated in younger children, cognitive-stress interactions predicted depression in adolescents but not in children.234,281

In addition to cognitive styles, other types of coping mechanisms, such as behavioral styles and problem-solving skills, have been examined in relation to pediatric depression.282,283,284 Earlier theories differentiated between problem-focused and emotion-focused coping. Problem-focused coping involves responses that act directly on the source of stress, whereas emotion-focused coping involves palliative measures to counter the negative emotions that arise from stressful situations. Recent models of coping proposed responses to stress that can be distinguished as voluntary or involuntary, and engaged or disengaged.282 Coping involves volitional and intentional responses to stress. Involuntary or automatic reactions to stress are, in part, a reflection of individual differences in temperament. Engaged coping includes problem-solving, cognitive restructuring, positive reappraisal, and distraction. In contrast, disengagement responses include avoidance, self-blame, emotional reaction, and rumination. Studies in children and adolescents indicated that higher levels of engaged coping and problem-focused coping are associated with lower levels of depressive symptoms. In contrast, disengagement, involuntary and emotion-focused coping are related to higher levels of depressive symptoms under stressful circumstances.230,282,284,285 Most of the research on coping has been cross-sectional, thereby limiting our ability to draw conclusions about the direction of the relationship between coping and depression.

**Summary and future directions**

In the past three decades, considerable advances have been made regarding our knowledge of the phenomenology and natural course of depression in children and adolescents. Basic epidemiologic and clinical research has also helped identify a number of risk factors associated with pediatric depression. There appears to be a complex interplay among genetic, neurobiological, cognitive, interpersonal, and environmental factors in concert with developmental challenges in the onset and maintenance of depression. Recent studies have emphasized the importance of gene-environment interactions in the genesis of depression. Time is another crucial factor, both in terms of windows of vulnerability when brain regions might be maximally sensitive to environmental influences and in the cascade of maturational events that lead to the unfolding of depression. Other factors, such as temperament/personality traits, cognitive styles and coping repertoires, moderate responses to stressful situations and precipitate depressive episodes.

Depression is likely to further compromise development by interfering with the achievement of key developmental tasks (eg, academic achievement, negotiating changes in family relationships, and establishing peer networks), resulting in the generation of additional stress, and perhaps even contributing to compromised neurobiological development and sensitization to future stress, depression, and other psychopathology. These dynamic processes may account, in part, for why early-onset depression tends to be recurrent throughout the life span and is also accompanied by other psychiatric problems and significant disability. The challenge for the field is to integrate the disparate findings across domains and to develop testable hypotheses with respect to clinical presentation, biopsychosocial processes, and clinical interventions. Effective interventions early in the course of the disorder will likely interrupt the “vicious cycle” and allow these youngsters to reach their full potential as adults. ❑

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Características, correlatos y evoluciones de los trastornos depresivos en niños y adolescentes

La enfermedad depresiva que aparece precozmente en la vida puede tener serias consecuencias funcionales y durante el desarrollo. Por lo tanto, la comprensión del trastorno durante esta etapa del desarrollo es crítica para determinar su etiología y su curso, como también para la generación de estrategias de intervención efectivas. Este artículo resume el conocimiento actual relacionado con la etiología, la fenomenología, los correlatos, el curso natural y las consecuencias de la depresión unipolar en niños y adolescentes. Utilizando la depresión del adulto como referente, los aspectos distintivos de la niñez y de la adolescencia son tomados en cuenta para una mejor comprensión de la depresión dentro del contexto del desarrollo. La información disponible sugiere que el cuadro clínico, los correlatos y el curso natural de la depresión son notablemente similares a lo largo de la vida. Sin embargo, hay importantes diferencias durante el desarrollo. Específicamente, el contexto familiar y psicológico en que ocurre la depresión en los muchachos está asociado con la variabilidad en la frecuencia y en la naturaleza de los síntomas depresivos y en las condiciones comórbidas entre los niños y adolescentes. También se han identificado diferencias madurativas en los correlatos neurobiológicos de la depresión. Estas diferencias durante el desarrollo pueden estar asociadas con la variabilidad observada en la respuesta clínica al tratamiento y en el curso longitudinal. La caracterización de las diferencias durante el desarrollo será útil para generar intervenciones más específicas y efectivas para los muchachos, lo que les permitirá alcanzar su potencial total cuando adultos.

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