

Prescribing Guidelines for Attention Deficit/Hyperactivity Disorder (ADHD)





Prescribing for Attention Deficit/Hyperactivity Disorder

This document was developed by Nationwide Children's Hospital in conjunction with Partners For Kids using evidence-informed clinical guidelines and expert opinion, where evidence is lacking, and is generally reflective of FDA approved indications and recommendations. It is designed to help primary care practitioners provide timely and effective treatment for children with mental health disorders. Information on cost is provided to aid in decision-making when appropriate. This document should not be considered a substitute for sound clinical judgment. Clinicians are encouraged to seek additional information if questions arise, as well as, refer to or consult with specialty behavioral health if therapeutic response is inadequate.

Additional information and resources are available through the Dayton Children's psychiatry team. To speak with one of the psychiatrists, call (937) 641-4385 or visit www.ChildrensDayton.org/Patients-Visitors/Services/Behavioral-Health-3/Programs-and-Services/Psychiatry.

Project ECHO (Extension for Community Healthcare Outcomes) is another resource that providers may enroll in. The Behavioral Health Primary Care ECHO series supports primary care providers in the assessment and management of behavioral health concerns for their patients. Sessions include short didactic presentations on commonly seen conditions in primary care, including depression, anxiety, ADHD, trauma, gender issues, substance use, and eating disorders, followed with behavioral health and pharmacological guidance through interactive, case-based learning.

To register for Project ECHO follow the link below and fill out the form:

https://www.NationwideChildrens.org/for-medical-professionals/education-and-training/echo/series/behavioral-health

For questions regarding enrollment in ECHO, please send an email to the address below:

BHPrimaryCareECHO@NationwideChildrens.org

Attention Deficit/Hyperactivity Disorder Overview

- ADHD is one of the most common pediatric behavioral health disorders affecting 9.4% of children and is characterized by hyperactivity, attention difficulties, and executive function deficits.
- Medication therapy is indicated for patients age 6 years or older. Behavioral therapy is recommended as first line treatment of pre-school aged children with ADHD (ages 4-5 years old).
- Behavioral therapy and, when indicated, academic support, should be considered in conjunction with medication therapy for patients greater than 6 years of age.

Diagnosis of Attention Deficit/Hyperactivity Disorder

- The Vanderbilt Assessment Scales is one of the most commonly used tools to diagnose and monitor ADHD in children and adolescents.
- The assessment asks parents and teachers about the child's behaviors within the past 6 months.
- Scoring exists for each section to reflect diagnosis of ADHD, oppositional-defiant disorder, conduct disorder, mood concerns, academic performance and classroom behavioral performance.
- Providers are encouraged to use the parent and teacher assessment versions for initial diagnosis, and the parent
 and teacher follow-up versions to monitor clinical response and adverse effects associated with medication and use
 clinical findings to optimize medication use for pediatric ADHD.
- Vanderbilt Assessment Scales are found at: https://www.nichq.org/resource/nichq-vanderbilt-assessment-scales

Pharmacogenomic Testing for ADHD

Although tests are available and covered by some insurers, there is limited clinically relevant data to support the use of testing in determining medication and dosage selections, and therefore, not recommended as part of standard care.

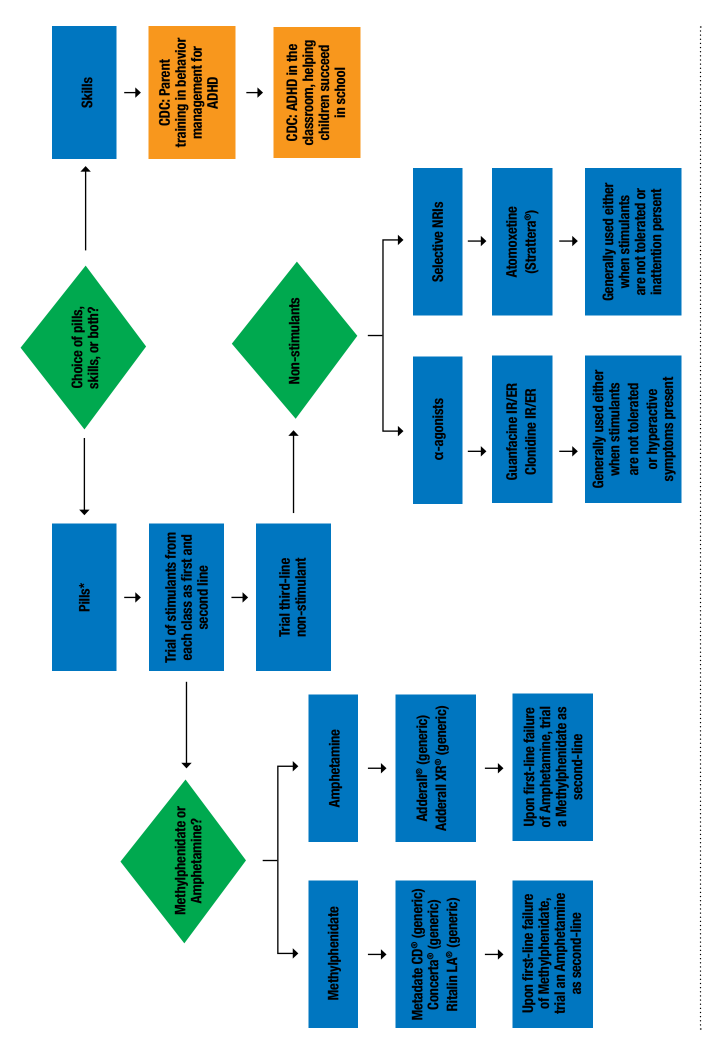
Treating Attention Deficit/Hyperactivity Disorder

