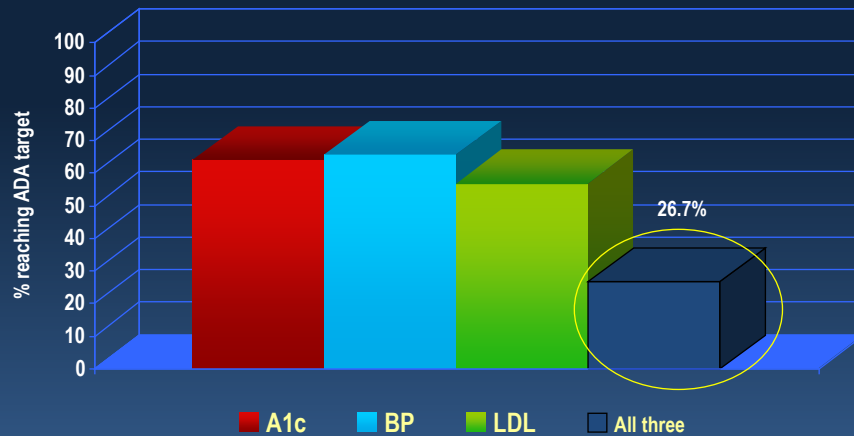

Lecture 1: 9:30– 10:30 a.m. PST

William Polonsky, PhD, CDCES, Presents:

Understanding and Addressing Problematic Adherence to Oral and Injectable
Cardiometabolic Medications

Patients Achieving Targets: 2014



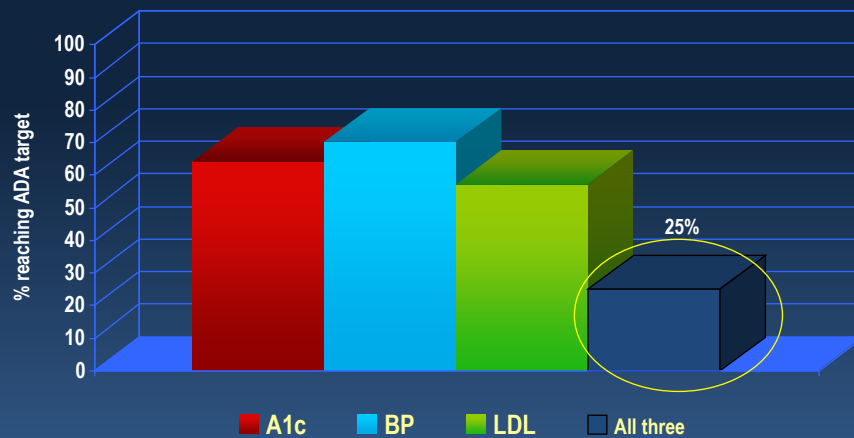
NHANES data: Ali et al, 2014

The Key Behavioral Contributor to Glycemic Control

Outcome: HbA1c (%)	Model 1: all self-care behaviours β	Model 2: all self-care behaviours + covariates β
General diet	0.04	0.06
Specific diet	-0.06	-0.04
Exercise	-0.10 ^a	-0.03
SMBG	0.03	-0.002
Medications	-0.14 ^b	-0.16 ^b

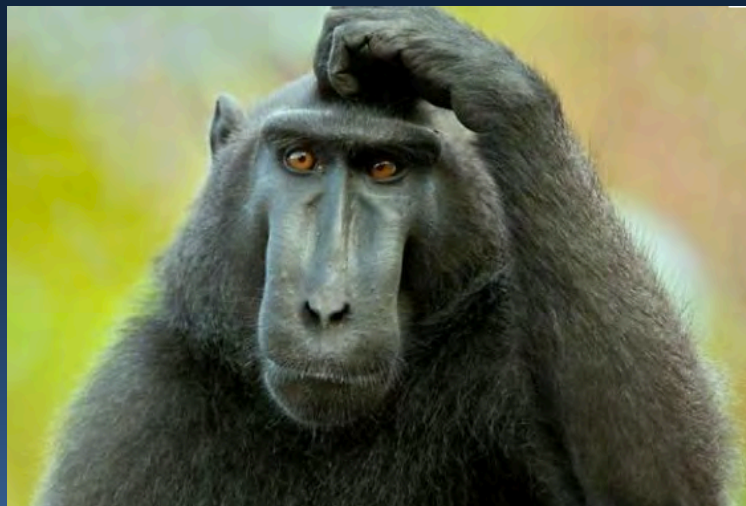
Osborn et al, 2016

Patients Achieving Targets: 2019

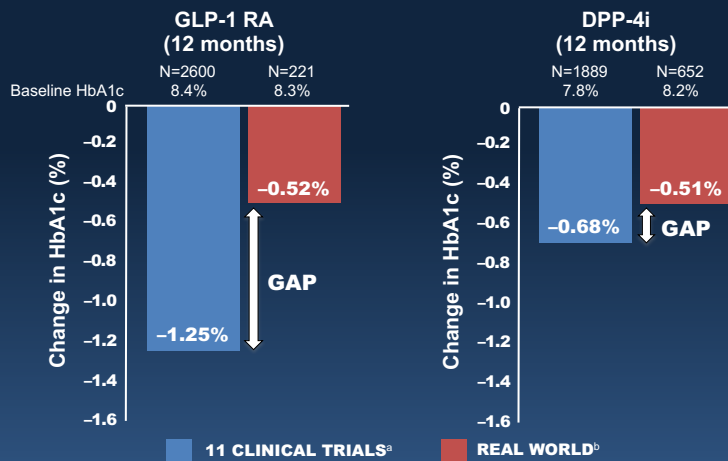


Kazemian et al, 2019

WHY AREN'T WE SEEING DRAMATIC IMPROVEMENTS?

