



WELCOME to the Obesity Care in All Ages ECHO

Session 1, Why Obesity is a Disease, May 13, 2025



Series Learning Objectives

- Describe obesity as a chronic disease, including evidence-based methods for evaluation and treatment
- Effectively communicate with patients about the health implications of obesity and its available treatment options
- Cultivate skills to effectively assess and treat patients with obesity in various care settings
- Identify when and how to refer patients to appropriate specialized obesity care services



Series Sessions

Date	Session Title
5/13/2025	Why Obesity is a Disease
6/10/2025	Approach to the Patient with Obesity
7/8/2025	Optimizing the Use of Lifestyle-based Obesity Care
8/12/2025	How to Use Anti-Obesity Medications Effectively (GLP-1 agonist)
9/9/2025	How to Use Anti-Obesity Medications Effectively (Non GLP-1 agonist)
9/23/2025	Approach to the Pediatric Patient with Obesity – AAP Clinical Practice Guidelines
10/7/2025	How to Use Endoscopic Therapy Effectively
10/21/2025	Pediatric Anti-Obesity Medications and Bariatric Surgery
11/4/2025	Metabolic-Bariatric Surgery: Who, When, Why, and Which One
11/18/2025	Improving Equitable Access to Obesity Care



Project ECHO (Extension for Community Healthcare Outcomes)

- All teach, all learn.
- ECHO is a telementoring model that uses virtual technology to support casebased learning and to engage the wisdom and experience of all attending.
- Highly Interactive.

Components of ECHO:





Today's Program

- Brief housekeeping
- Didactic: Why Obesity is a Disease Elizabeth Honigsberg, MD, MPH
- Role Play: Sarah Finn, MD and Abbey Berge-Clogston
- Discussion
- Summary
- Up Next



Housekeeping Notes

- Pre course survey: https://redcap.hitchcock.org/redcap/surveys/?s=EA47L8LEDJ43JTDN
- Raise virtual hand or enter comments in chat at any time. We will call on you when it works. Please mute otherwise.
- To protect individual privacy, please use non-identifying information when discussing cases.
- We will be recording the didactic part of these sessions. *Participating in these session is understood as consent to be recorded. Thank you!*
- Closed Captioning will be enabled during sessions
- Questions to ECHO Tech Support thru personal CHAT or ECHO@hitchcock.org



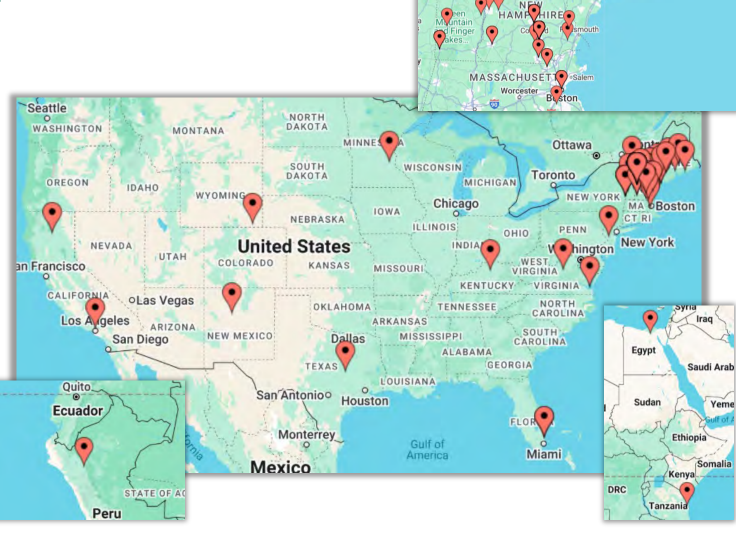
CME/CNE

- One hour of free CME/CNE is available for every session attended, up to 10 sessions.
- Track participation via <u>DH iECHO site</u>
- A link will be provided at the end of the course to submit your attendance and claim your CME/CNE



ECHO Participant Demographics Total Registrants: 193

Professional Identities		
Nurse		
Physician		
Dietitians and Nutritionists		
Administrator		
Behavioral Health Professional		
Physician Assistant/Medical Assistant		
Other healthcare professional		
Pharmacist		
Patient navigator/healthcare educator		
Child Development		



VERMONT

MAIN

Bar Harb



Core Panel

- Abigail Berge-Clogston Program Manager
- Amanda Boyd, MPH Health Coach, Certified Personal Trainer
- Auden McClure, MD, MPH Staff Physician, Pediatric Weight Center
- Charles Brackett, MD, MPH Staff Physician, General Internal Medicine
- Elaine Banerjee, MD, MPH Staff Physician, DH Weight Center
- Elizabeth Honigsberg, MD, MPH Staff Physician, DH Weight Center
- Hannah Brilling, RDN, LD Clinical Dietician
- Kimberly Dovin, MD Staff Physician, DH Weight Center
- Kristin Wheeler, RN Nurse, Weight Center
- Sarah Finn, MD Interim Section Chief, DH Weight Center



Echo Session 1 Why Obesity is a Disease.

Elizabeth Honigsberg MD MPH FACS DABOM May 13th, 2025



I have no financial interests or relationships to disclose.



There are four main objectives for today's discussion.

ASSESS	The current state of the obesity pandemic worldwide
UNDERSTAND	Obesity as a neurobiological/neuroendocrine disease
APPRECIATE	The multitude of factors that lead to the development of obesity
REVIEW	The various criteria for diagnosing the disease of obesity

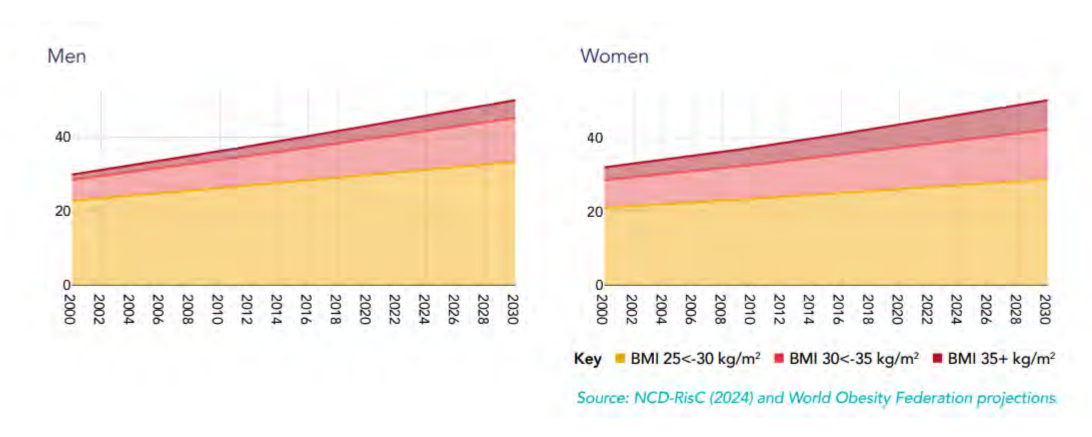


The current state of obesity worldwide.





Rates of men and women (20 years +) living with "high BMI" are increasing worldwide.

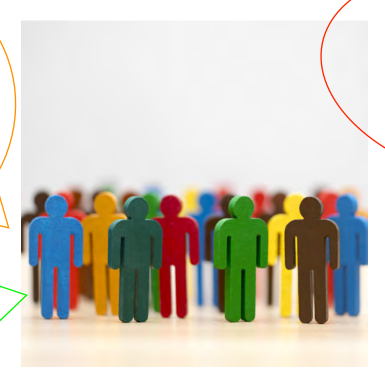




By 2030, THREE BILLION adults will have "high BMI", with 17% of men and 22% of women estimated to have BMI > 30 kg/m² (and the world is NOT prepared).

Yearly: 5
million/41
million adult
deaths
due to NCDs
related to high
BMIs

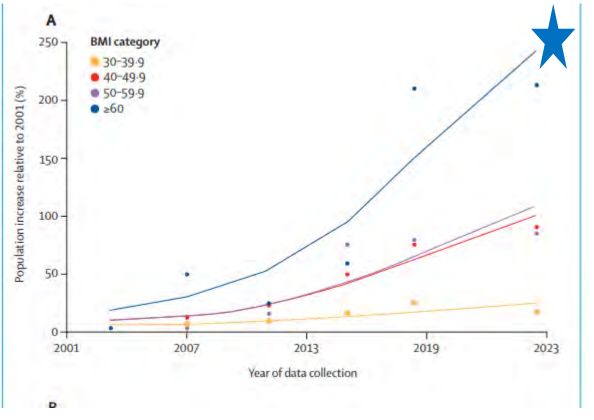
4 million deaths from T2DM, CVA, CAD, CA.



Obesity rates are doubling across the globe, TRIPLING in low-income countries.



In the United States, the prevalence of "normal" weight and overweight has declined since 2001, while all obesity categories have increased over this timeframe.



The largest relative increase of > 200%!



This global systemic failure to slow the obesity pandemic must end.



To do so, we must end:
 The misunderstanding
 The underinvestment
 The fragmentation
 The stigmatization



There is a fundamental misunderstanding about obesity...





THIS DOES <u>NOT</u> CAUSE OBESITY



THIS DOES <u>NOT</u> TREAT OBESITY





Both food intake and fat mass/set point are highly regulated by the brain.







Strikes (environmental factors, gene mutations, etc.) disrupt the regulation of energy balance in brain.



Energetic Homeostasis

Neurons express the orexigenic neuropeptides: AgRP, NPY

Neurons express anorexigenic peptides: POMC, CART -



Cross- talks between systems

Attention Systems Parietal, Visual cortices, Some areas of the frontal cortex Increase food in-take: increased activation to food cues during fed status.



Obesity

Hormones

Ghrelin, GLP-1, insulin, leptin, adiponectin, irisin, et al.



VTA and SN in the midbrain, Nucleus accumbens, Striatum, OFC 1

- Hyper-responsivity to food cues
- Lower availability of D2 receptors

Cognitive control

Mostly in the prefrontal cortex, particularly: Cingulate cortex, Inferior frontal cortex, Pre-SMA, DLPFC

- Inhibitory control
- Food motivation



Inflammation

in brain

Emotion systems

joy, anger, stress, etc.

Amygdala 🁚

Increase appetite: depress, anxiety,

Decrease appetite: fear, sadness,

Memory Systems

Hippocampus, Parahippocampal gyrus

BBB function

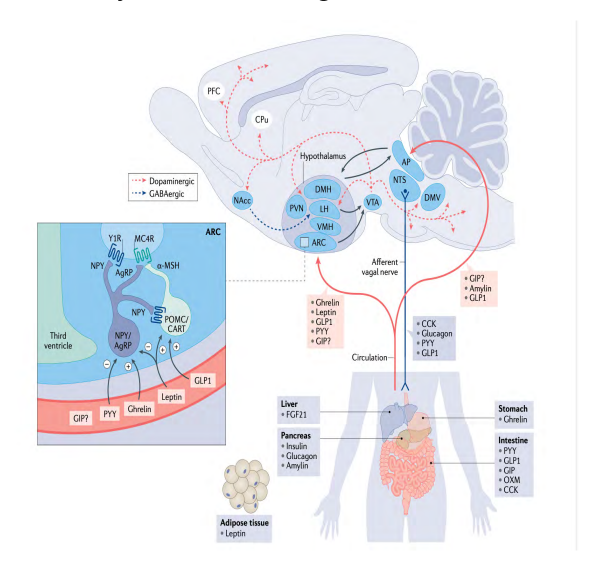
Increase food in-take: hippocampus 🎩

lesions or inflammation.





Various hunger and satiety hormones signal to the brain to affect food intake.





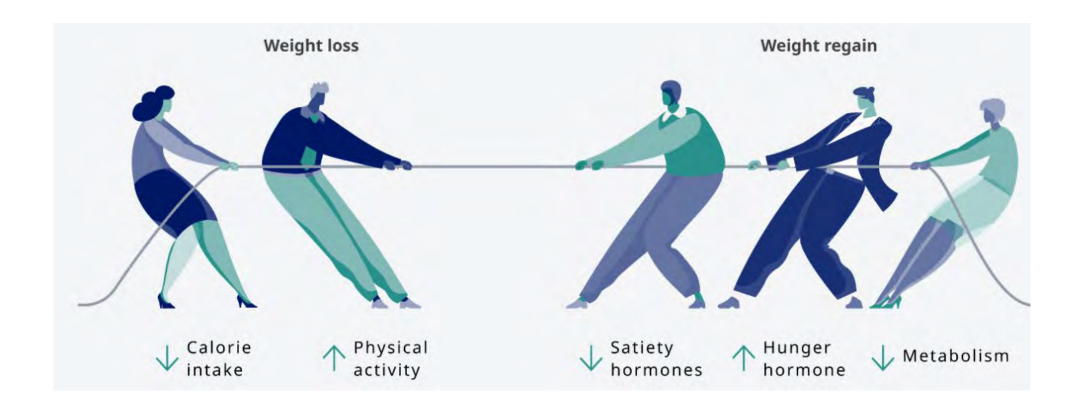
The brain sets AND defends a fat mass (set point) for everyone.



In obesity, that fat mass/set point is abnormally high.



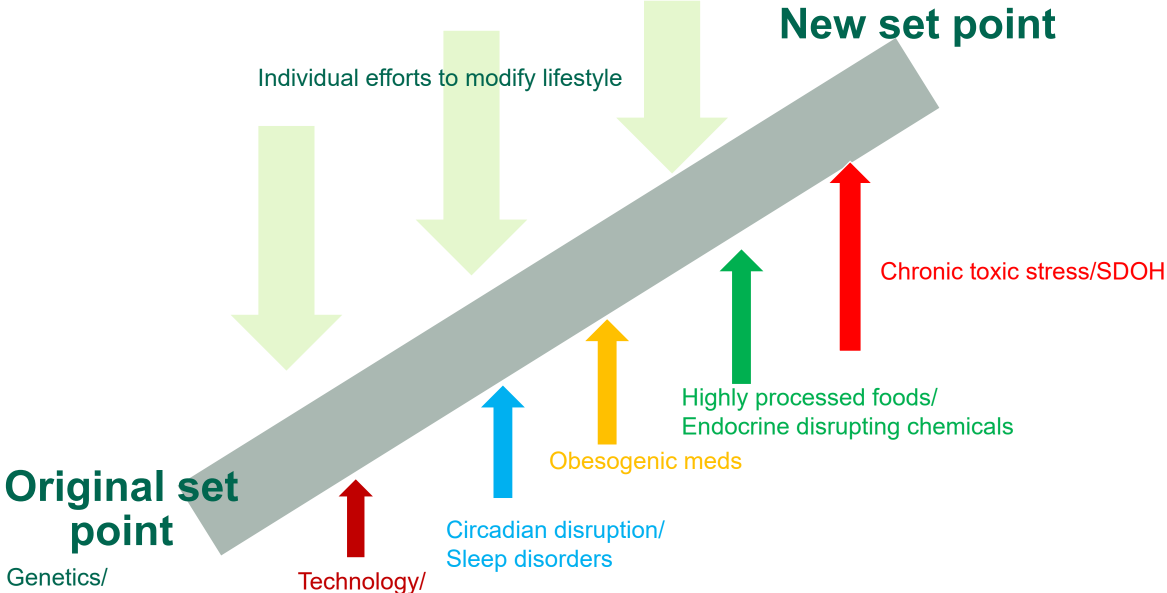
We have metabolically adapted to defend our fat mass.





What drives the development of obesity?





Sedentary lifestyle

Hormonal changes

Slide: adapted from Kaplan, L - Blackburn 2020.



The definition of obesity is evolving as is the diagnostic criteria.



WHO: abnormal or excessive fat accumulation that presents a risk to health.

CDC: BMI $> 30 \text{ kg/m}^2$

Obesity Medicine Association: A chronic, relapsing multi-factorial, neurobehavioral disease, wherein an increase in body fat promotes adipose tissue dysfunction and abnormal fat mass physical forces, resulting in adverse metabolic, biomechanical, and psychosocial health consequences.

The Lancet Commission 2025: provided explicit characterization of the illness intrinsically caused by excess adiposity and establish objective criteria for diagnosis.



OBESITY

Excess fat mass +/- abnormal distribution or function



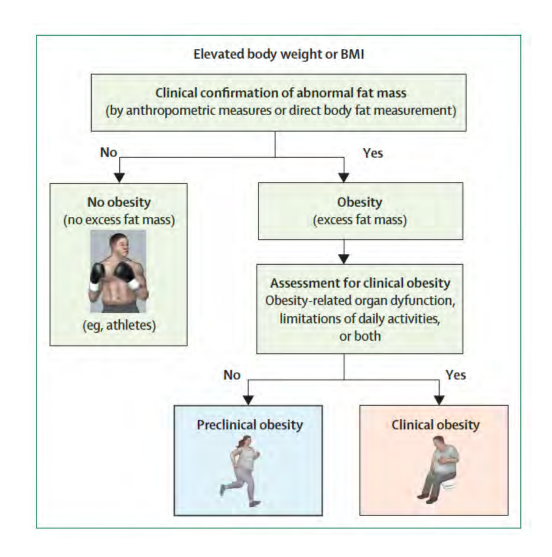
PRECLINICAL OBESITY

At increased risk of developing obesity related organ dysfunction, limitation of daily activities, or both



CLINCAL OBESITY

Chronic systemic illness with dysfunction of the tissues, organs, the entire individual





The objectives for today's session.

ASSESS	The worldwide pandemic of obesity continues to worsen and low and middle income countries are least prepared.
UNDERSTAND	Obesity is a chronic, relapsing neurobiological and neurohormal disease whereby the affected individual CANNOT lower the set point with diet and exercise alone.
APPRECIATE	Both internal and external factors contribute to the development of obesity
REVIEW	Diagnosis still largely relies on BMI, however criteria is changing to reflect the greater importance of metabolic/orthopedic/psychosocial health than BMI alone.



THANK YOU!



Role Play





WELCOME to the Obesity Care in All Ages ECHO

Session 2, Approach to the Patient with Obesity, June 10th, 2025



Today's Program

- Brief housekeeping
- Didactic: Approach to the Patient with Obesity Kimberly Dovin, MD
- Case Discussion
- Summary
- Up Next





APPROACH TO THE PATIENT WITH OBESITY

Kimberly Dovin, MD

Echo Series: Obesity Care in All Ages

Session #2

June 10, 2025



Goals

How to talk to patients about weight

Learn to take an obesity specific history

Understand the evaluation of obesity to identify complications



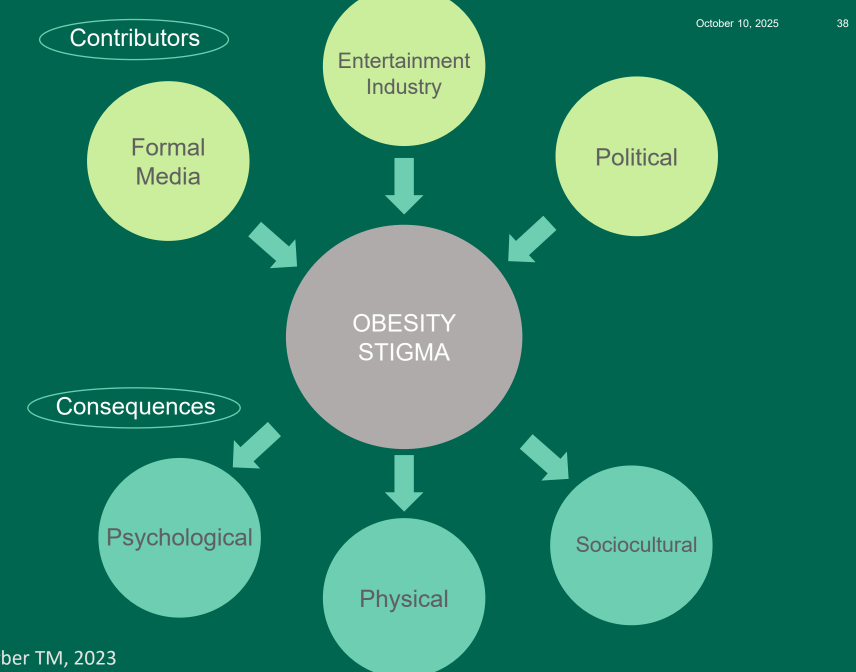
Obesity Stigma and Bias

"Society regularly regards [persons with obesity] not as innocent victims, but as architects of their own ill health, personally responsible for their weight problems because of laziness and overeating."

-Rebecca Puhl and Chelsea Heuer











Obesity Stigma - Medicine



- 2nd only to family in perceived bias
- Less time/discussion
- Less evaluation/screening







Lack of willpower

Laziness

Lack of motivation

Externalized Stigma

- Accusatory language
- Dismissiveness
- Gaps in healthcare

Internalized Stigma

- Disordered eating
- Avoidance of physical activity
- Depression/Anxiety
- Stress response

Weight (re)gain





Evaluation



Starting the Conversation



Take a weight history



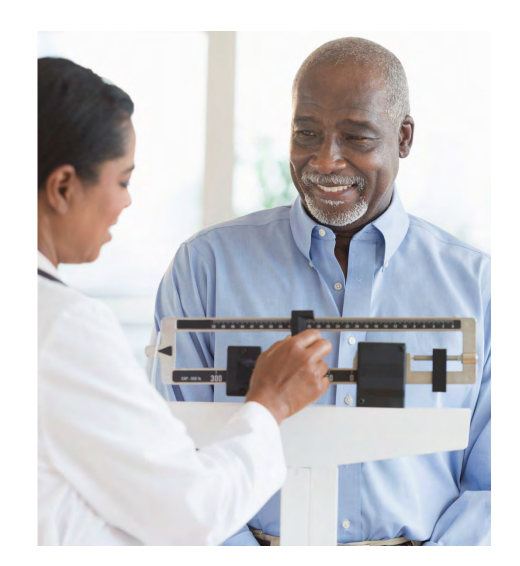
Assess symptoms and signs



Set Goals



(Re-)educate







Weight History

- "What is the story of your weight"
 - Did they have early childhood obesity (<5yo)
 - Stable adult weight?
 - Did they have any large gains and what might have been happening at that time?
 - Has it been gradual through adulthood?
 - How has excess weight impacted their life?





Symptoms of Obesity

- Pervasive thoughts of food
- Excess or no hunger
- Abnormal satiation/satiety
- Craving
- Pain or discomfort
- Difficulty with daily activities due to size
- Fatigue
- SOB
- Low body image







Evaluation

Physical Exam

- Gen: central, gynecoid, generalized adiposity.
- VS, Waist and Neck circumference
- HEENT: Mallampati? Moon facies?
- Neck: buffalo hump,thyroid?
- CV: evidence of arrythmia?
- Abd: hepatomegaly?

- Ext: edema, cuffing?
- Gait: antalgic?
- Skin: acanthosis, hidradenitis, acne, hirsutism, abdominal striae, tender subcutaneous nodules, intertrigo



Evaluation (continued)

Laboratory evaluation

- CBC, CMP
- TSH
- Lipid panel
- FBS, A1c
- Vitamin D

Complications

- Obesogenic medications
- MASLD/MASH Fib4 calculation
- OSA
- Eating disorders
- Contraindications to AOMs







Lipedema

Kruppa P, Georgiou I, et al PMID: 32762835; PMCID: PMC7465366.



1) thickened subcutis, soft, with small, palpable nodules, skin surface still smooth



Classification by stage 2) thickened subcutis, soft, some larger nodules, skin surface uneven



3) thickened subcutis, hardened, with large nodules, disfiguring fat deposition









Classification by morphology I) buttock II) thigh III) entire lower limb

* Type IV is often associated with type II or III.

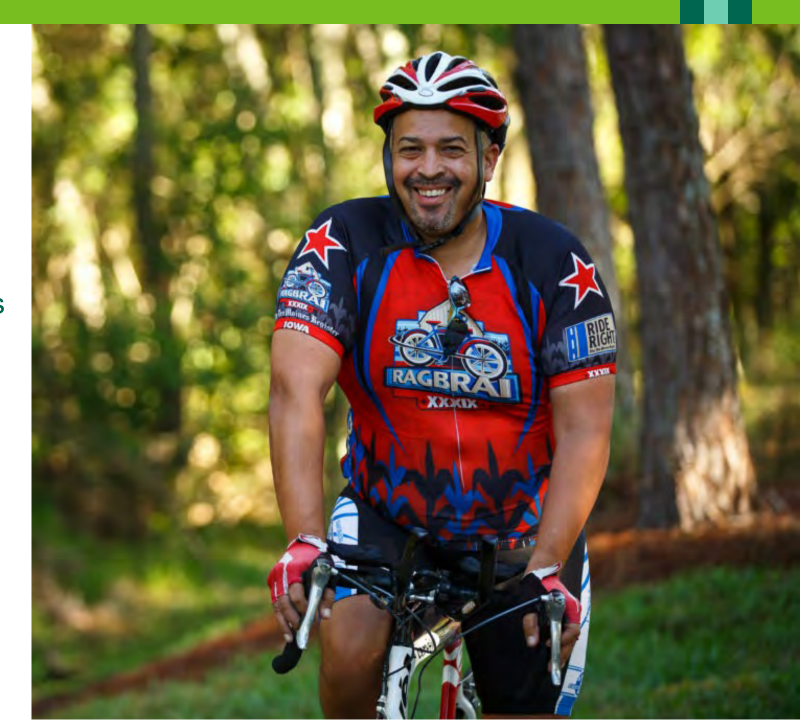
IV) arm*

V) leg



Goals of Treatment

- BMI < 25
- Improvement in complications
- Symptom Resolution
- QOL
- BMI <30?
- BMI >= 23







Summary – Evaluating the Patient with Obesity



Approach patients with compassion



Take a diseasespecific H&P



Set non-scale goals for treatment



- Kruppa P, Georgiou I, Biermann N, Prantl L, Klein-Weigel P, Ghods M. Lipedema-Pathogenesis, Diagnosis, and Treatment Options. Dtsch Arztebl Int. 2020 Jun 1;117(22-23):396-403. doi: 10.3238/arztebl.2020.0396. PMID: 32762835; PMCID: PMC7465366.
- Obesity Medicine Association. Pediatric Obesity Algorithm. https://obesitymedicine.org/resources/obesity-algorithm/. (Accessed = May 31, 2025)
- Pearl RL, Puhl RM, Himmelstein MS, Pinto AM, Foster GD. Weight Stigma and Weight-Related Health: Associations of Self-Report Measures Among Adults in Weight Management. Ann Behav Med. 2020 Nov 1;54(11):904-914. doi: 10.1093/abm/kaaa026. PMID: 32333673; PMCID: PMC7646152.
- Puhl RM, Brownell KD. Confronting and coping with weight stigma: an investigation of overweight and obese adults. Obesity (Silver Spring). 2006 Oct;14(10):1802-15. doi: 10.1038/oby.2006.208. PMID: 17062811.
- Puhl RM, Heuer CA. The stigma of obesity: a review and update. Obesity (Silver Spring). 2009 May;17(5):941-64. doi: 10.1038/oby.2008.636. Epub 2009 Jan 22. PMID: 19165161.
- Westbury, S., Oyebode, O., van Rens, T. *et al.* Obesity Stigma: Causes, Consequences, and Potential Solutions. *Curr Obes Rep* **12**, 10–23 (2023). https://doi.org/10.1007/s13679-023-00495-3





WELCOME to the Obesity Care in All Ages ECHO

Session 3, Optimizing the Use of Lifestyle-based Obesity Care, July 8th, 2025



Today's Program

- Brief housekeeping
- Didactic: Optimizing the Use of Lifestyle-based Obesity Care Shelby Sullivan, MD, FACG, FACG, DABOM
- Case Discussion
- Summary
- Up Next





Optimizing the use of Lifestyle-Based Obesity Care

Shelby Sullivan MD, FACG, FACG, DABOM

Director, Endoscopic Bariatric and Metabolic Program

Dartmouth-Hitchcock Medical Center and Geisel School of Medicine



Disclosure

The following planning committee member(s), speaker(s), author(s) or anyone in a position to control the content for this activity have reported the following financial relationship(s) with ineligible company(ies). All of the relevant financial relationships listed for these individuals have been mitigated.

Sarah Finn, MD ~ was a consultant to Harbor Capital (relationship has ended).



Disclosure:

Shelby Sullivan, MD ~ is a consultant to Allurion, Bioling, Pentax Medical, and Olympus Corporation. She also has grant/research support from Fractyl. Dr. Sullivan was a consultant to Fractyl (relationship has ended) and had grant/research support from Allurion (relationship has ended).

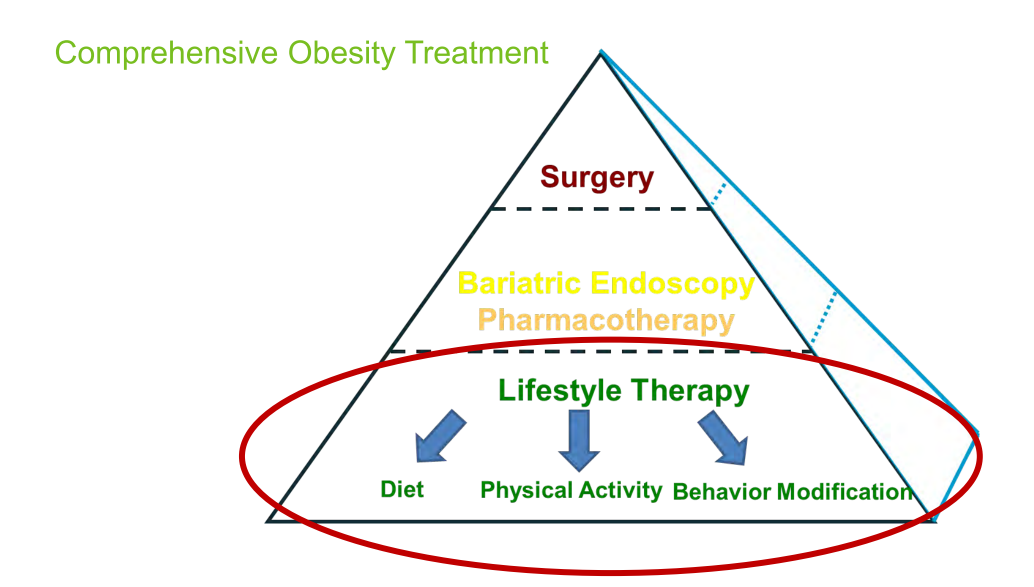
Other planning committee member(s), speaker(s), activity director(s), author(s) or anyone in a position to control the content for this activity have no relevant financial relationship(s) with any ineligible company(ies) to disclose.



Disclosures

- Shelby Sullivan, M.D. has financial interests to disclose.
- Research Support / Grants Last 24 Months
 - Allurion Technologies, Fractyl Laboratories
- Consulting / Employment Last 24 Months
 - Allurion Technologies, Fractyl Laboratories, Biolinq, Pentax, Olympus
 - Notes

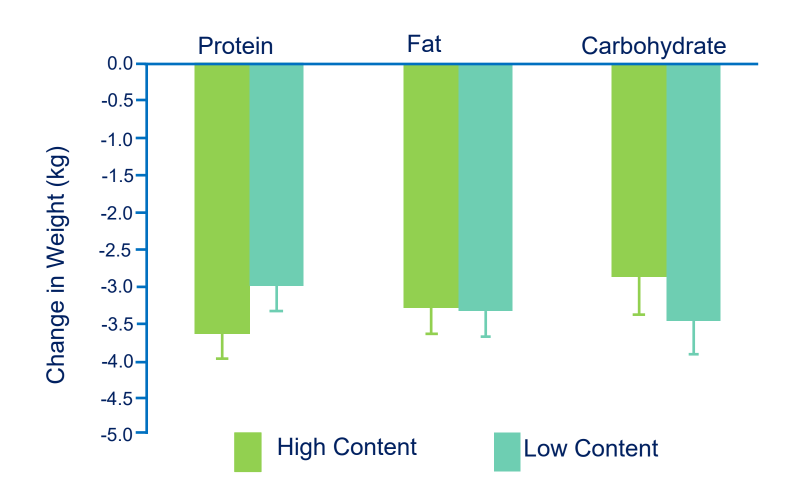






Comparison of Varying Macronutrient Composition: **Pounds Lost** Study

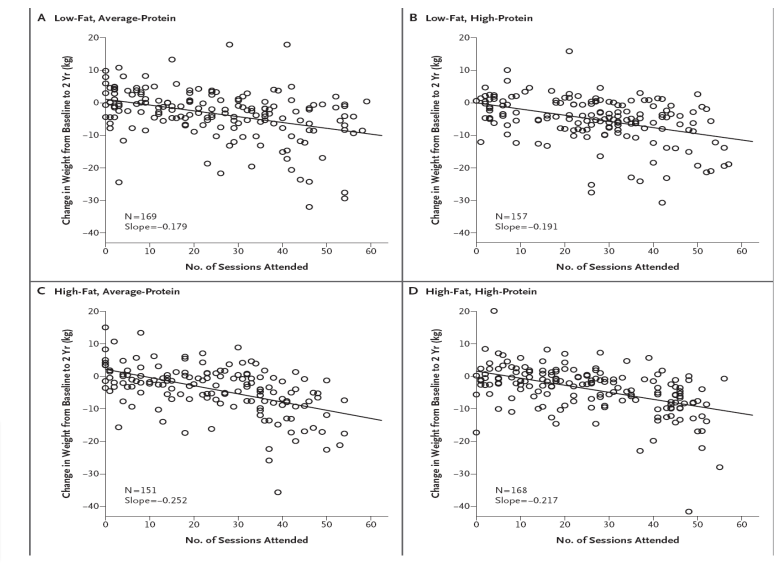
Differing macronutrient composition of the diet did not affect overall weight loss





Comparison of Varying Macronutrient Composition: **Pounds Lost** Study

The number of visits with the study team for lifestyle therapy was directly correlated with weight los





Lifestyle Therapy after Bariatric Surgery

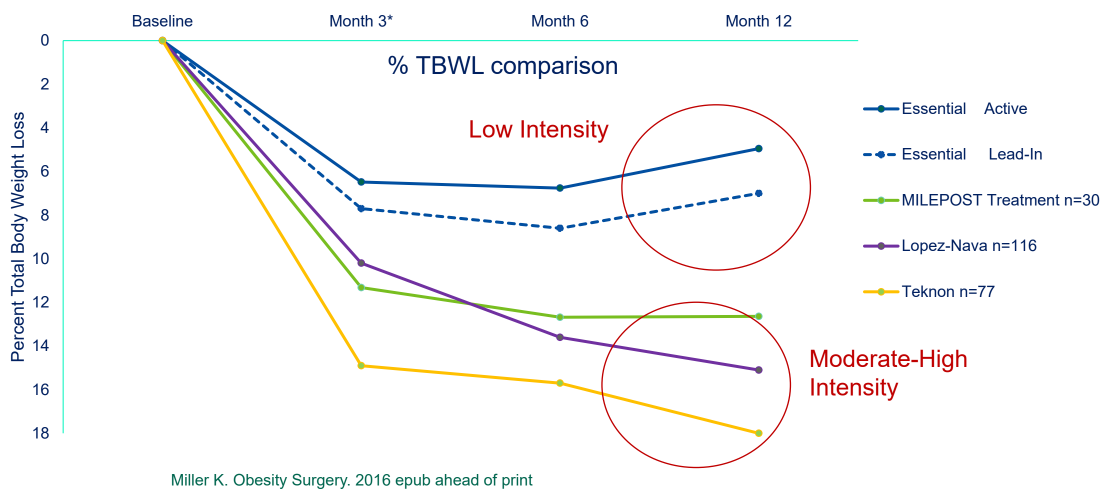
Table 3 Forest plot of standardized mean differences in a random-effects model for percentage of excess weight loss in treatment and control group patients 6–12 months after start of the intervention

Study or subgroup	Intervention			No intervention			Weight	Standardized mean differences IV,	Standardized mean differences IV, random 95% CI			andom,	
	M	SD	Total	M	SD	N		random, 95% CI	95% (1				
Kalarchian et al. (2011) (44)	5.8	3.5	18	0.9	3.2	18	20.1%	1.43 (0.69, 2.17)				-	
Nijamkin et al. (2012) (38)	79.6	15.5	72	63.8	14.2	72	23.5%	1.06 (0.71, 1.41)			4	F	
Papalazarou et al. (2010) (41)	76.4	4.1	15	57.5	4.1	15	13.6%	4.49 (3.07, 5.90)					-
Sarwer et al. (2012) (47)	26.1	1.5	41	23.5	1.5	43	22.3%	1.72 (1.21, 2.22)					
Tucker et al. (1991) (46)	55	15.9	17	48.8	17.9	15	20.5%	0.36 (-0.34, 1.06)					
Total (95% CI)			163			163	100.0%	1.60 (0.82, 2.38)				•	
Heterogeneity: Tau ² = 0.64; Ch	$i^2 = 31$	04, df	= 4 (P	< 0.000	001); [2	= 879	6	1 G. 625 C-51	1110	1		<u>+</u>	
Test for overall effect: $Z = 4.04$	(P < 0)	.0001)							-10	-5	0	5	1
									Favours	No Interve	ntion	Favours	Intervention

CI, confidence interval; df, degrees of freedom; M, mean; N, number of patients; SD, standard deviation.



Intensity of Lifestyle Therapy



Miller K. Obesity Surgery. 2016 epub ahead of print Lopez-Nava G. SOARD. 2015;11:861-865 Sullivan S. *Obesity*. 2017;25(2):294-301.



Diets with Data

Diet	Carb	Fat	Protein
LEARN	Moderate	Low	Normal
Atkins	Very Low	High	High
South Beach	Moderate	Moderate	Moderate
Paleo	Moderate	Low	High
Zone	Moderate	Moderate	Moderate
Pritikin/ Ornish	High	Very Low	Normal
Mediterrane an	Moderate	Moderate	Normal
Keto	Very Low	High	Normal

Common Themes

- Reduction in either the type or amount of food
- Reduce or eliminate sweets
- Reduce or eliminate sugar sweetened beverages
- Use whole grains when grain products are consumed

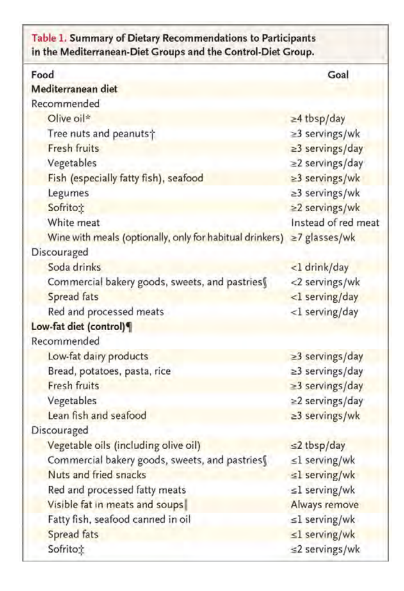


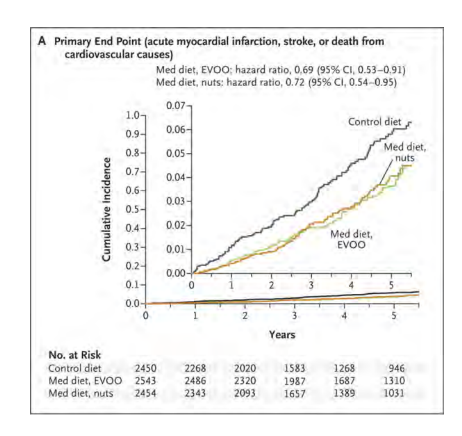
Network Meta-Analysis: Comparisons of Named Diet programs

		12-mo Weight Loss, kg					
	No diet	5.16	5.70	7.25	7.27		
	(6 mo: 0; 12 mo: 0) ^a	(2.68 to 7.63)	(4.14 to 7.35)	(5.33 to 9.25)	(5.26 to 9.34)		
, kg	6.07	LEARN	0.55	2.10	2.12		
	(4.23 to 7.84)	(6 mo: 0; 12 mo: 0.02) ^a	(-1.71 to 2.87)	(-0.20 to 4.47)	(-0.33 to 4.59)		
ht Loss	6.78	0.71	Moderate macronutrients	1.55	1.56		
	(5.50 to 8.05)	(-0.97 to 2.44)	(6 mo: 0; 12 mo: 0) ^a	(0.13 to 2.95)	(-0.17 to 3.30)		
no Weight	8.73	2.66	1.95	Low carbohydrate	0.02		
	(7.27 to 10.20)	(0.93 to 4.44)	(1.13 to 2.79)	(6 mo: 0.83; 12 mo: 0.48) ^a	(-1.78 to 1.79)		
9-mo	7.99	1.92	1.20	−0.74	Low fat		
	(6.01 to 9.92)	(-0.19 to 4.06)	(-0.42 to 2.79)	(−2.31 to 0.78)	(6 mo: 0.17; 12 mo: 0.50) ^a		

- > 59 Article with 7286 patients
- > Significant weight loss for both low-carb and low-fat diets
- > Difference between named diets was small

Mediterranean Diet: PREDIMED Study





Weight loss at 5 years:

- Control: -0.604 kg
- Med, EVOO: -0.88
 kg
- Med, Nuts: 0.188 kg
 HR primary Endpoint
- Med, EVOO: 0.69
- Med, Nuts: 0.72

Estruch R. NEJM. 2018;378:e34
Estruch R. Lancet Diabetes and Endocrinology.2019;7(5):e6-17



Meal Replacements and Odds of Achieving >5% and >10% TBWL at 1 year

	≥5% Weight Loss	≥10% Weight Loss
	OR [95% CI]	OR [95% CI]
MR diet vs diet only	2.83^*_{-} [1.37, 5.86] $l^2 = 40$	1.73 [0.92, 3.26] $I^2 = 0$
MR diet + support vs diet + support	1.49^*_{-} [1.08, 2.06] $l^2 = 44$	1.80^* [1.12, 2.87] $I^2 = 56$
MR diet + support vs diet only	2.83^*_{-} [1.37, 5.86] $l^2 = 25$	5.95^*_{-} [2.12, 16.67] $I^2 = 1$
MR diet + enhanced support vs diet + support	4.32^* [3.01, 6.20] $I^2 = 0$	6.63^* [4.01, 10.94] $I^2 = 0$
MR diet + support vs minimal control	4.03^*_{-} [1.87, 8.69] $l^2 = 82$	8.32^*_{-} [2.02, 34.16] $I^2 = 93$

Astbury NM. Obesity Reviews. 2019;20(4):569-587



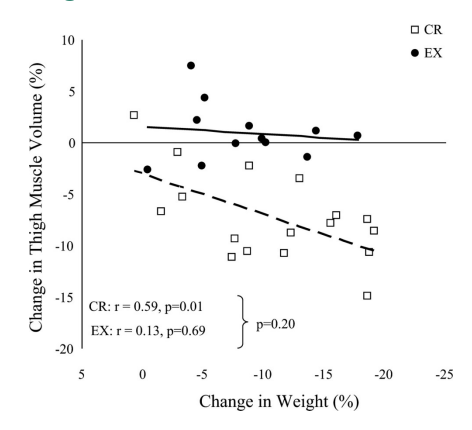
Long-term Calorie Goals

- Based on start weight, gender, level of physical activity
- Goal for 1-2 pound weight loss per week
 - 500 kcal/day deficit = 1 pound per week
 - 750-1000 kcal/day deficit = 2 pounds per week
- Estimates for BMI 30-40 kg/m²:
 - 1200-1500 kcal/day women
 - 1500-1800 kcal/day men
- Comparison Gastric Bypass
 - 500-970 kcal/ day in the first 3 months
 - 870-1420 kcal/day at the end of the first year

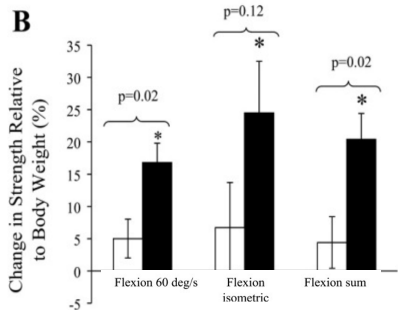


Exercise Preserves Lean Muscle Tissue

Muscle Thigh Volume

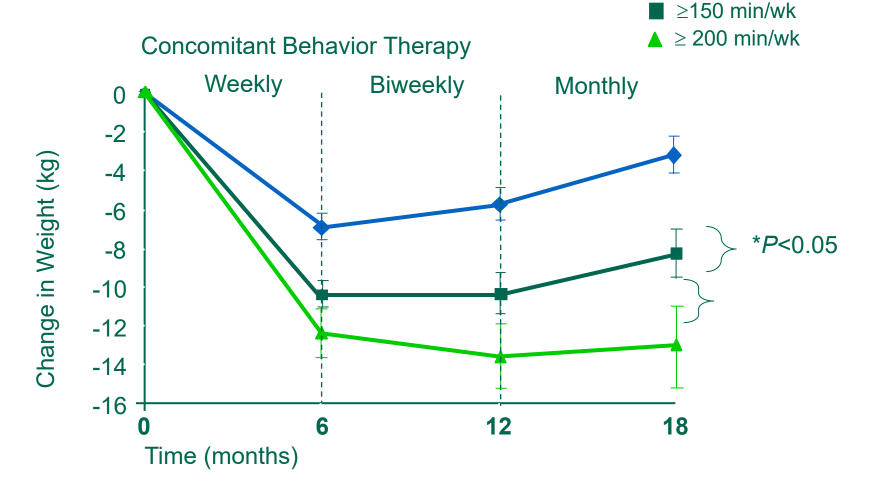


Strength Relative to Body Weight





Physical Activity
Is Necessary for
Weight Loss
Maintenance

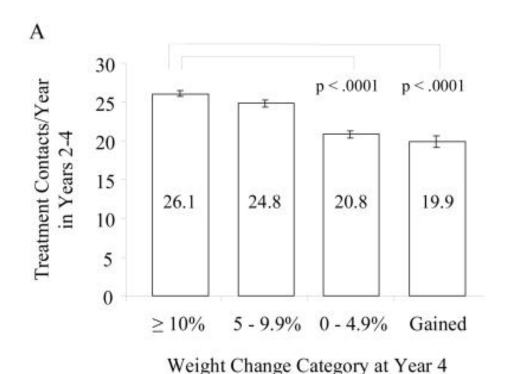


< 150 min/wk

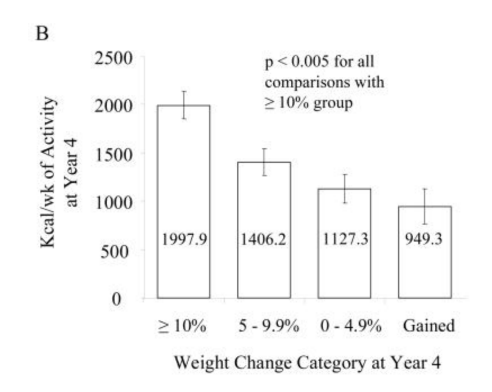


Predictors of Success in Lifestyle Therapy

Treatment Contacts



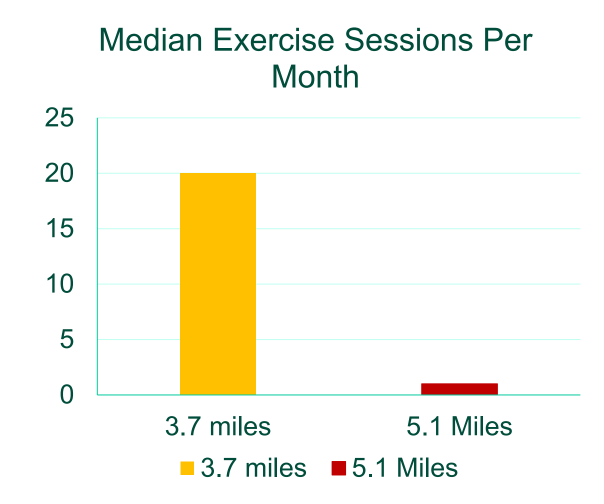
Exercise





Gym Use and Distance From Home

- Data collected from 7.5 million mobile devices by the data firm Dstillery
- Difference of only 1.4 miles between going to the 5 times a week vs one time per month
- Summary even small barriers will reduce exercise





Exercise for Weight Loss and Weight Maintenance

American Heart Association/American College of Cardiology/The Obesity Society Guidelines and American Diabetes Association Guidelines

Weight loss and Adults with type I and type II diabetes

- ≥150 minutes per week moderate intensity (brisk walk)
- Equal to ≥ 30 minutes/day most days of the week

Weight maintenance

- 200-300 minutes per week moderate intensity
- 40-60 minutes/day most days of the week

Strength and flexibility

- Recommended as a consideration by the obesity guidelines
- ADA Guidelines Recommend 2-3 sessions per week

Jensen, MD. Obesity. 2014;22(2):S5-S39 Diabetes Care. 2022;45(Supplement_1):S60-S82

Department of Health and Human Services 2018 Physical Activity Guidelines

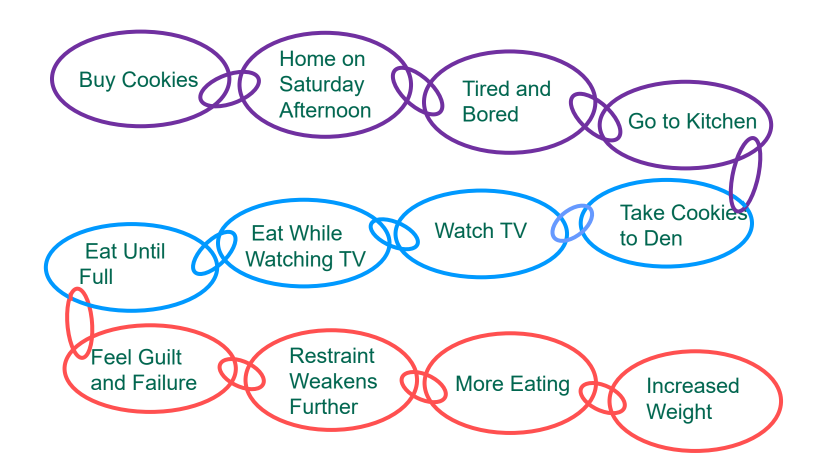
Age	Aerobic Activity	Muscle Strengthening			
6-17	60 minutes of moderate or vigorous physical activity (PA)/day including at least 3 days ofvigorous PA/wk	3 days/week and included as part of the 60 minutes of daily PA. Also include bone-loading activity			
18-64	150-300 minutes of moderate PA/wk, 75 minutes of vigorous PA/wk or equivalent combination spread throughout the week	Muscle strengthening activities at moderate or greater intensity (all major muscle groups) on 2 or more days/wk			
65+	Same as adults, or be as active as abilities and health conditions allow	Same as adults, but include balance training and combination activities (strength and aerobic training together)			
All Ages	Sit Less. Move More				

Piercy K. JAMA. 2018;320(19):2020-2028





Breaking the Obesity Behavior Chain





Pillars of Behavior Modification



Klein, S. Gastroenterology. 2002;123(3):882-932 Wadden, TA. Circulation. 2012;125(9):1157-70



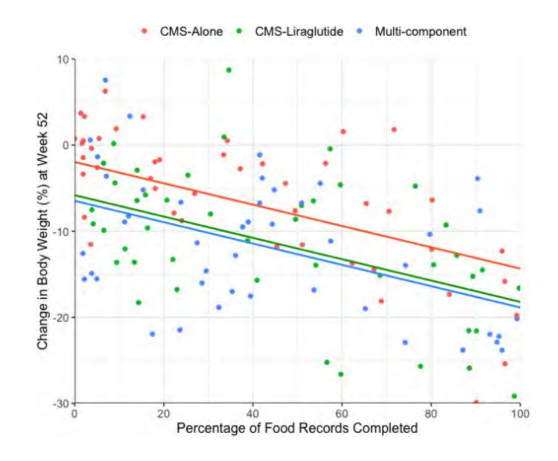
Self-Monitoring: Independently Associated with Weight loss

Systematic review of 22 studies

 More frequent and complete self-monitoring of food intake, exercise and body weight was consistently associated with more weight loss

Post hoc analysis of a randomized controlled trial of 3 arms: intensive behavioral therapy (IBT) alone vs IBT + liraglutide 3.0 mg/d vs IBT vs liraglutide 3 mg/d + meal replacements

- In a linear regression model controlled for treatment group, only adherence to self-monitoring predicted weight loss at 52 weeks
- Patients who completed 100% of their food records lost 12.4 percentage points more than those who completed 0%

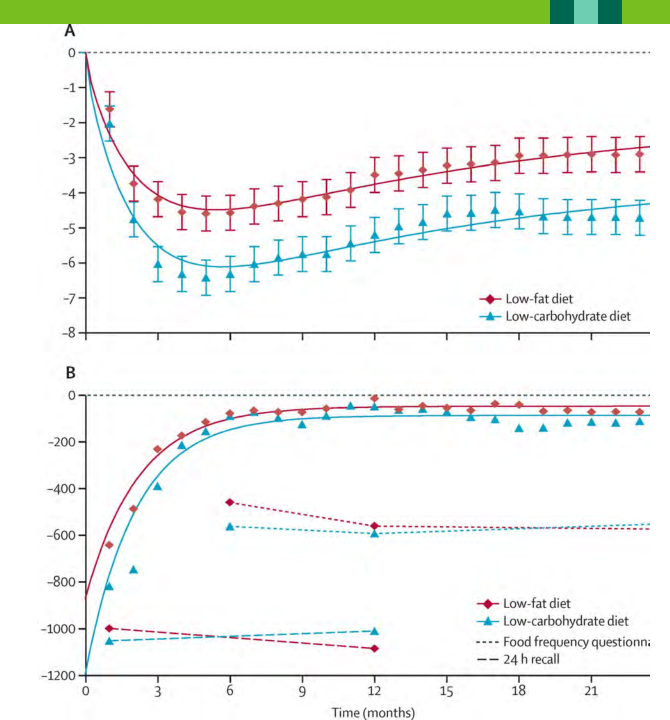




Use of Food and Activity Logs

- Cue the patient on what they have already eaten in the day
- Help the patient be more mindful with food choice
- Help identify patterns
- Food logs and in person 24-hour recalls are not accurate for total energy intake
 - Very hard to estimate energy intake
 - Should only be used as a tool to guide choices

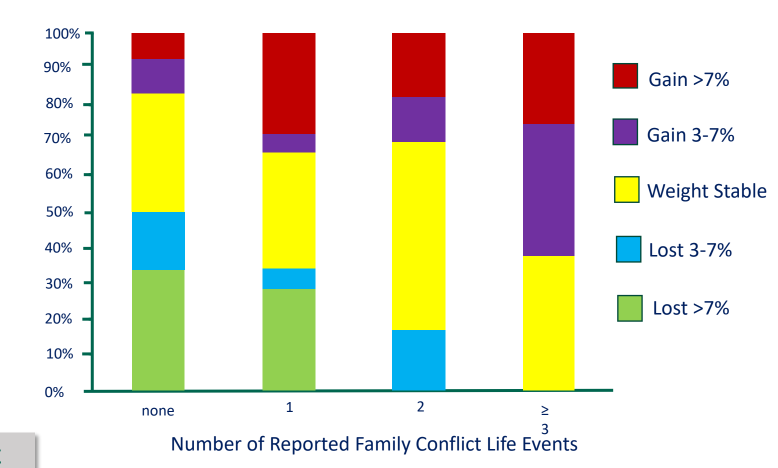
Freedhoff Y. The Lancet. 2016;338(10047):849-851





Factors That Can Derail Lifestyle Therapy

- Stress
 - Home related
 - Work related
- Lack of Sleep
 - Not getting to bed on time
 - OSA
 - Insomnia
- Physical Injuries
- Food insecurity



May need referrals for management



Delivering Lifestyle Therapy

- Trained Interventionist
 - Dietitian, nurse, psychologist, behavior coach, exercise professional, physician (Billing may be limited to physician, dietitian, psychologist)
- In person and over the telephone may yield similar results, less via internet only
- Intensity of lifestyle intervention does matter
 - High Intensity = 14 or more "visits" in 6 months*
 - Moderate Intensity = 6 to 13 "visits" in 6 months*
 - Difference between high and moderate intensity is about 5% TBWL
- Does not appear to be a difference in weight loss between individual sessions and group sessions





Five Steps to Facilitate Behavior Change



- 2
- Review when, where, and how behaviors will be performed
- Have patient keep record of behavior change
 - 4

Review progress at next treatment visit

5

Congratulate patient on successes (do not criticize shortcomings)



Goal Setting

Specific

Names a specific action or behavior

Measurable

• The goal is made so that it can be measured

Attainable

The goal can be reasonable attained

Relevant

The goal is relevant to the desired behavior change

Time-Based

• The goal has a deadline for accomplishment



What Do I Do for My Patients Without a Dietitian or Health Coach?

Diet Recall

- Everything consumed (liquid and solid)
- You can skip this for time, but ask about snacking and meals out

Set calorie goal with typical ranges (BMI <40 kg/m2), calculate for higher BMI

- 1200-1500 kcal/day women,
- 1500-1800 kcal/day men,

Set Exercise Goal:

- 150 min/week moderate intensity exercise for weight loss
- 200-300 min per week for weight maintenance
- 2-3 sessions of strength/resistance exercise per week

Discuss barriers and goals at every visit

At least monthly follow-up for the first 6 months

Reduce

- Added sugars
- Processed grains
- Animal fat (except fatty fish)

Protein in moderation – 16-24% of calories (80-120 gm/day for most patients)

For MASLD or CVD patients: discuss Mediterranean diet Increase

- · Non-starchy Vegetables/Fruit
- Unsaturated vegetable oil (olive oil)
- Nuts (but limit total number of servings)

Use meal replacements to help achieve dietary goals

Jensen, MD. Obesity. 2014;22(2):S5-S39 Webb, VL. Gastroenterology. 2017;152(7):1752-1764 Mozaffarian D. Obesity. 2025. epub ahead of print

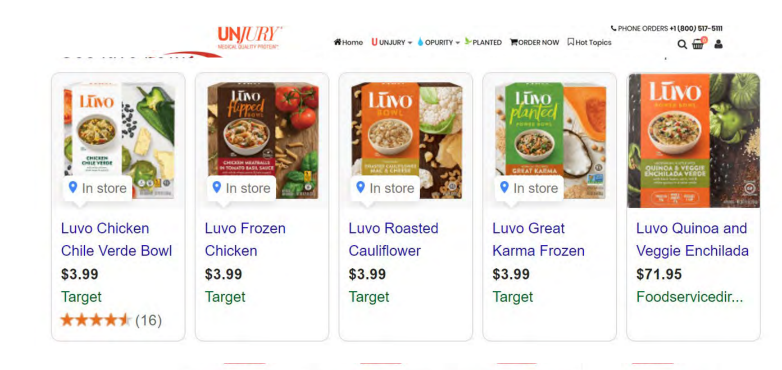


Resources for Exercise

- https://www.mayoclinic.org/healthy-lifestyle/fitness/in-depth/strength-training/art-20046031
- https://www.acefitness.org/resources/everyone/exerciselibrary/?srsltid=AfmBOopUPYpKGr7eraaZG6q4TdBJkKKPdVwCueAXo3MezSgDf4zWxcw
- Apple Fitness App
- Multiple other exercise apps

Meal Replacements

- Types
 - Shakes
 - Bars
 - Frozen Entrée
 - Patient self purchase
 - Carry in the office
- Calorie Controlled
- Stimulus controlled





Conclusions

- Lifestyle therapy alone achieves only modest weight loss
- Lifestyle therapy maximizes weight loss with all adjunctive therapies
 - Anti-obesity medications
 - Endoscopic Bariatric Therapies
 - Bariatric Surgery
- Components
 - Diet
 - Exercise
 - Behavior Modification
- Can be done in a primary care practice if time is limited, focus on one goal at a time





WELCOME to the Obesity Care in All Ages ECHO

Session 4, How to Use Anti-Obesity Medications Effectively (GLP-1 agonist) - August 12th, 2025



Today's Program

- Brief housekeeping
- Didactic: How to Use Anti-Obesity Medications Effectively (GLP-1 agonist) Elaine Banerjee, MD, MPH; Sarah Finn, MD
- Case Discussion
- Summary
- Up Next



Speaker Slides: Obesity Treatment with Glucagon-like Peptide (GLP-1) Receptor Agonists

Elaine S Banerjee, MD, MPH



Disclosures

I have no relevant conflicts of interest to disclose



Acknowledgements

 I appreciate the work of the ECHO team and the DH Weight Center team on this panel, especially Dr Sarah Finn and Dr Minda Gowarty for some the slides and content of this presentation



Objectives

- By the end of this presentation, participants should be able to:
 - Identify the indications, contra-indications, and side effects of GLP-1 RA
 - Apply evidence-based pharmacological management to develop personalized care plans for patients with obesity
 - Prescribe and manage GLP-1 RA treatment for obesity
- Of note, GLP-1 RA are not the only medications for the treatment of obesity and non-GLP-1 medications will be discussed in our next session





"I shouldn't need a medication..."









GLP-1 Medications

Obesity

- Semaglutide (Wegovy)
- Tirzepatide (Zepbound)
- Liraglutide (Saxenda)

Diabetes

- Semaglutide (Ozempic, Rybelsus)
- Tirzepatide (Mounjaro)
- Liraglutide (Victoza)
- Dulaglutide (Trulicity)





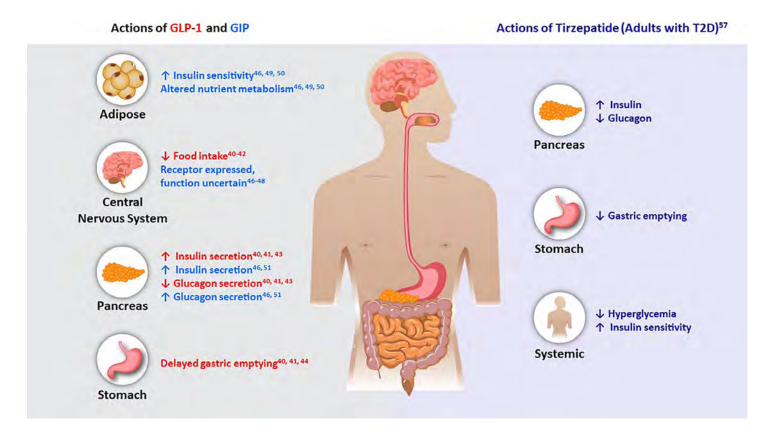
Indications

- Obesity
 - BMI \geq 30 kg/m²
 - BMI ≥ 27 kg/m², AND serious complication of obesity
- Diabetes
- Zepbound: moderate-to-severe OSA
- Wegovy: CAD, PAD, CVA
- Not FDA approved: MASLD, HFpEF





Mechanism



De Block, C et al. Tirzepatide for the treatment of adults with type 2 diabetes: an endocrine perspective. Diabetes Obesity and Metabolism 1.2023. 25: (1). 3-17.





Mechanism







Side-effects,
Contraindications, and
Precautions



Side-effects of weight loss

- Gallstones and their complications which may result in gallstone pancreatitis
- Hair loss
- Gout flares
- Loss of lean body mass
- Loose skin



Side Effects – Medication-related

- GI*
 - Nausea 16-44%
 - Constipation 3-24%
 - Diarrhea 9-30%
 - Abdominal pain 6-20%
 - Vomiting 5-24%
 - Belching, GERD, & Flatulence 1-7%
- Fatigue 5-11%
- Headache 1-17%
- Hypoglycemia semaglutide and tirzepatide 1-6%, liraglutide 2-28%
- Injection site reaction 3-14%

UpToDate Lexidrug/Semglutide. UpToDate Lexidrug/Tirzepatide. UpToDate Lexidrug/Liraglutide. Accessed Aug 3, 2025.





Are they too new for us to know all the side effects?

- Exenatide was approved for diabetes in 2005
- Liraglutide was approved for diabetes in 2010 and for obesity in 2014
- Semaglutide was approved for diabetes in 2017 and for obesity in 2021
- Tirzepatide was approved for diabetes in 2021, for obesity in 2023, and for obstructive sleep apnea in 2024





Contraindications

- Personal or family hx of medullary thyroid cancer or MEN-2
- Pregnancy

UpToDate Lexidrug/Semglutide. UpToDate Lexidrug/Tirzepatide. UpToDate Lexidrug/Liraglutide. Accessed Aug 3, 2025. Cesta CE, Rotem R, Bateman BT, et al. Safety of GLP-1 receptor agonists and other second-line antidiabetics in early pregnancy. *JAMA Intern Med*. Doi: 10.10001/jamainternmed.2023.6663





Precautions

- Known gallstones
 - Consider ursodiol
- Pancreatitis
- Tirzepatide & oral contraceptives



Effectiveness



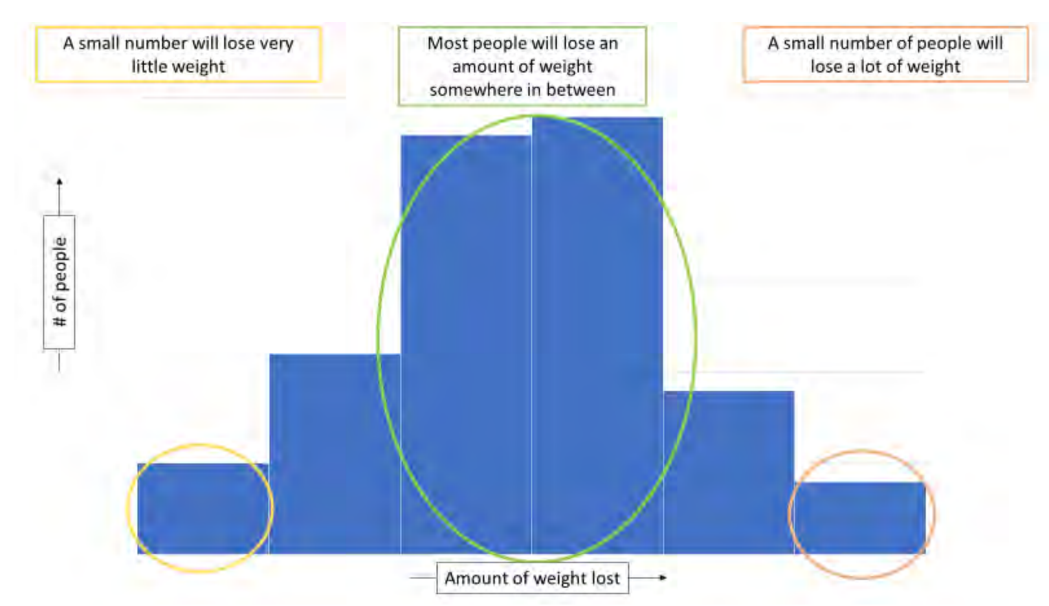


Effectiveness – Weight - Summary

	Average Total Body Weight Loss	Difference from Lifestyle
Lifestyle	3-5%	
Liraglutide	8-11%	5-6%
Semaglutide	15-17%	10-13%
Tirzepatide	21-25%	18-20%











Effectiveness – Weight - Semaglutide

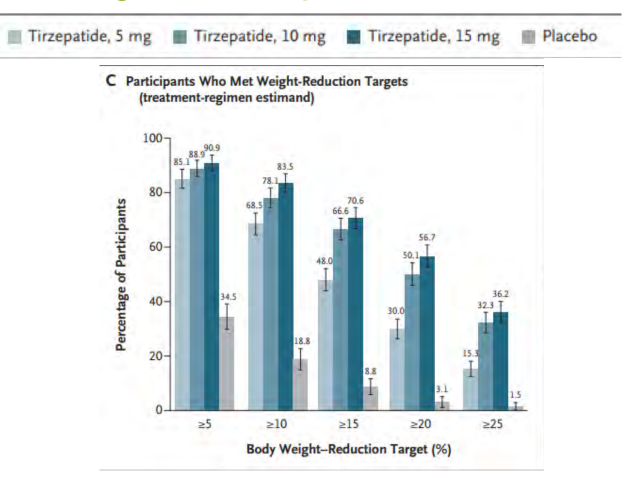
	Percentage achieving 5% weight loss	Percentage achieving 10% weight loss	Percentage achieving 15% weight loss	Percentage achieving 20% weight loss
Semaglutide (N=2,366)	86.4%	71.9%	54.6%	34.8%
Placebo (N=1,222)	37.5%	16.4%	7.4%	2.8%
Risk Ratio (95% CI)	2.2 (1.8,2.8)	4.1 (3.1,5.7)	7.1 (4.8,10.3)	11.9 (8.3,16.9)

Qin W, et al. Efficacy and safey of semaglutide 2.4mg for weight loss in overweight or obese adults without diabetes: An updated systematic review and meta-analysis including the 2-year STEP 5 trial. *Diabetes Obes Metab.* 2024;26:911-923.





Effectiveness – Weight - Tirzepatide



Jastreboff AM, et al. Tirzepatide once weekly for the treatment of obesity. N Engl J Med 2022;387:205-16.



Effectiveness – Quality of Life

Parameter	Tirzepatide 5 mg (N = 630)	Tirzepatide 10 mg (N = 636)	Tirzepatide 15 mg (N = 630)	Placebo (N = 643)
IWQOL-Lite-CT				
Total Score				
Baseline	64.2 (0.9)	61.9 (0.9)	63.0 (0.9)	63.2 (1.0)
Change from baseline to week 72	18.6 (0.6)	21.2 (0.6)	22.6 (0.6)	10.5 (0.7)
Change from baseline difference vs. placebo (95% CI), p value	8.1 (6.3 to 9.9)***	10.7 (8.9 to 12.5)***	12.1 (10.3 to 13.9)***	4
Physical Composite Score				
Baseline	64.0 (1.0)	61.5 (1.0)	62.7 (1.0)	63.3 (1.1)
Change from baseline to week 72	16.8 (0.7)	19.5 (0.7)	20.8 (0.7)	9.7 (0.7)
Change from baseline difference vs. placebo (95% CI), p value	7.2 (5.2 to 9.2)***	9.9 (7.9 to 11.9)***	11.1 (9.1 to 13.1)***	8
Physical Function Composite Score				
Baseline	64.4 (1.0)	61.9 (1.0)	63.3 (1.0)	64.0 (1.1)
Change from baseline to week 72	17.8 (0.7)	20.7 (0.7)	21.8 (0.7)	10.1 (0.8)
Change from baseline difference vs. placebo (95% CI), p value	7.7 (5.6 to 9.8)***	10.7 (8.6 to 12.8)***	11.7 (9.6 to 13.8)***	
Psychosocial Composite Score				
Baseline	64.3 (1.0)	62.1 (1.0)	63.2 (1.0)	63.2 (1.0)
Change from baseline to week 72	19.6 (0.7)	22.1 (0.7)	23.6 (0.7)	11.0 (0.7)
Change from baseline difference vs. placebo (95% CI), p value	8.7 (6.7 to 10.6)***	11.2 (9.3 to 13.1)***	12.7 (10.7 to 14.6)***	Ų

Gudzeune KA, et al. Association between weight reduction achieved with tirzepatide and quality of life in adults with obesity: Results from the SURMOUNT-1 study. *Diabetes Obes Metab.* 2025;27:539-550.



Dosing & Titration





Dosing & Titration – General Principles

- Start at the lowest dose
- Titrate based on effect and tolerance
- Most people will need to get to the 3rd or 4th dose for effect
- Message after the 3rd shot at each dose
 - Weight
 - Side effects



Dosing & Titration - Semaglutide

Wegovy

- •0.25mg weekly for ≥ 4 weeks
- •0.5mg weekly for ≥ 4 weeks
- •1mg weekly for ≥ 4 weeks
- •1.7mg weekly for ≥ 4 weeks
- •2.4mg weekly

Ozempic

- •0.25mg weekly ≥ 4 weeks*
- •0.5mg weekly ≥ 4 weeks*
- •1mg weekly ≥ 4 weeks
- •2mg weekly†
- *0.25/0.5mg adjustable dose pen with 2mg medication in the pen
- † 1mg and 2mg pens are adjustable by counting clicks to get lower doses



Dosing & Titration - Tirzepatide

Zepbound & Mounjaro

- •2.5 mg weekly for ≥ 4 weeks
- •5 mg weekly for ≥ 4 weeks
- •7.5 mg weekly for ≥ 4 weeks
- •10 mg weekly for ≥ 4 weeks
- •12.5 mg weekly for ≥ 4 weeks
- •15 mg weekly



Dosing & Titration - Liraglutide

Saxenda

- •0.6mg daily for 1 week
- •1.2mg daily for 1 week
- •1.8mg daily for 1 week
- •2.4mg daily for 1 week
- •3mg daily

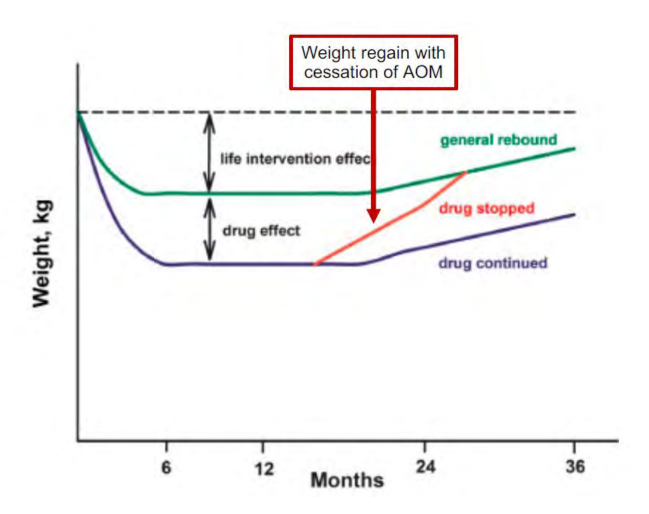
Victoza

- •0.6mg daily for 1 week
- •1.2mg daily for 1 week
- •1.8mg daily for 1 week





Duration





Insurance coverage

- Medicare does not cover any medications for obesity
 - Wegovy for CAD with hx of MI
 - Zepbound for moderate-to-severe OSA
 - \$2,000/year limit on out of pocket costs for medications
- Medicaid state dependent
 - VT no coverage for obesity, but may cover zepbound for OSA
 - NH varies but some plans will cover GLP-1 RA for obesity
- Federal BCBS tier 3 with high copay





Insurance coverage

- Marketplace generally not covered
- Employer-based plans are employer dependent
- Most plans will cover a GLP-1 RA for Diabetes
 - Many plans have restrictions on which medication and other treatments
 - Generally not covered for pre-diabetes or insulin resistance



Troubleshooting



Troubleshooting: Managing GI side effects

- Constipation
 - Hydration & fiber (increasing gradually)
 - Osmotic laxatives (miralax)
 - Stimulant laxatives only if needed
- Diarrhea generally self limited

- Nausea
 - Small meals, eating something first thing in the morning
 - Ginger
 - Ondansetron if needed
- Belching
 - Ginger, papaya enzyme

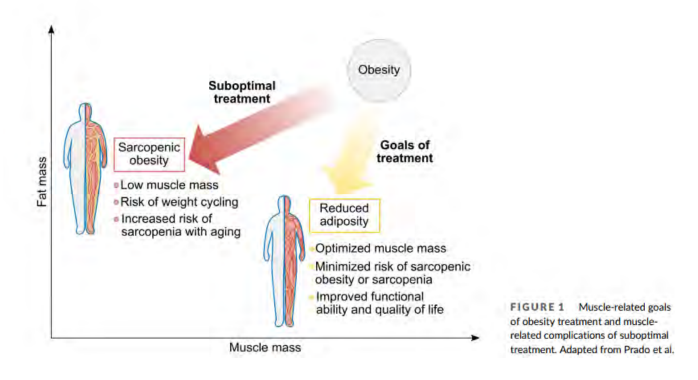


Common Side Effect Pattern with GLP1 Medications





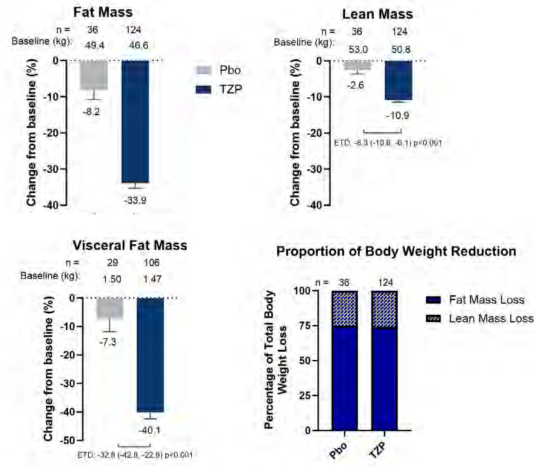
Troubleshooting: Limiting loss of lean body mass



Mechanick JI. Strategies for minimizing muscle loss during use of incretin-mimetic drugs for treatment of obesity. *Obesity Reviews*. 2025;26:e13841.



Troubleshooting: Limiting loss of lean body mass



Look M, et al. Body composition changes during weight reduction with tirzepatide in the SURMOUNT-1 study of adults with obesity or overweight. *Diabetes Obes Metab.* 2025;27:2720-2729.



Troubleshooting: Limiting loss of lean body mass

- Nutrition
 - Protein
 - 0.8g 1.5g/kg body weight
 - Bariatric: 1.5g/kg ideal body weight
 - Joslin: 1-1.5g/kg adjusted body weight (IBW + 0.25xEBW)
 - Micronutrients
- Physical activity especially resistance exercise
- Possible future role for bimagrumab, apiegromab, or cagrilintide

Mechanick JI. Strategies for minimizing muscle loss during use of incretin-mimetic drugs for treatment of obesity. *Obesity Reviews*. 2025;26:e13841.

Wilkinson TJ, et al. Preservation of healthy lean body mass and function during weight loss. *Clinical Obesity*. 2024:14:e12683





WELCOME to the Obesity Care in All Ages ECHO

Session 5, How to Use Anti-Obesity Medications Effectively (1 AOM non glp 1 agonist)

- September 9th, 2025



Today's Program

- Brief housekeeping
- Didactic: How to Use Anti-Obesity Medications Effectively (1 AOM non glp 1 agonist)
- Sarah Finn, MD
- Case Discussion
- Summary
- Up Next



Obesity Treatment with Anti obesity Medicine (oral agents/non GLP1 agonist)

Sarah Finn, MD



Disclosures

I have no relevant conflicts of interest to disclose



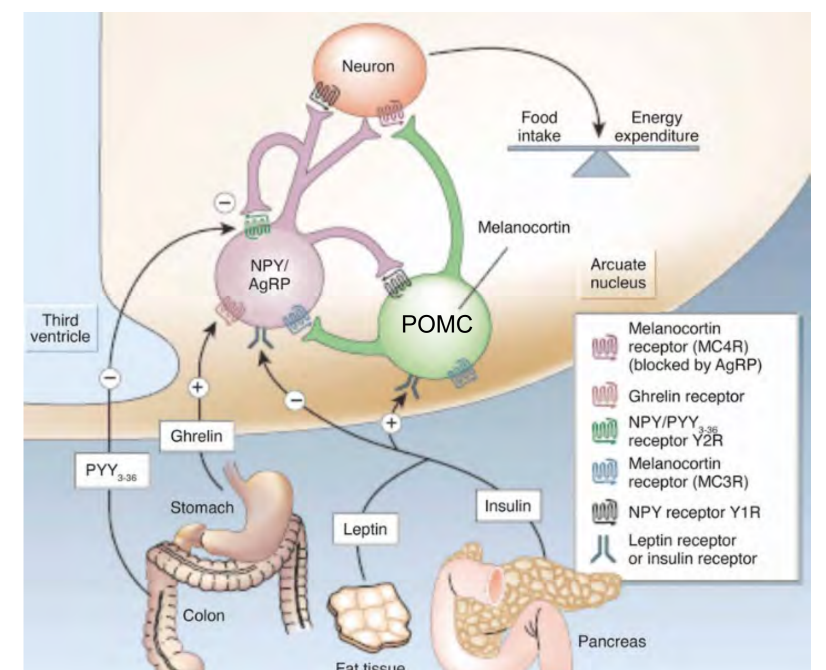
Objectives

By the end of this presentation, participants should be able to:

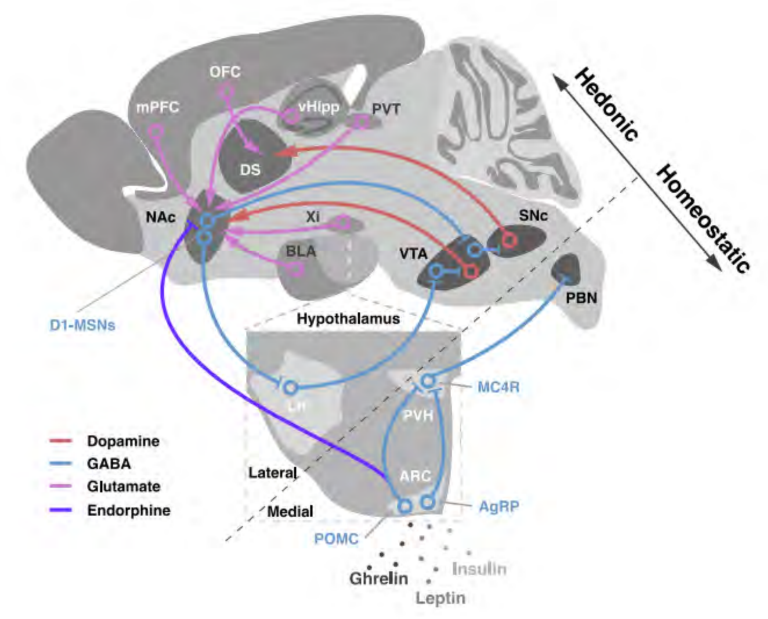
- 1. Understand mechanism of action of non GLP1 anti obesity medications
- 2. Understand short- and long-term benefits
- 3. Understand risks and management of complications
- 4. Variability of response and management of suboptimal initial response, recurrent weight gain and complications











Stuber GD, Schwitzgebel VM, Luscher C. The neurobiology of overeating. *Neuron.* 2025;113(11):1680-1693.

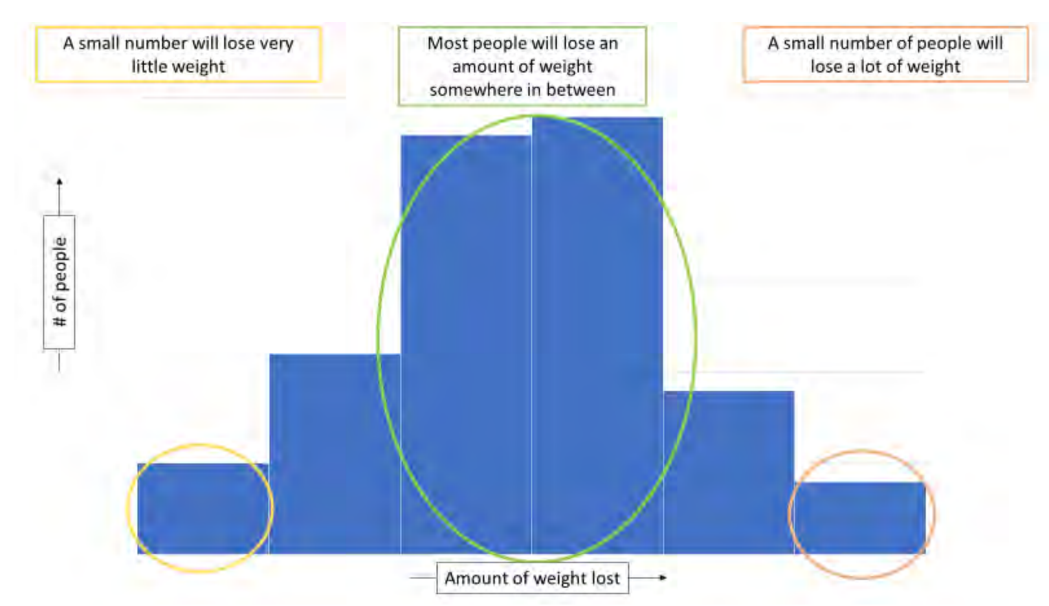


Current AOM Pharmacotherapy

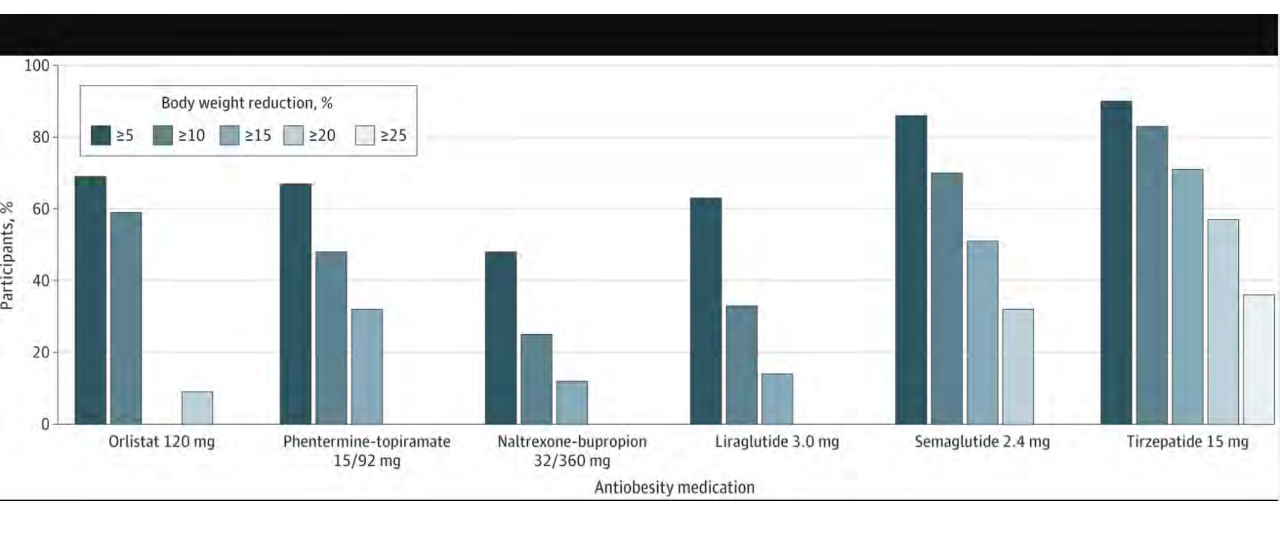
IIII LICARIT			
	FDA Approved	Off Label	
	Phendimetrazine Benzphetamine Diethylpropion	Metformin Dapaglifloxin Diabetes	
First generation	Phentermine		
Second generation	Orlistat Phentermine/Topiramate Naltrexone/Bupropion	Topiramate (seizures/migraines) Zonisamide (seizures/migraines) Bupropion (depression) Naltrexone (addiction)	
	Liraglutide 3.0 mg		
Third generation	Semaglutide 2.4 mg Tirzepatide	Liraglutide 1.8 mg Semaglutide Diabetes Tirzepatide Exanatide Dulaglutide	
Monogenic obesity	Setmelanotide		















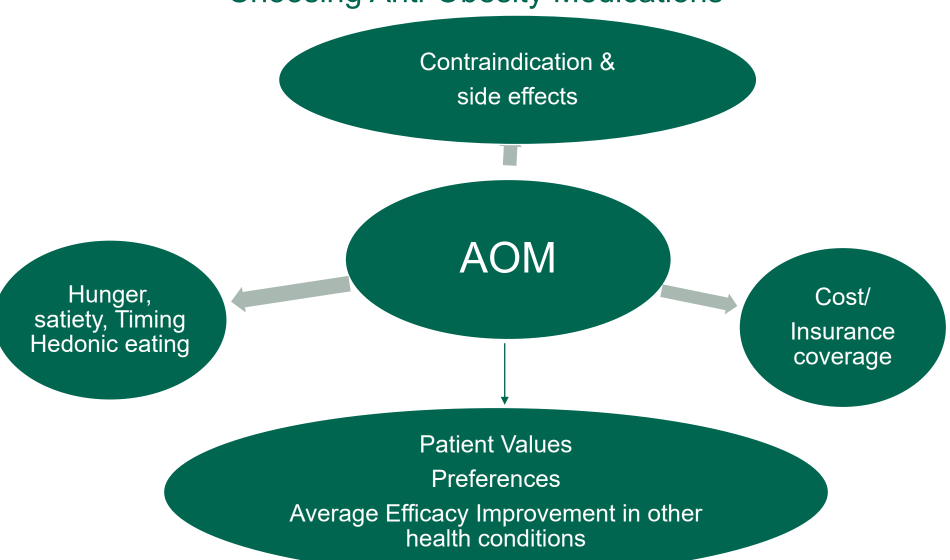
	Orlistat	Phentermine	Qsymia (phentermine/topiramate)	Contrave (bupropion/naltrexone)
Conditions to avoid in	Bariatric surgery Liver disease Pregnancy/breast feeding	Seizures Glaucoma, ESRD Pregnancy/breast feeding	Seizures Glaucoma Kidney stones ESRD on HD Pregnancy/breast feeding	Seizures Glaucoma Lower doses with CKD/Liver disease Pregnancy/breast feeding
% TBW loss	2.9-6.1 KG (6-8 lbs.)	5; 3.6kg (~8lbs)	7-9.8	5-7
Contraindications	Malabsorption syndromes	concurrent stimulant, Substance use disorder	lack of highly-effective pregnancy prevention	untreated bipolar disorder, active opioid use,
Adverse effects	Diarrhea Flatus with discharge Fecal urgency Fatty stool (Steatorrhea) Oily evacuation Fecal incontinence Rare liver injury	Jitteriness, tremor, increased BP or HR, dry mouth, insomnia, constipation	Same as phentermine plus dizziness, abnormal taste, paresthesia, kidney stones	Nausea, constipation, insomnia, dry mouth, sweating, headaches, increased BP







Choosing Anti-Obesity Medications



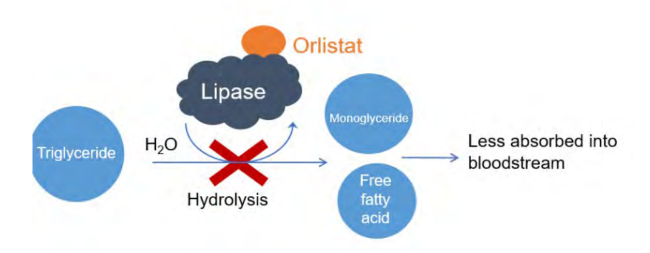




Orlistat

Mechanism of Action

- Reversible inhibitor of gastrointestinal lipases reducing lipid absorption
- (25-30% of fat calories are not absorbed)
- Reinforces avoidance of energy dense foods





Clinical Trials Orlistat 120 mg TID

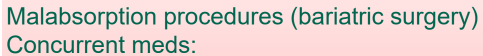
Authors Participants	Average KG weight loss
Hutton, Fergusson et al. 10, 631	6.1 kg vs 2.6 kg placebo
Davidson MH, Hauptman, J et al. 3, 305	5.8 kg vs 3 kg placebo
Suyog, J, et al. 80	4.65 kg vs 2.5 kg placebo
Rissanen A, Lean MEJ (XENDOS trial) 2,550	-2.4 KG weight changes treatment difference -0.4% treatment improvement in A1c vs placebo

Igel LI, Kumar RB, Saunders KH, Aronne LJ. Practical Use of Pharmacotherapy for Obesity. Gastroenterology. 2017 May;152(7):1765-1779. doi: 10.1053/j.gastro.2016.12.049. Epub 2017 Feb 10. PMID: 28192104.











Cyclosporine

Coumadin

Anti-epileptics

Antiretroviral Agents

Levothyroxine



High cholesterol Diabetes Constipation

Diarrhea

Flatus with discharge



Fatty stool (Steatorrhea)

Fecal incontinence

Rare liver injury





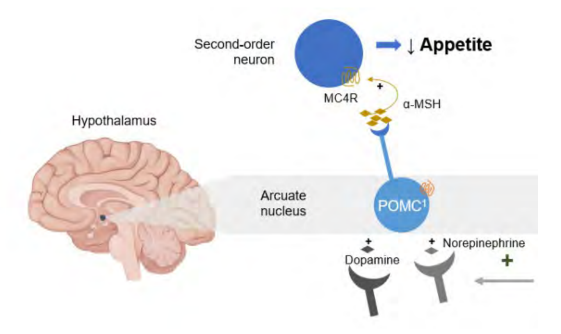
Phentermine

Mechanism of Action

 Increases release of norepinephrine (and dopamine and serotonin to lesser degree) in hypothalamus

Enhance appetite suppression via central adrenergic pathways, reduced food

consumption







Phentermine



Prior heart attack, Aneurysm, stroke Seizure

Uncontrolled HTN

Tachyarrhythmia Hyperthyroidism (untreated) Severe anxiety, Bipolar DO

Glaucoma

Pregnancy/breastfeeding



Lack of co-morbid conditions High hunger/cravings



Insomnia
Headache
Constipation
Irritability
Eye pain (monitor)
Increased BP, HR
Dizziness

Jitteriness, Tremor



Phentermine

Dosing

- Phentermine (Lomaira) 4 mg 1-3 times daily (start ½ tablet) before meals
- Phentermine (Adipex) 15mg daily capsule (can start ¼ tablet 37.5mg tab)
- Approved for short term use (3 months); long term therapy is recommended by experts
- Schedule IV Controlled substance
- Check state requirements for controlled substance requirements; for NH: Yearly PDMP and in office visit

Patient Monitoring

- Avoid caffeine, energy drinks, decongestants
- Monitor BP and HR at home
- Suicidal ideation or worsening mood/anxiety



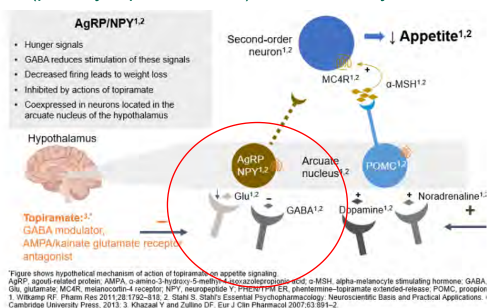
Phentermine/Topiramate

Mechanism of Action

- Phentermine
 - Increases release of norepinephrine (and dopamine and serotonin to lesser degree) in hypothalamus
 - Enhance appetite suppression via central adrenergic pathways, reduced food consumption
- Topiramate

GABA receptor modulator (post synaptic neurons) carbonic anhydrase inhibition,

glutamate antagonism



The Scientific Exchange for U.S. Healthcare Professionals: FORWARD: Focus on Obesity Education Curriculum (2023) available at: Obesity FORWARD Resources for HCPs | Scientific Exchange (scientific-exchange.com) (Accessed:8.15.2025)



Clinical Trials Phentermine/Topiramate

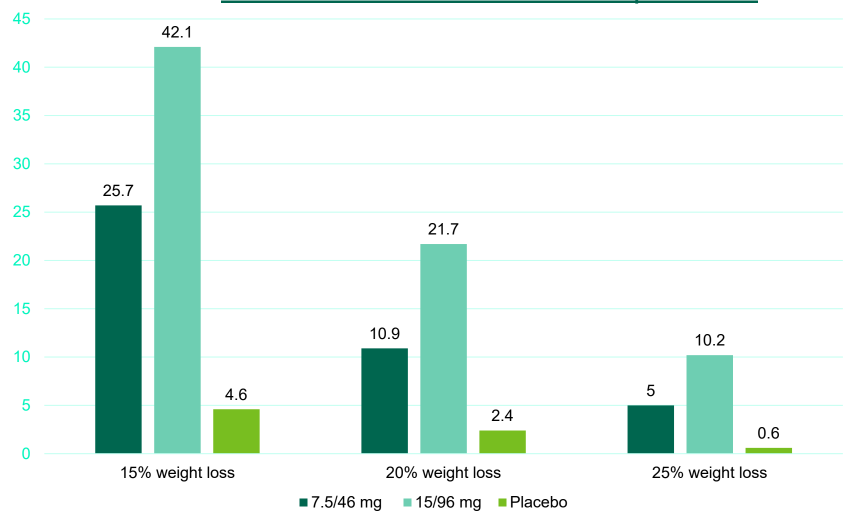
Trial Name Participants	Average Percent Total body weight loss
EQUIP 1267	14%
EQUATE 756	9.2%
CONQUER 2487	12.4%
SEQUEL 676	10.5% 2 years

Igel LI, Kumar RB, Saunders KH, Aronne LJ. Practical Use of Pharmacotherapy for Obesity. Gastroenterology. 2017 May;152(7):1765-1779. doi: 10.1053/j.gastro.2016.12.049. Epub 2017 Feb 10. PMID: 28192104.

^{*} No Cardiovascular outcomes trial completed



Clinical trials Phentermine/Topiramate







Phentermine + Topiramate XR (Qsymia)



Uncontrolled HTN

Tachyarrhythmia

Hyperthyroidism (untreated)

Severe anxiety

Glaucoma

Kidney stones (calcium phosphate)

Pregnancy or breast feeding

CKD stage IV/V



Lack of co-morbid conditions

High hunger/cravings

Women on birth control

Depression not well treated

Paresthesia

Memory issues/forgetfulness

Taste distortion

Dry mouth

Dizziness

Constipation

Insomnia

Serious but rare: hypokalemia** pay attention to nonpotassium sparing diuretics







Phentermine/Topiramate (Qsymia)

Dosing

- Qsymia: 3.75/23mg x 14d then 7.5/46mg, 15/92mg
- Phentermine 8 mg daily and topiramate 25-50 mg daily (more affordable)
- Schedule IV Controlled substance
- Check state requirements for controlled substance requirements; for NH: Yearly PDMP and in office visit

Patient Monitoring

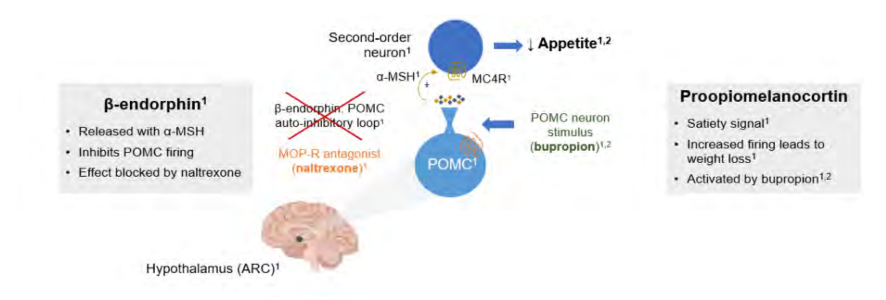
- If on diuretics-monitoring potassium before and during treatment
- Monitor metabolic acidosis higher doses/long term exposure
- Suicidal ideation or anxiety/depression
- Evaluate pregnancy status prior to use in patients who can become pregnant.
- Patients who can become pregnant should have a negative pregnancy test prior to and monthly during therapy.
- Effective contraception should be used during treatment.
- Irregular bleeding may occur with use of combination oral contraceptives; efficacy of contraception may be dependent upon dose.



Naltrexone/Bupropion (Contrave)

Mechanism of Action

- Opioid receptor antagonist dopamine agonist/norepinephrine-dopamine reuptake inhibitor
- Increased satiety, appetite suppression







Clinical Trials Naltrexone + Bupropion (Contrave)

52 weeks multicentered RCT	% Total weight loss
COR-I	6.1% vs. 1.3% placebo -4.8%
COR-II	7.1% vs 2% placebo 5%-48% 10%-27% 15%-10%
COR-BMOD	9% vs 5% placebo
COR-DM	> 5%- vs 1.8% placebo

^{*} No Cardiovascular outcomes trial completed; No dedicated RCT for prevention of weight regain or long -term mainten





Naltrexone/ Bupropion (Contrave)

Seizure disorder

Severe anxiety

Uncontrolled headaches

Heart disease

Uncontrolled high blood pressure

Chronic opioid or methadone use

Drug or alcohol withdrawal

Anorexia nervosa or bulimia nervosa

Glaucoma

Liver failure (reduce dose)

Kidney disease (reduce dose)



Type 2 Diabetes

Emotional eater

Cravings for food and addictive behaviors related to food

Quitting tobacco

Quitting alcohol

Depression

Nausea

Vomiting

Constipation

Stomach upset

Trouble sleeping

Headache

Dry mouth

Dizziness

Worsening mood, suicidality

Sweating





Naltrexone + Bupropion (Contrave)

Dosing

Week 1: 1 tablet in am

Week 2: 1 tablet in am and 1 tablet in pm

Week 3: 2 tablets in am and 1 tablet in pm

Week 4 and on: 2 tablets in am and 2 tablets in pm

....

Bupropion SR 100 mg tablets

Week 1: 1 tablet in the am

Week 2: 1 tablet in the am and 1 tablet in the pm

Week 3: 2 tablets in the am and 1 tablet in the pm

Week 4 and on: 2 tablets in am and 2 tablets in pm

*You can change from 2 100 mg tablets to 200 mg tablet

Naltrexone 50 mg tablet

Week 1: 1/4 tablet in the am Week 2: 1/2 tablet in the am

Week 3: ½ tablet in the am and ¼ tablet in the pm

Patient Monitoring

- Monitor BP and HR weekly for a month
- Worsening depression/anxiety
- Can reduce side effects by reducing dose





Treatment Recommendations

Do not delay treatment; be proactive in prevention of obesity complications

Starting treatment

- BMI ≥ 27 with complications
- BMI ≥ 30

12-16 weeks if less than 3-5% of body weight change course (discontinue, change dose, alternate medication, etc.)

Use in combination with lifestyle and behavioral change!

Maintenance Medication recommended life long

Monitor every 6, 12 months Reassess (exercise, diet, psychological, health issues, concomitant medications)



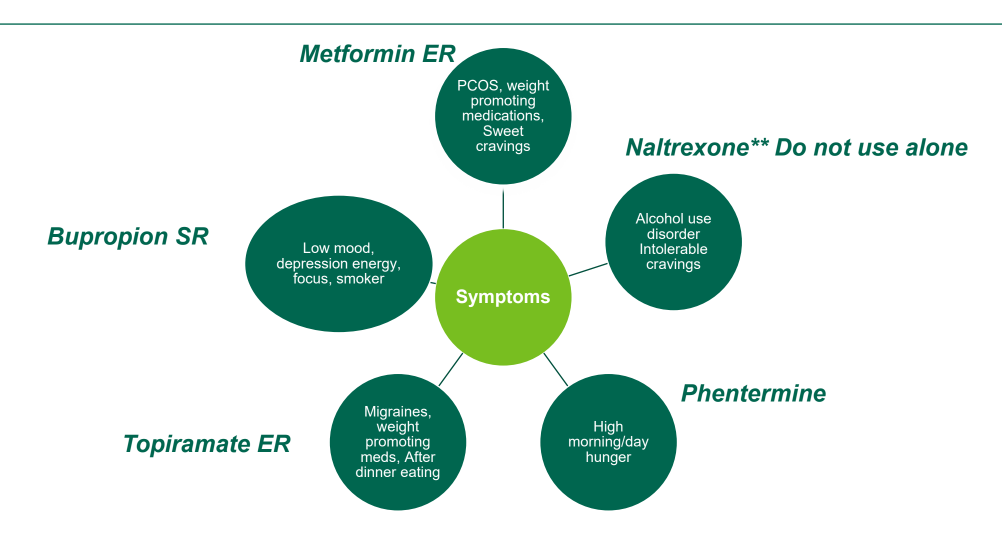


Shared Decision making on Treatment Goals

- In addition to weight loss, focusing on health metrics, reduction in cardiometabolic risk, improvement, remission or resolution of adiposity related complications, maintenance of weight loss, management of symptoms of obesity (appetite cravings), improvement of quality of life.
- Medications help prevent weight regain and maintain weight loss after health changes alone
- Personalized to meet individual values, preferences and treatment goals, safe, effective, culturally acceptable and affordable for long-term adherence
- If BMI still in obesity range, but patients complications have reduced, good QOL, and less amount of ASE on medication, that should be goal weight and tell the patient that!



"Off label" AOM that are affordable and tailored for combination treatment







Ms. Suarez 45 year old woman, weight 230, BMI 35, waist circumference 40 inches. BP 125/76

Insomnia: Amitriptyline 250 mg QHS

Anxiety: Clonazepam 1.5 mg PO BID

PCOS: Fasting insulin 36 uU/ml, A1c 6.0 on metformin

HTN: controlled on lisinopril

Kidney stones: Calcium oxalate





You she is already on metformin to help with psychotropic induced weight gain in addition to prediabetes. She would like to lose more weight and metformin "isn't' working any more" because her hunger has come back. Labs: A1c 5.6. Patient would like to lose more weight and she does not have coverage for GLP1 agonist.

What are your next steps?

- A. Tell her she has already lost 6% of her body weight and she doesn't need any further intervention. Stop metformin as it isn't working
- B. Continue metformin and add phentermine 4 mg titrate to 8 mg
- C. Continue metformin and start naltrexone titration
- D. Stop metformin and start on phentermine 8 mg



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Combination AOM Pearls

- Goals, reasonable expectations on treatment discussion at every visit!
- You can safely add phentermine (and Wellbutrin) in a patient with well controlled HTN even if they need their medication adjusted
- Both phentermine/Wellbutrin have overlapping adrenergic activity and using combination may result in higher risk of anxiety, lower threshold for seizures
- When an agent is ineffective for monotherapy (without serious ASE), it may be reintroduced later in a combination resulting in good weight loss





• You add phentermine and titrate up. She has some constipation, but it improves with water. Over a period of 10 months, she loses another 25 pounds. Her total weight loss is 41 pounds (Total 21.25%) and final weight 190 pounds (BMI 29.3). She wants to lose more weight, but also doesn't want to be on 'medications forever'.

How do you counsel her?



Mr. Williams 32 year old weight, Vitals: 260 lbs, BMI 38 BP 116/76 HR 83

PMH:

- Mild OSA, uses CPAP
- OA: ibuprofen
- Depression: Zoloft
- Migraines: propranolol

He has high hunger during the day, no emotional eating. He would like GLP1 agonist, but no coverage. He is concerned of taking a medication twice a day. What would you do for this patient?

- A) Start Phentermine/Topiramate
- B) Start Naltrexone/Bupropion
- C) Start Metformin
- D) Start Wellbutrin



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Mr. Williams follows up with you after being on phentermine/topiramate and has lost 15 lbs. (5% body weight) in 3 months. His headaches are much improved. He notes he is having tingling in his hands and feet, but it's not bothering him to much. He would like to increase his medication. You increase him to 11.25/69 x 2 weeks then 15 mg phentermine/ 96 mg of topiramate. He follows up with you in 3 months and has lot additional 20 pounds (total 35 lbs. lost, 13.46% of his body weight), but notes that he feels "foggy" and has been more forgetful.

What would be your next steps?

- A. Slowly titrate off Phentermine/Topiramate
- B. Reduce phentermine/topiramate back to 11.25/69 mg dose
- C. Stop Phentermine/Topiramate, start Naltrexone/Bupropion
- D. Continue phentermine, stop Topiramate



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Phentermine/Topiramate PEARLS

- When patients with moderate/high dose of topiramate develop cognitive side effects, reducing the dose to a lower, but still effective dose can allow the continued use of the medication
- Taper down phentermine/Topiramate slowly to minimize side effects
 - Combo step down or alternative every other day x 7 days to off
 - Two separate drugs, decrease dose of topiramate
- Recommend trial of ER versions
- Calcium phosphate stones are associated with topiramate



Ms. Reed is a 38 year old, BMI 40 waist circumference 43, BP 145/89 HR 92

PMH:

DM with gastroparesis: A1c 7.8, metformin, glipizide

HTN: Lisinopril

PCOS: OCP

Depression: untreated

Back pain: untreated

She feels she is an emotional eater with food noise 'all day'. What medication would you try first?

- A) Tirzepatide 2.5 mg and titrate up
- B) Qsymia 3.75/23 and titrate up
- C) Naltrexone/bupropion



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- C) Naltrexone/bupropion



Naltrexone/bupropion Pearls

- Monitor BP and HR weekly for a month
- Monitor depression/anxiety
- •Can reduce side effects by reducing dose back to week 2 or 3 of titration
- •Use reduced doses (week 2) for CKD/Liver disease



• You see Mrs. Emmel for obesity follow up and you would like to start GLP1 agonist. She has had gastric bypass in 2018 with 15% total body weight regain and no concerns for starting GLP1 agonist. Insurance came back with denial saying patient has tried and failed 'making' a patient try orlistat before prescribing GLP1 agonist. Patient is worried about side effects as used Alli (over the counter) pre-surgery and had explosive diarrhea.

What can you say to the patient and insurance company:

- A) Orlistat is contraindicated in malabsorption syndromes
- B) We need to try orlistat first
- C) Orlistat may help your constipation
- D) sorry I can't prescribe any made for you



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WELCOME to the Obesity Care in All Ages ECHO

Session 6, Approach to the Pediatric Patient with Obesity - AAP Clinical practice guidelines
- September 23rd, 2025



Today's Program

- Brief housekeeping
- Didactic: Approach to the Pediatric Patient with Obesity AAP Clinical practice guidelines
- Auden McClure, MD, MPH
- Case Discussion Christine Arsnow
- Summary
- Up Next



Approach to the Pediatric Patient with Obesity

The American Academy of Pediatrics Clinical Practice Guidelines

Auden McClure, MD MPH

Co-director, Pediatric Weight and Lipid Program

The Weight Center, Center for Digestive Health



Dr McClure has no financial conflicts of interest to disclose



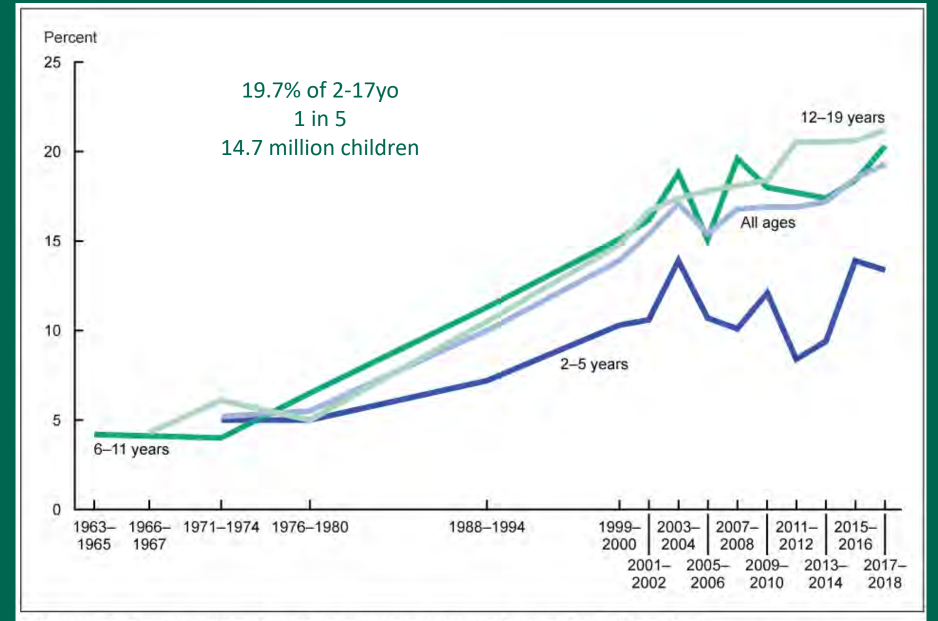
Objectives for today

Define obesity as a disease

Provide an overview of pediatric obesity medicine using the American Academy of Pediatrics Clinical Practice Guidelines



Rising Prevalence of Pediatric Obesity

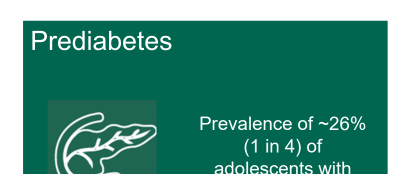


NOTE: Obesity is body mass index (BMI) at or above the 95th percentile from the sex-specific BMI-for-age 2000 CDC Growth Charts.

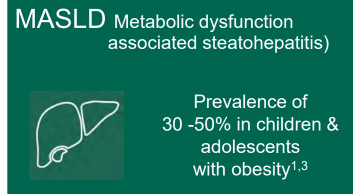
SOURCES: National Center for Health Statistics, National Health Examination Surveys II (ages 6–11), III (ages 12–17); and National Health and Nutrition Examination Surveys (NHANES) I–III, and NHANES 1999–2000, 2001–2002, 2003–2004, 2005–2006, 2007–2008, 2009–2010, 2011–2012, 2013–2014, 2015–2016, and 2017–2018.

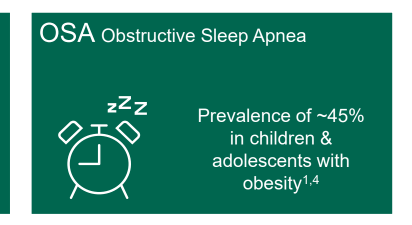


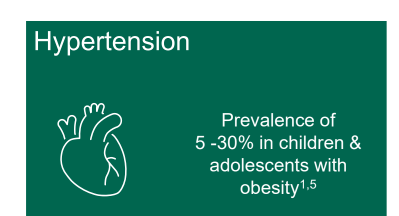
Complications of Obesity

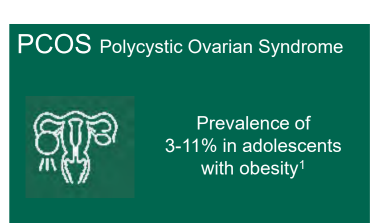


obesity^{1,2}















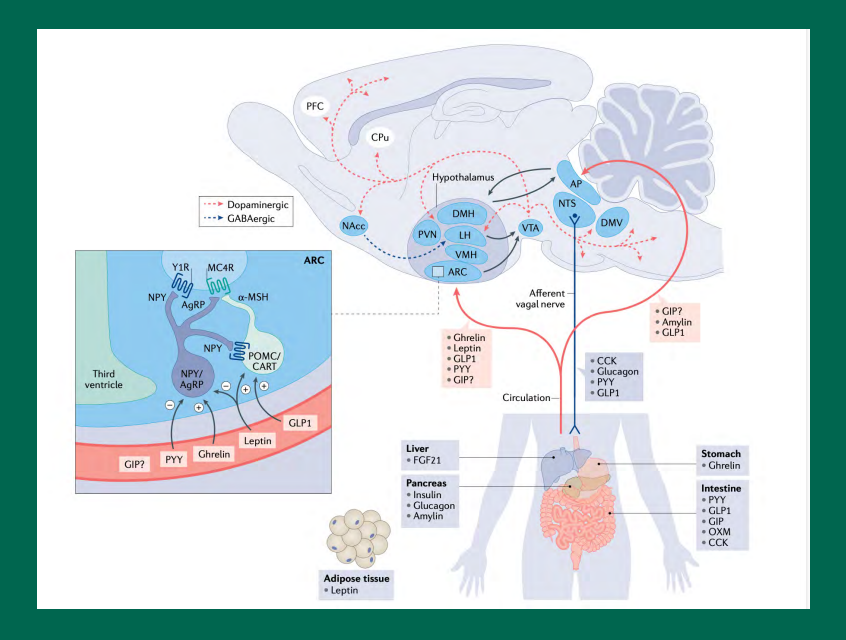
OMA Definition of Obesity

Chronic, relapsing, multi-factorial, neurobehavioral disease, wherein an increase in body fat promotes adipose tissue dysfunction and abnormal fat mass physical forces, resulting in adverse metabolic, biomechanical, and psychosocial health consequences



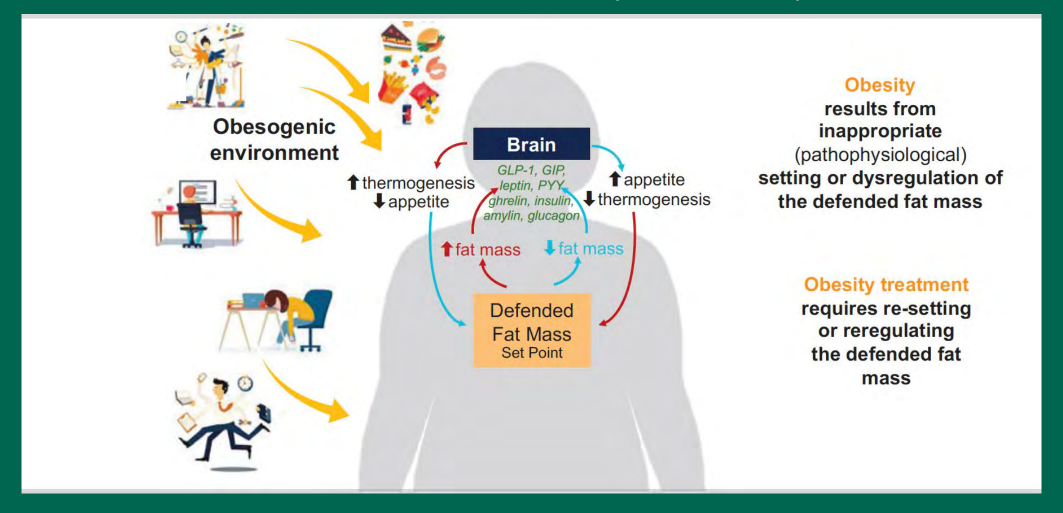
Neurohormonal Regulation of Weight

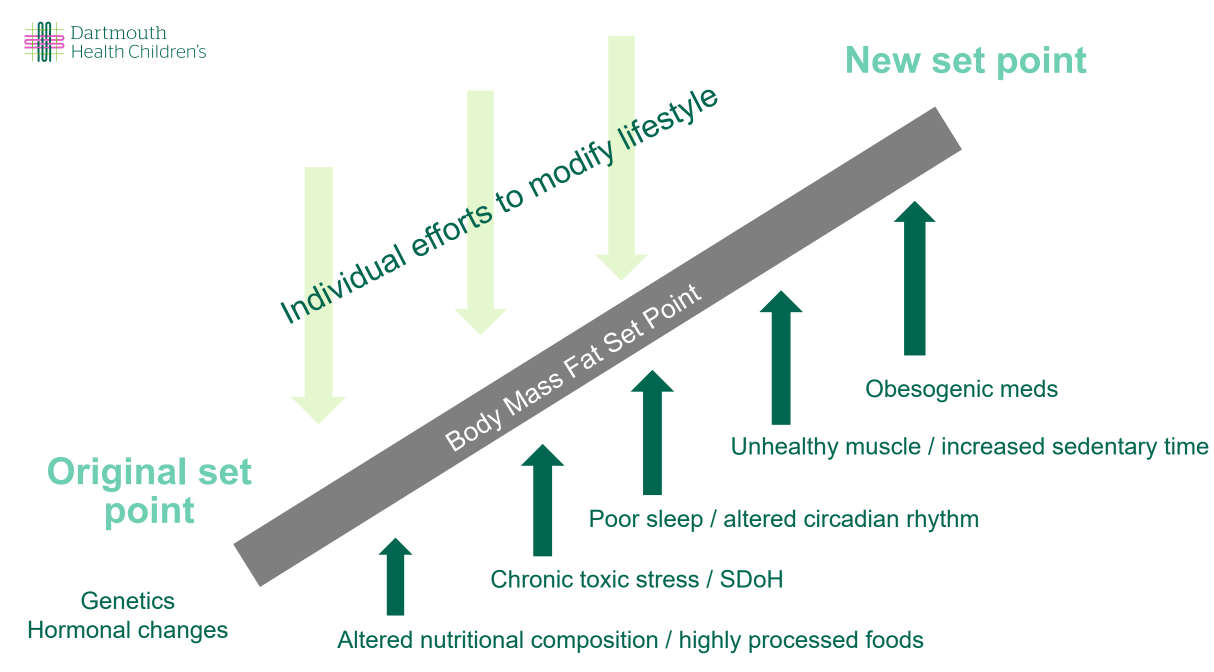
Complex interplay of hormones which regulate energy balance and determine set point (defended fat mass)



In obesity that set point is too high

Concept of Defended Fast Mass (Set Point)







What Obesity Is



Caused by countless factors (many/ most of which are not within the control of the individual) that collectively facilitate weight gain over time



Doggedly persistent, particular when surfacing early in life:

- If obesity surfaces in childhood, it is probably a particularly aggressive form of the disease
- >85% of youth with obesity will grow up to be adults with obesity





Altered

nutritional

composition

Highly

processed

diet

Durable Anti-Obesity Therapy (lifestyle, medication, surgery)



rhythms

Obesogenic Environment / drivers of obesity

sedentary time



Some Important Points to Consider

- Obesity is a complex and chronic disease!
 - Disease of abnormal physiology resulting from multi-factorial causes, including genetics, environment, and social determinants of health
 - Long term management will be necessary, and treatment response may vary

It is not

- a "choice," lack of willpower, or poor parenting
- a simple balance between calories in and calories out is much more complex



What is our job?

- Identification (Diagnosis)
 - Not just obesity as defined by BMI, but the disease of obesity
 - Identify complications such as insulin resistance, pre-diabetes, lipid abnormalities, fatty liver disease, hypertension, PCOS, etc.
- Understand the prognosis of each issue
 - Families can't make treatment decisions if they do not understand the prognosis and treatment options
- Discuss an initial (individualized) management plan
- Treat when able, refer when appropriate



Treatment Goals

- Goal is to <u>durably</u> reduce excess adiposity
- While concurrently treating obesity-related complications
- Watching and waiting is no longer an appropriate option

- For adults, even 5-10% weight loss is associated with improved cardiometabolic outcomes
- For children, goal is BMI reduction > 10%, even below 85-95th percentile, if complications persist

CLINICAL PRACTICE GUIDELINE Guidance for the Clinician in Rendering Pediatric Care



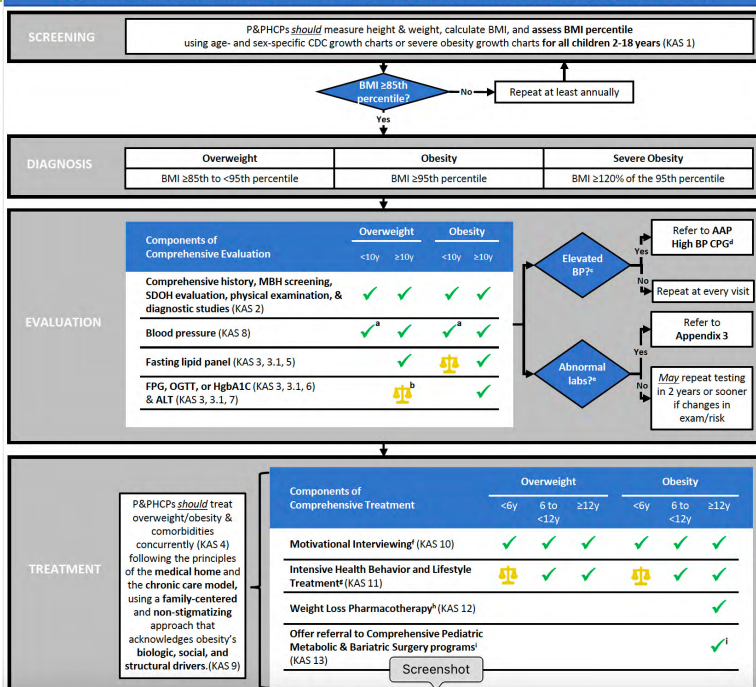
Clinical Practice Guideline for the Evaluation and Treatment of Children and Adolescents With Obesity



Clinical Practice Guidelines

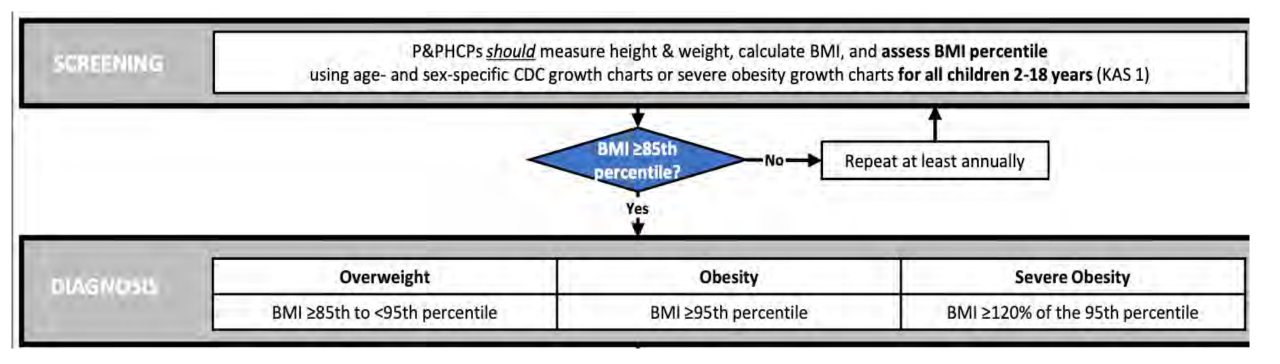
- Full Report
- Executive Summary
- AAP Decision Support Tools
 - Algorithm
 - Key Action Statements
 - Consensus
 Recommendations
- Found at AAP Institute for Childhood Healthy Weight

Algorithm for the Evaluation and Treatment of Children and Adolescents with Overweight and Obesity



Algorithm for Screening, Diagnosis, Evaluation, and Treatment of Pediatric Overweight and Obesity https://www.aap.org/en/patient-care/institute-for-healthy-childhood-weight/clinical-practice-guideline-for-the-evaluation-and-treatment-of-pediatric-obesity/

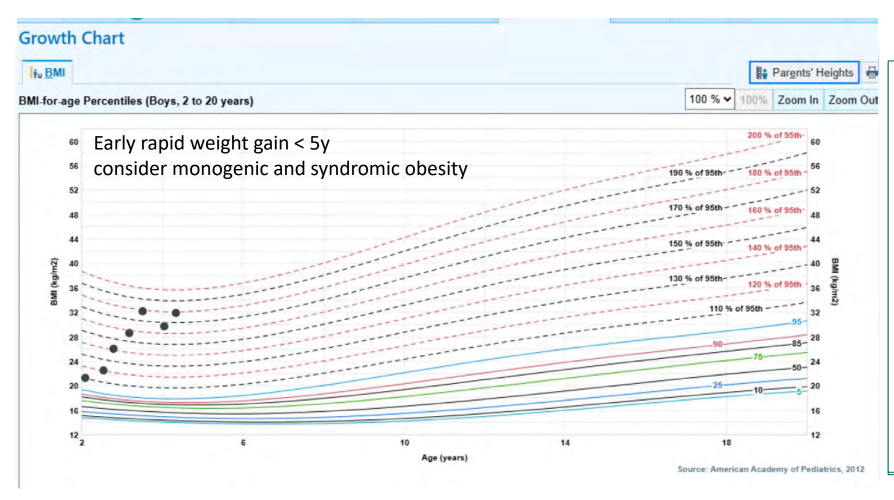




US Preventive Services Task Force Grade B evidence for screening for obesity using BMI



Extended Growth Charts



Class I obesity

BMI ≥ 95th percentile to < 120% of 95th percentile

Class 2 obesity

BMI \geq 120% to < 140% of 95th percentile or BMI \geq 35 to <40 kg/m²

Class 3 obesity

BMI \geq 140% of 95th percentile or BMI \geq 40 kg/m²

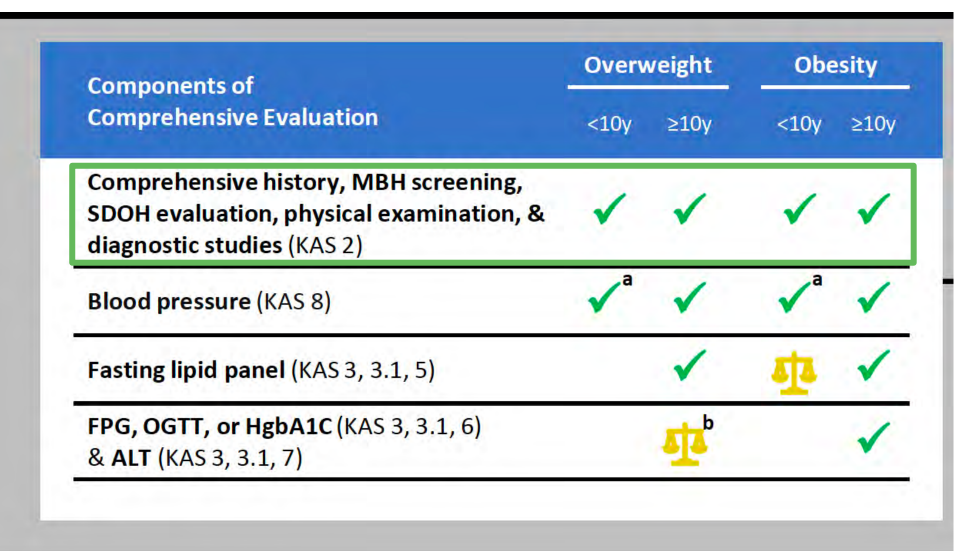






Screening for obesity-related complications

EVALUATION





Focused History

In children with overweight/ obesity evaluate for obesity-related co-morbidities:

- Assess individual, structural, and contextual risk and protective factors related to healthy behavior and healthy weight, including:
 - Medical history: chief complaint/history of present illness, <u>review of systems</u>, medication history, <u>family history</u>
 - Social determinants of health
 - Individual/family lifestyle behavior history
 - Mental and behavioral health, psychosocial consequences of living with obesity
- Physical exam
- Diagnostic studies



Acanthosis nigricans



Communicating with families

- Partner with families use respect, trust, open and objective communication
 - Avoid labeling by using patient first language (child with obesity)
 - Use words that are perceived as neutral by parents, teens, children
- Parents know their children best listen, with compassion
- Acknowledge that they have been working hard to address obesity
- Explain the multifactorial nature of obesity
 - What we can influence = obesity pillars (diet, activity, sleep, stress, meds)
 - What we can't, like our genetics or family history
- Share the story of obesity pathophysiology and target treatment to that



Treat obesity and co-morbidities concurrently

EVALUATION

Comprehensive Evaluation

Comprehensive history, MBH screening, SDOH evaluation, physical examination, & diagnostic studies (KAS 2)

Blood pressure (KAS 8)

Overweight Obesity

<10y ≥10y

✓10y ≥10y

✓10y

Volume 140, Issue 3

September 2017



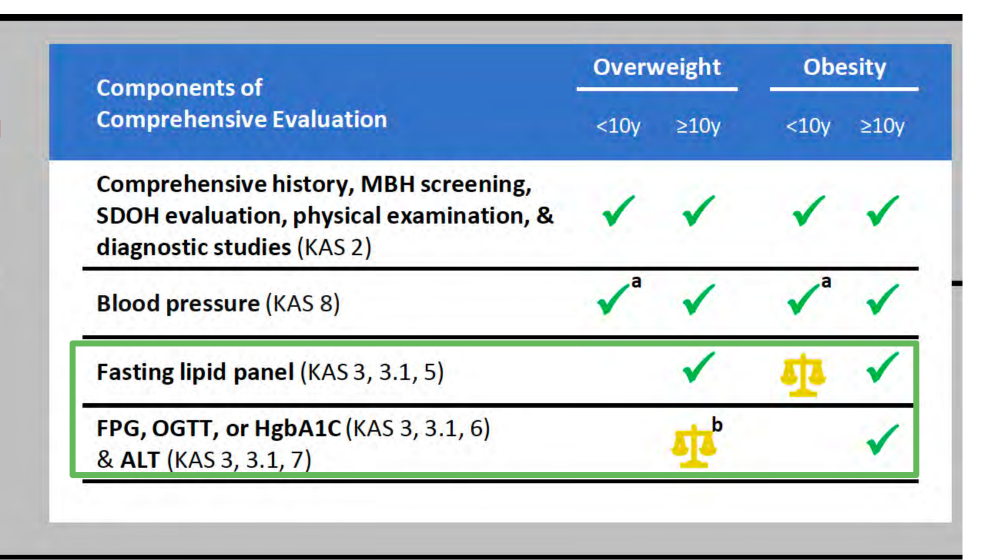
FROM THE AMERICAN ACADEMY OF PEDIATRICS | CLINICAL PRACTICE GUIDELINE | SEPTEMBER 01 2017

Clinical Practice Guideline for Screening and Management of High Blood Pressure in Children and Adolescents ⊘



Treat obesity and co-morbidities concurrently

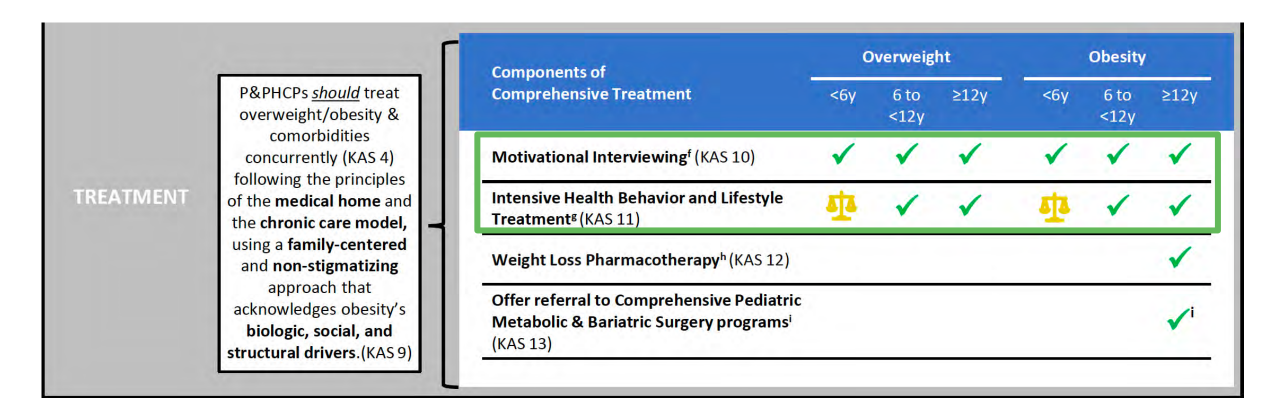
EVALUATION





Infancy (0-24 months)	Toddler (Age 2-4 years)	Early Childhood (Age 5-9 years)	Puberty (Age 10-14 years)	Adolescent (Age 15-18 years)
Weight>Length	BMI ≥ 95 th percentile Or ≥ 85 th percentile with 2 or more risk factors (24-48 months)	BMI ≥ 95 th percentile Or ≥ 85 th percentile with 2 or more risk factors	BMI ≥ 95 th percentile Or ≥ 85 th percentile with 2 or more risk factors	BMI ≥ 95 th percentile Or ≥ 85 th percentile with 2 or more risk factors
	 Fasting Blood Glucose and/or HbA1c Fasting Lipid Panel/Non fasting if fasting not feasible ALT Consider 25 OH Vitamin D, Consider iron studies BP annually if ≥ 3 years 			
- Consider Sleep Study - Consider Uric Acid - Consider fasting serum insulin				
Abnormal labs results for which additional testing is recommended: LDL ≥130; TG ≥100 (<10 years) or 130 (≥10 years); Prediabetes: HgbA1C ≥5.7 –6.4; FBS 100-125, OGTT 140-199; T2DM: FPG ≥126mg/dL, OGTT ≥200, HgbA1C ≥6.5; ALT≥2x upper limit of normal (≥52 males / ≥44 females) - Consider Urine Microalbumin/Creatinine ratio - Consider C-peptide, hs-CRP				

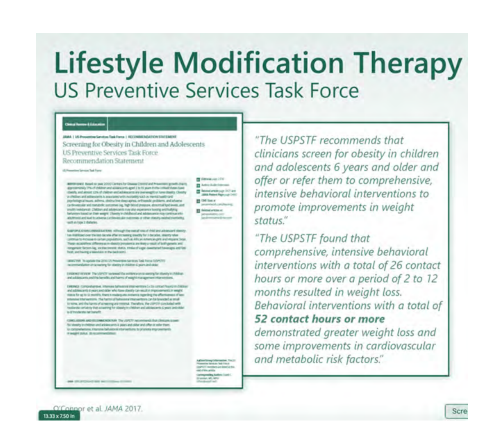






Intensive Health Behavior and Lifestyle Treatment

- Patient and family working with multidisciplinary team
 - Provider with training in obesity medicine
 - Other professionals with behavior and lifestyle and mental health expertise
- Focus on nutrition, activity, sleep, reduction of sedentary time, mental health, parenting skills...
- Longitudinal more effective with more contact hours
- Healthcare or community setting community partnerships are key, offer what is feasible





We have the tools

- For children and adolescents there may be a potential window of opportunity to prevent/treat inflammation and changes in hypothalamus that drive insulin resistance and increased set point
 - Health behavior and lifestyle change is a foundation
 - Switching out obesity promoting medications when possible
 - When appropriate, offering concurrent <u>adjunctive</u> treatment and / or referral to tertiary care for:
 - IHBLT
 - Weight loss medications
 - Metabolic & bariatric surgery



TREATMENT

P&PHCPs should treat overweight/obesity & comorbidities concurrently (KAS 4) following the principles of the medical home and the chronic care model, using a family-centered and non-stigmatizing approach that acknowledges obesity's biologic, social, and structural drivers.(KAS 9)

The Rationale For Obesity Pharmacotherapy

- Filling the treatment gap
- Ability to target underlying biological pathways regulating energy balance
- Potential for enhancement of weight loss maintenance
- · Potential to scale up





Series Sessions

Date	Session Title	
5/13/2025	Why Obesity is a Disease	
6/10/2025	Approach to the Patient with Obesity	
7/8/2025	Optimizing the Use of Lifestyle-based Obesity Care	
8/12/2025	How to Use Anti-Obesity Medications Effectively (GLP-1 agonist)	
9/9/2025	How to Use Anti-Obesity Medications Effectively (1 AOM non glp 1 agonist)	
9/23/2025	Approach to the Pediatric Patient with Obesity – AAP Clinical Practice Guidelines	
10/7/2025	How to Use Endoscopic Therapy Effectively	
10/21/2025	Pediatric Anti-Obesity Medications and Bariatric Surgery	
11/4/2025	Metabolic-Bariatric Surgery: Who, When, Why, and Which One	
11/18/2025	Improving Equitable Access to Obesity Care	

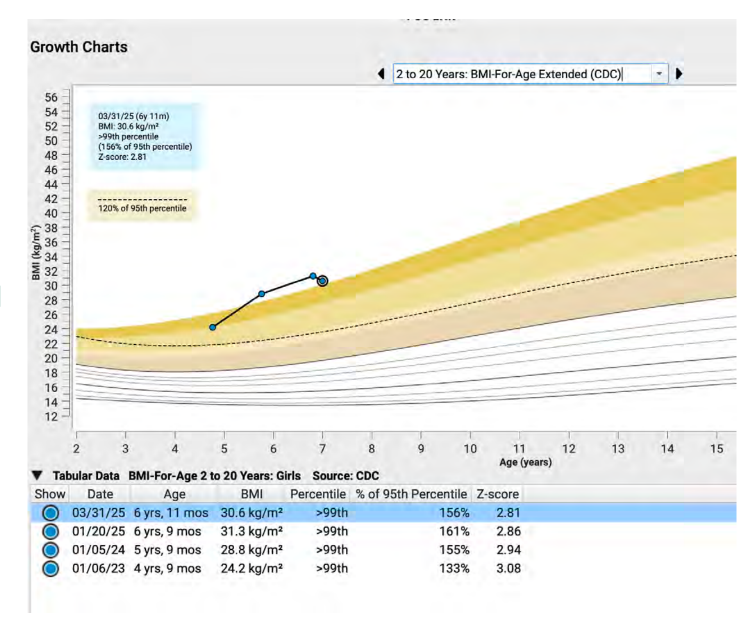


CASE DISCUSSION



Patient Overview

- Patient: 6-year-old female with obesity since toddlerhood
- Parental Concern: Mother prefers weight not discussed in front of child





Diet History

- What do you like to eat?: Hot dogs, chicken fingers, plain noodles, ramen, bagels
- Patient refuses all fruits and vegetables, including cucumbers, apples, strawberries, carrots
- Beverages:
 - At mom's: Water, milk, chocolate milk
 - At dad's: Soda, water, chocolate milk



Social History

• Family: Divorced parents, limited communication

- What do you like to do?:
 - Riding bikes
 - Playing at the park
 - "Doing moves" (demonstrates a cool dance move)



Clinical Data

Measurement

Weight

Height

BMI

Blood Pressure

AST / ALT

Hemoglobin A1c

Total Cholesterol

Triglycerides

HDL / LDL

Value

117 lbs

52 inches

30.6 kg/m2 (>99th

percentile)

100/80

17 / 24

5.3%

150

85

51/82





Discussion Question

How can we provide nutrition guidance to children with obesity in a way that:

- Avoids shame
- Protects self-esteem
- Encourages ongoing trust in medical care?





CASE #2

Presentation

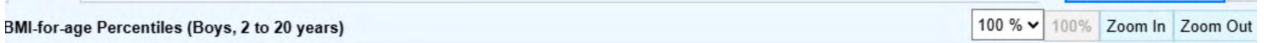
13 yo with BMI 130th% of 95th percentile seen with concern for weight.

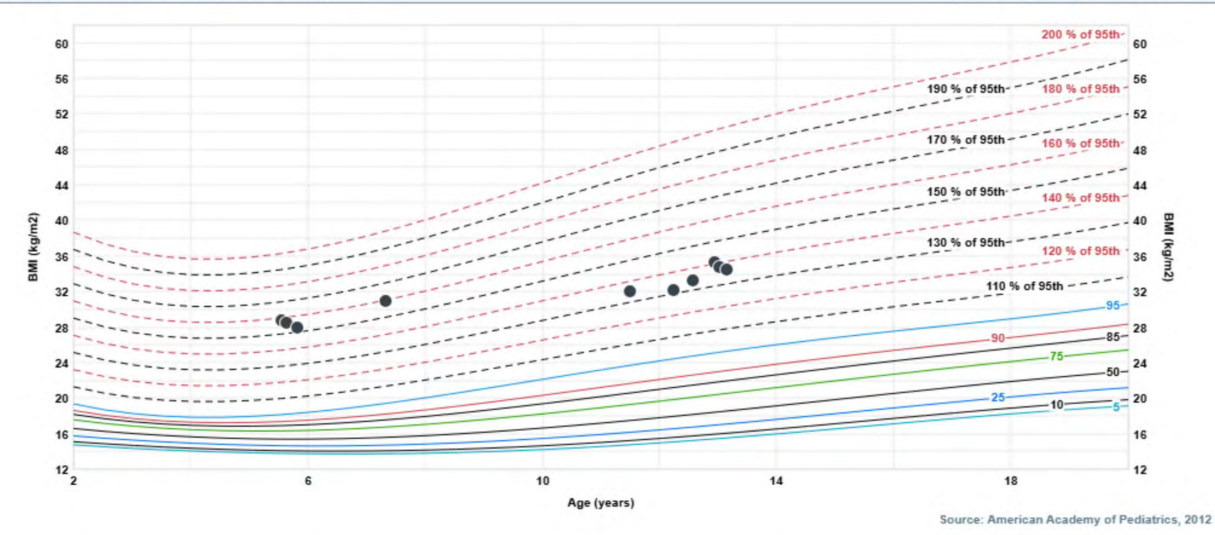
- Social History lives with parents and sister, parents work, supportive household
- Family history
 - Obesity on mom's side, MGM with Type 2 DM and MASH
 - Mom says she works hard to stay healthy weight –no prediabetes, but had GDM
 - No Fhx of bariatric surgery or AOM use. Dad and sister are thin, "can eat anything"
- Past medical history / ROS generally healthy, no asthma, constipation, headaches, snoring, mental health concerns. No medications

Growth Chart











Please Share Anything Else you feel is relevant to understanding the case

Weight history

 Parents say patient has always struggled with weight, even with intensive high school football, his weight drops some but then rebounds. Teen expresses frustration with being so active and not losing weight, being "stuck". Would like to try a medication

Review of health behaviors

- Mom says they have a very healthy diet at home but Caleb is much pickier than her other children, tends to be hungry all the time. Likes 1-2 vegetables, likes fruit but doesn't often choose it. Tends toward pasta, pizza, burgers, also snack foods which they have at home since younger sister needs to gain weight. They don't have soda at home but he drinks sweet tea, and energy drinks, which he buys with his own money
- Very active with football in fall, baseball in spring. No winter sport





Question(s) raised:

- What would be the next step in addressing weight concern in this teen?
- What evaluation would you do?
- What would your initial step in treatment be?



Evaluation

Labs

- A1C 5.7
- ALT 50 / AST 36
- Lipids with HDL 33 / Trig 215 / LDL 79

Abdominal Ultrasound

• Diffuse increased echogenicity suggestive of steatosis. Normal gallbladder



CASE #2

Presentation

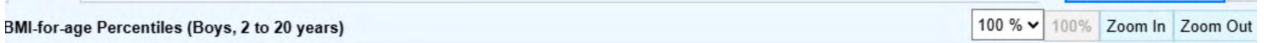
13 yo with BMI 130th% of 95th percentile seen with concern for weight.

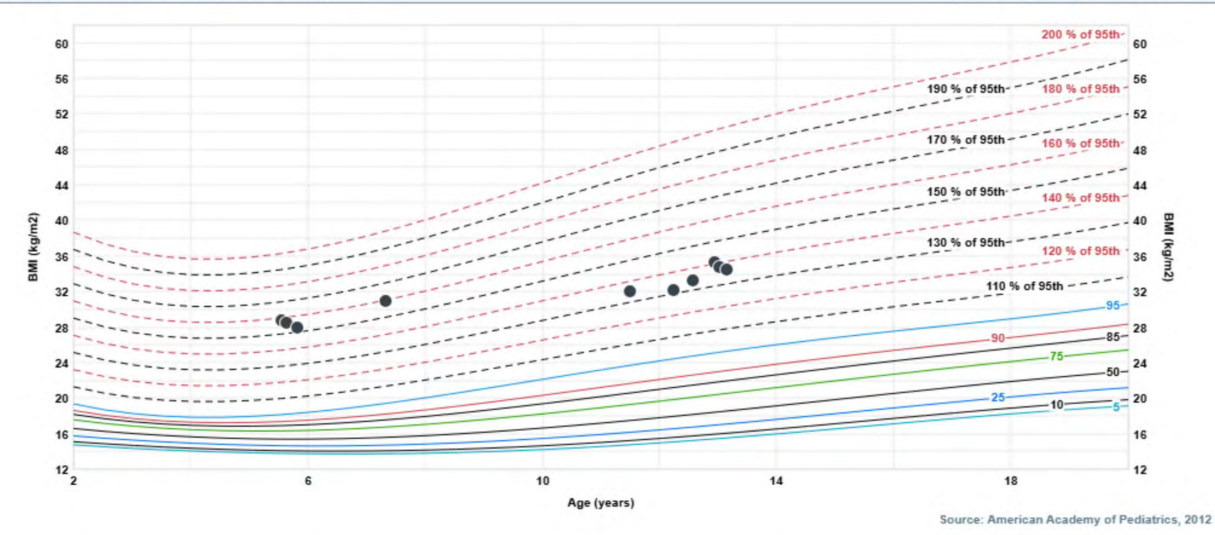
- Social History lives with parents and sister, parents work, supportive household
- Family history
 - Obesity on mom's side, MGM with Type 2 DM and MASH
 - Mom says she works hard to stay healthy weight –no prediabetes, but had GDM
 - No Fhx of bariatric surgery or AOM use. Dad and sister are thin, "can eat anything"
- Past medical history / ROS generally healthy, no asthma, constipation, headaches, snoring, mental health concerns. No medications

Growth Chart











Please Share Anything Else you feel is relevant to understanding the case

Weight history

 Parents say patient has always struggled with weight, even with intensive high school football, his weight drops some but then rebounds. Teen expresses frustration with being so active and not losing weight, being "stuck". Would like to try a medication

Review of health behaviors

- Mom says they have a very healthy diet at home but Caleb is much pickier than her other children, tends to be hungry all the time. Likes 1-2 vegetables, likes fruit but doesn't often choose it. Tends toward pasta, pizza, burgers, also snack foods which they have at home since younger sister needs to gain weight. They don't have soda at home but he drinks sweet tea, and energy drinks, which he buys with his own money
- Very active with football in fall, baseball in spring. No winter sport



Question(s) raised:

- What would be the next step in addressing weight concern in this teen?
- What evaluation would you do?
- What would your initial step in treatment be?



Evaluation

Labs

- A1C 5.7
- ALT 50 / AST 36
- Lipids with HDL 33 / Trig 215 / LDL 79

Abdominal Ultrasound

• Diffuse increased echogenicity suggestive of steatosis. Normal gallbladder





WELCOME to the Obesity Care in All Ages ECHO

Session 7, How to use Endoscopic Therapy Effectively

- October 7th , 2025





How to Use Endoscopic Therapy Effectively

Shelby Sullivan MD, FACG, FACG, DABOM

Director, Endoscopic Bariatric and Metabolic Program

Dartmouth-Hitchcock Medical Center and Geisel School of Medicine

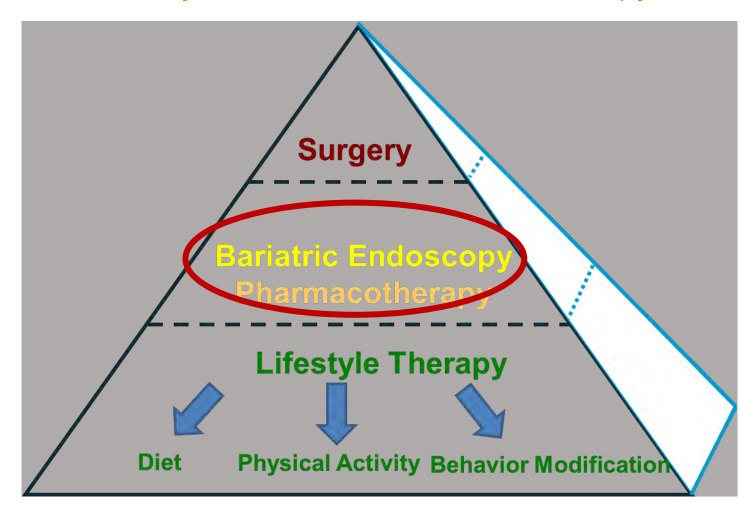


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 - Allurion Technologies, Fractyl Laboratories
- Consulting / Employment Last 24 Months
 - Allurion Technologies, Fractyl Laboratories, Biolinq, Pentax, Olympus



Comprehensive Obesity Treatment: Bariatric Endoscopy





What are is Endoscopic Therapy for Obesity?

Procedures and Devices that are performed or placed in the GI tract for weight loss or glycemic control



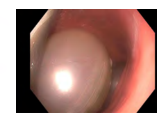
















Suturing and Plication Devices







Bariatric and Metabolic Endoscopy





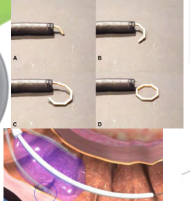




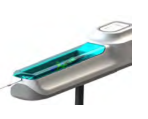




Small Bowel Therapies



Weight-Loss **Independent Effects**







Orbera Balloon approved for weight loss



Space Occupying Devices

Suturing and Plication Devices Overstitch are specifically approved for endoscopic sleeve gastroplasty and revision of bariatric surgery for obesity treatment

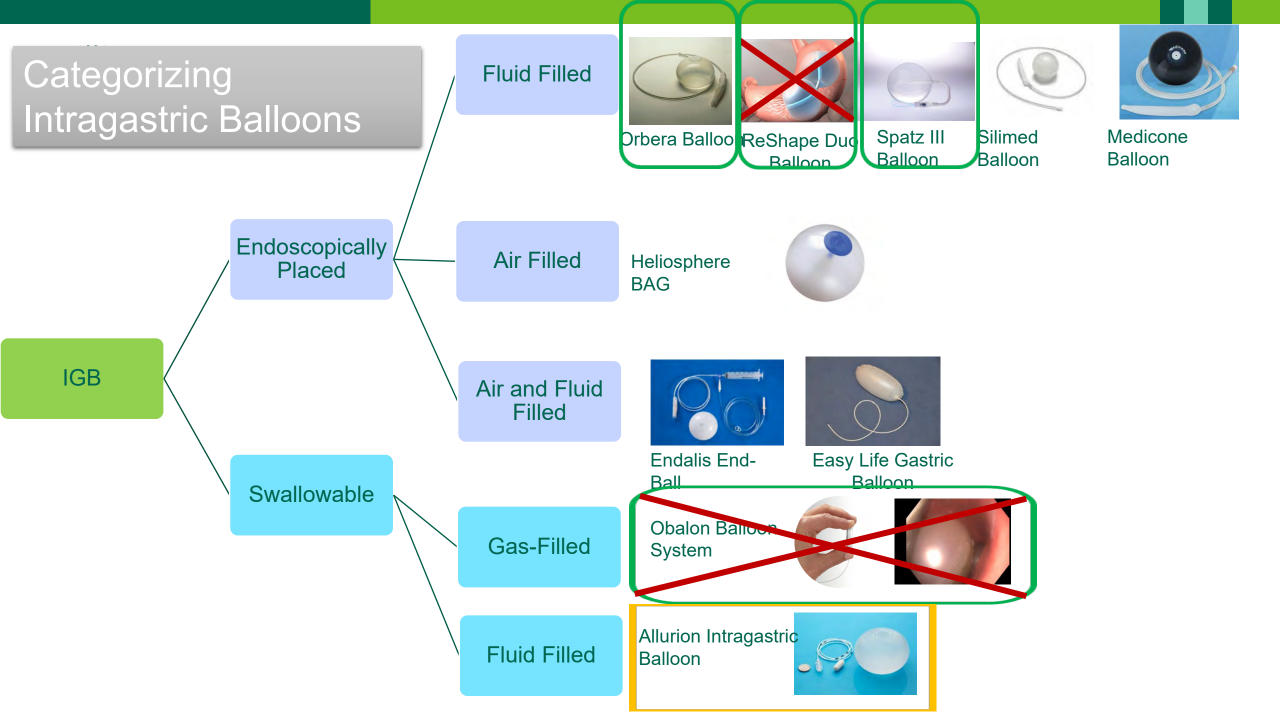
Bariatric Endoscopy Non-IGB Gastric Devices

Small Bowel Therapies FDA Approved And Current availability in the US

Spatz 3 Balloon approved for weight loss

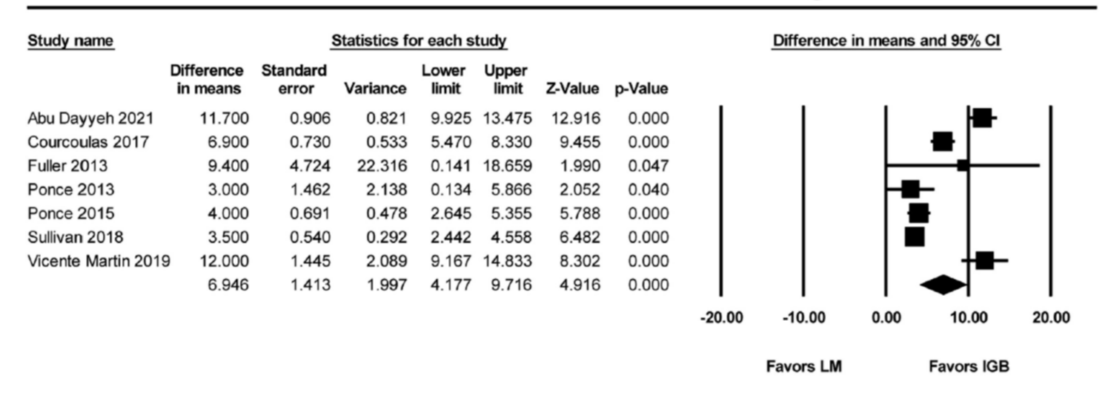


Intragastric Balloons



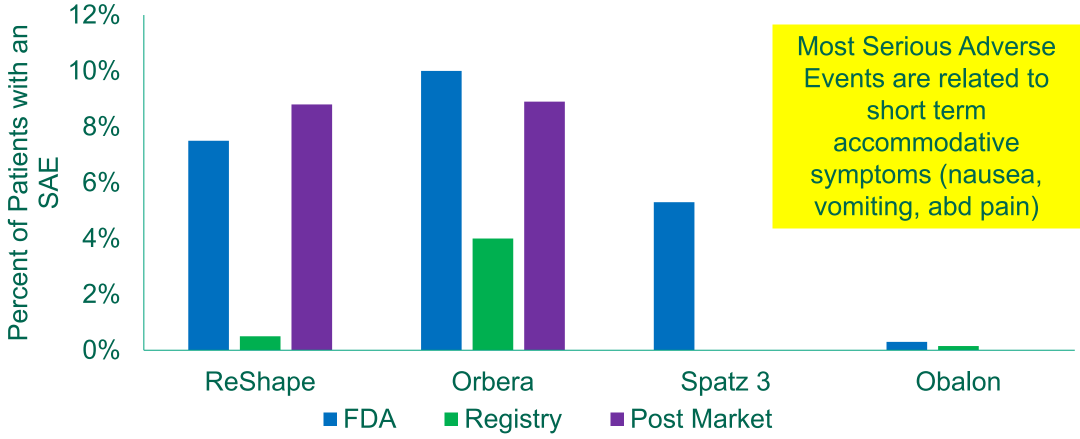
ASGE/ESGE EBMT Guideline: Placebo Subtracted IGB Efficacy

Mean Difference in %TWL at 6-8 Months Following IGB Placement





Intragastric Balloon: Serious Adverse Events in the US



Ponce J. Surgery for Obesity and Related Diseases. 2015;11(4):874-881 Agnihotri A. Clinical Gastroenterology and Hepatology 2018;16:1081–1088

Courcoulas A. *Int J Obes*. 2017;41:427-433 Vargas EJ. Clinical Gastroenterology and Hepatology 2018;16:1073–1080 Sullivan S. Surgery for Obesity and Related Diseases. 2018; 14(12):1876-188 Moore R. Surgery for Obesity and Related Diseases. 2019 Mar;15(3):417-423

https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfpma/pma_pas.cfm?c_id=3557&t_id=538679 Moore R. Obesity Surgery. 2020 Nov;30(11):4267-4274 Abu Dayyeh B. The Lancet. 2021;398:1965-1973

ASGE/ESGE EBMT Guideline: IGB

Recommendation	Strength of Recommendati on	Evidence
In adults with obesity, the ASGE-ESGE suggests the use of an IGB plus LM over LM alone	Conditional	Moderate
In adults with obesity undergoing IGB placement, the ASGE- ESGE suggests the use of antiemetics periprocedurally	Conditional	Very Low
In adults with obesity undergoing IGB placement, the ASGE- ESGE suggests the use of pain medication periprocedurally	Conditional	Very Low
In adults with obesity undergoing IGB placement, the ASGE- ESGE suggests the use of proton pump inhibitors while the IGB is in place	Conditional	Very Low

Jirapinyo P. GIE. 2024;99(6):867-885



AGA Clinical Practice Guidelines on Intragastric Balloons in the Management of Obesity	Strength of Rec	Quality of evidence
1. In individuals with obesity seeking a weight-loss intervention who have failed a trial of conventional weight-loss strategies, AGA suggests the use of IGB therapy with lifestyle modification over lifestyle modification alone. a, b	Conditional	Moderate
2. In individuals with obesity undergoing IGB therapy, AGA recommends moderate- to high-intensity concomitant lifestyle modification interventions to maintain and augment weight loss.	Strong	Moderate
3. In individuals undergoing IGB therapy, AGA recommends prophylaxis with PPIs.	Strong	Moderate
4. In individuals undergoing IGB therapy, AGA suggests using the intraoperative anesthetic regimens associated with the lowest incidence of nausea along with perioperative antiemetics. AGA suggests a scheduled antiemetic regimen for 2 week after IGB placement. [©]	Conditional	Low
5. In individuals undergoing IGB therapy, AGA suggests against perioperative laboratory screening for nutritional deficiencies.	Conditional	Low
6. AGA suggests daily supplementation with 1–2 adult dose multivitamins after IGB placement.	Conditional	Very low
7. After IGB removal, AGA suggests subsequent weight-loss or maintenance interventions that include dietary interventions, pharmacotherapy, repeat IGB, or bariatric surgery.		



Patient Selection

- Failed lifestyle therapy
- ➤ BMI FDA labelling is 30-40 kg/m2, no difference in safety or percent total body weight loss outside of that range
- Interested in a removal device
- Willing to abstain from NSAIDs and not on anticoagulation
- Willing to take PPI during balloon implantation
- Willing to participate in lifestyle therapy

Contra-indications

- Prior foregut surgery
- Cirrhosis
- Esophageal stricture
- Large hiatal hernia
- Need for anticoagulation
- History of PUD with unknown cause

Cautions

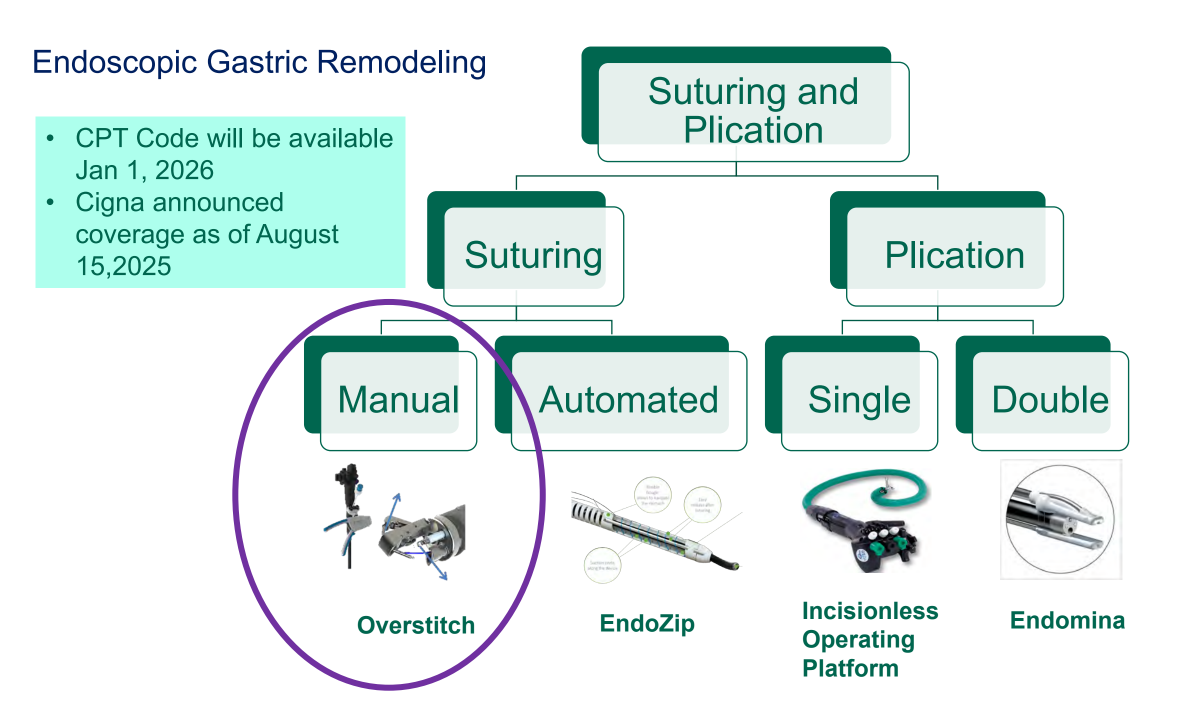
- Gastroparesis
- Poorly controlled GERD

Considerations

- More accommodative symptoms with fluid filled compared with gas-filled
- Gas-filled not currently available



Endoscopic Gastric Remodeling



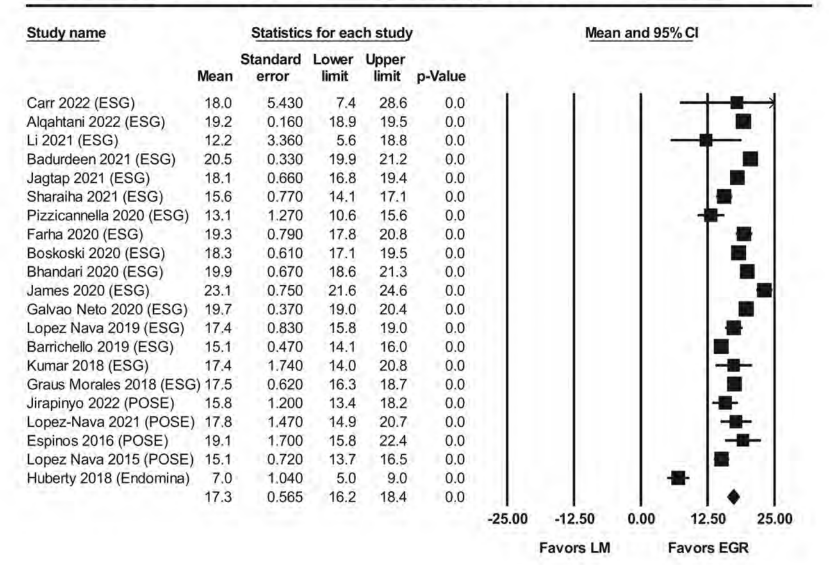


Endoscopic Gastric Remodeling

- Endoscopic Sleeve Gastroplasty
- Overstitch Device
- Most common EGR procedure in the US and around the world



%TWL at 12 Months Following EGR (Observational Studies Only)

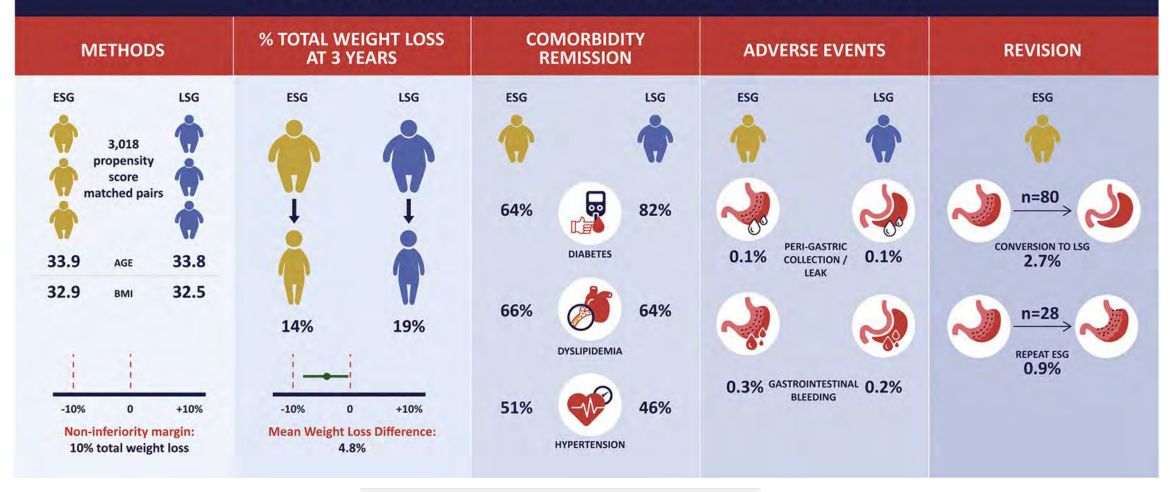


American Society for Gastrointestinal Endoscopy–European Society of Gastrointestinal Endoscopy guideline on primary endoscopic bariatric and metabolic therapies for adults with obesity Jirapinyo P. GIE. 2024. epub 4.19.24

Recommendation	Strength of Recommendation	Quality of Evidence
In adults with obesity, the ASGE-ESGE suggests the use of Endoscopic Gastric Remodeling (EGR) plus LM over LM alone	Conditional	Moderate
In adults with obesity undergoing EGR, the ASGE-ESGE suggests the use of antiemetics periprocedurally	Conditional	Very Low
In adults with obesity undergoing EGR, the ASGE-ESGE suggests the use of pain medication periprocedurally	Conditional	Very Low
In adults with obesity undergoing EGR, the ASGE-ESGE suggests the use of prophylactic antibiotics periprocedurally	Conditional	Very Low
In adults with obesity undergoing EGR, the ASGE-ESGE suggests the use of short-term proton pump inhibitors while the IGB is in place	Conditional	Very Low

Endoscopic Gastroplasty vs Laparoscopic Sleeve Gastrectomy:

A Non-Inferiority Propensity Score Matched Comparative Study



Follow-up:

ESG 78%-87% (85% at 3 years) LSG 71-87% (71% at 3 years)

Alqahtani AR. GIE. 2022;96:44-50

Cost Effectiveness Analysis: Semaglutide vs ESG

No. of months	Strategy	Costs, \$ Cumulati	For no	n-domir	nance w	ith (\$/QALY)	NMB, \$	EMI
50	No semaglutide or ESG	NA	FC(~ ~ ~ ~ 4 .	- 4 4 6 9 6	20	NA	37.8
12	ESG	17 229	an ES	J COST (of \$1636	00, 165	54 996	32.2
12	Semaglutide	11.742	Sama	alutida	must oo	ct	60 255	32.9
24	ESG	19 685	Sema	giuliae	must co	Sl	127 288	32.2
24	Semaglutide	22848	\$3591 or less with 20% or 1584		or '584	123 216	32.9	
36	ESG	19 685	ψυσυτι) 1033 V	VILIT 20 /	0 01	202 853	31.6
36	Semaglutide	33 688	less dro	n out o	ver 5 ve	ars 1580	186 515	32.9
48	ESG	19 685		p out o		, ai o	275 691	31,7
48	Semaglutide	43814	24129	2.92	-0.04	-617831	247 653	32.9
60	ESG	19685	awara.	3.66	0.00	0	345.854	31.7
60	Semagliitide	53 268	33 583	3.60	-0.06	-595 532	306 632	33.0

Abbreviations: BMI, body mass index (calculated as weight in kilograms divided by height in meters squared): ESG, endoscopic sleeve gastroplasty; ICER, incremental costeffectiveness ratio; NA, not applicable; NMB, net monetary benefit; QALY, qualityadjusted life-year,



Patient Selection

- Failed lifestyle therapy
- ➤ BMI FDA labelling is 30-50 kg/m², but current guidelines are BMI 27-29.9 kg/m² with co-morbidity or 30 kg/m² and above
- Interested in a semi-permanent procedure
- Willing to comply with post procedure diet
- Willing to participate in lifestyle therapy
- Can abstain from NSAIDs and anticoagulants during the post-op period

Contra-indications

- Cirrhosis
- Esophageal stricture
- Large hiatal hernia
- Need for surveillance of gastric mucosa (history of advanced dysplasia or gastric cancer, genetic diseases)

Cautions

- Gastroparesis
- Poorly controlled GERD
- Family history of gastric cancer

Considerations

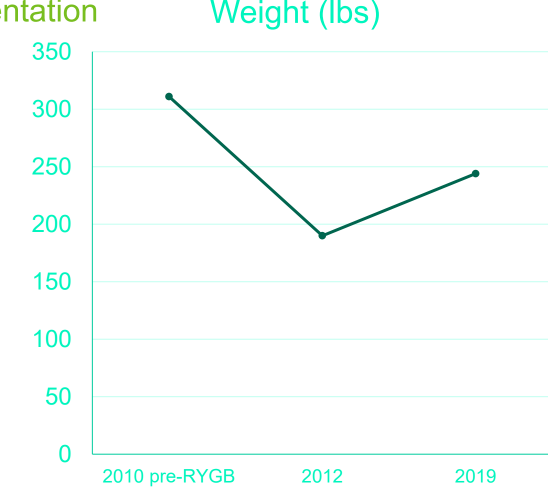
Less accommodative symptoms than the endoscopically placed fluid filled intragastric balloons



Revision after Bariatric Surgery

Weight Regain after RYGB: Case Presentation

- 47 yo woman with history of class III obesity, migraines and RA
- RYGB in 2010 at weight of 311 lbs
- Lost 121 lbs (39% TBWL), weight nadir 190
 lbs
- Started regaining weight in 2014
- Regained 54 lbs (44.6% weight regain) by July 2019
- Reports being able to eat 3 times as much at a meal compared with the first few years after surgery, does not feel restriction



Argon Plasma Coagulation (APC) Resurfacing Technique

- Low flow, high watts
- 2 to 3 rings on the gastric side of the anastomosis
- BID PPI therapy (and sucralfate) and liquid diet for 45 days after procedure
- Repeat every 8-10 weeks until stoma is reduced



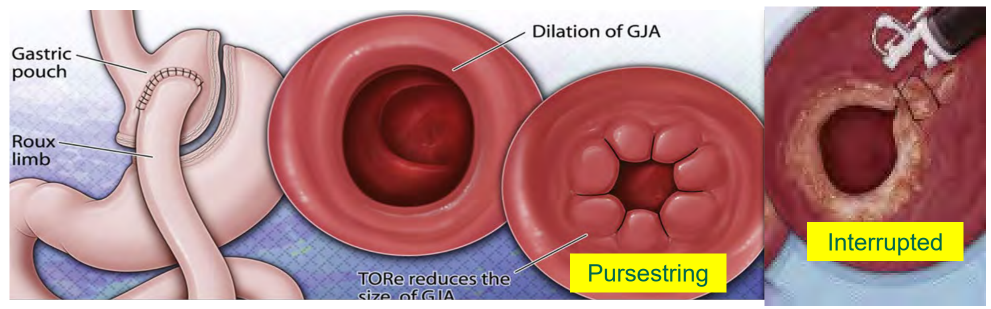




Storm AC. Gastrointestinal Endoscopy Clinics of North America. 2017;27:233-244

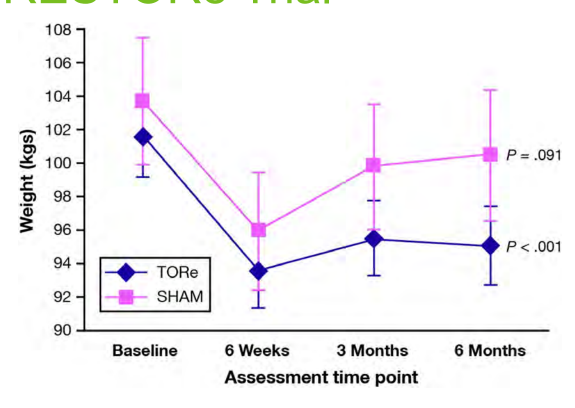
Transoral Outlet Reduction (TORe) Technique

- Prep of the gastric side of the anastomosis with APC or ESD
- Types of suturing
 - Interrupted
 - Purse-String



Vargas EJ. Surgical Endoscopy. 2018:32:252-259
Jirapinyo P. Gastrointestinal Endoscopy. 2020;91:1067-1073
Jaruvongvanich V. Gastrointestinal Endoscopy. 2020;92(6):1164-1175

Endoscopic Transoral Outlet Reduction (TORe): RESTORe Trial



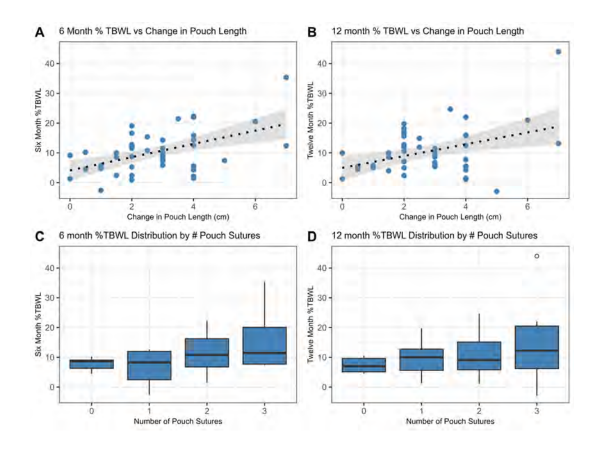
- Randomized Sham Controlled trial
 - TORe n=50
 - Sham control n=27
- Non-full thickness suturing system
- APC to the anastomosis
- Achieved weight loss or weight maintenance
 - Active 96% and Sham control 77.8%, p=0.019

TORe Data from University of Colorado

Post – Procedure Weight Loss Outcomes				
Group	6-month %TBWL	12-month %TBWL		
All (intention-to-treat)	10.2 ± 6.9%	10.3 ± 7.7%		
All (per-protocol)	11.3±7.6% (n=29)	12.2±9.2% (n=26)		
Purse-String	12.3±8.5% (n=21)	13.5±9.2% (n=21)		
Non-Purse String	8.7±3.7% (n=8)	7.0±7.9% (n=5)		

Two Serious Adverse events:

- Perforation in a patient not compliant with medications or diet
- Bleeding in a patient getting heparin with dialysis



Meyers MH. *Journal of Gastrointestinal Surgery.* 2023;27:1587-1593.

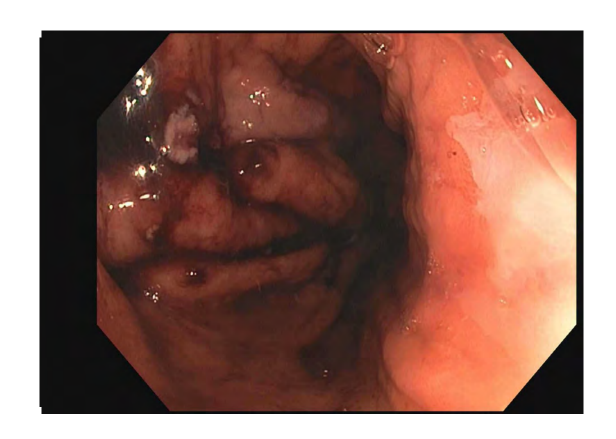
Comparison to Surgical Revision: Retrospective Matched Cohort

	Endoscopic (n = 31)	Surgical (n = 31)	<i>P</i> value
Age, y	48.6 (9.5)	48.1 (8.4)	.84
Female gender, n (%)	26 (83.9)	26 (83.9)	1.0
Time since RYGB, y	9.1 (3.2)	6.7 (4.1)	.01
Total weight loss from RYGB (%)	40.8 (8.7)	37.1 (12.7)	.2
Weight regain (%)	52.8 (34.1)	55.3 (29.5)	.77
Weight (kg)	111.1 (29.3)	114.6 (30.3)	.65
BMI (kg/m²)	40.5 (9.4)	41.5 (9.1)	.68
Adverse Events (%)	2 (6.5)	9 (29)	0.043
Serious Adverse Events	0 (0)	6 (19.4)	0.024



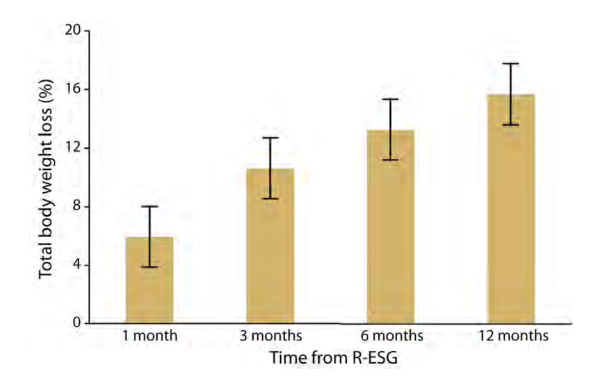
Sleeve Revision

- 59 yo woman with h/o class 2 severe obesity
 - Sleeve gastrectomy in 2008
 - 45 lbs weight loss (19% TBWL)
 - 107% weight regain
- Weight loss medications
 - Saxenda no significant weight loss
- Endoscopic sleeve revision performed
 - 13.2% TBWL at 7 months follow-up



Endoscopic Sleeve Revision

- Retrospective study
 - N=82
 - Weight regain: 27.9 \pm 20.7 kg
 - Weight at revision: 128.2 ± 57.5 kg
- Revision
 - Endoscopic sleeve gastroplasty within the dilated sleeve
- Adverse events:
 - Narrowed GEJ treated with dilation



Patient Selection

- Weight loss failure:
 - Weight regain of at least 20% of the weight lost
 - <50% excess weight loss from the time of surgery</p>
- Interested in a permanent procedure
- Willing to comply with post procedure diet
- Willing to participate in lifestyle therapy
- Can abstain from NSAIDs and anticoagulants during the post-op period

- Contra-indications
 - Cirrhosis
 - Esophageal stricture
 - Large hiatal hernia
- Cautions
 - Poorly controlled GERD
- Considerations
 - RYGB Any medical issues that could be in the small bowel or remnant stomach should be evaluated before revision
 - SG moderately controlled GERD or risk for Barrett's esophagus.



Conclusions

- Lifestyle therapy alone achieves only modest weight loss
- Lifestyle therapy maximizes weight loss with all adjunctive therapies
 - Anti-obesity medications
 - Endoscopic Bariatric Therapies
 - Bariatric Surgery
- Components
 - Diet
 - Exercise
 - Behavior Modification
- Can be done in a primary care practice if time is limited, focus on one goal at a time