

## Obesity: The Role of Pharmacotherapy

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## Disclosures: Professor John B Dixon

Apollo Endosurgery	Consultant
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BUPA	Research Support
I-Nova	Consultant
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Nestle Health Science	Consultant
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Novartis	Advisory board and speaker fees

## Pharmacotherapy for obesity AFP July 2017



Phung Qing Lee, John Dixon

Lee, P. C. and J. Dixon (2017). "Pharmacotherapy for obesity." *Aust Fam Physician* **46**(7): 472-477.

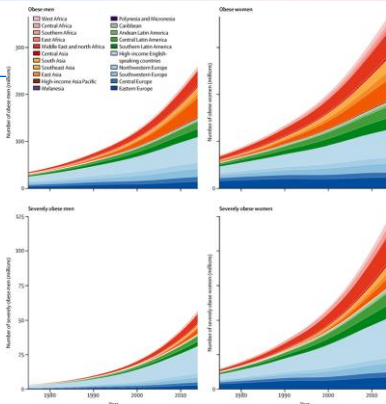


Global & Regional  
Obesity

High income  
English  
speaking  
countries

BLUE

Global & Regional  
Severe obesity



## ANNALS OF THE NEW YORK ACADEMY OF SCIENCES Issue: Five Year in Diabetes and Obesity

### Managing obesity in primary care practice: a narrative review

Raymond Carvajal,<sup>1</sup> Thomas A. Wadden,<sup>1</sup> Adam G. Tzai,<sup>2</sup> Katherine Pfock,<sup>1</sup>  
and Caroline H. Moran<sup>1</sup>

Clinically meaningful weight loss results are not  
achieved in the primary care setting— 1-3% at 6  
months to 2 years

These quite intensive combination behavioural  
programs struggle to achieve 5% or 5kg weight loss

Achieving 5% is currently not achievable for the  
majority at 12 months and less likely at 2-years

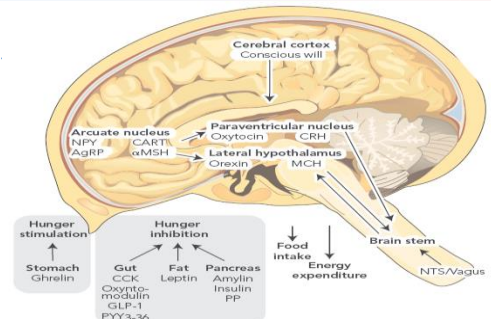
Carvajal et al. *Ann NY Acad Sci* 2013;1281:191-206.



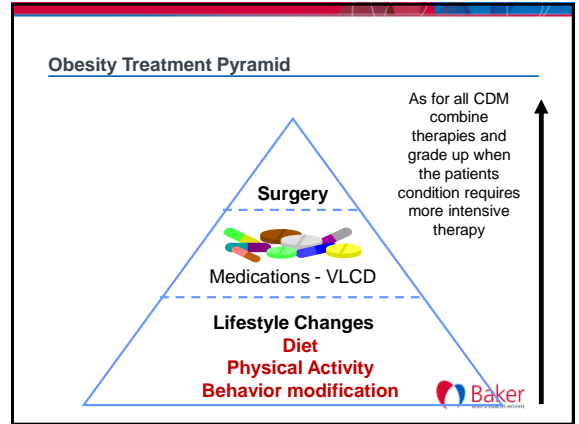
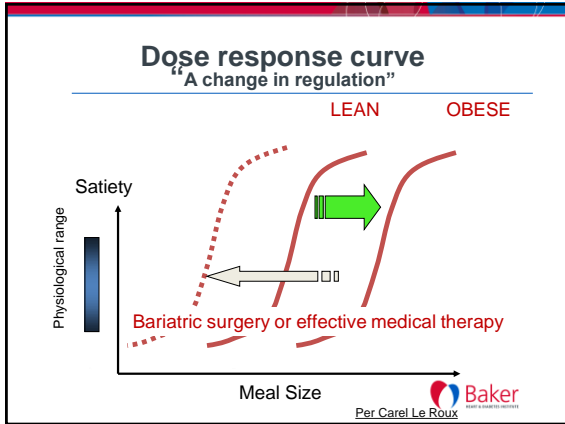
## Benefits of modest weight loss 3-10%