Medication

- Long-acting stimulant medications are generally preferred for school-age children.
- Start with a first line medication from the methylphenidate or dextroamphetamine-amphetamine class.
- Maximize dosing of one agent before moving to the next. If ineffective at maximal dosing or side effects develop, switch classes within first line options, then move to second or third line medication, if needed.
- Maximize dosing of long-acting stimulant before adding an immediate release formulation medication.
- Before considering a stimulant medication, obtain cardiac history, including sudden cardiac death in first degree relative under age 50, history of congenital heart defect, or conduction defect.

Behavioral Therapy

- Parent Training in Behavioral Management: Parent Training in Behavior Management for ADHD | CDC
- Classroom Training: ADHD in the Classroom | CDC



Refer to medication charts at the end of the guideline for a listing of preferred and non-preferred agents and clinical pearls, including information regarding alternative formulations such as crushable tablets, capsules to be opened, liquids or patch.

Long-acting Stimulant Conversion Guide

At times, prescribers may need to switch patients from one stimulant to another due to various reasons including patient tolerability and insurance preference/formulary changes. This guide serves as a resource to aid in decision-making for stimulant dose conversions. This guide should not be considered a substitute for clinical judgment, and all patients should be monitored closely for clinical and adverse effects.

General Recommendation:

• Concerta® (methylphenidate ER) and Vyvanse® (lisdexamfetamine) are uniquely dosed. The table below provides an initial dose which may require additional titration.

Dextroamphetamine/ amphetamine ER (Adderall® XR)	Methylphenidate ER (Ritalin® LA or Metadate® CD)	Methylphenidate ER (Concerta®)	Dexmethylphenidate (Focalin XR®)	Lisdexamfetamine (Vyvanse®)	Methylphenidate XR liquid (Quillivant XR®)
N/A	N/A	N/A	N/A	10 mg	N/A
5 mg	10 mg	N/A	5 mg	20 mg	10 mg
10 mg	20 mg	18 mg	10 mg	30 mg	20 mg
15 mg	30 mg	27 mg	15 mg	40 mg	30 mg
20 mg	40 mg	36 mg	20 mg	50 mg	40 mg
25 mg	50 mg	5.4 mg	25 mg	60mg	50 mg
30 mg	60 mg	54 mg	30 mg	70 mg	60 mg

Patient-related Considerations for ADHD Medication							
Patient-related Considerations	Recommendation						
Appetite suppression	 Eat protein rich breakfast prior to administration Schedule meals and provide regular snacks and drinks Monitor height and weight 						
Difficulty swallowing	 Consider alternate medication form: Capsule (refer to medication table to determine which can be opened and sprinkled) Chewable tablet Liquid 						
Insomnia	 If long duration of stimulant action, ensure early morning administration or change to shorter duration stimulant Encourage good sleep hygiene habits 						
Abdominal pain	Take with meals						
Headache	Increase hydration Schedule meals						
Tachycardia and chest pain	 Consider dose reduction Switch to a different stimulant or a non-stimulant Consider cardiology consult with EKG 						
Concern for abuse and/or diversion	Consider a prodrug form of a stimulant, tamper resistant stimulant, or non-stimulant						
Flat affect or mood lability	 Consider dose reduction Switch to a different stimulant or non-stimulant 						

Stimulant and Non-Stimulant Medications for Treatment of Attention Deficit/Hyperactivity Disorder

The table below contains options covered on the Unified Preferred Drug List (UPDL) by Ohio Medicaid Managed Care Plans.