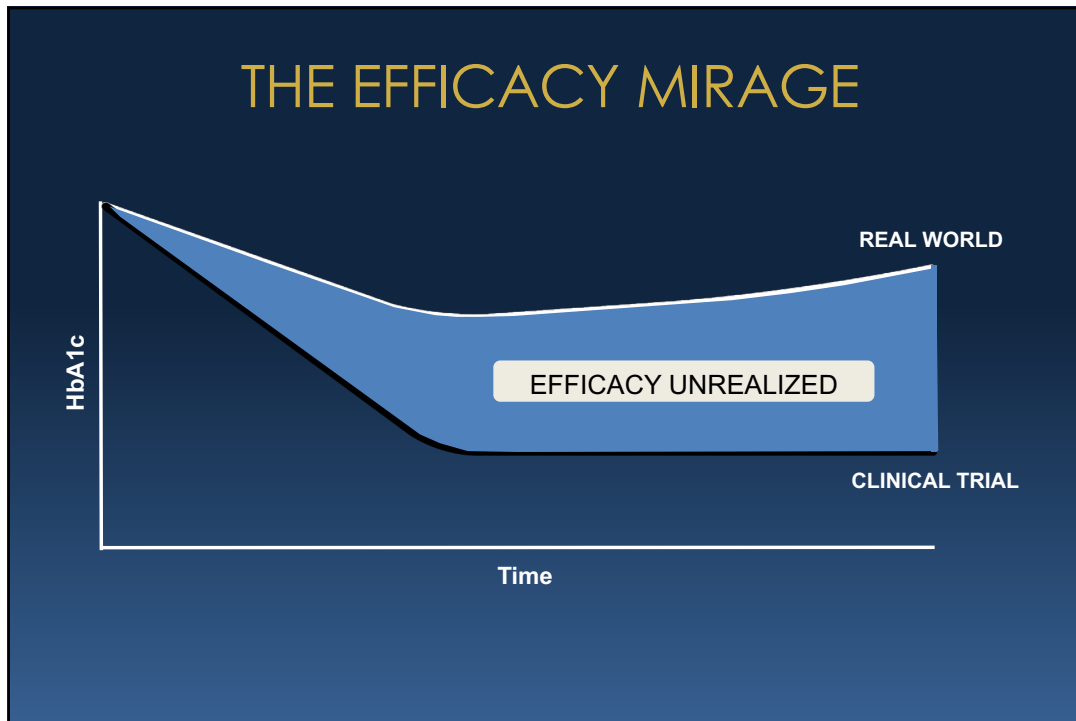


CLINICAL TRIAL RESULTS LOOK GOOD, BUT...

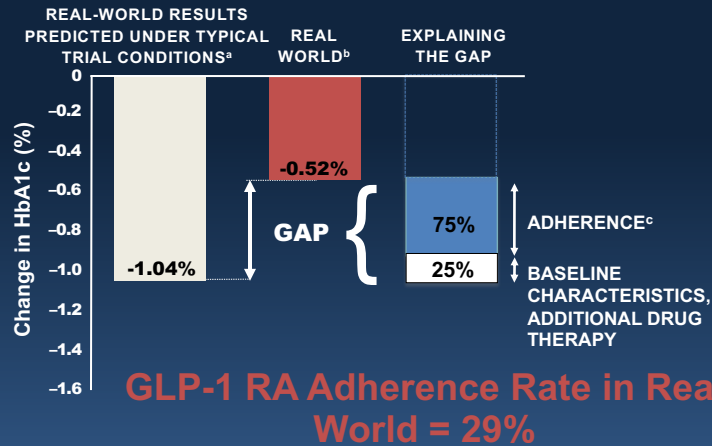


^aIdentified 11 pivotal randomized controlled trials with published change in HbA1c (7 GLP-1 RA [2600 patients] and 4 DPP-4i [1889 patients]).
^bOptum/Humedica SmartFile database (2007-2014) was used (GLP-1 RA 221 patients; DPP-4i 652 patients). Change in HbA1c measured from drug initiation to 365±90 days later.
 Carls et al, 2017

THE EFFICACY MIRAGE

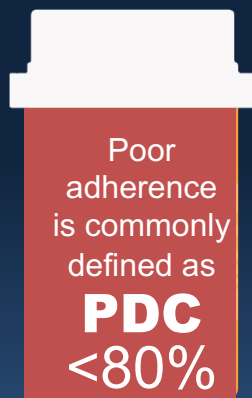


POOR ADHERENCE IS THE KEY



RCT, randomized clinical trial.
^aLinear regression model fitted to estimate the change in HbA1c 1 year after initiating GLP-1 RA or DPP-4i based on baseline and treatment characteristics.
^bOptum/Humedica SmartFile database (2007-2014) was used (GLP-1 RA 221 patients; DPP-4i 652 patients). Change in HbA1c measured from drug initiation to 365±90 days later. ^cMedical adherence classified as poorly adherent if percentage of days covered (PDC) <80%.
 Carls GS et al. 76th ADA Scientific Sessions. June 10–14, 2016. New Orleans, LA. Poster 117-LB.

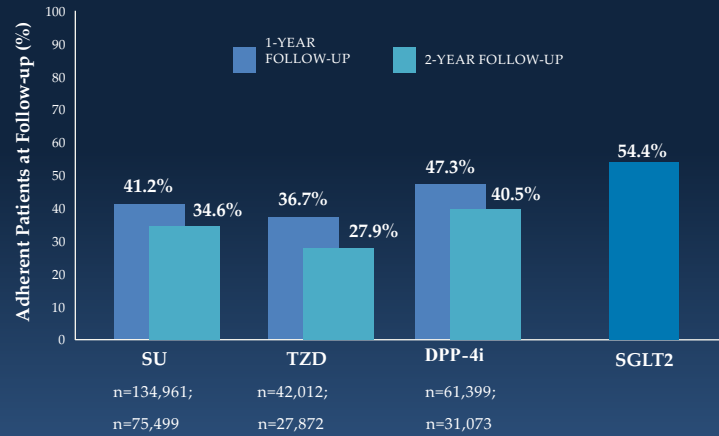
DEFINING POOR ADHERENCE



- Proportion of days covered
- Typically measured after first refill
- PDC doesn't account for
 - Prescriptions that are never filled at all¹
 - What the patient actually takes

PDC, proportion of days covered.
 1. Fischer MA et al. *J Gen Intern Med.* 2010;25:284-290.

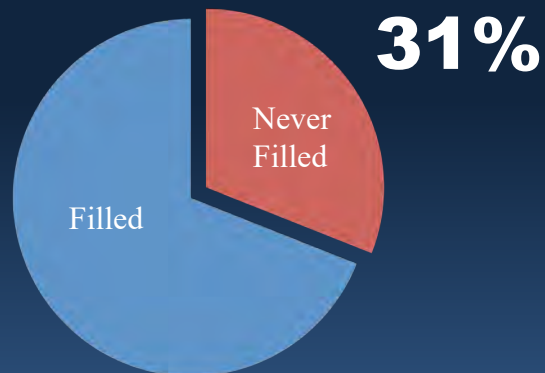
Adherence Rates for T2D Agents



PDC, proportion of days covered; SU, sulfonylurea; TZD, thiazolidinedione.
 Retrospective claims analysis of 238,372 patients with T2D with at least 1 prescription claim for a DPP-4i, SU, or TZD from January 1, 2009 to January 31, 2012.
 Adherence defined as PDC \geq 0.8. Farr AM, et al. *Adv Ther.* 2014;31:1287-1305.
 Symphony PTD Data Set; Nov 2016 – Sep 2017 - Baseline characteristics of the total cohort (N=6,086,767, No of Claims=62,224,558)

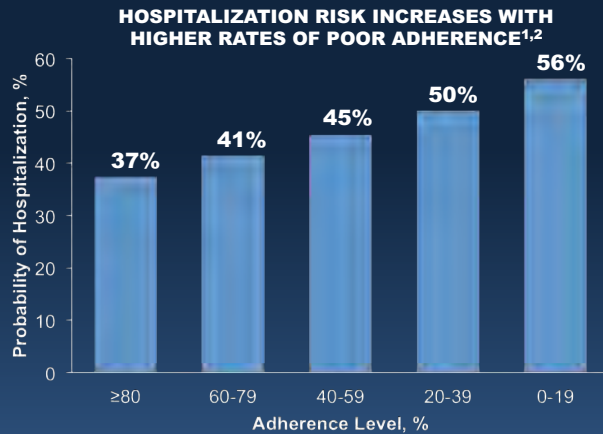
TRACKING NEW E-PRESCRIPTIONS FOR DIABETES MEDICATIONS

AMONG 75,589 INSURED PATIENTS IN THE FIRST YEAR OF A COMMUNITY-BASED E-PRESCRIBING INITIATIVE



Fischer MA et al. *J Gen Intern Med.* 2010;25:284-290.

IMPACT OF POOR ADHERENCE



Data was provided by a large, Medicare supplemental (MarketScan) database from July 1, 2009 to June 30, 2014. There were 123,235 patients with T2D aged ≥65 who received glucose-lowering agents. Comparisons between adherent (defined as PDC ≥80%) and poorly adherent (PDC <80%) were all statistically significant at $P < 0.001$.¹

1. Boye KS et al. 76th ADA Scientific Sessions, June 10–14, 2016. Poster 1221-P. 2. Ho PM et al. *Arch Intern Med.* 2006;166:1836-1841.

73%

increased risk
of all-cause
mortality
due to poor
adherence to oral
hypoglycemics²

Poor adherence
defined as PDC <0.8

INTERVENTION STRATEGIES TO ADDRESS MEDICATION ADHERENCE

- Written medication instructions
- Enhancing HCP adherence skills
- Goal setting
- Stimuli/prompts to take medications
- Enhancing support from significant others
- Special packaging of medications
- Self-monitoring of medication adherence
- Habit analysis and intervention

Conn and Rupar, 2017

INTERVENTION STRATEGIES TO ADDRESS MEDICATION ADHERENCE

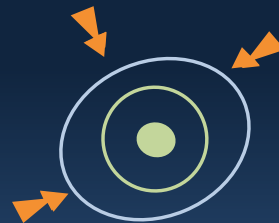
- Medication side effect management
- Feedback about medication adherence
- Medication calendars
- Enhancing patient self-management skills
- Providing consequences/rewards for adherence
- Motivational interviewing
- Stress management

Conn and Rupar, 2017

EFFECTIVENESS OF CURRENT INTERVENTION STRATEGIES

Review of 771 RCTs indicate that effects are modest (Cohen's d):

- Overall: 0.29
- Behavioral strategies: 0.33
- Addressing habits: 0.37
- No behavioral strategies: 0.28



“Much room remains for improvement.”

Conn and Rupar, 2017

WHAT ARE WE MISSING?



THE PROBLEM: FORGETFULNESS?



THE SOLUTION: ADDRESS FORGETFULNESS?



Gadkari and McHorney *BMC Health Services Research* 2012, **12**:98
<http://www.biomedcentral.com/1472-6963/12/98>

 BMC
Health Services Research

RESEARCH ARTICLE

Open Access

Unintentional non-adherence to chronic prescription medications: How unintentional is it really?

Abhijit S Gadkari* and Colleen A McHorney

“Patient’s medication beliefs, especially perceived need for medication and perceived medication affordability, were strong predictors of unintentional non-adherence.”