Obesity complication	Weight loss required for therapeutic benefit (%)	Notes	References
Diabetes (prevention)	3-10	Maximum benefit at 10%	DPP Research Group, 2009 (63) Garvey et al., 2014 (64)
Hypertension	5 to >15	Blood pressure still decreasing at >15%	Wing et al., 2011 (65)
Dyslipidemia	3 to >15	Triglycerides still decreasing at >15%	Wing et al., 2011 (65)
Hyperglycemia (elevated A1C)	3 to >15	A1C still decreasing at >15%	Wing et al., 2011 (65)
NAFLD	10	Improves steatosis, inflammation, and mild fibrosis	Asy et al., 2007 (66) Dixon et al., 2004 (67) Patel et al., 2009 (68)
Sleep apnea	10	Little benefit at 5%	Foster et al., 2009 (69) Winslow et al., 2012 (70)
Osteoarthritis	5-10	Improves symptoms and joint stress mechanics	Christensen et al., 2007 (71) Felton et al., 1992 (72) Aaboe et al., 2011 (73)
Stress incontinence	5-10		Burgio et al., 2007 (74) Subak et al., 2009 (75)
Gastroesophageal reflux disease	5-10 in women; 10 in men		Singh et al., 2013 (76) Tobias, 2011 (77)
Polycystic ovary syndrome	5-15 (>10 optimal)	Lowers androgens, improves ovulation, and increases insulin sensitivity	Panidis et al., 2008 (78) Norman et al., 2002 (79) Moran et al., 2013 (80)

Cefalu WT, Bray GA, Home PD, et al. *Diabetes care*. Aug 2015;38(8):1567-1582.



AgRP = agouti-related peptide. CART = cocaine and amphetamine-regulated transcript. CCK = cholecystokinin. CRH = corticotropin-releasing hormone. GLP-1 = glucagon-like peptide. MCH = melanin-concentrating hormone. αMSH = alpha melanocyte-stimulating hormone. NPY = neuropeptide Y. NTS = nucleus of the tractus solitarius. PP = pancreatic polypeptide. PYY = peptide YY.



### Mechanisms

Reduce energy intake  
Nutrient malabsorption

**Suppress appetite by effects on the central control of energy balance**

Increase energy expenditure

- Stimulate brown fat
- Non-exercise activity thermogenesis (NEAT)
- Generate metabolic inefficiency

**Aim: To reset energy balance**

Baker

### Indications for weight management pharmacotherapy

BMI >30 kg/m<sup>2</sup>, or those with a  
BMI of 27–30 kg/m<sup>2</sup> with obesity-related risks and complications.

Lower BMI thresholds (BMI >27 kg/m<sup>2</sup>, or BMI >25 kg/m<sup>2</sup> with obesity-related complications) should be considered in Aboriginal and Torres Strait Islander and Asian populations.

Baker

### The importance of a stopping rule

There is no point

- in taking a drug that is not effective
- in continuing a drug that produces unacceptable side-effects
- in taking drug if it increases net risk of future disease

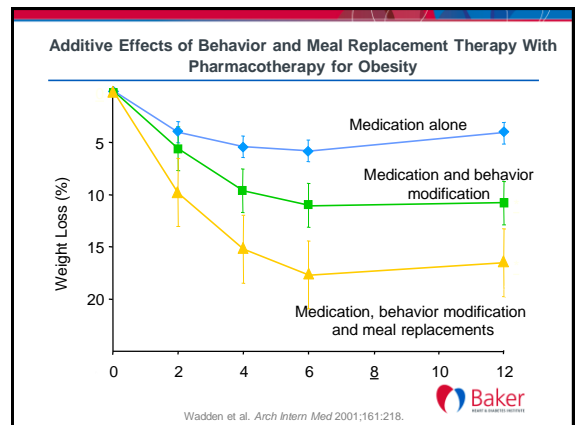
There is no point

- in stopping an effective drug if well tolerated and reduces risk

3-months on the full dose is usually a sufficient time to assess effectiveness

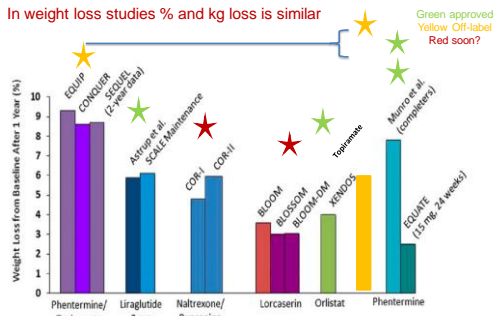
In chronic disease management we often combine therapy for greater efficacy so we need additional time if we increase therapy when synergy is expected

Baker



## Comparison of weight loss medications at 1-year (placebo subtracted – ITT – LOCF)

In weight loss studies % and kg loss is similar



I am going to focus on the use of the 4 drugs we have available in Australia today

Orlistat (Xenical, Alli)

Phentermine (Duromine, Metermine)

Topiramate (Many) Off Label for weight loss

Liraglutide (Saxenda)

How to use them and combine them?



## Orlistat 120mgs tds (oral)

### ACTION

Inhibits intestinal lipase - reduces fat absorption by 30%.

### EVIDENCE

In a one-year clinical study, weight loss of 9 kg was reported in the orlistat group, compared with 5-6kg for the placebo-treated group

XENDOS (XENical) in the Prevention of Diabetes in Obese Subjects) study, at four years, it reduced the risk of developing diabetes by 37.3% compared with the lifestyle intervention alone

Torgerson JS, Hauptman J, Boldrin MN, Sjöström L. XENical in the prevention of diabetes in obese subjects (XENDOS) study: A randomized study of orlistat as an adjunct to lifestyle changes for the prevention of type 2 diabetes in obese patients. Diabetes Care 2004;27(1):155-61.



## Orlistat prevention of diabetes study XENDOS

4-year, double-blind, prospective study

3,305 randomized to lifestyle changes plus either orlistat 120 mg or placebo, three times daily

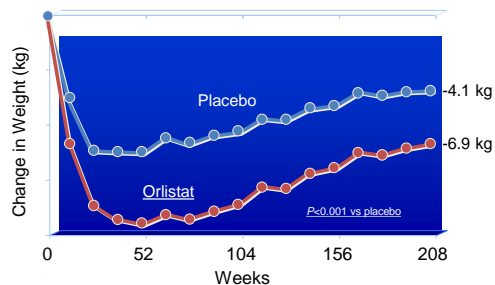
BMI  $\geq 30$  kg/m<sup>2</sup> and normal (79%) or impaired (21%) glucose tolerance (IGT)

Primary endpoints were time to onset of type 2 diabetes and change in body weight.

Torgerson JS. Et al. Diabetes Care 2004;27:155-61.



## Effect of Long-term Orlistat Therapy on Body Weight



Torgerson et al. Diabetes Care 2004;27:155



## Results at 4 years

52% completed in the treatment group compared with 34% of placebo recipients ( $P < 0.0001$ )

Diabetes incidence 9.0% with placebo and 6.2% with Orlistat a risk reduction of 37.3% ( $P = 0.0032$ )

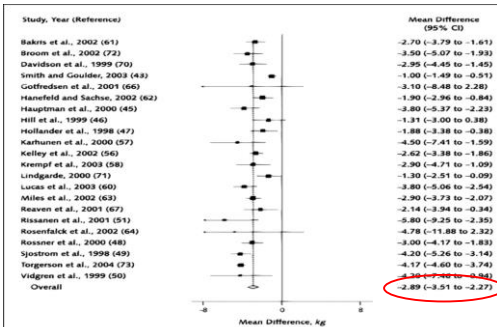
Orlistat plus lifestyle changes resulted in a greater weight loss and reduction in the incidence of type 2 diabetes.

The latter restricted to the IGT group

Torgerson JS. Et al. Diabetes Care 2004;27:155-61.



### Weight loss with orlistat versus placebo at 12 months



Li, Z. et al. Ann Intern Med 2005;142:532-546



### Side effects – related to fat malabsorption

- Steatorrhea, oily spotting, bloating and flatulence with discharge, faecal incontinence
  - Effects attenuated on a low fat diet
- Oxalate kidney stones – associated with fat malabsorption
- Fat-soluble vitamin deficiency – in the long term
  - Supplement and use caution with patients on warfarin
- This drug is well tolerated in the long term and has beyond weight loss benefits in those with type-2 diabetes

Jacob S, Rabbia M, Meier MK, Hauptman J. Orlistat 120 mg improves glycaemic control in type 2 diabetic patients with or without concurrent weight loss. Diabetes Obes Metab 2009;11(4):361-71.



### All other medications act centrally to reduce energy intake

When asked “What is the effect of the drug?” obese patients treated with anti-obesity drugs offer a wide variety of answers such as:

“I don’t eat as much.”

“I can stop eating.”

“I don’t graze all day and night.”

“I’m not hungry as soon as I stop eating.”

“I’m normal” (in respect to eating).

### 3 - Factor eating questionnaire

Improved cognitive restraint

Lower levels disinhibition

Reduced hunger



### Phentermine

15mgs, 30mgs, 40mgs (oral)

Prepared with a slow release resin – single daily dosage

Approved in the 1960s this drug has been the most commonly used weight management drug in Australia and the United states for decades

#### ACTION

Phentermine is a sympathomimetic amine with significant anorectic activity in animal models.

It’s appetite suppressant effect is generally considered to be exerted through the hypothalamus, but it is not certain that this is the only effect related to weight loss.

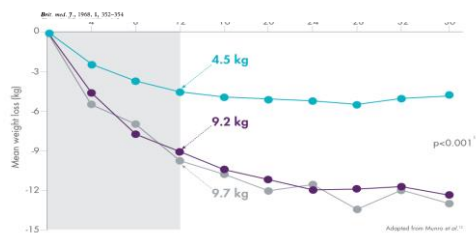
#### EVIDENCE

Differences in weight loss at 6 months of 3.6 – 4.5 kg or %WL for phentermine-treated patients vs placebo



### Comparison of Continuous and Intermittent Anorectic Therapy in Obesity

J. F. MUNRO\* M.B., M.R.C.P.E.D.; A. C. MCCOISNE\* M.B., Ch.B.  
ELIZABETH M. WILSON\* M.B., B.S.; L. J. P. DUNCAN\* M.B., B.Sc., F.R.C.P.E.D.



- Placebo once-daily (continuous regimen) + 1,000 Cal/day dietary advice, n=25
- Duramine™ 30 mg once-daily (continuous regimen) + 1,000 Cal/day dietary advice, n=17
- Duramine™ 30 mg once-daily for 4 weeks alternating with placebo once-daily for 4 weeks (alternate regimen) + 1,000 Cal/day dietary advice, n=22



### Phentermine Contraindications and Precautions

Unstable hypertension, history of heart disease, hyperthyroidism, anxiety disorders, Hx of Drug & Alcohol abuse, Major psychiatric illness, pregnancy, breast feeding, MAOIs, and glaucoma

- Caution with combined use with SSRI’s, ergot drugs, and clomipramine



## Phentermine

It is primarily a sympathomimetic  
 It's effects on dopamine and serotonin are trivial  
 Therefore it has little or no addictive potential  
 While it may be expected in some to raise blood pressure there is no clear evidence that it does  
 No evidence of increased CV risk  
 There is generally the expected fall in BP associated with weight loss



## Phentermine Treatment

Start with Duromine 15 mg/day.  
 Most adult patients tolerate 30 mg/day some may need 40mg  
 Evaluate for adverse effects.  
 Evaluate for effectiveness  
 Titrate dose to effectiveness

- Weight loss
- Eating behavior – smaller meals – satiety – hunger - control

Tachyphylaxis with lower effect  
 Higher doses can be used – up to 40 mg

I recommend taking the dose in the morning



## Side Effects

### Common

- Dry mouth - usually tolerable
- Insomnia – typically fades quickly
- Increased energy
- Feeling anxious / palpitations
- Other – e.g. constipation

Warn patients of these common early issues  
 They usually resolve spontaneously

### Less Common

- Impotence, decreased sex drive
- Irritability
- Mood elevation

If phentermine is effective and there are no adverse effects it can be continued



## When may phentermine be continued on a long term basis?

- low-to-intermediate cardiovascular risk with no evidence of serious cardiovascular disease
- no serious psychiatric disease or history of substance abuse
- no clinically significant increase in pulse or blood pressure while taking phentermine
- close monitoring – monthly during dose escalation and at least every three months thereafter
- efficacy–safety stopping rule is followed

Phentermine is now US-FDA approved on a long term basis with topiramate.  
 Studies out to 2-years at this stage

Lee, P.C. and J. Dixon (2017). "Pharmacotherapy for obesity." Aust Fam Physician 46(7): 472-477.



## Topiramate

Approved for seizures in 1996  
 Approved for migraine prevention in 2004  
 Mono-therapy not approved for obesity

### Doses

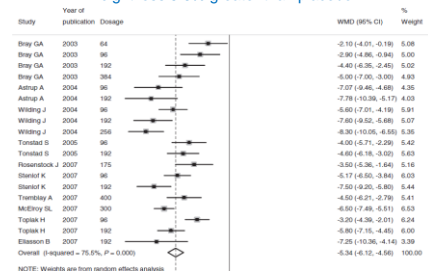
- Epilepsy: 400 mg/day
- Migraine prevention: 25 - 100 mg/day
- Obesity: 25 – 100 mg/day

Starting Rx 12.5 or 25 mg/day, titrate dose slowly



## Meta-analysis of Topiramate RCTs

Weight loss 5.3% greater than placebo



Verrotti, A. et al. (2011). "Topiramate-induced weight loss: a review." Epilepsy Res 95(3): 189-199.



## Topiramate – Side effects

Paraesthesia, dry mouth, altered taste sensation, constipation, dizziness, insomnia, fatigue, somnolence

Cognitive effects may include psychomotor slowing, decreased concentration and attention, memory impairment, and language difficulties

Common mild dose related and usually resolve

Major mood change - suicidal thoughts or ideation

Rare rapid onset serious and drug induced

Acute Myopia & Angle Closure Glaucoma

Increased risk of oral clefts if taken during pregnancy in first trimester

Warn and manage risk

Contraindications: Pregnancy, Glaucoma, Kidney Stones



## Using Topiramate Alone

Start with 25 mg/day; best given at night first

Stay at 25 mg/day at least 2 weeks

Evaluate for ASEs, cravings, binge eating

If marked improvement stay at 25 mg/day

If no ASEs consider increase to 50 mg/day

If appropriate increase to 75mg/day and 100mg/day (maximum)

If ASEs either reduce dose or stop depending on the nature of the ASE

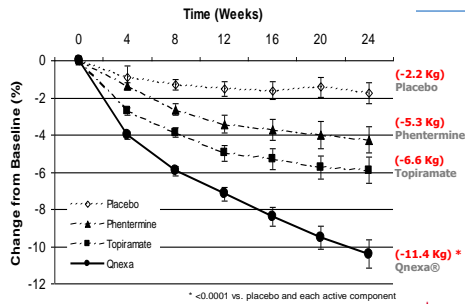
If no cravings or binge eating look for weight loss +/- changes in eating behavior.

I recommend taking the dose at night



## Qsymia®: 24 Week Weight Loss

Topiramate and Phentermine (ITT)

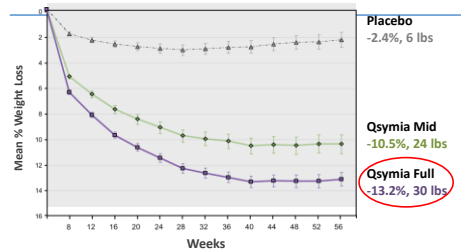


<http://lr.vivus.com/releaseDetail.cfm?ReleaseID=407933>



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## Qsymia : Weight Loss Over Time (Completer Population)



	Placebo	Mid	Full
Patients	564	344	634
Completers (% of randomized)	57%	69%	64%

1. Statistically greater number of patients completing study on Qnexa vs. placebo,  $p < 0.0001$   
 \* Data from patients that completed 56 weeks on treatment



## Liraglutide 3mg for weight management (TGA approved December 2015)

Liraglutide 3.0 mg

Liraglutide is a human glucagon-like peptide (GLP-1) analogue, with 97% amino acid sequence homology to endogenous human GLP-1

Like endogenous GLP-1, liraglutide binds to and activates the GLP-1 receptor (GLP-1R)

GLP-1 is a physiological regulator of appetite and calorie intake

GLP-1 is hormone secreted from the distal gut in response to a meal  
 Slows gastric emptying = sense of fullness  
 Satiety

Central action to reduce hunger and provide prolonged satiety

Saxenda® Approved Product Information, December 2015



## Weight loss with liraglutide 3.0 mg

Significantly greater and clinically meaningful weight loss in a wide range of patients

Obesity and prediabetes patient population <sup>1,2</sup>	8.0% at week 56 compared with 2.6% in the placebo group ( $p < 0.0001$ )
Type 2 diabetes population <sup>1,3</sup>	6.0% at week 56 compared with 2.0% in the placebo group ( $p < 0.0001$ )
Obstructive sleep apnoea population <sup>1</sup>	5.7% at week 32 compared with 1.6% in the placebo group ( $p < 0.0001$ )

Treatment arm = Liraglutide 3.0 mg plus diet and exercise; Placebo = diet and exercise alone. Data are for patients in the full analysis set, with last observation carried forward. Changes from baseline are estimated mean weight loss

1. Saxenda® Approved Product Information, December 2015. 2. Pi-Sunyer X et al. N Engl J Med 2015;373:11-22.  
 3. Davies MJ et al. JAMA. 2015;314(7):687-699.



## Liraglutide weight-loss review



Patients who lost at least 5% of baseline weight after 16 weeks of treatment on average lost 11.2% of their baseline weight at week 56



Saxenda® Approved Product Information, December 2015. Pi-Suryer X et al. N Engl J Med 2015;373:11-22.



## Summary of liraglutide 3.0 mg safety profile

GI side effects are common	Most episodes of GI events were mild to moderate, transient and the majority did not lead to discontinuation of therapy
Dehydration	Potential risk of dehydration in relation to GI side effects
Gallbladder-related events	Associated with above average weight loss
Low rates of pancreatitis	Mild grade and of short duration Half the liraglutide-associated pancreatitis cases were associated with gallstones

Pi-Suryer X et al. N Engl J Med 2015;373:11-22. Saxenda® Approved Product Information, December 2015.



Drug	Starting dose	Available doses	Weight loss versus placebo (% or kg)	Side effects	Contraindications
Phentermine	15 mg	15, 30, 40 mg	3.6-4.5 kg at six months	Dry mouth, insomnia, agitation, constipation, and tachycardia	Severe hypertension, cardiovascular disease, glaucoma, history of drug or alcohol abuse, monoamine oxidase inhibitors, selective serotonin reuptake inhibitor use, pregnancy
Orlistat	120 mg TDS	120 mg	2.9-3.4% at one year	Statorrhoea, oily spotting, flatulence with discharge, faecal incontinence, fat-soluble vitamin malabsorption	Pregnancy
Liraglutide	0.6 mg	0.6-3.0 mg	5.4% at one year	Nausea, vomiting, diarrhoea, constipation Rare: Pancreatitis, cholecystitis	Severe renal or hepatic insufficiency, pregnancy, past history of pancreatitis and major depression or psychiatric disorder
<b>Off-label pharmacotherapy (not approved by Therapeutic Goods Administration for weight loss)</b>					
Topiramate	12.5 mg mane	25, 50, 100 mg	3.4-5.0 kg	Paresthesia, dry mouth, constipation, altered taste sensation, insomnia, dizziness, cognitive effects Rare: Closed angle glaucoma, depression or suicidal ideation	Glaucoma, renal stones, pregnancy (if used for weight loss)
Phentermine (Phe)/topiramate (Top)	Phe: 15 mg mane Top: 12.5 mg mane	Phe: 15 mg Top: 12.5, 25, 50, 100 mg	5.0-6.6% at one year	Side effects of phentermine and topiramate	Contraindications to phentermine or topiramate

Lee, P. C. and J. Dixon (2017). "Pharmacotherapy for obesity." Aust Fam Physician 46(7): 472-477.



## Where and how do we use medications?

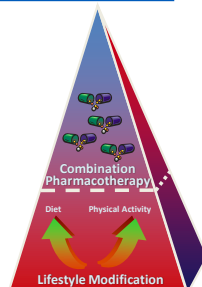
If you cannot provide the behavioural program out source it

- To support weight loss and weight maintenance
- Reduce weight regain
  - Diets, VLCD, balloons, surgery, or any other .....
- The action plan always has short and long-term aims
- Combine medications to enhance effect and reduce individual doses
- Start low and go slow with the dosage
- Remember the stopping rule



## Obesity Treatment of the future

As for dysregulation of blood pressure and blood glucose we will need combination drug therapy with lifestyle interventions to successfully manage clinically severe obesity



Adapted from source: www.obesityonline.org