Drug	Initial Daily Dose ¹	Titration Recommendation	Max Daily Dose	Strengths Available	Avg Cost Per Script ²	Clinical Pearls		
	Stimulants							
Dextroamphet- amine-Amphet- amine Immediate Release (Adderall®)	2.5-5 mg	Increase by 2.5-5 mg weekly	40 mg	5; 7.5; 10; 12.5; 15; 20; 30 mg	\$36	3:1 ratio dextro- to levo- amphetamine ratio. ³ Tablet can be crushed. Duration 4-6 hours.		
Dextroamphet- amine-Amphet- amine Long-Act- ing (Adderall XR®)	5-10 mg	Increase by 5-10 mg weekly	30-60 mg	5; 10; 15; 20; 25; 30 mg	\$35	3:1 ratio dextro- to levo-amphetamine ratio. ³ Capsule can be opened and sprinkled. Duration 10-12 hours.		
Methylpheni- date Immediate Release (Ritalin®)	5 mg	Increase by 5-10 mg/day weekly	60 mg	5; 10; 20 mg	\$24	Tablet can be crushed. Duration 4 hours.		
Methylphenidate Long-Acting (Ritalin LA®)	10-20 mg	Increase by 10 mg/ day weekly	60 mg	Brand: 10; 20; 30; 40 mg Generic: 10; 15; 20; 30; 40; 50; 60 mg	\$67	50% is immediate release and 50% is extended release. Capsule can be opened and sprinkled. Duration 8-10 hours.		
Methylphenidate Long-Acting (Concerta®)	18 mg	Increase by 18 mg weekly	54 mg (<13y) 72 mg (≥13y)	18; 27; 36; 54 mg	\$51	22% is immediate release and 78% is extended release. Tablet cannot be crushed. Duration 10-12 hours.		
Methylphenidate Long-Acting (Metadate CD®)	20 mg	Increase by 10-20 mg/day weekly	60 mg	10; 20; 30; 40; 50 mg	\$72	30% is immediate release and 70% is extended release. Capsule can be opened and sprinkled. Duration 8-10 hours.		
Dexmethylpheni- date Long-Acting (Focalin XR®)	0.5 mg	Increase by 5 mg/ day weekly	30 mg	5; 10; 15; 20; 25; 30; 35; 40 mg	\$76	50% is immediate release and 50% is extended release. Capsule can be opened and sprinkled. Duration 10-12 hours. When switching from methylphenidate, reduce dose by half.		
Methylphenidate Long-Acting (Quillivant XR®)	20 mg	Increase by 10-20 mg/day weekly	60 mg	25 mg/ 5mL as 60; 120; 150; 180mL	\$339	Long-acting oral suspension. Duration 12 hours.		

Methylphenidate Long-Acting (QuilliChew ER®)	10-20 mg	Increase by 10,15, or 20 mg/day weekly	60 mg	20; 30; 40 mg	\$447	Long-acting chewable tablet. 30:70 mixture of immediate:delayed release. Duration 8 hours.
Lisdexamfetamine (Vyvanse®)	30 mg	Increase by 10-20 mg/day at 3-7 day intervals	70 mg	Capsule: 10; 20; 30; 40; 50; 60; 70 mg Chewable tab- let: 10; 20; 30; 40; 50; 60 mg	\$402	Pro-drug metabolized to 100% dextroamphetamine. Decreased risk of abuse. Available in capsule and chewable tablet, which are interchangeable on mg-mg basis. Capsule can be opened and dissolved in liquid, then immediately ingested. Duration 10-12 hours.

Non-Stimulants							
Guanfacine Extended Release (Intuniv®)	1 mg	Increase by 1 mg/ day weekly	4 mg	1; 2; 3; 4 mg	\$22	Not equivalent to immediate-release guanfacine. Take at the same time each day. Do not administer with high-fat meal. Tablet cannot be opened or crushed. Monitor blood pressure. Taper when discontinuing.	
Atomoxetine (Strattera®)	<70kg: 0.5 mg /kg >70kg: 40 mg	Increase by 5-10 mg weekly	1.4 mg/ kg or 100 mg	10; 18; 25; 40; 60; 80; 100 mg	\$110	Must be taken daily. Takes 2 weeks to attain maximum efficacy. Cannot be opened or crushed. Black box warning for an increased risk of suicidal ideation; balance risk with clinical need. Bolded warning of liver damage; decrease dose in hepatic impairment.	

Bolded medications are available generically.

¹Dosing is for school-aged children. Medication treatment in preschool-aged children should be considered after a trial of behavioral intervention.

²Cost based on generic drug when available using average 30-day strength and dosing without insurance.

³Contains a combination of d-amphetamine and I-amphetamine. More potent release of dopamine occurs with d-amphetamine, resulting in more symptom reduction for hyperactivity/impulsivity, but more appetite suppression. More potent release of norepinephrine occurs with I-amphetamine, resulting in more symptom reduction for inattention, but less CNS excitation and more cardiovascular adverse effects.

The table below contains information on options not currently covered on the Ohio Medicaid Unified Preferred Drug List (UPDL), and as such, these medications will require a prior authorization or step therapy request.

	Stimulants						
Drug	Initial Daily Dose ¹	Titration Recommendation	Max Daily Dose	Strengths Available	Avg Cost Per Script ²	Clinical Pearls	
Dextroamphetamine- Amphetamine Long-		Increase by 12.5 mg increments weekly	25 mg	12.5; 25; 37.5; 50 mg	\$369	Approved for children 13 years and older. Capsule can be opened and sprinkled. Duration 16 hours.	
Acting (Mydayis®)		increments weekly		30 mg		See package insert for mg conversion to mixed amphetamine salts.	
Dextroamphetamine Extended Release (Dexedrine® Spansule®)	5 mg	Increase by 5mg/day weekly	40 mg	5; 10; 15 mg	\$844	Extended release capsule. Swallow capsule whole. Duration 3-5 hours.	
Dextroamphetamine Immediate Release (Zenzedi®)	5 mg	Increase by 2.5-5mg/ day weekly	40 mg	Brand: 2.5; 5; 7.5; 10; 15; 20; 30 mg Generic: 5; 10 mg	\$264	Immediate release tablet. Can be crushed. Duration 4-6 hours. Generic available in only 5 mg and 10 mg strengths.	
Dextroamphetamine Immediate Release (ProCentra®)	5 mg	Increase by 10 mg/day weekly	40 mg	5 mg/mL	\$304	Short acting oral solution. Duration 4-6 hours.	
Methylphenidate Long-Acting (Aptensio XR®)	10 mg	Increase by 10 mg/day weekly	60 mg	10; 15; 20; 30; 40; 50; 60 mg	\$396	40% is immediate release and 60% is extended release. Capsule can be opened and sprinkled.	
						Duration 8-12 hours.	
Methylphenidate Long-Acting (Cotempla XR-ODT®)	17.3 mg	Increase by 17.3 mg/ day weekly	51.8 mg	8.6; 17.3; 25.9 mg	\$534	Long-acting orally disintegrating tablet. Duration 8-12 hours.	
Methylphenidate Long-Acting (Daytrana®)	10 mg	Increase to next transder- mal patch size no more frequently than every week	30 mg	10; 15; 20; 30 mg	\$504	Transdermal system. Apply for 9 hours. Strength of patch is how much medicine is delivered in a day. Duration 10-12 hours. May cause skin irritation.	
Methylphenidate Long-Acting (Jornay PM®)	20 mg	Increase by 20 mg/day weekly	100 mg	20; 40; 60; 80; 100mg	\$485	Take in the evening between 6:30-9:30pm. If converting from another methylphenidate formulation, discontinue previous formulation and titrate Jornay PM® using initial schedule. Capsules can be opened and sprinkled.	

Non-Stimulants							
Viloxazine (Qelbree®)	6yo- 11yo: 100mg >12yo: 200mg	6-11yo: increase by 100mg/day weekly >12yo: increase by 200mg/day weekly	400 mg	100; 150; 200 mg	\$358	Must be taken daily. Takes 2 weeks to attain maximum efficacy. Cannot be opened or crushed. Black box warn- ing for an increased risk of suicidal ideation; balance risk with clinical need. Bolded warning of liver damage; decrease dose in hepatic impairment.	

Bolded medications are available generically.

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²Cost based on generic drug when available using average 30-day strength and dosing without insurance.

³Contains a combination of d-amphetamine and l-amphetamine. More potent release of dopamine occurs with d-amphetamine, resulting in more symptom reduction for hyperactivity/impulsivity, but more appetite suppression. More potent release of norepinephrine occurs with l-amphetamine, resulting in more symptom reduction for inattention, but less CNS excitation and more cardiovascular adverse effects.

Note: Drug information is compiled from data at Lexicomp Online, online.lexi.com. Prices are references and actual cost may vary based on drug strength, quantity and other factors. Average cost per script is based on tablet and/or capsule formulations for a one-month supply. In general, liquid formulations are more expensive and not as widely covered compared to tablet and capsules. Please refer to the specific medication's package insert for the most up to date information.

References:

- 1. Subcommittee on Attention-Deficit/Hyperactivity Disorder, Steering Committee on Quality Improvement and Management; ADHD: Clinical Practice Guideline for the Diagnosis, Evaluation, and Treatment of Attention-Deficit/ Hyperactivity Disorder in Children and Adolescents. Pediatrics November 2011; 128 (5): 1007–1022. 10.1542/ peds.2011-2654
- 2. Parent Training in Behavioral Management for ADHD. Parent Training in Behavior Management for ADHD | CDC. 01/14/2022
- 3. ADHD in the Classroom: Helping Children Succeed in School. ADHD in the Classroom | CDC. 01/14/2022

Partners For Kids is the oldest and largest pediatric accountable care organization in the United States. It was founded 25 years ago by Nationwide Children's Hospital and has improved the health of millions of children in south central and southeastern Ohio.

PartnersForKids.org