Gadkari and McHorney, 2012

MEDICINE AND SOCIETY

Debra Malina, Ph.D., Editor

Beyond Belief — How People Feel about Taking Medications for Heart Disease

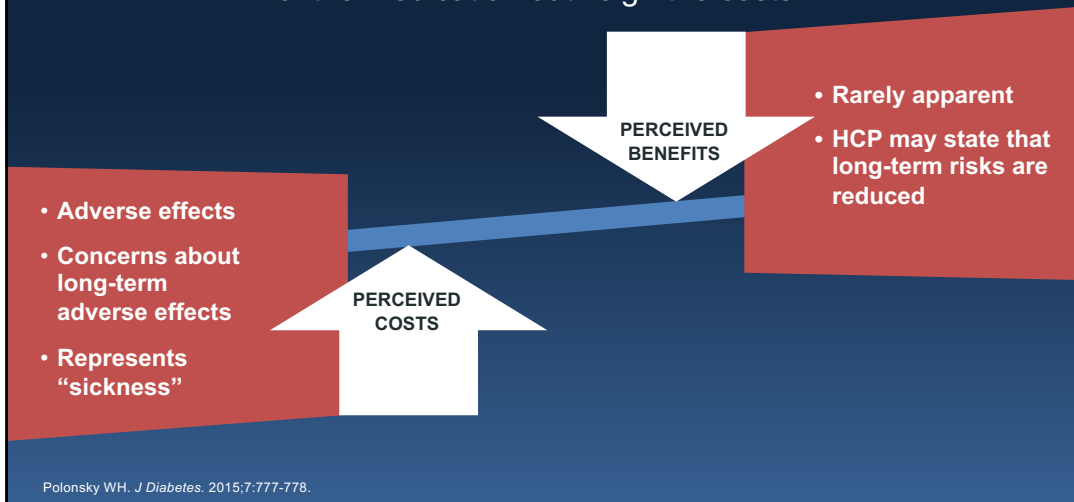
Lisa Rosenbaum, M.D.

“It’s our job to help patients live as long as possible free of CVD complications. Although most patients share that goal, we don’t always see the same pathways to get there. I want to believe that if patients knew what I know, they would take their medicine. What I’ve learned is that if I felt what they feel, I’d understand why they don’t.”

Rosenbaum, 2015

MEDICATION BELIEFS

Perceived worthwhileness: Does the patient believe the benefits of the medication outweigh the costs?



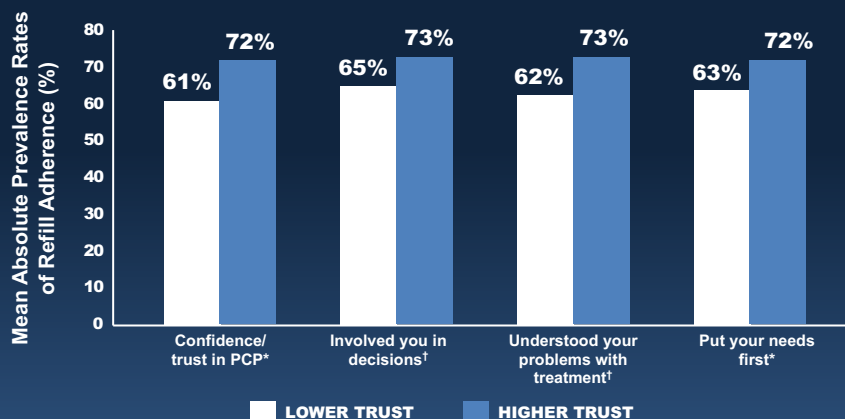
PERCEIVED TREATMENT INEFFICACY



Lack of tangible benefits contributes to discouragement and poor adherence

1. Polonsky WH. *J Diabetes*. 2015;7:777-778. 2. Polonsky WH, Skinner TC. *Clin Diabetes*. 2010;28(2):89-92.

LACK OF PHYSICIAN TRUST



Differences in prevalence of poor refill adherence for any cardiometabolic medication in a cohort of 9377 patients with diabetes. Respondents were classified as poorly adherent when they had no medication supply for >20% of the observation time.

*Trust is defined using 2 items from the Trust in Physicians Scale (TIPS) modified to match the 4-point Consumer Assessment of Healthcare Providers and Systems (CAHPS) scale options during the preceding 12 months. †Shared decision-making was determined using 2 items from the Interpersonal Processes of Care (IPC) instrument during the preceding 12 months.

Ratanawongsa N et al. *JAMA Intern Med*. 2013;173:210-218.

Association Between Primary Care Practitioner Empathy and Risk of Cardiovascular Events and All-Cause Mortality Among Patients With Type 2 Diabetes: A Population-Based Prospective Cohort Study

Hajira Dambha-Miller, MRCGP,
PhD^{1,2}

Adina L. Feldman, PhD²

Ann Louise Kinnonth, FRCGP,

ABSTRACT

PURPOSE To examine the association between primary care practitioner (physician and nurse) empathy and incidence of cardiovascular disease (CVD) events and all-cause mortality among patients with type 2 diabetes.

Dambha-Miller et al, 2019

Assessing Your HCPs' Empathy

How good was your HCP at:

1. making you feel at ease
2. letting you tell your story
3. really listening
4. being interested in you as a whole person
5. fully understanding your concerns
6. showing care and compassion
7. being positive
8. explaining things clearly
9. helping you to take control
10. making a plan of action with you

Dambha-Miller et al, 2019

HCP Empathy and Mortality Outcomes

- 10-year follow up of patients with newly diagnosed T2D:
- “those reporting better experiences of empathy in the first 12 months after diagnosis had a significantly lower risk **(40% to 50%)** of all-cause mortality over the subsequent 10 years vs. those who experienced low practitioner empathy.”

Dambha-Miller et al, 2019

WHY DO PATIENTS FEEL THIS WAY?

- Threatening patients with medication
 - *“If you can ’t make some positive changes, then we’ll have no choice but to put you on more medication, and perhaps even start insulin.”*
- Underlying messages
 - More medication should be avoided **at all costs**
 - You have failed
 - You are to be punished

SO WHAT TO DO?



SO WHAT TO DO?



1. Ask correctly

- *“Any problems taking those medications?”*
- vs.
- *“What’s one thing about taking your medications that’s been challenging?”*

SO WHAT TO DO?



1. Ask correctly

2. **Forgetfulness**

- “Aside from forgetting, what else is tough about taking your meds?”
- Anchoring strategies

SO WHAT TO DO?



1. Ask correctly

2. Forgetfulness

3. **Patient-provider trust and collaboration**

- Listen, listen, listen

SO WHAT TO DO?

1. Ask correctly
2. Forgetfulness
3. Patient-provider trust
4. **Talk about beliefs about diabetes and medications**



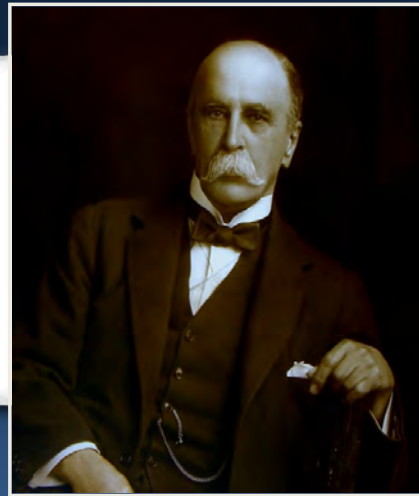
Challenging Harmful Beliefs

1. Taking your medications is one of the most powerful things you can do to positively affect your health
2. Your medications are working even if you can't feel it
3. Needing more medication isn't your fault
4. More medication doesn't mean you are sicker, less medication doesn't mean you are healthier
5. Emphasize the potential long-term gains

Diabetes and Your Health

“To live a long and healthy life, develop a chronic disease and take care of it.”

- Sir William Osler



CONCLUSIONS

Poor medication adherence:

- ... explains a great deal of the lack of glycemic progress over the past decade
- ... is commonly an *attitudinal* issue, not just a behavioral issue.
- ... is best addressed by considering the patient's perspective, and encouraging a two-way conversation about the perceived pro's and con's of the medication.

Thanks for Listening!

Critical Psychosocial Issues in Diabetes

Web-based video modules

UC San Diego
SCHOOL OF MEDICINE



Home

Modules

The **Critical Psychosocial Issues in Diabetes** web-based program is a series of video modules designed to examine psychosocial issues in diabetes, provide a brief review of the research literature, clarify how and why the problems manifest themselves among patients with diabetes, and put forward practical solutions for the busy healthcare professional.

The American Diabetes Association published its first Psychosocial Position Statement in December, 2016, recognizing the important

www.behavioraldiabetes.org

Lecture 2: 10:30 – 11:45 a.m. PST

Jeremy H. Pettus, MD, Presents:

A Focus on Time in Range,
Unmet Needs and Modern Management of Type 1 Diabetes

To Be Discussed...

- Incidence and pathophysiology
- Demographics of T1D in the U.S.
- A1c and time in range (TIR)
- Overview of pumps and CGM devices
- Interpreting CGM downloads in ~ 30 secs.
- Identifying and addressing common problems
- New insulin and glucagon formulations
- Advances in hybrid and closed AP

TCOYD

Prevalence of T1D Is Increasing!

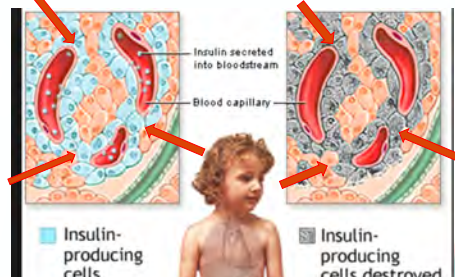


- 40,000 people diagnosed each year in U.S.²
- 110 people are diagnosed with T1D each day
- By 2040 there will be 5 million people with T1D

¹ T1D Exchange T1D population based on company research
² American Diabetes Association

TCOYD

Type 1 is an Autoimmune Disease: The Immune System Attacks Healthy Beta Cells



Natural Progression is months to a few years.

August 6, 2019

Teplizumab Gets Breakthrough Status for Type 1 Diabetes Prevention

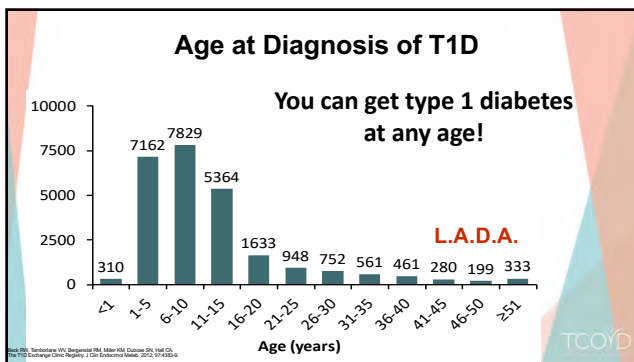
Steve Duffy

Type 1 Diabetes TrialNet

The Food and Drug Administration (FDA) has granted Breakthrough Therapy designation to teplizumab (PRV-031; Provention Bio), an anti-CD3 monoclonal antibody, for the prevention or delay of clinical type 1 diabetes (T1D) in individuals at risk of developing the disease.

BREAKTHROUGH DESIGNATION

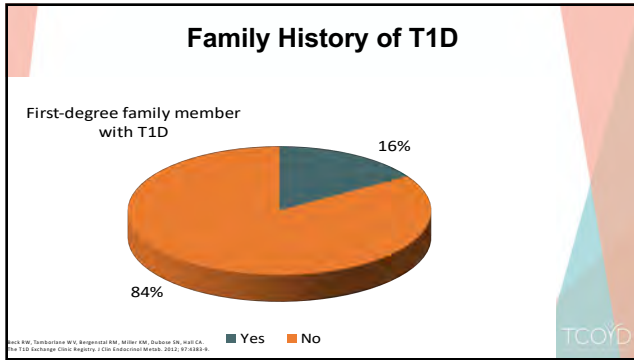
The designation was based on data from a recent Teplizumab is an investigational anti-CD3 monoclonal antibody



Latent Autoimmune Diabetes in Adults (L.A.D.A.)

- o The most missed diagnosis in diabetes
- o Type 1 diabetes can occur at any age
- o Slower beta-cell destruction (may respond briefly to oral agents)
- o Typically does not have features of the Metabolic Syndrome
- o Blood test positive for type 1 diabetes (GAD auto antibodies)

TCOYD

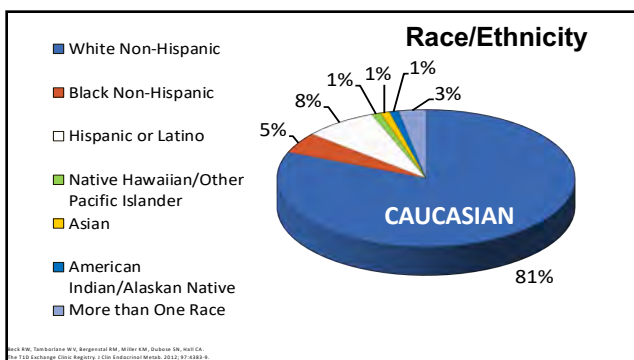


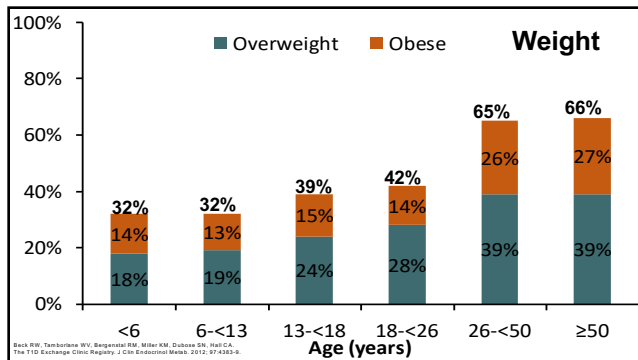
Risk of Developing Type 1 vs Type 2

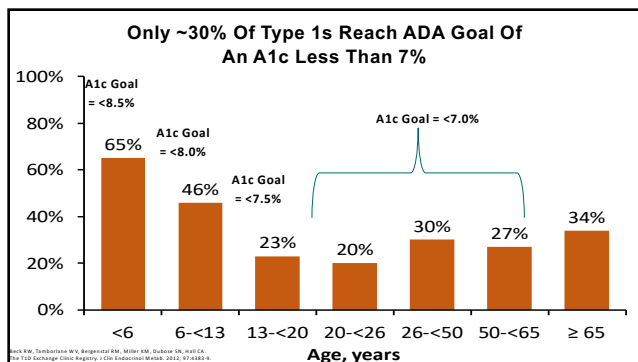
General Population	0.3%	8-11%
If you have a sibling with T1D	4%	~30%
If your mother has T1D	2-3%	~30%
If your father has T1D	6-8%	~30%
If you have an identical twin with T1D	~50%	100%

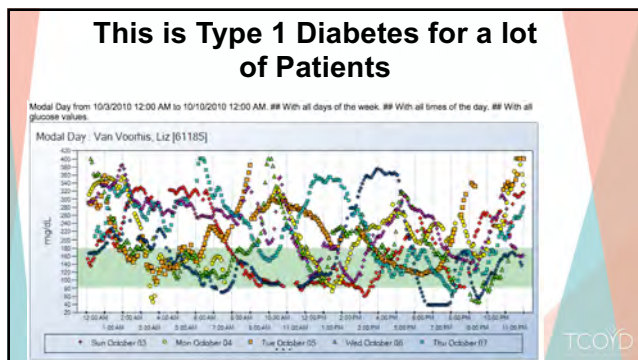
Beck RW. Being control of your diabetes: a patient-oriented course. J Clin Endocrinol Metab. 2012; 97:1038-41.

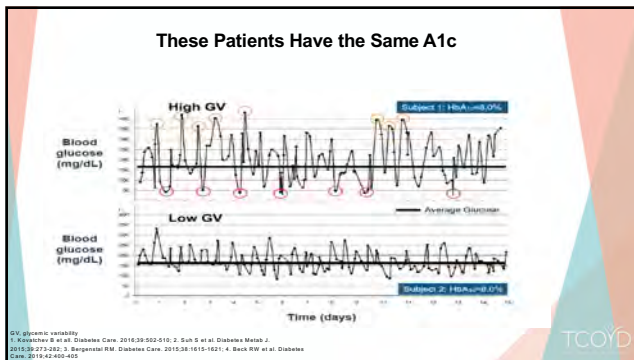
TCOYD



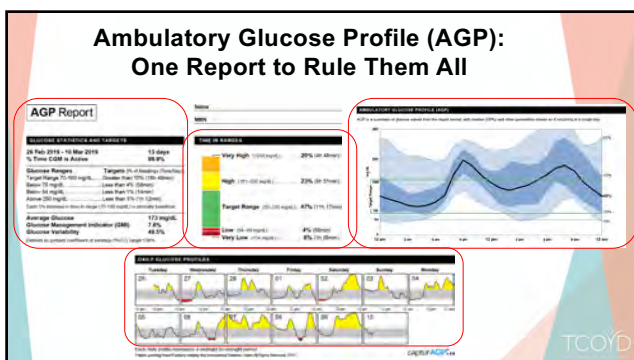


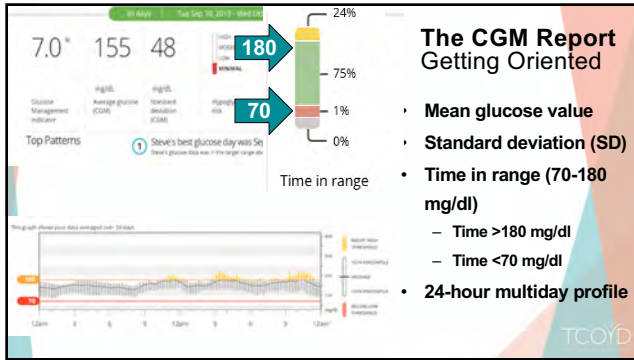


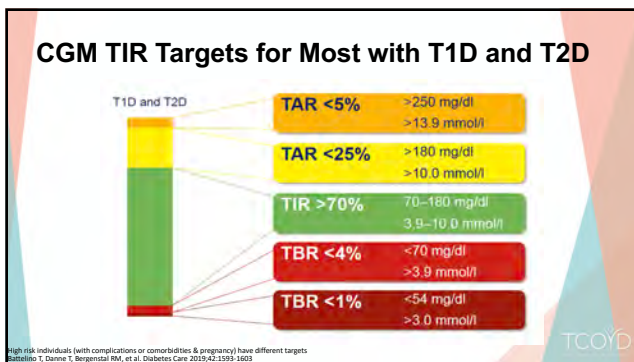


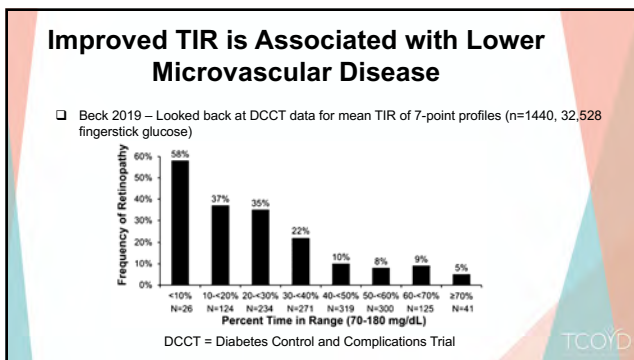


- ### Provider CGM Overview
1. Review CGM download **together** with the patient, explain what you are observing
 2. Look at average glucose and predicted A1c
 3. Look at time in range and start with time hypoglycemic (goal < 5%)
 4. Look at total time in ideal range (goal > 70%)
 5. Look at 24 hour day to see when highs and lows occur
 6. Look at individual days to tease out those problem areas
 7. Review **alert settings** on the CGM. Especially if the significant other looks exhausted and has alarm PTSD
- TCOYD









Options to Connect Directly to Smart Phone/Smart Watch

- Last 10 days
- No calibration
- No finger sticks
- Predictive low alert
- Medicare approved



TCOyD



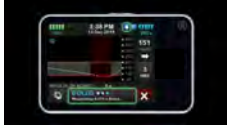
CGM System



- Requires calibration
- Predictive low alerts
- Requires high alerts
- 6-day wear
- Need to confirm with fingerstick when dosing insulin
- No sharing capabilities

TCOyD

How Does Control IQ Keep You in Range?



Delivers	Ensures an automatic correction bolus if glucose is predicted to be above 180 mg/dL.
Increases	Increases basal insulin delivery if glucose is predicted to be above 160 mg/dL.
Maintains	Maintains actual Personal Profile settings.
Decreases	Decreases basal insulin delivery if glucose is predicted to be below 112.5 mg/dL.
Stops	Stops basal insulin delivery if glucose is predicted to be below 70 mg/dL.

TCOYD

The NEW ENGLAND JOURNAL of MEDICINE

ESTABLISHED IN 1812 OCTOBER 21, 2019 VOL. 381 NO. 18

Six-Month Randomized, Multicenter Trial of Closed-Loop Control in Type 1 Diabetes

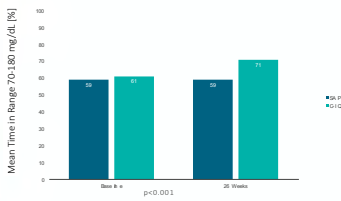


Improved "Time in Range"
Reduced A1c
Reduced Hypoglycemia

TCOYD

Results

Primary Outcome Time in Range 70-180 mg/dL*



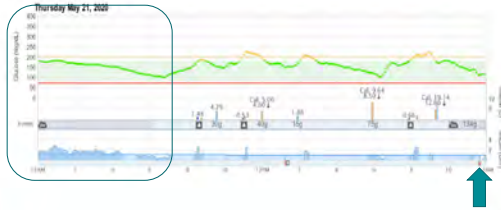
2.6 hours

average additional time per day that Control-IQ participants spent in range compared to SAP users*

*As measured by CGM

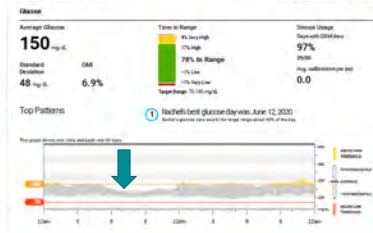
TCOYD

Basal Rate Modulation Overnight to Improve Control...



TCOYD

AP Systems Very Effective Overnight



TCOYD

Hybrid-Closed Loop System

- Auto-adjusts basal rate when in auto mode
- Target blood sugar: 120mg/dl
- Mealtime boluses required
- Sensor (needs frequent calibration to stay in auto mode)



TCOYD

DIY Looping Hybrid Closed Loop NOT FDA Approved

- Basal rate modulator
- Communicates with certain CGM devices (no calibration needed)
- Always in auto mode
- Still need to enter carbs and give correction doses

Smart Pens: Same Software Programs as Pumps

- I:Carb ratio
- Correction factor
- Insulin log
- Cloud-based

Example of a Bionic Pancreas

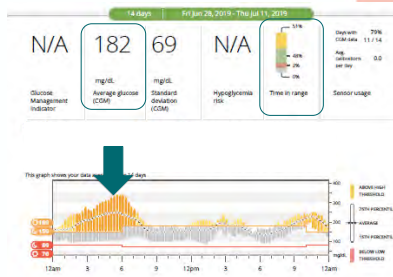
2 ports for insulin and glucagon

Let's Practice Example Cases

TCOYD

Case 1: Sam

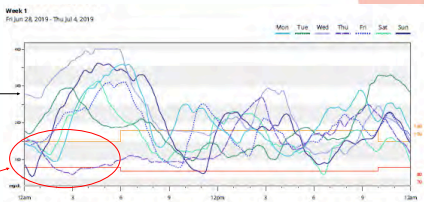
- Quick Interpretation**
- Adult with T1D
 - A1c ~8%
 - High variability
 - Minimal lows
 - Most glycemc burden overnight



TCOYD

Case 1 Cont...

Rises around 1AM without coverage, may be a learned behavior based on experience when no snacking



- To address overnight fall in glucose, we reduced the basal 20%
- Eliminates the need for "mandatory" bedtime snack
- Over time, increased time in range

ENDOCRINE SOCIETY

TCOYD

How Do you Know if the Basal Does is “Right”?

- Check blood sugar when there is no insulin boluses in the system and no carbohydrates from last meal (e.g. 2-4 AM) and compare to morning blood sugar
- Be on the lookout for variable bedtimes
- If ≥ 30 mg/dL rise in glucose raise basal insulin dose
- If ≥ 30 mg/dL fall in glucose decrease basal insulin dose

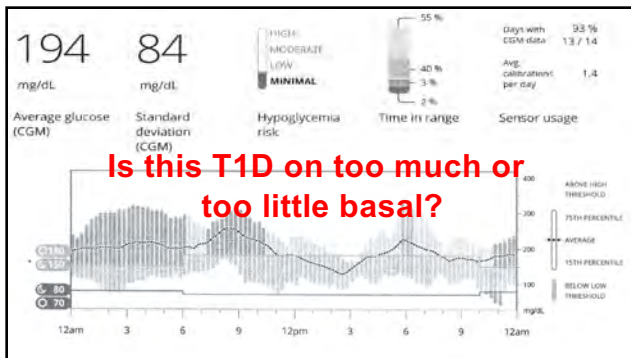
ENDOCRINE SOCIETY

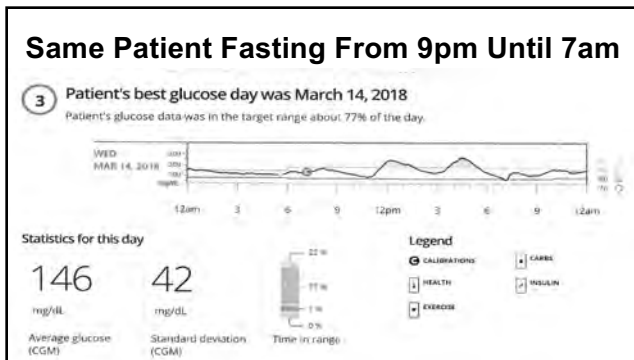
TCOYD

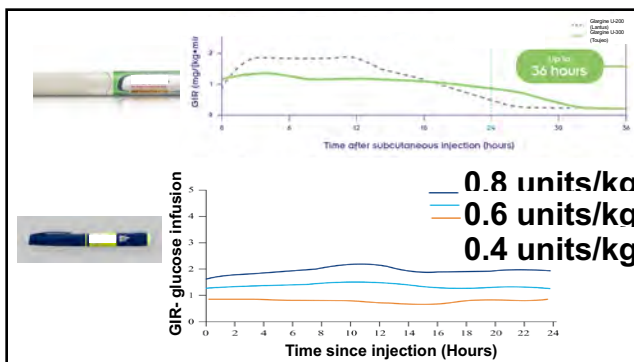
Physiologic Basal



TCOYD







Case 1 Learning Points

- Type 1 diabetes does not require a midnight snack
- Nighttime highs SHOULD NOT reflex to increasing basal dose
- To determine if the issue is basal or bolus related, do "basal testing" as discussed
- Often, nighttime highs need to be addressed with more insulin before bed rather than changes to basal
- Newer basal insulins (Glargine U-300, Degludec U-00/U-200) are more consistent, have more flexible dosing, and less hypoglycemia

TCOYD

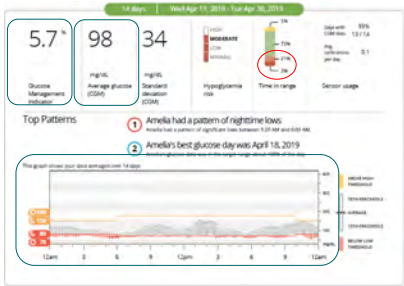
Case 2: Amelia

• Amelia is a 57 yo female with Type 1 diabetes since age 2

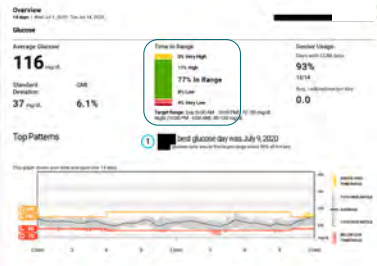
• Was told she needed tight glucose control to avoid complications

• Has since had a fear of Hyperglycemia and prefers to "Ride low"

• Currently on insulin pump with CGM



- Switched pumps
- Episodes of hypoglycemia markedly reduced...



Suspensions to Reduce Hypoglycemia



New Formulations of Glucagon

Nasal Glucagon



Pre-Filled Syringe



Auto-injector Pen



TCOYD

Case 2 Learning Points

- A “good” A1c doesn’t mean good control
- When you see a low A1c, look immediately at percent hypos
- Make sure these patients are on a CGM with alarms turned ON!
- Hybrid closed loop systems can help reduce hypoglycemia
- ALL type 1 patients MUST have glucagon available with loved ones trained on how to use

TCOYD

Case 3: Brian

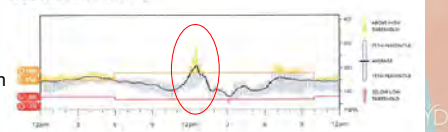


Quick Interpretation

- Overall glucose just slightly below goal
- Low variability
- