

American Academy of Child and Adolescent Psychiatry

AACAP is pleased to offer Practice Parameters as soon as they are approved by the AACAP Council, but prior to their publication in the *Journal of the American Academy of Child and Adolescent Psychiatry (JAACAP)*. This article may be revised during the *JAACAP* copyediting, author query, and proof reading processes. Any final changes in the document will be made at the time of print publication and will be reflected in the final electronic version of the Practice Parameter. AACAP and *JAACAP*, and its respective employees, are not responsible or liable for the use of any such inaccurate or misleading data, opinion, or information contained in this iteration of this Practice Parameter.

PRACTICE PARAMETER FOR THE ASSESSMENT AND TREATMENT OF CHILDREN AND ADOLESCENTS WITH EATING DISORDERS

ABSTRACT

This Practice Parameter reviews evidence-based practices for the evaluation and treatment of eating disorders in children and adolescents. Where empirical support is limited, clinical consensus opinion is utilized to supplement systematic data review. The Parameter focuses on the phenomenology of eating disorders, comorbidity of eating disorders with other psychiatric and medical disorders, and treatment in children and adolescents. Since the database related to eating disorders in younger patients is limited, relevant literature drawn from adult studies is included in the discussion. **Key Words:** eating disorders, anorexia nervosa, bulimia nervosa, food avoidance, binge eating, treatment.

DEVELOPMENT AND ATTRIBUTION

This Parameter was developed by James Lock, MD, PhD, Maria C. La Via, MD, and the American Academy of Child and Adolescent Psychiatry (AACAP) Committee on Quality Issues (CQI): Heather J. Walter, MD, MPH and Oscar G. Bukstein, MD, MPH, co-chairs; and Christopher Bellonci, MD, R. Scott Benson, MD, Regina Bussing, MD, Allan Chrisman, MD, Tiffany R. Farchione, MD, John Hamilton, MD, Munya Hayek, MD, Helene Keable, MD, Joan Kinlan, MD, Nicole Quiterio, MD, Carol Rockhill, MD, Ulrich Schoettle, MD, Matthew Siegel, MD, and Sandra Stock, MD.

The AACAP Practice Parameters are developed by the AACAP CQI in accordance with American Medical Association (AMA) policy. Parameter development is an iterative process between the primary author(s), the CQI, topic experts, and representatives from multiple constituent groups, including the AACAP membership, relevant AACAP committees, the AACAP Assembly of Regional Organizations, and the AACAP Council. Details of the Parameter development process can be accessed on the

AACAP web site. Responsibility for Parameter content and review rests with the author(s), the CQI, the CQI Consensus Group, and the AACAP Council.

The AACAP develops both patient-oriented and clinician-oriented Practice Parameters. Patient-oriented Parameters provide recommendations to guide clinicians toward best assessment and treatment practices. Recommendations are based on the critical appraisal of empirical evidence, when available, and clinical consensus, when evidence is unavailable, and are graded according to the strength of the empirical and clinical support. Clinician-oriented Parameters provide clinicians with the information, stated as principles, needed to develop practice-based skills. Although empirical evidence may be available to support certain principles, principles are primarily based on clinical consensus. This Parameter is a patient-oriented Parameter.

The primary intended audience for the AACAP Practice Parameters is child and adolescent psychiatrists; however, the information contained therein may also be useful for other medical or mental health clinicians.

The authors wish to acknowledge the following experts for their contributions to this Parameter: Deborah Katzman, MD; Guido Frank, MD; Daniel Le Grange, PhD; Jennifer Hagman, MD; Jennifer Couturier, MD; and Wendy Spettigue, MD.

Jennifer Medicus served as the AACAP staff liaison for the CQI.

This Practice Parameter was reviewed at the Member Forum at the AACAP Annual Meeting in October 2012.

From November 2013 to February 2014, this Parameter was reviewed by a Consensus Group convened by the CQI. Consensus Group members and their constituent groups were as follows: Heather Walter, MD, MPH, co-chair; Christopher Bellonci, MD, Regina Bussing, MD, and R. Scott Benson, MD (CQI); Jennifer Couturier, MD and Wendy Spettigue, MD (topic experts); Adelaide Robb, MD (AACAP Committee on Research); Timothy Brewerton, MD and Michael Enenbach, MD (AACAP Assembly of Regional Organizations); and Jennifer S. Saul, MD and Laurence L. Greenhill, MD (AACAP Council).

This Practice Parameter was approved by the AACAP Council on July 1, 2014.

This Practice Parameter is available on the Internet (www.aacap.org).

Disclosures: James Lock, MD, PhD serves or has served on the Advisory Board for the Center for Discovery, the Global Foundation for Eating Disorders, and the National Eating

Disorders Association; serves or has served as a consultant for the Training Institute for Eating Disorders in Children and Adolescents; receives or has received grant support from the Davis Foundation, the Global Foundation for Eating Disorders, and the National Institutes of Health; and receives or has received royalties from Guilford Press and Oxford University Press. Maria La Via receives or has received grant support from the National Institute of Mental Health. Oscar Bukstein, MD, MPH, co-chair, receives royalties from Routledge Press. Heather Walter, MD, MPH., co-chair, has no financial relationships to disclose. Disclosures of potential conflicts of interest for all other individuals named above are provided on the AACAP web site on the Practice Parameters page.

Correspondence to the AACAP Communications Department, 3615 Wisconsin Ave, NW, Washington, DC, 20016.

© 2014 by the American Academy of Child and Adolescent Psychiatry.

INTRODUCTION

This Practice Parameter provides an evidence-based approach to the evaluation and treatment of eating disorders in children and adolescents, including specifically Anorexia Nervosa (AN), Bulimia Nervosa (BN), Binge Eating Disorder (BED), and Avoidant Restrictive Food Intake Disorder (ARFID). The parameter will not address feeding problems in infancy (e.g., failure to thrive), Pica, Rumination Disorder, Purging Disorder, or the evaluation and treatment of obesity. Evaluation and treatment of eating disorders in children and adolescents is complex and often requires specific expertise and relevant clinical experience. This Practice Parameter is designed to help child psychiatrists to accurately assess and effectively treat children and adolescents with eating disorders. This parameter may also provide useful information for other medical and mental health professionals because the treatment of eating disorders commonly requires consultation and involvement with other experts in addition to child psychiatrists.

METHODOLOGY

The recommendations in this practice parameter were developed after searching literature including PubMed/Medline and employing the relevant medical subject headings (MeSH terms) “eating disorders”, adding limits “child: 6 – 12 years” and “adolescent: 13-18 years”, “clinical

trial”, and a time period from 1985 to 2011 inclusive (yields 497 citations); Cochrane, employing the relevant medical subject headings (MeSH terms) “eating disorders”, adding limits “clinical trials” (yields 646 citations); and PsycINFO, employing the term “eating disorders”, adding limits “child: 6 – 12 years” and “adolescent: 13-17 years”, “clinical trial”, and a time period from 1985 to 2011 inclusive (yields 112 citations). In addition, the bibliographies of book chapters and treatment guideline articles were reviewed; and lastly, colleagues were asked for suggested source materials.

The online search was narrowed on PubMed/Medline using delimiters and filters such as English language only, human subjects, and using the Boolean operator ‘AND’, ‘OR’, and ‘NOT’ to include the following search terms: family therapy, comorbid, treatment outcome, psychopharmacology, and eating disorder not otherwise specified to reduce citations to 141. Similarly, the online search was narrowed on Cochrane by searching clinical trials, and using the Boolean operator ‘AND’ and ‘OR’ to include the following search terms: anorexia, bulimia, child, adolescent, and family based therapy to reduce citations to 17. Finally, the online search was narrowed on PsycINFO, by using the Boolean operator ‘AND’ and ‘OR’ to include the following search terms: anorexia and bulimia. The subject of this search was further specified by including treatment outcomes, family therapy and clinical trials to reduce citations to 69 results.

For this practice parameter, we hand culled 91 publications for examination based on their relevance to clinical practice. In addition, 19 more recent references for 2012-13 were identified by expert and member reviews.

HISTORICAL REVIEW

The first comprehensive description of a condition resembling AN was provided by Richard Morton in 1689, which he called *nervous consumption*.¹ While there is evidence in ancient history, usually in the context of religious beliefs, of clinical problems similar to AN,² it was not until 1874 when Sir William Gull in England and Charles Lasegue in France coined the terms *Anorexia Nervosa* and *Anorexia Hysterique*, respectively, to describe the symptoms of self-starvation and weight preoccupations associated with AN³. It was not until this time that theories related to etiology and treatment began to evolve. Both Gull and Lasegue suggested that families likely contributed to the disorder, but Jean-Martin Charcot directly blamed families and advocated complete separation of affected individuals from their families whose influence he

viewed as “pernicious.”⁴ Thus, treatment for AN from the late nineteenth century to the present has often included hospitalization and long separations from family members, where patients are treated exclusively by professionals.^{5,6} Psychoanalytic approaches suggested that affected individuals suffered from a range of unconscious problems including pregnancy fears, primary narcissism, and confusion between body and emotion.⁷ Hilde Bruch conceptualized AN as a disorder of suppression and neglect in childhood, leading to food refusal and the formation of AN symptoms for self-assertion. She advocated for individually-oriented psychodynamic therapy for the patient to promote autonomy and independence from parents and families.^{8,9} Patients were often treated in psychiatric hospitals and in the 1980’s until the 2000’s, inpatient hospital-based and specialized residential treatment programs became more prevalent for patients with AN.¹⁰ Salvador Minuchin’s pioneering work in structural family therapy with psychosomatic disorders suggested that families could be important in treatment, despite the prior practices.¹¹ Subsequently, researchers at the Institute of Psychiatry and Maudsley Hospital in London developed a form of family therapy that was specifically designed to utilize parental skills to disrupt the maintaining behaviors of AN.¹² Over the past 10 years a substantial database supports including families in the treatment of adolescents with AN.¹³

BN was first included in DSM-III in 1980 following clinical descriptions of patients with binge eating and purging by Boskand-Lodahl (bulimarexia)^{14,15} and Russell who called it “an ominous variant” of AN.¹⁶ Although, few studies have investigated treatment for BN in adolescents, many intervention studies of adults with BN have demonstrated the effectiveness of cognitive behavioral therapy (CBT) for this disorder.¹⁷ In addition, antidepressant medications and interpersonal psychotherapy (IPT) are effective in adults with BN.¹⁸ The diagnosis of Binge Eating Disorder (BED) is now included in DSM-5.¹⁹ BED is understudied in children and adolescents, but appears to be rarer in younger patients than in adults.²⁰ Effective treatments for adults with BED include CBT, IPT, and medications.^{21,22} Pilot studies support the use of IPT for adolescents with BED.²³ Another new diagnosis in DSM-5 is ARFID.¹⁹ This is a disorder found principally in children. In ARFID, food or eating is avoided usually leading to low weight, but is not associated with shape or weight concerns, or intentional efforts to reduce weight.²⁴

ANOREXIA NERVOSA (AN)

Clinical Presentation and Course

DSM-5 criteria for AN include: restriction of energy intake leading to low body weight; fear of gaining weight or behavior that interferes with weight gain; and self-evaluation unduly influenced by weight and body shape.²⁵ Denial of the seriousness of malnutrition is also a common symptom, especially in younger patients.²⁶ There are two subtypes of AN, a restricting type and a binge-eating/purging type. In DSM-5, amenorrhea is no longer required to meet diagnostic criteria.²⁷ DSM-5 suggests that clinicians rate the level of severity of AN (mild to extreme) in adults based on current BMI; in children and adolescents, severity is based on age and gender norms according to BMI percentiles.^{26,28,29} A BMI below the 10th percentile is considered to be consistent with the degree of malnutrition associated with AN.^{30,31} Alternatively, if longitudinal growth charts are available, deviations from individual growth trajectories can be observed.³² At present, there is little scientific basis for using weight as a marker of severity in children and adolescents so it remains to be seen how useful this approach will be in defining severity.

There is evidence that AN symptoms may be expressed differently in childhood and adolescence, as compared to adulthood.³³ Children and adolescents are often incapable of verbalizing abstract thoughts; therefore, behaviors such as food refusal that leads to malnutrition may manifest as non-verbal representations of emotional experiences. As a result, parental reports about the child's behavior are critical, as self-report is often unreliable due to a lack of insight, minimization, and denial by the child or adolescent.³⁴ Children and adolescents with AN are less likely than adults to engage in binge eating and purging behaviors.³⁵

Children and adolescents with AN most often present for psychiatric evaluation after a pediatrician or other medical provider suspects an eating disorder based on the patient's unexpected weight loss or failure to gain expected weight. Patients with AN often develop weight concerns and subsequent behavioral change directed toward weight loss 6 -12 months prior to the full clinical diagnosis. The rate of weight loss typically escalates in the last few weeks prior to referral, prompting parents to seek a medical evaluation.³⁶ Patients sometimes report initial drive for thinness, but often claim that they are trying to eat less, avoid fattening foods, and exercise more for health reasons. Other young patients deny body image or weight concerns at assessment and insist they just "aren't hungry," or complain of abdominal discomfort. It is important, however, not to infer fear of weight gain or weight and shape concerns on the basis of developmental immaturity alone; instead, the presence of behaviors

indicating avoidance of “fattening” foods or indicating fear of weight gain (such as repeated weighing, pinching skin) may confirm the diagnosis. Typically, caloric reduction increases over time as food choices become more limited (usually elimination of protein, fats, and sweets). As these dieting behaviors persist, the patient focuses more exclusively on weight and dieting to the exclusion of friends and family. Academic and athletic pursuits usually continue and sometimes become more compulsive and driven. Patients may have a compulsion to stand or move, and may be exercising secretly, or may no longer have the energy to over-exercise. They may dress in baggy clothing or layers, and complain of feeling cold. Some patients may drink water excessively, while others may restrict fluid intake. Patients often appear withdrawn, depressed, and anxious. They, usually, remain cognitively intact until more severe malnutrition develops. In some instances, compensatory behavior such as purging develops, but this is usually later in the course of the disorder in younger patients.³⁵

Long-term adult follow-up studies of AN suggest rates of chronicity, defined as having AN greater than five years, of 7-15% and mortality of 5-7%, although mortality as high as 18% has been reported in some samples.³⁷ Death is most often secondary to medical complications of starvation (50%) or suicide (50%).³⁸ The prognosis for adolescents with AN is better than in adult populations.^{39,40}

Epidemiology

The prevalence of AN in the United States is reported to be 1-2% among females, while the prevalence in teenage girls has been reported to be between 0.3- 0.7%.^{41,42} One study examining the rates of AN between 1939 and 1984 found evidence that the incidence of AN increased in females age 10-24.^{43,44} Little is known about the prevalence of AN in males, although a ratio of one male case to every ten females is commonly cited,⁴⁵ while some estimates are as high as 1:1.^{46,47} Studies of males with eating disorders are limited, though some studies suggest that incidence or detection rates of AN in males are increasing.^{46,48,49} The prevalence of sub-threshold AN is estimated to be 1.5% in adolescent females and 0.1% in adolescent males.^{42,50} There are also only sparse data related to the racial and ethnic distribution of AN, but recent studies suggest that the disorder may be less common in persons of African origin.⁵¹ Further, only three studies have assessed the incidence rates of AN in children 14 years old or younger.^{43,52} The first study reported incidence rates of 9.2-25.7 females per 100,000 per year in

10-14 year olds and 11.9-69.4 females per 100,000 per year in 15-19 year olds. The second study reported incidence rates of 25.7 females and 3.7 males per 100,000 of the population per year in 10-14 year olds and 69.4 females and 7.3 males per 100,000 of the population per year in the 15-19 year old range.⁴³ Peak incidence occurs at 14-18 years of age,⁵³ with rare cases presenting after 25 years of age.⁴³ The third study found an overall incidence of eating disorders of 3:100,000 between 5-13 years of age. Incidence rates were found to rise with age to 4.5:100,000 by age 11 and to 9.5:100,000 by age 12. Of these, 80% had AN or a restrictive eating problem similar to AN, but not meeting full diagnostic criteria.⁵⁴

Etiology and Risk Factors

The etiology of AN is likely multifactorial and precipitated by the interaction of several risk factors including biological, psychological, environmental/cultural, and sociological. Family aggregation studies demonstrate that AN occurs at about five times the expected rates in affected families. Twin studies find heritability estimates ranging from approximately 30% to 75% in AN.⁵⁵ Recent twin studies show an interaction between genes and developmental processes in children and adolescents who develop AN. One study⁵⁶ found that for 11-year old twins, genetic influences were marginal, but in 17-year old twins, heritability was high.⁵⁷ One possible explanation for this finding is that hormonal changes during adolescence could mediate gene expression during puberty.^{58,59}

Temperament and personality type are also risk factors associated with AN,^{55,60} and perfectionistic, obsessive, and avoidant personality features are likely heritable.^{61,62} Studies also suggest that specific cognitive features are associated with AN, including cognitive rigidity and a bias toward detail information processing^{63,64} Some have characterized these cognitive features as an endophenotype because they are present in patients after recovery and in unaffected siblings.⁶⁵⁻⁶⁷ Picky eating early in life has also been associated with later development of AN.⁶⁸ Developmental challenges associated with adolescence such as autonomy, self-efficacy, and intimacy are found in patients with AN, though it is uncertain whether this is a cause or a result of the disorder.^{8,69} Psychosocial factors associated with the development of AN include societal pressures related to thinness and appearance that can trigger extreme dieting in vulnerable individuals.⁷⁰⁻⁷² In addition, some studies suggest that certain activities such as ballet, gymnastics, wrestling, and modeling may increase risk for eating disorders because of the role of

appearance and/or weight in performance.^{73,74} The westernization of culture, with an emphasis on the “thin ideal,” is also associated with AN in some studies, particularly in Asian culture and immigrants to Europe and the Americas.⁷⁵⁻⁷⁷

Differential Diagnosis and Comorbidity

There are many possible causes of weight loss, loss of appetite, and refusal to eat. Differential diagnosis for AN includes ARFID and Rumination Disorder when these disorders have resulted in low weight. Any medical or psychiatric illness that leads to changes in appetite, weight loss, or changes in food intake likely increases the risk for the development of an eating disorder. In addition, chronic infection, thyroid disease, Addison’s disease, inflammatory bowel disease, connective tissue disorders, cystic fibrosis, peptic ulcer disease, disease of the esophagus, celiac disease, infectious diseases, disease of the small intestine, diarrhea, diabetes mellitus, and occult malignancies can lead to weight loss, appetite loss, and refusal to eat.⁷⁸ Many of these can be ruled out with a thorough history and physical exam along with laboratory studies.

Further complicating the diagnosis of AN is the potential presence of other significant psychiatric comorbid conditions. In adolescents with AN, results from the National Comorbidity Survey Replication Adolescent Supplement (NCS-A) found that the lifetime rate of comorbidity with at least one other psychiatric disorder is 55.2%.⁷⁹ Psychiatric comorbidity in adults with AN includes depression, social anxiety, separation anxiety, obsessive compulsive disorder (OCD), generalized anxiety and substance abuse; while avoidant, dependent, obsessive-compulsive or passive-aggressive personality disorders are also often diagnosed.⁸⁰⁻⁸² AN and OCD share obsessional preoccupations, such as over eating, food, weight, and shape obsessions, and compulsive behaviors, such as restricting and counting calories, over exercising, checking behaviors, meal time rituals, making it difficult to differentiate these two disorders. When the preoccupations and compulsions emerge at the same time and are focused on food, eating, and weight, the diagnosis, however, is likely to be AN. Additionally, comorbid conditions, particularly anxiety disorders, are often present before AN develops and persists after recovery.⁸³⁻⁸⁵ Specific phobias, particularly those such as fear of swallowing, can lead to weight loss and be confused with AN. Similarly, dysphoric thoughts, low self-esteem, and guilt are found in both depression and AN. With depression these features encompass common aspects of

life, while in AN these thoughts are focused more specifically on eating and weight-related concerns.⁸⁶

BULIMIA NERVOSA (BN)

Clinical Presentation and Course

DSM-5 criteria for BN include: recurrent binge-eating where a binge is defined as consuming a very large amount of food in a discrete period of time, such as within two hours, and a sense of loss of control over eating during that episode; recurrent compensatory behavior, such as vomiting, fasting, exercise, laxative use, diuretic use, taking diet pills; and self-evaluation unduly influenced by weight and body shape. The binge-eating and compensatory behaviors both occur, on average, at least once a week for three months, and do not occur exclusively during AN.^{19,25} The severity of BN, mild to extreme, is based on the frequency of compensatory behaviors. Compensatory behaviors distinguish BN from BED. Patients with BN are often within a normal or high normal weight range for age, gender, and height. Secrecy, shame and guilt tend to accompany BN in adolescents, and in turn, these emotions may negatively impact the age-appropriate development of adolescents by interfering with social and interpersonal processes.^{87,88} Males with BN appear to be more likely than females to present with over exercise and steroid use.⁸⁹ Risk for BN is increased in males who participate in sports such as wrestling, gymnastics, diving, and long distance running where weight and appearance can affect performance.⁴⁹ There is also a limited database suggesting that homosexual males may be more at risk than their heterosexual peers for the development of BN.^{90,91}

There are challenges in diagnosing BN in childhood and adolescence because of developmental differences between younger patients and adults.³³ Data suggests that loss of control may be a more valid marker than calories consumed in terms of determining if an eating episode should be characterized as binge eating.⁹² Binge-eating may also occur less frequently in younger patients because children do not have access to, or control over, foods in the same way that adults with BN do.^{33,35} Similar limitations on abstract thinking, self-expression, and minimization as described for adolescent AN are relevant for adolescent BN as well; thus, parental interviews and other collateral reports may be essential to obtain an accurate history.³³

The short-term course of BN is marked by fluctuating symptoms with varying cycles of remission and exacerbations. Typically individuals with BN have exhibited symptoms for nearly

five years before seeking treatment.⁹³ Outcome in BN is highly variable. 50% of those with BN who receive treatment are symptom free 5-10 years later, while the other 50% continue to exhibit eating disorder symptoms. One long-term follow-up study suggests, however, that the number of women who continue to meet full criteria for BN declines as the duration of follow-up increases.⁹⁴ A history of low self-esteem, childhood obesity, and personality disorders predict a longer duration of illness.⁹⁵ Self-injury, substance abuse, and other impulsive and risk-taking behaviors are common in adults and older teens seeking treatment, but appear to be less common in younger adolescents with BN.⁹⁶

Epidemiology

The incidence of BN is increasing in urbanized areas and countries undergoing rapid Westernization.^{41,97} Between 1-2% of adolescent females and 0.5% of adolescent males are estimated to meet DSM IV criteria for BN.⁹⁸⁻¹⁰⁰ The estimated male to female ratio is 1:10 though some estimates are as high as 1:3.^{47,90} BN typically begins in adolescence between 14-22 years of age^{42,101} and for some arises after an episode of AN. BN is rarely diagnosed in children and young adolescents, although older patients presenting for treatment often pinpoint the onset of their illness to early adolescence.³⁵

Etiology and Risk Factors

The etiology of BN is multifactorial, including biological, psychological and sociological factors. BN occurs more often in first degree relatives of those with eating disorders than in the general population.¹⁰² Twin studies show a higher concordance rate in monozygotic twins than in dizygotic twins with heritability threshold estimates ranging from 60% to 83%.¹⁰³⁻¹⁰⁵ Family factors, abuse, posttraumatic stress disorder (PTSD), impulsive personality traits, and perfectionistic temperament have also been suggested to be risk factors for BN.^{95,102,106-111} In addition, social pressures to be thin, body dissatisfaction, dieting, and negative affect are associated with the development of BN.^{73,74,77} Dieting to try to accomplish a thin ideal physique is thought to lead to physical and psychological starvation.⁹³ Physiological starvation then leads to hunger and promotes binge eating when food becomes available. Guilt associated with binge eating and fear of weight gain leads to purging behavior and increased dieting. Thus, a cycle of deprivation, binge eating, and purging is established.

Differential Diagnosis and Comorbidity

Differential diagnosis for BN includes AN, binge/purge type, sub-threshold BN, Purging Disorder, BED, major depressive disorder, central nervous system (CNS) tumors, Kleine-Levin Syndrome, Kluver-Bucy Syndrome, and gastrointestinal pathology, such as obstruction and gastroparesis. Several psychiatric disorders are highly associated with BN, including affective disorders, anxiety disorders, substance use disorders and personality disorders.^{84,93,112} In adolescent BN, a recent study found a lifetime psychiatric comorbidity rate of 88%, with a majority of adolescents having at least one comorbid psychiatric illness.⁷⁹ In addition, significant rates of suicidal ideation (53%), plans (26%), and attempts (35%) were found among adolescents with BN.

BINGE EATING DISORDER (BED)

Clinical Presentation and Course

BED is characterized, as is BN, by recurrent binge eating episodes accompanied by a sense of loss of control over eating during the episode. In BED, these episodes are associated with at least three of the following: eating more rapidly, eating until uncomfortably full, eating when not hungry, eating alone because of embarrassment about the amount of food consumed, and feelings of disgust, depression, or guilt. According to DSM-5, binge eating episodes need to occur, on average, at least once a week for three months, and must be associated with marked distress, to meet diagnostic severity levels. BED is distinguished from BN in part because the binge eating episodes are not associated with inappropriate compensatory behaviors. In addition, to meet criteria for BED, binge eating episodes cannot occur exclusively during the course of AN or BN.

There are similar challenges in diagnosing BED and BN in childhood and adolescence because of developmental differences between younger patients and adults.³³ A sense of being out of control when eating is likely more important than eating an objectively large amount of food in younger patients⁹² because younger patients often cannot gain access to food as easily as adults.³⁵ For these reasons, a clinical concern about BED in children and adolescents should consider using a lower threshold for the rate and duration of binge eating episodes.³³ A suggested rate of once per month, over the previous three month period, was recommended by a consensus

group of experts in child and adolescent eating disorders.³³ In addition, as with AN and BN, children and adolescents are limited in their abstract thinking ability and self-expression. They may also minimize any discomfort or shame they experience when binge eating. Thus, parental interviews and other collateral reports are often necessary for making a definitive diagnosis.³³ BN usually occurs in patients of normal weight or slightly overweight; BED often occurs in overweight and obese individuals. In BN, binge eating is considered to be a response to restriction of food intake,¹¹³ whereas in BED, binge eating occurs in the context of overall chaotic and unregulated eating patterns.¹¹⁴

Epidemiology

BED may be the most common eating disorder, affecting 3.5% of females and 2% of males among adults.⁴⁷ Rates of BED in children and adolescents are estimated to be 2.3% in adolescent females and 0.8% in adolescent males. Prevalence of sub-threshold BED is estimated to be 2.3% in female adolescents and 2.6% in male adolescents.⁴² Some studies suggest that among obese adolescents, about 1% meet criteria for BED, while 9% have objective binge episodes.²⁰ A high percentage of adults and adolescents that present with BED are moderately overweight or obese. Prevalence rates of BED are higher among women enrolled in weight-control programs compared to those who are not.¹¹⁵

Etiology and Risk Factors

Comparatively, little is known about the specific risk factors for BED in children and adolescents, as it is a relatively rare disorder in younger populations. Onset of binge eating typically begins in late adolescence or early adulthood, often after a period of significant dieting or weight loss. Some studies suggest that restrictive dieting, pressure to be thin, body dissatisfaction, emotional eating, low self-esteem, and poor social support are risk factors for the onset of adolescent binge eating.¹¹⁶ Other studies suggest increased eating related to specific psychopathology, depressed mood, and increased anxiety are associated with the initiation of binge eating in adolescents.¹¹⁷

Differential Diagnosis and Comorbidity

Differential diagnosis includes BN, AN, and sub-threshold BED, Night Eating Syndrome, and Nocturnal Sleep Related Eating Disorder. Medical reasons for binge eating should be considered including CNS tumors, Kleine-Levin Syndrome, Kluver-Bucy Syndrome, Prader-Willi Syndrome, and gastrointestinal pathology. In adults, BED is associated with significant comorbid psychopathology, including depressive disorders, anxiety disorders, PTSD, impulse control disorders, substance use disorders, and personality disorders.^{50,118}

AVOIDANT RESTRICTIVE FOOD INTAKE DISORDER (ARFID)

Clinical Presentation

Diagnostic criteria for ARFID^{24,119,120} includes food restriction or avoidance without shape or weight concerns or intentional efforts to lose weight that results in significant weight loss, nutritional deficiencies, and is associated with disturbances in psychological development and functioning. Some patients present with highly selective eating, neophobia (the fear of new things) related to food types, or hypersensitivity to food texture, appearance, and taste.¹²⁰⁻¹²³ For some patients, fear of swallowing or choking contributes to food avoidance; a specific event can sometimes be identified as triggering that fear. ARFID also applies to individuals who have a lack of interest in eating or who have low appetite.

Epidemiology

There are no epidemiological studies available for this new diagnosis.

Etiology and Risk Factors

Specific etiologic risk factors for ARFID are unknown. Patients with autism spectrum disorders (ASD) frequently display selective eating patterns. Anxiety disorders and anxious traits, as well as depressive symptoms, often pre-date the development of ARFID. Neglect, abuse, and developmental delays may increase the risk for chewing and spitting associated with ARFID.

Differential Diagnosis

ARFID can be confused with AN, but distinguishing features include a lack of fear of weight gain in ARFID, no shape and weight concerns in ARFID, and no specific focus on weight

loss in ARFID. It is essential to obtain collateral history from parents who will usually indicate no avoidance of high calorie food in ARFID. The patients are aware that they are low weight and may express a wish to eat more and gain weight, but their anxiety and fear prevents them from actually consuming enough.²⁴ ARFID can sometimes be confused with ASD, other neurodevelopmental disorders, and anxiety disorders.

OTHER SPECIFIED FEEDING OR EATING DISORDERS

In previous iterations of the DSM, an eating disorder not otherwise specified (EDNOS) was the broad diagnostic category that encompassed atypical and subthreshold presentations of AN, BN, BED and other atypical eating problems. In DSM-5 many of the patients who would formally have been diagnosed with EDNOS will now meet the revised criteria for AN, BN, BED and ARFID. Remaining patients will be diagnosed under the general grouping of “other specified feeding or eating disorders” and will include specified diagnostic groupings such as atypical AN, which meets all criteria for AN, except that despite significant weight loss, the individual’s weight remains within or above the normal range; BN, with symptoms of low frequency and/or limited duration; BED, with symptoms of low frequency and/or limited duration; purging disorder, recurrent purging to influence body shape or weight in the absence of binge eating; and night eating syndrome, recurrent episodes of eating at night that leads to impairment and/or distress. Finally, there is a diagnosis called “unspecified feeding and eating disorder” for cases in which the clinician chooses not to specify the reason that criteria are not met for a specific feeding or eating disorder, and includes presentations in which there is insufficient information to make a more specific diagnosis. The rationale for this reorganization in DSM-5 was based in part on studies that suggested that many younger patients with AN and BN would receive more appropriate clinical diagnoses based on DSM-5 criteria.¹²²

Epidemiology, Etiology, and Risk Factors

The epidemiology of this new grouping of specified eating disorders in DSM-5 is unknown, particularly in children and adolescents. For sub-threshold AN and BN cases, the same etiologic and risk factors for full syndrome disorders are likely pertinent.

Differential Diagnosis

The differential diagnosis of disorders likely to be found in this group includes AN and BN, as well as the range of medical and psychological problems that may cause eating and weight changes in children and adolescents. The most common psychological problems include changes in appetite related to depression, anxiety and phobia secondary to choking, difficulty swallowing, gastric upset, gastrointestinal motility problems, and fear of vomiting and food intake after viral gastroenteritis. Common medical problems include diabetes mellitus, celiac disease, irritable bowel syndrome, and allergies. In addition to these types of presentations, a clinical syndrome associated with women athletes has been described and is called the Female Triad Syndrome. The Female Triad Syndrome consists of three elements: low dietary energy availability from disordered eating, amenorrhea, and low bone density. While not a specific eating disorder, presentations consistent with this triad should be considered at high risk for an eating disorder and monitored closely for worsening medical problems and eating disorder symptoms. Studies suggest that among female high school athletes, 18.2% met criteria for disordered eating, 12.5% met criteria for menstrual irregularity, and 21.8% met criteria for low bone mass.^{124,125}

EVIDENCE BASE FOR PRACTICE PARAMETERS

In this parameter, recommendations for best assessment and treatment practices are stated in accordance with the strength of the underlying empirical and/or clinical support, as follows:

- Clinical Standard [CS] is applied to recommendations that are based on rigorous empirical evidence, such as meta-analyses, systematic reviews, individual randomized controlled trials, and/or overwhelming clinical consensus;
- Clinical Guideline [CG] is applied to recommendations that are based on strong empirical evidence, such as non-randomized controlled trials, cohort studies, case-control studies, and/or strong clinical consensus;
- Clinical Option [OP] is applied to recommendations that are based on emerging empirical evidence, such as uncontrolled trials or case series/reports, or clinical opinion, but lack strong empirical evidence and/or strong clinical consensus;
- Not Endorsed [NE] is applied to practices that are known to be ineffective or contraindicated.

The strength of the empirical evidence is rated in descending order as follows:

- Randomized, Controlled Trial [rct] is applied to studies in which subjects are randomly assigned to two or more treatment conditions;
- Controlled Trial [ct] is applied to studies in which subjects are non-randomly assigned to two or more treatment conditions;
- Uncontrolled Trial [ut] is applied to studies in which subjects are assigned to one treatment condition;
- Case Series/Report [cs] is applied to a case series or a case report.

RECOMMENDATIONS

Recommendation 1: Mental health clinicians should screen all child and adolescent patients for eating disorders (CS).

All preteen and adolescent patients should be asked about eating patterns and body satisfaction. In addition, height and weight should be obtained and plotted on growth curves. If there is a concern about the ratio of height to weight or evidence of growth failure, referral for further evaluation is warranted. The Eating Disorder Examination-Questionnaire (EDE-Q), Eating Disorder Inventory (EDI) and Eating Attitudes Test (EAT) are validated, short, self-report measures that can be useful screening instruments for eating disorders.¹²⁶⁻¹²⁸ The Kids' Eating Disorder Survey (KEDS), ChEDE-Q, the EDI-C, and the Child-Eating Attitudes Test (CHEAT) are all validated measures for use in younger children.¹²⁹

Recommendation 2: A positive screening should be followed by a comprehensive diagnostic evaluation, including laboratory tests and imaging studies as indicated (CS).

Evaluation of a child or adolescent who screens positive for an eating disorder should include complete psychiatric evaluation and physical examination. A comprehensive evaluation, in addition to components typically included in a psychiatric assessment, should include evaluation of weight and height, changes in body weight, menstrual history in females, body image concerns, presence of dieting and calorie counting, as well as the amount, type and frequency of exercise. A registered dietician can be helpful in assessing specific dietary intake. Percent of weight loss and rapidity of weight loss should also be evaluated.¹³⁰ In addition, binge eating or purging behaviors should be assessed including frequency, amount consumed, and

duration of the behaviors. Evaluation of comorbid symptoms including depressive symptoms, anxiety symptoms, obsessional thoughts, self-injurious behavior and suicidality is also necessary. Screening for psychotic symptoms and drug use is also recommended. Inquiry into label-checking and self-weighing can be helpful if other symptoms of body image preoccupation are denied. Obtaining a history from parents is critical to corroborate symptoms.²⁶ Percentile body mass indexes (BMIs) are necessary to estimate levels of malnutrition in children and adolescents given the inaccuracy of standard BMI calculations in this age group.³⁰ Methods for calculating percentile BMIs are available using the Centers for Disease Control (CDC) age and gender growth charts.^{31,131} Efforts should be made to obtain longitudinal growth charts and deviations from individual growth trajectories should be evaluated.

The best characterized and most commonly used structured interview for assessing disordered eating behaviors and eating-related psychopathology is the Eating Disorder Examination (EDE).^{132,133} The EDE is reliable for patients down to 12 years of age. A child version for evaluating children and adolescents under 14 years of age is also available.¹³⁴ There is also a self-report version (EDE-Questionnaire), which takes about five minutes to complete and is reliable in adolescents.^{128,135,136} It is noteworthy that for adolescent AN, studies suggest that minimization and denial may produce unusually low scores on the EDE despite the clinical presence of clear behaviors consistent with AN.²⁶ In this instance, parental report using the EDE interview provides scores similar to clinician ratings.³⁴ In addition to the EDE, several other commonly utilized questionnaires are available, including the EAT,¹²⁶ CHEAT,¹³⁷ and (EDI).¹³⁸

For adolescents with BN, the EDE appears to provide a good measure of eating disordered behaviors and constructs, even without parent report.^{34,132} Another measure specific to BN that may be considered is the Bulimia-Test-Revised (BULIT-R). Several assessment instruments are relevant for AN, BN, and BED, such as EDE, CHEAT,^{126,127} EAT, and EDI.^{132,137} A critical review of the strengths of these various instruments is available.¹³⁹

In those patients where there is evidence of malnutrition or purging behaviors, initial laboratory testing typically includes a complete blood count, chemistry profile including electrolytes, blood urea nitrogen, creatinine, glucose, and liver functions including aspartate aminotransferase (AST) and alanine aminotransferase (ALT). These tests also can be used to monitor treatment. A thyroid stimulating hormone (TSH) test should be ordered to rule out underlying thyroid dysfunction. Further blood testing should include measuring calcium,

magnesium, phosphate, total protein, albumin, erythrocyte sedimentation rate (ESR), amylase as some studies suggest that elevations of amylase provides evidence that the patient is vomiting, B12, and lipid profiles; in females, luteinizing hormone (LH), follicle stimulating hormone (FSH), and estradiol levels should also be tested. If indicated, a pregnancy test using beta human chorionic gonadotropin (B-hCG) should also be considered to evaluate amenorrhea.

Electrocardiograms are often necessary to further evaluate bradycardia and risk of cardiac arrhythmias. Dual-energy x-ray absorptiometry (DEXA) of bone should initially be conducted on females with amenorrhea lasting more than six months and yearly if amenorrhea persists.¹⁴⁰ All males with significant weight loss should also have a DEXA scan. Results of these data provide patients and families with guidance about the clinical impact of starvation on physical health and growth. Over time, improvements in these physical health parameters can help benchmark clinical progress. For example, for females with AN, normalization of estrogen levels can be a marker of healthy weight. At the same time, in more chronically ill patients with eating disorders, there can be a normalization or laboratory values that can suggest better health than is present.

Recommendation 3: Severe acute physical signs and medical complications need to be treated (CS).

Adverse physical effects of eating disorders include cardiac arrhythmias, bradycardia, hypotension, hypothermia, dehydration, congestive heart failure, kidney failure, pancreatitis, amenorrhea or irregular menses, low bone mineral density, neurological and cognitive impairments, delay in growth or growth impairment, pubertal delay or interruption, hormonal imbalances, and fluid and electrolyte abnormalities. In addition, clinical signs of malnutrition include hair loss, lanugo hair, dry skin, dependent edema, muscle weakness, and cramps.⁷⁸ Most physical abnormalities are reversible with adequate diet and restoration of a healthy weight. In children and adolescents with longstanding AN, or other low weight eating and feeding disorders, some clinical abnormalities may be irreversible, including growth impairment, decreased bone density, structural brain changes and infertility.⁷⁸ Patients admitted for malnutrition should be monitored for potential re-feeding syndrome during the initial re-nourishment phase.¹⁴¹

Frequent purging associated with BN, or Purging Disorder, may cause parotid swelling, calluses on the dorsum of the hand (Russell's sign) from teeth scraping on top of the hand when using fingers to stimulate regurgitation, erosion of dental enamel, or serious fluid and electrolyte disturbances, such as hypokalemia.¹⁴¹ These problems may lead to orthostatic hypotension and syncope. Esophageal tears from excessive erosion of throat tissue can lead to serious and difficult to control bleeding. Binge-eating can cause both gastric and esophageal rupture.¹⁴² The mortality rate is estimated to be about 1% in BN patients, but more recent studies suggest that this may be an underestimate.^{143,144}

Indications for medical hospitalization of children and adolescents have been published by the American Academy of Pediatrics and the Society of Adolescent Health and Medicine.^{78,145} These include severe abnormality of heart rate (bradycardia and orthostatic heart rate changes), blood pressure (orthostatic hypotension), body temperature (hypothermia), electrolyte abnormalities, and severe malnutrition.^{78,146} These standards were developed primarily in relation to physiologic impacts secondary to malnutrition and purging; they are, however, reasonable indicators of medical compromise for all child and adolescent eating disorders.

In the context of hospitalization for weight restoration, some data suggests that the use of nasogastric tube feeding is more efficient than other approaches in promoting weight gain.^{147[cs],148[cs]} The long term benefits of this approach, however, have not been established and the clinical need for the approach not substantiated.⁷⁸

Recommendation 4: Psychiatric hospitalization, day programs, partial hospitalization programs, and residential programs for eating disorders in children and adolescents should be considered only after outpatient interventions have been unsuccessful (CG).

There is no evidence that psychiatric hospitalization for eating disorders is more effective than outpatient treatment.^{149[rct],150[rct]} A few uncontrolled studies suggest that residential and day treatment^{151[ut]} may be useful, but no studies have been randomized or have compared residential and day treatment^{152[ut]} to outpatient treatment in adolescents.^{153[ut],154[ut]} Negative impacts of such programs include separation of the developing child from family, friends, and community. Nonetheless, such intensive programs are sometimes clinically necessary because of poor response to, or the lack of, availability of appropriate specialty outpatient treatment. In those instances, these negative impacts can be mitigated by keeping length of stay short, using the

lowest safe level of care, involving families in programming, and utilizing highly expert and experienced staff.

Recommendation 5: Treatment of eating disorders in youth usually involves a multi-disciplinary team that is developmentally aware, sensitive, and skilled in the care of children and adolescents with eating disorders (CS).

Eating disorders affect the psychological and physical health of children and adolescents with these illnesses.¹⁵⁵ It is therefore usually necessary to have a team of practitioners who work together. The team typically consists of a psychotherapist, pediatrician, and dietician. There may also be a role for medication management by a child psychiatrist who may play a dual role as both psychotherapist and psychopharmacologist. The physician tasked with ongoing medical monitoring ensures the patient is safe for outpatient treatment according to the guidelines provided by the American Academy of Pediatrics and the Society of Adolescent Health and Medicine.^{78,145} The dietician may be a consultant to the pediatrician or therapist about any nutritional concerns. Some dieticians may provide direct consultation to families and patients, depending on the overall treatment approach being utilized. Studies do not support the use of nutritional counseling as a sole treatment for eating disorders.^{156[rect],157[rect]}

Recommendation 6: Outpatient psychosocial interventions are the initial treatment of choice for children and adolescents with eating disorders (CS).

There is a limited database of empirically supported treatment in AN.^{158,159} Outpatient psychosocial interventions studied for adolescent AN include family and individual therapies. Six RCTs, including a total of 323 adolescent participants, have evaluated family therapy for adolescents with AN.^{160[rect]-165[rect]} The findings from these studies and a meta-analysis of RCTs¹⁵⁰ suggested that family therapy, particularly Family-Based Treatment (FBT),¹³ sometimes referred to as Maudsley Family Therapy,¹⁶⁶ is effective and superior to comparison individual therapies. FBT is an outpatient form of family therapy. FBT consists of between 10-20 family meetings over a 6-12 month treatment course.¹³ FBT empowers parents to take charge of the weight restoration of their child by taking action to disrupt symptoms of self-starvation and over exercise. Once the child is able to eat independently without parental supervision and has reached a normal weight, the treatment briefly focuses on developmental issues of adolescence.

There is evidence that this approach is both highly efficient^{163[rct]} and likely decreases the need for hospitalization.^{167[ut]} Although individual therapy was not as effective as family therapy in these studies,^{165[rct]} individual approaches were nonetheless beneficial and could be offered for patients where FBT is not an acceptable or tenable option.^{164[rct]} In particular, Adolescent Focused Therapy (AFT), which is an individual therapy focused on individuation and self-efficacy, was found to be useful.^{168[rct]} AFT encourages the adolescent to manage her own eating and weight gain through the relationship with the therapist. In addition, the main focus of AFT is to encourage an increased awareness and tolerance of emotions, particularly negative ones.¹⁶⁸ CBT for adolescent AN was used in one study^{150[rct]} and found to be the most cost-effective approach compared to treatment as usual or hospitalization.^{169[rct]} More recent studies suggest that a new form of expanded CBT (CBT-E), may be useful for adolescents with AN, BN, and patients who do not meet full criteria for these disorders, though randomized studies have not yet been conducted.^{170[ut]}

Although there are a number of RCTs examining treatment for adults with BN, there are only two RCTs examining treatment for adolescent BN including a total of 165 adolescent participants. Schmidt and colleagues compared FBT and a self-help version of CBT for 85 adolescents (ages 13-20) who met full or partial DSM-IV BN criteria.^{171[rct]} CBT begins with psycho-education about BN followed by cognitive and behavioral exercises designed to change maladaptive eating-related thoughts and behaviors. CBT initially focuses on normalizing eating patterns to reduce excessive hunger and binge eating. Use of self-monitoring through a diet record is central to this aspect of CBT for BN. Once a normal pattern of eating is established, cognitive distortions and behavioral experiments are encouraged to challenge beliefs and fears.¹⁷² One study of adolescents with BN comparing CBT group therapy with FBT found that the CBT group had a significantly lower rate of binge-eating compared to those in the FBT group at the end of treatment, although no statistically significant difference was found between treatments at the six-month follow-up.^{171[rct]} Le Grange and colleagues compared FBT and individual supportive psychotherapy (SPT) in adolescents (ages 12 to 19) who met full or partial DSM-IV BN criteria.^{173[rct]} Results of this study indicated that FBT was more effective than SPT at both end-of-treatment and six-month follow-up. Additionally, FBT reduced binge eating/purging rates in significantly shorter time when compared to SPT. Other psychosocial interventions have been examined and found useful in adults with BN, including interpersonal psychotherapy (IPT)¹⁷⁴

and dialectical behavioral therapy (DBT).^{175[rect]} In one multi-site study of adults with BN that compared CBT to IPT, CBT resulted in a shorter time to abstinence from binge eating and purging, but at follow-up there were no differences in outcome between the two treatments.^{176[rect]} IPT has not been studied in adolescent BN, but it is effective for adolescent depression.^{177[rect]} DBT has been modified for use with adolescent BN, but only case reports are available.^{178[cs]} DBT has also been gaining an evidence base for adolescents with multiple diagnoses who have affect dysregulation and self-harming behaviors, and has been applied to adults with eating disorders¹⁷⁹ and comorbid borderline personality disorder. The results suggested that DBT is worthy of further study.

Treatment of BED in adults has received considerable attention in research studies.^{163[cs],165[rect],166[rect],167[rect],168 [ut]} To date, studies in adults suggest that CBT,^{21[rect]} IPT,^{21[rect]} and DBT^{180[rect]} are effective. In adolescents with BED, preliminary studies support the use of IPT, but BED has otherwise been relatively unexamined in younger patients.^{23[cs]}

For those children and adolescents with ARFID, there are no empirical studies to guide treatment. For the most part, these disorders require individualized behavioral plans to address the specific eating problem, but use of CBT and family interventions may be helpful. For example, gradual desensitization procedures are often helpful along with behavior reinforcement plans for many of these problems.^{121[cs]} When these problems are severe enough to lead to medical instability or severe malnutrition, hospitalization may be needed.^{78[cs]}

Recommendation 7: The use of medications, including complementary and alternative medications, should be reserved for comorbid conditions and refractory cases (CG).

Medications for adults with AN have been tried in case series and small pilot studies. To date, the results of these studies are not encouraging.^{181[cs]} Selective serotonin reuptake inhibitors (SSRIs) were initially thought to be helpful to prevent relapse, but larger scale studies did not support this.^{182[rect]} No systematic studies of SSRIs have been conducted in adolescents with AN. More recently, atypical antipsychotics have been examined in small studies^{183[cs]} and small RCTs in adults^{184[rect]-186[rect]} because of their potential effects on weight, anxiety, and obsessive thinking.¹⁸⁷ Feasibility and acceptability of medication is a major problem because of fear of weight gain in adolescents.^{188[rect]} A recent pilot RCT for adolescents with AN found few benefits to adding risperidone to standard treatment, though the medication appeared to be well

tolerated.^{189[rct]} Another small trial combined treatment as usual with either placebo or olanzapine over a 10 week period and found no differential benefit with the addition of olanzapine.^{186[rct]} A small randomized study of quetiapine, compared to treatment as usual found some evidence of greater improvements in weight and eating related thinking in the group randomized to quetiapine, but there were no statistically significant differences between groups.^{190[rct]}

Medications for BN have been studied almost exclusively in adults with the disorder.¹⁷ Studies suggest that antidepressants are feasible, acceptable, and effective in adults with BN. Fluoxetine has been the most widely studied, and has been found to decrease urges to binge and purge. Fluoxetine is FDA approved for the treatment of BN, but the dose of fluoxetine in these studies was considerably higher than typically needed for depression (e.g., 60 mg per day). One small study of antidepressants such as SSRIs in adolescents found that the medications were feasible and acceptable with few side effects in this age group.^{191[ut]} CBT, however, appears to be superior to antidepressants and current recommendations support the use of medications for patients who refuse CBT or who do not have an optimal response to CBT.¹⁹² Adult studies suggest that for depressed patients with BN, the combination of CBT and SSRIs is the most effective approach. Although some medications appear to be useful for adults with BED, none have been studied in adolescents.

For comorbid conditions, guidelines for the specific condition should be followed. When patients are starved, however, lower levels of available serotonin may limit the effectiveness of SSRIs and other antidepressants until weight is at least partially restored. In addition, it is important to note that obsessionality and depressed mood often improve with weight gain alone.^{193[cs]}

PARAMETER LIMITATIONS

AACAP Practice Parameters are developed to assist clinicians in psychiatric decision making. These Parameters are not intended to define the sole standard of care. As such, the Parameters should not be deemed inclusive of all proper methods of care or exclusive of other methods of care directed at obtaining the desired results. The ultimate judgment regarding the care of a particular patient must be made by the clinician in light of all of the circumstances presented by the patient and his or her family, the diagnostic and treatment options available, and available resources.

Disclosures: James Lock, MD, PhD serves or has served on the Advisory Board for the Center for Discovery, the Global Foundation for Eating Disorders, and the National Eating Disorders Association; serves or has served as a consultant for the Training Institute for Eating Disorders in Children and Adolescents; receives or has received grant support from the Davis Foundation, the Global Foundation for Eating Disorders, and the National Institutes of Health; and, receives royalties from Guilford Press and Oxford University Press. Maria C. La Via, MD received research funding from the National Institute of Mental Health. Oscar Bukstein, MD, MPH, co-chair, receives royalties from Routledge Press. Heather Walter, MD, MPH, co-chair, has no financial relationships to disclose.

Treatment	Treatment Targets	Evidence-Base	Recommendations
Family-Based Treatment ¹⁹⁴	Family therapy supports parental management of eating and related behavior until adolescent demonstrates improvement.	Six randomized controlled trials ¹⁶⁰⁻¹⁶⁵ support efficacy for AN. Superiority over other treatments unclear. Two randomized controlled trials supporting usefulness for BN. ^{171,195}	Useful for most cases of short duration AN and BN in young patients.
Adolescent Focused Therapy	Individual therapy targets autonomy and self-efficacy in the context of adolescent development.	This treatment has been the subject of two RCTs ^{164,165} where it performed worse than FBT for AN, but was still effective.	Useful for adolescents with AN when FBT is not feasible.
Cognitive Behavior Therapy	Individually focused therapy targets adolescent management of behaviors and distorted cognitions associated with AN and BN.	This treatment has been the subject of one RCT and one case series for adolescents with BN. ^{171,172} No published studies of CBT for adolescent AN.	Adolescent version of CBT may be appropriate for use with BN.

Interpersonal Psychotherapy ¹⁷⁴	Interpersonal Psychotherapy (IPT) focuses on changing problematic interpersonal relationships that trigger or maintain eating disorder symptoms.	Two randomized controlled trials ^{21,176} in adults with BN and BED support the use of IPT for these disorders. Preliminary studies suggest that IPT is useful for adolescents with BED. ^{23, 196}	Useful for cases of BN and BED as an alternative to CBT. ²¹
Antidepressants	Symptoms of obsessionality, anxiety, and depression in AN and BN. Targets binge eating and purging in BN. ¹⁹⁷	One uncontrolled trial ¹⁹¹ suggests antidepressants are tolerated and may be helpful for adolescent BN.	Useful for comorbid disorders. Use as a second line treatment for adolescent BN.
Atypical Antipsychotics	Body image distortion, weight gain fears, and anxiety related to AN.	Case series data and three pilot RCTs. ^{186,189,190} Insufficient evidence to suggest efficacy for use in AN.	Useful for comorbid conditions. Further study needed to determine efficacy for core symptoms of AN.

1 REFERENCES

- 2 1. Morton R. *Phthisiologia: Or, a Treatise of Consumptions*. London: Smith & Walford;
3 1694.
- 4 2. Saraf M. Holy anorexia an anorexia nervosa: Society and concept of disease. *The*
5 *Pharos*. 1998;61:2-4.
- 6 3. Gull W. Anorexia Nervosa (apepsia hysterica, anorexia hysterica). *Transactions of the*
7 *Clinical Society of London*. 1874;7:222-228.
- 8 4. Silverman J. Charcot's comments on the therapeutic role of isolation in the treatment of
9 anorexia nervosa. *Int J Eat Disord*. 1997;21:295-298.
- 10 5. McKenzie JM. Hospitalization for anorexia nervosa. *Int J Eat Disord*. 1992;11:235-241.
- 11 6. Zhao Y, Encinosa W. Hospitalizations for eating disorders from 1999-2006. In: Quality
12 AfHRA, ed. Vol April 20092009.
- 13 7. Thoma H. *Anorexia Nervosa*. New York: International Universities Press; 1967.
- 14 8. Bruch H. *Eating Disorders: Obesity, Anorexia Nervosa, and the Person Within*. New
15 York: Basic Books; 1973.
- 16 9. Bruch H. *The Golden Cage: The Enigma of Anorexia Nervosa*. Cambridge, MA:
17 Harvard University Press; 1978.
- 18 10. Meads C, Gold L, Burls A. How effective is outpatient compared to inpatient care for
19 treatment of anorexia nervosa? A systematic review. *European Eating Disorders*
20 *Review*. 2001;9:229-241.
- 21 11. Minuchin S, Rosman B, Baker I. *Psychosomatic Families: Anorexia Nervosa in Context*.
22 Cambridge, MA: Harvard University Press; 1978.
- 23 12. Dare C, Eisler I. Family therapy for anorexia nervosa. In: Cooper I, Stein A, eds. *The*
24 *Nature and Management of Feeding Problems in Young People*. New York: Harwood
25 Academics; 1992:146-160.
- 26 13. Lock J, Le Grange D. *Treatment Manual for Anorexia Nervosa: A Family-Based*
27 *Approach*. 2nd ed. New York: Guilford Press; 2013.
- 28 14. Boskind-Lodahl M. Cinderella's stepsisters: a feminist perspective on anorexia nervosa.
29 *Signs*. 1976;2:342-356.
- 30 15. Boskind-Lodahl M, White W. The definition and treatment of bulimarexia in college age
31 women-a pilot study. *Journal of the American College Health Association*. 1978;27:84-
32 97.
- 33 16. Russell G. Bulimia nervosa: An ominous variant of anorexia nervosa. *Psychological*
34 *Medicine*. 1979;9:429-448.
- 35 17. Mitchell J, Agras WS, Wonderlich S. Treatment of bulimia nervosa: Where are we and
36 where are we going? *Int J Eat Disord*. 2007;40:95-101.
- 37 18. Mitchell J, Agras W, Crow S, et al. Stepped care and cognitive-behavioural therapy for
38 bulimia nervosa :randomised trial. *Br J Psychiatry*. 2011;198:391-397.
- 39 19. Association AP. *Diagnostic and Statistical Manual of Mental Disorders*, . Arlington, VA:
40 American Psychiatric Association; 2013.
- 41 20. Decaluwe V, Braet C. Prevalence of binge eating in obese children and adolescents
42 seeking weight loss treatment. *International Journal of Obesity*. 2003;27:404-409.
- 43 21. Wilson G, Wilfley D, Agras W, Bryson S. Psychological treatments of binge eating
44 disorder. *Arch Gen Psychiatry*. 2010;67:94-101.

- 1 22. Wilfley D, Crow S, Hudson JI, et al. Efficacy of subutramine for the treatment of binge
2 eating disorder: a randomized multicenter placebo-controlled double-blind study.
3 *American Journal of Psychiatry*. 2008;165:51-58.
- 4 23. Tanofsky-Kraff M, Wilfley D, Young J, et al. A pilot study of interpersonal
5 psychotherapy for preventing excess weight gain in adolescent girls at-risk for obesity.
6 *IJED*. 2010;43:701-706.
- 7 24. Bryant-Waugh R, Kreipe R. Avoidant/Restrictive Food Intake Disorder in DSM-5.
8 *Psychiatric Annals*. 2012;42:402-405.
- 9 25. American Psychiatric Association. *Diagnostic and Statistical Manual of Mental*
10 *Disorders, 5th edition: DSM-5*. Washington D.C.: American Psychiatric Association;
11 2013.
- 12 26. Couturier J, Lock J. Denial and minimization in adolescent anorexia nervosa. *Int J Eat*
13 *Disord*. 2006;39:175-183.
- 14 27. Roberto C, Steinglass J, Walsh BT. The clinical significance of amenorrhea as a
15 diagnostic criterion for anorexia nervosa. *Int J Eat Disord*. 2008;41:559-563.
- 16 28. Hebebrand J, Himmelman G, Hesecker H, Schaefer H, Remschmidt H. The use of
17 percentiles for the body mass index in anorexia nervosa: diagnostic, epidemiological, and
18 therapeutic considerations. *Int J Eat Disord*. 1996;19:359-369.
- 19 29. Zhang A, Lai, HuicChan J. Comparison of the use of body mass index percentiles and
20 percentage of ideal body weight to screen for malnutrition in children with cystic fibrosis.
21 *American Journal of Clinical Nutrition*. 2004;80:982-991.
- 22 30. Hebebrand J, Wehmeier P, Remschmidt H. Weight criteria for diagnosis of anorexia
23 nervosa. *Am J Psychiatry*. 2000;157:6.
- 24 31. Le Grange D, Doyle P, Swanson S, Ludwig K, Glunz C, Kreipe R. Calculation of
25 expected body weight in adolescents with eating disorders. *Pediatrics*. 2012;129:e438-
26 e446.
- 27 32. Company MLI. 1983 Metropolitan height and weight tables. *Stat Bull Metropolitan Life*
28 *Insurance Company*. 1983;64:1-9.
- 29 33. Workgroup, Adolescents fCoEDiCa. Classification of child and adolescent eating
30 disturbances. *Int J Eat Disord*. 2007;40(S117-S122).
- 31 34. Couturier J, Lock J, Forsberg S, Vanderheyden D, Lee HY. The addition of a parent and
32 clinician component to the eating disorder examination for children and adolescents. *Int J*
33 *Eat Disord*. 2007;40:472-475.
- 34 35. Peebles R, Wilson J, Lock J. How do children and adolescents with eating disorders
35 differ at presentation, *Journal of Adolescent Health*. *Journal of Adolescent Health*.
36 2006;39:800-805.
- 37 36. Pinhas L, Morris A, Crosby R, Katzman D. Incidence and age specific presentation of
38 restrictive eating disorders in children: a Canadian surveillance program study. *Archives*
39 *of Pediatrics and Adolescent Medicine*. 2011;165:895-899.
- 40 37. Steinhausen H. The outcome of anorexia nervosa in the 20th century. *Am J Psychiatry*.
41 2002;159:1284-1293.
- 42 38. Arcelus J, Mitchell A, Wales J, Nielsen S. Mortality rates in patients with anorexia
43 nervosa and other eating disorders. *Archives of General Psychiatry*. 2011;68:724-731.
- 44 39. Fichter M, Quadflieg N, Hedlund S. Twelve-year course and outcome predictors of
45 anorexia nervosa. *Int J Eat Disord*. 2006;39:87-100.

- 1 40. Herperz-Dahlmann B, Muller B, Herperz S, Heussen N, Hebebrand J, Remschmidt H.
2 Prospective 10-year follow-up in adolescent anorexia nervosa--course, outcome,
3 psychiatric comorbidity, and psychosocial adaptation. *J Child Psychol Psychiatry*.
4 2001;42:603-612.
- 5 41. Hoek H, Hoeken Dv. Review of prevalence and incidence of eating disorders. *Int J Eat*
6 *Disord*. 2003;34:383-396.
- 7 42. Swanson J, SJ C, Le Grange D, Swendsen J, Merikangas K. Prevalence and correlates of
8 eating disorders in adolescents: Results from the National CoMorbidity Survey
9 Replication Adolescent Supplement. *Archives of General Psychiatry*. 2011;68:714-723.
- 10 43. Lucas AR, Beard CM, O'Fallon WM. 50-year trends in the incidence of anorexia nervosa
11 in Rochester, Minn: A population-based study. *American Journal of Psychiatry*.
12 1991;148:917-929.
- 13 44. Keski-Rahkonen A, Hoek H, Susser ES, et al. Epidemiology and course of anorexia
14 nervosa in the community. *Am J Psychiatry*. 2007;164:1259-1165.
- 15 45. Norris M, Apsimon M, Harrison M, et al. An examination of medical and psychological
16 morbidity in adolescent males with eating disorders. *Eating Disorders*. 2012;20:405-415.
- 17 46. Braun D, Sunday S, Huang A, Halmi CA. More males seek treatment for eating
18 disorders. *Int J Eat Disord*. 1999;25:415-424.
- 19 47. Hudson JI, Hiripi E, Pope HG, Kessler RC. The prevalence and correlates of eating
20 disorders in the national comorbidity survey replication. *Biological Psychiatry*.
21 2007;61:348-358.
- 22 48. Strober M, RK F, Lampert C, Diamond J, WH K. Males with anorexia nervosa: a
23 controlled study of eating disorders in first degree relatives. *IJED*. 2001;29:263-269.
- 24 49. Lock J. Fitting Square Pegs into Round Holes: Males with Eating Disorders. *Journal of*
25 *Adolecent Health*. 2008;44:99-100.
- 26 50. Hudson JI, Hiripi E, Pope HG J, RC K. The prevalence and correlates of eating disorders
27 in the National Comorbidity Survey Replication. *Biol Psychiatry*. 2007;61(3):348-358.
- 28 51. Hoek H, van Harten PN, Hermans KM, Katzman MA, Matroos GE, Susser ES. The
29 incidence of anorexia nervosa on Curacao. *Am J. Psychiatry*. 2005;162:748-752.
- 30 52. Joergensen J. The epidemiology of eating disorders in Fyn Country, Denmark, 1977-
31 1986. *Acta Psychiatrica Scandinavica*. 1992;85:30-34.
- 32 53. Halmi K, Brodland G, Loney J. Progress in anorexia nervosa. *Annals of Internal*
33 *Medicine*. 1973;78:907-909.
- 34 54. Nicholls D, Lynn R, Viner R. Childhood eating disorders: British national surveillance
35 study. *Br J Psychiatry*. 2011;198:295-301.
- 36 55. Bulik C, Sullivan P, Tozzi F, Furberg H, Lichtenstein P, Pedersen N. Prevalence,
37 heritability and prospective risk factors for anorexia nervosa. *Archives of General*
38 *Psychiatry*. 2006;63:305-312.
- 39 56. Klump KL, McGue M, Iacona W. Age differences in genetic and environmental
40 influences on eating attitudes and behaviors in preadolescent female twins. *Journal of*
41 *Abnormal Psychology*. 2000;109:239-251.
- 42 57. Klump KL, Burt SA, McGue M, Iacona W. Changes in genetic and environmental
43 influences on disordered eating across adolescence: A longitudinal twin study. *Archives*
44 *of General Psychiatry*. 2007;64:1409-1415.
- 45 58. Klump KL, McGue M, Iacona W. Differential heritability of eating attitudes and
46 behaviors in prepubertal versus pubertal twins. *IJED*. 2003;33:287-292.

- 1 59. Klump KL, Perkins P, Burt SA, McGue M, Iacona W. Puberty moderates genetic
2 influences on disordered eating. *Psychological Medicine*. 2007;37:627-634.
- 3 60. Klump K, Bulik CM, Pollice C, et al. Temperament and character in women with
4 anorexia nervosa. *J Nerv Ment Dis*. 2000;188:559-567.
- 5 61. Cassin S, von Ramson L. Personality and eating disorders: A decade in review. *Clin*
6 *Psycho Rev*. 2005;25:895-916.
- 7 62. Wade T, Tiggemann M, Bulik CM, Fairburn CG, Wray N, Martin N. Shared
8 temperament risk factors for anorexia nervosa: A twin study. *Psychosomatic Medicine*.
9 2008;70:239-244.
- 10 63. Tchanturia K, Davies H, Harrison A, et al. Poor cognitive flexibility in eating disorders:
11 Examining the evidence. *PLoS ONE*. 2012;7:e28331.
- 12 64. Lopez C, Tchanturia K, Stahl D, Treasure J. Central coherence in eating disorders: a
13 systematic review. *Psychol Med*. 2008;38:1075-1084.
- 14 65. Holliday J, Tchanturia K, Landau S, Collier D. Is Impaired Set-Shifting an
15 Endophenotype of Anorexia Nervosa? *Am J Psychiatry*. 2005;162:2269-2275.
- 16 66. Treasure J. Getting beneath the phenotype of anorexia nervosa: the search for viable
17 endophenotypes and genotypes. *La revue de psychiatrie*. 2007;52:212-219.
- 18 67. Fitzpatrick K, Lock J, Darcy A, Colburn D, Gudorf C. Neurocognitive processes in
19 adolescent anorexia nervosa. 2012:12 JUN 2012 | DOI: 2010.1002/eat.22027.
- 20 68. Marchi M, Cohen P. Early childhood eating behaviors and adolescent eating disorders. *J*
21 *Am Acad Child Adolesc Psychiatry*. 1990;29:112-117.
- 22 69. Crisp AH. Anorexia Nervosa as flight from growth: Assessment and treatment based on
23 the model. In: Garner DM, Garfinkel P, eds. *Handbook of treatment for eating disorders*.
24 New York: Guilford; 1997:248-277.
- 25 70. Attie I, Brooks-Gunn J. Development of eating problems in adolescent girls: A
26 longitudinal study. *Developmental Psychology*. 1989;25:70-79.
- 27 71. Killen JD, Taylor CB, Hayward C, al e. Pursuit of thinness and onset of eating disorder
28 symptoms in a community sample of adolescent girls: a three year prospective analysis.
29 *Int J Eat Disord*. 1994;16:227-238.
- 30 72. Investigators M. Risk factors for the onset of eating disorders in adolescent girls: results
31 of the McKnight longitudinal risk factor study. *Am J Psychiatry*. 2003;160:248-254.
- 32 73. Anderson-Fye, Becker AE. Sociocultural Aspects of Eating Disorders. In: Thompson J,
33 ed. *Handbook of Eating Disorders and Obesity*. Hoboken, New Jersey: John Wiley &
34 Sons; 2004:565-589.
- 35 74. Stice E. Sociocultural influences on body image and eating disturbance. In: Fairburn CG,
36 Brownell K, eds. *Eating and Weight Disorders and Obesity: A Comprehensive*
37 *Handbook*. Second ed. New York: Guilford Press; 2002:103-107.
- 38 75. Lake A, Staiger P, Glowinski H. Effects of western culture on women's attitudes to eating
39 and perceptions of body shape. *Int J Eat Disord*. 2000;27:83-89.
- 40 76. Wildes J, Emery R. The roles of ethnicity and culture in the development of eating
41 disturbance and body dissatisfaction: A meta-analytic review. *Clinical Psychology*
42 *Reviv*. 2001;21:521-551.
- 43 77. Gunewardene A, Huon G, Zheng R. Exposure to westernization and dieting: a cross-
44 cultural study. *Int J Eat Disord*. 2001;29:289-293.

- 1 78. Golden N, Katzman D, Kreipe R, et al. Eating disorders in adolescents: position paper of
2 the Society for Adolescent Medicine: Medical Indications for Hospitalization in an
3 Adolescent with an Eating Disorder. *J Adolesc Health*. 2003;33:496-503.
- 4 79. Swanson J, Crow S, Le Grange D, Swendsen J, Marikangas K. Prevalence and correlates
5 of eating disorders in adolescents. *Archives of General Psychiatry*. 2011;68:714-713.
- 6 80. Godart N, Flament M, Perdereau F, Jeammet P. Comorbidity between eating disorders
7 and anxiety disorders: A review. *Int J Eat Disord*. 2002;32:253-270.
- 8 81. Kaye W, Bulik CM, Thonton L, et al. Anxiety disorders comorbid with bulimia and
9 anorexia nervosa. *Am J Psychiatry*. 2004;161:2215-2221.
- 10 82. Strober M. The association of anxiety disorders and obsessive compulsive personality
11 disorder with anorexia nervosa: evidence from a family study with discussion of
12 nosological and neurodevelopmental implications. *Int J Eat Disord*. 2007;40:S46-S51.
- 13 83. Bulik CM, Sullivan PF, Fear J, Joyce PR. Eating disorders and antecedent anxiety
14 disorders: a controlled study. *Acta Psychiatrica Scandinavica*. 1997;96:101-107.
- 15 84. Godart NT, Flament MF, Lecrubier Y, Jeammet P. Anxiety disorders in anorexia nervosa
16 and bulimia nervosa: comorbidity and chronology of appearance. *European Psychiatry*.
17 2000;15:38-45.
- 18 85. Silberg J, Bulik CM. The developmental association between eating disorder symptoms
19 and symptoms of depression and anxiety in juvenile twin girls. *Journal of Child
20 Psychology and Psychiatry*. 2005;46:1317-1326.
- 21 86. Eisenberg M, Wall M, Neumark-Sztainer D. Muscle enhancing behaviors among
22 adolescent girls and boys. *Pediatrics*. 2012;130:1019-1026.
- 23 87. Le Grange D, Lock J. Bulimia nervosa in adolescents: Treatment, eating pathology, and
24 comorbidity. *South African Psychiatry Review*. 2002;August:19-22.
- 25 88. Le Grange D, Loeb KL, Orman S, Jellar C. Bulimia nervosa: a disorder in evolution?
26 *Archives of Pediatrics and Adolescent Medicine*. 2004;158:478-482.
- 27 89. Eisenberg M, Neumark-Sztainer D, Paxton S. Five-year change in body satisfaction
28 among adolescents. *Journal of Psychosomatic Research*. 2004;61:521-527.
- 29 90. Carlat DJ, Camargo CA, Jr., Herzog DB. Eating disorders in males: a report on 135
30 patients. *Am J Psychiatry*. 1997;154(8):1127-1132.
- 31 91. Russell C, Keel P. Homosexuality as a specific risk factor for eating disorders in men. *Int
32 J Eat Disord*. 2002;31:300-306.
- 33 92. Marcus M, Kalarchian M. Binge eating in children and adolescents. *Int J Eat Disord*.
34 2003;34 Suppl:S47-57.
- 35 93. Fairburn CG, Cooper Z, Doll H, Norman P, O'Connor M. The natural course of bulimia
36 nervosa and binge eating disorder in young women. *Archives of General Psychiatry*.
37 2000;57:659-665.
- 38 94. Wilson GT, Fairburn CG, Agras WS, Walsh BT, Kraemer H. Cognitive behavior therapy
39 for bulimia nervosa: time course and mechanism of change. *J Clin Consulting
40 Psychology*. 2002;70:267-274.
- 41 95. Fairburn CG, Welch SL, Doll HA, Davies BA, O'Connor ME. Risk factors for bulimia
42 nervosa. A community-based case-control study. *Arch Gen Psychiatry*. 1997;54(6):509-
43 517.
- 44 96. Jacobi C, Hayward C, de Zwaan M, Kraemer H, Agras W. Coming to terms with risk
45 factors for eating disorders: application of risk terminology and suggestions for a general
46 taxonomy. *Psychological Bulletin*. 2004;130:19-65.

- 1 97. Crowther C, Arney M, Luce K, Dalton G, Leahey T. The point prevalence of bulimia
2 nervosa from 1990-2004. *Int J Eat Disord.* 2008;41:491-497.
- 3 98. Flament M, Ledoux S, Jeammet P, Choquet M, Simon Y. A population study of bulimia
4 nervosa and subclinical eating disorders in adolescence. In: Steinhausen H, ed. *Eating*
5 *Disorders in Adolescence: Anorexia and Bulimia Nervosa.* New York: Brunner/Mazel;
6 1995:21-36.
- 7 99. Fairburn CG, Beglin SJ. Studies of the epidemiology of bulimia nervosa. *Am J*
8 *Psychiatry.* 1990;147(4):401-408.
- 9 100. Ahs F, Furmark T, Michelgard A, et al. Hypothalamic blood flow correlates positively
10 with stress-induced cortisol levels in subjects with social anxiety disorder. *Psychosomatic*
11 *Medicine.* 2006;68:859-862.
- 12 101. Stice E, Agras WS. Predicting onset and cessation of bulimic behaviors during
13 adolescence: A longitudinal grouping analysis*. *Behavior Therapy.* 1998;29(2):257-276.
- 14 102. Strober M, Humphrey L. Family contributions to the etiology and course of anorexia
15 nervosa and bulimia nervosa. *J Consulting Clin Psychol.* 1987;55:654-659.
- 16 103. Bulik C, Sullivan PF, Wade T, Kendler KS. Twin studies of eating disorders: a review.
17 *IJED.* 2000;27:1-20.
- 18 104. Fairburn CG, Cowen PJ, Harrison PJ. Twin studies and the etiology of eating disorders.
19 *Int J Eat Disord.* 1999;26(4):349-358.
- 20 105. Bulik CM, Sullivan PF, Kendler KS. Heritability of binge eating and broadly defined
21 bulimia nervosa. *Biological Psychiatry.* 1998;44:1210-1218.
- 22 106. Welch SL, Fairburn CG. Sexual abuse and bulimia nervosa: three integrated case control
23 comparisons. *Am J Psychiatry.* 1994;151(3):402-407.
- 24 107. Fairburn CG, Doll HA, Welch SL, Hay PJ, Davies BA, O'Connor ME. Risk factors for
25 binge eating disorder: a community-based, case-control study. *Arch Gen Psychiatry.*
26 1998;55(5):425-432.
- 27 108. de Zwaan M, Mitchell JE, Seim HC, et al. Eating related and general psychopathology in
28 obese females with binge eating disorder. *Int J Eat Disord.* 1994;15(1):43-52.
- 29 109. Dansky B, Brewerton T, O'Neil P, Kilpatrick D. The National Women's Study:
30 Relationship of crime victimization and PTSD and Eating Disorders. *IJED.* 1997;21:213-
31 228.
- 32 110. Brewerton T. Eating disorders, trauma, and comorbidity: Focus on PTSD. *Eating*
33 *Disorders.* 2007;15:285-304.
- 34 111. Mitchell K, Mazzeo S, Schlesinger M, Brewerton T, Smith B. Comorbidity and partial
35 and subthreshold PTSD in men and women with eating disorders in the National
36 Comorbidity Survey-Replication Study. *IJED.* 2012;45:307-315.
- 37 112. Herzog DB, Keller MB, Sacks NR, Yeh CJ, Lavori PW. Psychiatric comorbidity in
38 treatment-seeking anorexics and bulimics. *J Am Acad Child Adolesc Psychiatry.*
39 1992;31(5):810-818.
- 40 113. Fairburn CG. *Cognitive Behavioral Therapy and Eating Disorders.* New York: Guilford;
41 2008.
- 42 114. Wilfley D, Schwartz M, Spurrell E, Fairburn C. Using the Eating Disorder Examination
43 to identify the specific psychopathology of binge eating disorder. *IJED.* 2000;27:259-
44 269.
- 45 115. Striegel-Moore RH, Wilson GT, Wilfley DE, Elder KA, Brownell KD. Binge eating in an
46 obese community sample. *Int J Eat Disord.* 1998;23(1):27-37.

- 1 116. Stice E, Presnell K, Spangler D. Risk factors for binge eating onset in adolescent girls: A
2 2-year prospective investigation. *Health Psychology*. 2002;21:131-138.
- 3 117. Glasofer D, Tanofsky-Kraff M, Eddy K, et al. Binge eating in overweight treatment
4 seeking adolescents. *Journal of Pediatric Psychology*. 2007;32:95-105.
- 5 118. Wilfley D, Friedman M, Douchis J, et al. E. Comorbid psychopathology in binge eating
6 disorder: relation to eating disorder severity at baseline and following treatment. *Journal
7 of Consulting & Clinical Psychology*. 2000;68:641-649.
- 8 119. Bryant-Waugh R, Markham L, Kreipe R, Walsh BT. Feeding and eating disorders in
9 childhood. *IJED*. 2010;43:98-111.
- 10 120. Kreipe R, Palomaki A. Beyond picky eating: Avoidant/Restrictive Food Intake Disorder.
11 *Curr Psychiatry Rep*. 2012;14:421-431.
- 12 121. Nicholls D, Randall D, Lask B. Selective eating: Symptom disorder or normal variant?
13 *Clin Child Psychology and Psychiatry*. 2001;6:257-270.
- 14 122. Bryant-Waugh R, Nicholls D, eds. *Diagnosis and classification of disordered eating in
15 childhood*. New York: Guilford Press; 2011. Le Grange D, Lock J.. In (eds): . . . , eds.
- 16 123. Faith M, ed *Development of child taste and food preferences: The role of exposure*. New
17 York: Oxford Press; 2010. Agras WS, ed. *The Oxford Handbook of Eating Disorders*.
- 18 124. Nichols J, Rauh M, Lawson M, Barkai H. Prevalence of the female athlete triad
19 syndrome among high school athletes. *Arch Pediatr Adolesc Med*. 2006;160:137-142.
- 20 125. Thein-Nissenbaum J, Carr K. Female athlete triad syndrome in the high school athlete.
21 *Phys Ther Sport*. 2011; August 12(3):108-115.
- 22 126. Garner DM, Olmsted M, Bohr Y, Garfinkel P. The Eating Attitudes Test: Psychometric
23 features and clinical correlates. *Psychol Med*. 1982;12:871-878.
- 24 127. Garner D, Olmstead M, Polivy J. Development and validation of a multidimensional
25 eating disorder inventory for anorexia nervosa and bulimia. *IJED*. 1983;2:15-34.
- 26 128. Passi V, Bryson S, Lock J. Assessment of eating disorders in adolescents with anorexia
27 nervosa: Self-report versus interview. *Int J Eat Disord*. 2003;33:45-54.
- 28 129. Childress A, Brewerton T, Hodges E, Jarrell M. The Kids Eating Disorder Survey
29 (KEDS): A study of middle school students. *J Am Acad Child Adolesc Psychiatry*.
30 1993;32:843-850.
- 31 130. Lock J, Peebles R. Predictors of Serious Medical Complications in Adolescent Females
32 Hospitalized for Eating Disorders. Paper presented at: Academy of Eating
33 Disorders 2010; Salzburg, Austria.
- 34 131. Control CfD. *CDC Growth Charts for the United States: Development and Methods*.
35 Atlanta, GA: Center for Disease Control; 2002.
- 36 132. Cooper Z, Fairburn CG. The Eating Disorder Examination: A semi-structured interview
37 for the assessment of the specific psychopathology of eating disorders. *International
38 Journal of Eating Disorders*. 1987;6:1-8.
- 39 133. Cooper Z, Cooper PJ, Fairburn CG. The validity of the eating disorder examination and
40 its subscales. *British Journal of Psychiatry*. 1989;154:807-812.
- 41 134. Bryant-Waugh R, Cooper P, Taylor C, Lask B. The use of the Eating Disorders
42 Examination with children: A pilot study. *Int J Eat Disord*. 1996;19:391-397.
- 43 135. Fairburn CG, Beglin SJ. Assessment of eating disorders: interview or self-report
44 questionnaire? *Int J Eat Disord*. 1994;16(4):363-370.
- 45 136. Binford R, Le Grange D, Jellar C. EDE and adolescent bulimia nervosa: Interview or
46 self-report? *Int J Eat Disord*. 2005;37:44-49.

- 1 137. Maloney M, McQuire J, Daniels S. The reliability testing of a children's version of the
2 Eating Attitudes Test. *J Am Acad Child Adolesc Psychiatry*. 1988(27):541-543.
- 3 138. Garner DM, Olmsted M, Polivy J. Development and validation of a multidimensional
4 eating disorder inventory for anorexia nervosa and bulimia. *Int J Eat Disord*. 1983;2:15-
5 34.
- 6 139. Anderson D, Lundgren J, Shapiro J, Paulosky C. Assessment of eating disorders.
7 *Behavior Modification*. 2004;28:763-782.
- 8 140. Katzman D. Medical Complications in Adolescents with Anorexia Nervosa: A Review of
9 the Literature. *Int J Eat Disord*. 2005;37:S52-S59.
- 10 141. Golden NH. Eating disorders in adolescence and their sequelae. *Best Pract Res Clin*
11 *Obstet Gynaecol*. Feb 2003;17(1):57-73.
- 12 142. Rome ES, Ammerman S. Medical complications of eating disorders: an update. *J Adolesc*
13 *Health*. Dec 2003;33(6):418-426.
- 14 143. Fichter MM, Quadflieg N. Six-year course of bulimia nervosa. *Int J Eat Disord*. Dec
15 1997;22(4):361-384.
- 16 144. Crow S, Peterson C, Swanson S, et al. Increased mortality in bulimia nervosa and other
17 eating disorders. *Am J Psychiatry*. 2009;166:1342-1346.
- 18 145. Pediatrics AAO. Policy Statement: Identifying and treating eating disorders. *Pediatrics*.
19 2003;111:204-211.
- 20 146. Pediatrics AAO. Identifying and Treating eating disorders: Policy statement. *Pediatrics*.
21 2003;111:204-211.
- 22 147. Robb A, Silber T, Orrell-Valente J, et al. Supplemental Nocturnal Nasogastric Refeeding
23 for Better Short-Term Outcome in Hospitalized Adolescent Girls With Anorexia
24 Nervosa. *American Journal of Psychiatry*. 2002;159:1347-1353.
- 25 148. Silber T, Robb A, Orrell-Valente J, Ellis N, Valadez-Meltzer A, Dadson M. Nocturnal
26 nasogastric refeeding for hospitalized boys with anorexia nervosa. *Developmental and*
27 *Behavioral Pediatrics*. 2004;25:415-418.
- 28 149. Crisp AH, Norton K, Gowers S, et al. A controlled study of the effect of therapies aimed
29 at adolescent and family psychopathology in anorexia nervosa. *Br J Psychiatry*.
30 1991;159:325-333.
- 31 150. Gowers S, Clark A, Roberts C, et al. Clinical effectiveness of treatments for anorexia
32 nervosa in adolescents. *Br J Psychiatry*. 2007;191:427-435.
- 33 151. Olmsted M. Day hospital treatment of anorexia nervosa and bulimia nervosa. In: Fairburn
34 CG, Brownell K, eds. *Eating Disorders and Obesity: A Comprehensive Review*. New
35 York: Guilford Press; 2002:330-334.
- 36 152. Gerlinghoff M, Backmund H, Franzen U. Evaluation of a day treatment programme for
37 eating disorders. *European Eating Disorders Review*. 1998;6:96-106.
- 38 153. Brewerton T, Costin C. Treatment results of anorexia nervosa and bulimia nervosa in a
39 residential treatment program. *Eating Disorders*. 2011;19:117-131.
- 40 154. Brewerton T, Costin C. Long-term outcome of residential treatment for anorexia nervosa
41 and bulimia nervosa. *Eating Disorders*. 2011;19:132-144.
- 42 155. Lock J. *The Oxford Handbook of Child and Adolescent Eating Disorders: Developmental*
43 *Perspectives*. New York, New York: Oxford University Press; 2012.
- 44 156. Hall A, Crisp AH. Brief psychotherapy in the treatment of anorexia nervosa: Outcome at
45 one year. *Br J Psychiatry*. 1987;151:185-191.

- 1 157. Pike K, Walsh BT, Vitousek K, Wilson GT, Bauer J. Cognitive-Behavioral Therapy in
2 the Posthospitalization Treatment of Anorexia Nervosa. *American Journal of Psychiatry*.
3 2004;160:2046-2049.
- 4 158. Berkman N, Lohr K, Bulik C. Outcomes of eating disorders: A systematic review of the
5 literature. *IJED*. 2007;40:283-309.
- 6 159. Hay P. A systematic review of evidence for psychological treatments in eating disorders:
7 2005-2012. *IJED*. 2013;46:462-469.
- 8 160. Russell GF, Szmukler GI, Dare C, Eisler I. An evaluation of family therapy in anorexia
9 nervosa and bulimia nervosa. *Arch Gen Psychiatry*. 1987;44(12):1047-1056.
- 10 161. Le Grange D, Eisler I, Dare C, Russell G. Evaluation of family treatments in adolescent
11 anorexia nervosa: a pilot study. *Int J Eat Disord*. 1992;12(4):347-357.
- 12 162. Eisler I, Dare C, Hodes M, Russell G, Dodge E, Le Grange D. Family therapy for
13 adolescent anorexia nervosa: the results of a controlled comparison of two family
14 interventions. *J Child Psychol Psychiatry*. 2000;41(6):727-736.
- 15 163. Lock J, Agras WS, Bryson S, Kraemer H. A comparison of short- and long-term family
16 therapy for adolescent anorexia nervosa. *Journal of the American Academy of Child and*
17 *Adolescent Psychiatry*. 2005;44:632-639.
- 18 164. Lock J, Le Grange D, Agras WS, Moye A, Bryson S, Jo B. A randomized clinical trial
19 comparing family based treatment to adolescent focused individual therapy for
20 adolescents with anorexia nervosa. *Archives of General Psychiatry*. 2010;67:1025-1032.
- 21 165. Robin A, Siegal P, Moye A, Gilroy M, Dennis A, Sikand A. A controlled comparison of
22 family versus individual therapy for adolescents with anorexia nervosa. *J Am Acad Child*
23 *Adolesc Psychiatry*. 1999;38(12):1482-1489.
- 24 166. Eisler I. The empirical and theoretical base of family therapy and multiple family day
25 therapy for adolescent anorexia nervosa. *Journal of Family Therapy*. 2005;27:104-131.
- 26 167. Wallis A, Rhodes P, Kohn M, Madden S. Five-years of family based treatment for
27 anorexia nervosa: the Maudsley Model at the Children's Hospital at Westmead. *Int J*
28 *Adolesc Med Health*. 2007;19:277-283.
- 29 168. Fitzpatrick K, Moye A, Hostee R, Le Grange D, Lock J. Adolescent Focused Therapy for
30 Adolescent Anorexia Nervosa. *Journal of Contemporary Psychotherapy*. 2010;40:31-39.
- 31 169. Byford S, Barrett B, Roberts C, et al. Economic evaluation of a randomised controlled
32 trial for anorexia nervosa in adolescents. *Br J Psychiatry*. 2007;191:436-440.
- 33 170. Dalle Grave R, Calugi S, Doll H, Fairburn C. Enhanced cognitive behavioral therapy for
34 adolescents with anorexia nervosa: An alternative to family therapy? *Behav Res Ther*.
35 2013;51:R9-R12.
- 36 171. Schmidt U, Lee S, Beecham J, et al. A randomized controlled trial of family therapy and
37 cognitive behavior therapy guided self-care for adolescents with bulimia nervosa and
38 related conditions. *Am J Psychiatry*. 2007;164:591-598.
- 39 172. Lock J. Adjusting Cognitive Behavioral Therapy for Adolescent Bulimia Nervosa:
40 Results of a Case Series. *American Journal of Psychotherapy*. 2005;59:267-281.
- 41 173. Le Grange D, Crosby RD, Rathouz PJ, Leventhal BL. A randomized controlled
42 comparison of family-based treatment and supportive psychotherapy for adolescent
43 bulimia nervosa. *Archives of General Psychiatry*. 2007;64(9):1049.
- 44 174. Fairburn CG. Interpersonal psychotherapy for bulimia nervosa. In: Garner DM, Garfinkel
45 P, eds. *Handbook of Treatment for Eating Disorders, Second Edition*. New York:
46 Guilford Press; 1997.

- 1 175. Safer DL, Telch C, F., Agras WS. Dialectical behavior therapy for bulimia nervosa. *Am J*
2 *Psychiatry*. 632-634 2001(158).
- 3 176. Agras WS, Walsh BT, Fairburn CG, Wilson GT, Kraemer HC. A multicenter comparison
4 of cognitive-behavioral therapy and interpersonal psychotherapy for bulimia nervosa.
5 *Archives of General Psychiatry*. 2000;57:459-466.
- 6 177. Mufson L, Weissman MM, Moreau D, Garfinkel R. Efficacy of interpersonal
7 psychotherapy for depressed adolescents. *Arch Gen Psychiatry*. 1999;56:573-579.
- 8 178. Safer D, Couturier J, Lock J. Dialectical Behavior Therapy modified for adolescent binge
9 eating disorders: a case report. *Cognitive Behavioral Practice*. 2007;14:157-167.
- 10 179. Federici A, Wisniewski L, Ben-Porath D. Description of an intensive dialectical behavior
11 therapy program for multidagnostic clients with eating disorders. *Journal of Counseling*
12 *and Development*. 2012;90:330-338.
- 13 180. Safer DL, Robinson AH, Jo B. Outcome from a Randomized Controlled Trial of Group
14 Therapy for Binge Eating Disorder: Comparing Dialectical Behavior Therapy Adapted
15 for Binge Eating to an Active Comparison Group Therapy. *Behavior Therapy*.
16 2010;41:106-120.
- 17 181. Attia E, Mayer L, Killory E. Medication response in the treatment of patients with
18 anorexia nervosa. *Journal of Psychiatric Practice*. 2001;7:157-162.
- 19 182. Walsh BT, Kaplan AS, Attia E, et al. Fluoxetine after weight restoration in anorexia
20 nervosa: a randomized clinical trial. *JAMA*. 2006;295:2605-2612.
- 21 183. La Via M, Gray J, WH K. Case reports of olanzapine treatment of anorexia nervosa. *Int J*
22 *Eat Disord*. 2000;27:363-366.
- 23 184. Bissada H, Tasca G, Barber A, Bradwejn J. Olanzapine in the treatment of low body
24 weight and obsessive thinking in women with anorexia nervosa: a randomized, double-
25 blind, placebo controlled trial. *Am J Psychiatry*. 2008;165:1281-1288.
- 26 185. Mondraty N, Birmingham C, Touys W, Sundakov V, Chapman L, Beumont P.
27 Randomized controlled trial of olanzapine in the treatment of cognitions in anorexia
28 nervosa. *Australas Psychiatry*. 2005;13:72-75.
- 29 186. Kafantaris V, Leigh E, Hertz S. A placebo controlled pilot study of adjunctive olanzapine
30 for adolescents with anorexia nervosa. *Journal of Child and Adolescent*
31 *Psychopharmacology*. 2011;21:207-2012.
- 32 187. Brewerton T. Anti-psychotic agents in the treatment of anorexia nervosa:
33 Neuropsychopharmacologic rationale and evidence from controlled trials. *Current*
34 *Psychiatry Reports*. 2012;14:398-405.
- 35 188. Norris R, Spettigue W, Buchholz A, Henderson K, Obeid N. Factors influencing research
36 drug trials in adolescents with anorexia nervosa. *Eating Disorders*. 2010;18:210-217.
- 37 189. Hagman J, Gralla J, Sigel E, et al. A double-blind, placebo-controlled study of
38 risperidone for the treatment of adolescents and young adults with anorexia nervosa: a
39 pilot study. *American Journal of Child and Adolescent Psychiatry*. 2011;50:915-924.
- 40 190. Court A, Mulder C, Kerr M, et al. Investigating the effectiveness, safety, and tolerability
41 of quetiapine for anorexia nervosa in young people: a pilot study. *Journal of Psychiatric*
42 *Research*. 2010;44:1027-1034.
- 43 191. Kotler L, Devlin B, Davies M, Walsh BT. An open trial of fluoxetine in adolescents with
44 bulimia nervosa. *Journal of Child and Adolescent Psychopharmacology*. 2003;13:329-
45 325.

- 1 192. Mitchell JE, Raymond N, Specker S. A review of the controlled trials of
2 pharmacotherapy and psychotherapy in the treatment of bulimia nervosa. *Int J Eat*
3 *Disord.* 1993;14(3):229-247.
- 4 193. Meehan K, Loeb K, Roberto C, Attia E. Mood change during weight restoration in
5 patients with anorexia nervosa. *IJED.* 2006;39:587-589.
- 6 194. Lock J, Le Grange D, Agras WS, Dare C. *Treatment manual for anorexia nervosa: A*
7 *family-based approach.* New York: Guilford Publications, Inc.; 2001.
- 8 195. Le Grange D, Crosby R, Rathouz P, Leventhal B. A randomized controlled comparison
9 of family-based treatment and supportive psychotherapy for adolescent bulimia nervosa.
10 *Archives of General Psychiatry.* 2007;64:1049-1056.
- 11 196. Rieger E, Van Buren D, Bishop M, Tanofsky-Kraff M, Welch R, Wilfley D. An eating
12 disorder-specific model of interpersonal therapy (IPT-ED): Causal pathways and
13 treatment implications. *Clin Psychol Rev.* 2010;30:400-410.
- 14 197. Group: FBNS. Fluoxetine in the treatment of bulimia nervosa. *Archives of General*
15 *Psychiatry.* 1992;49:139-147.
16
17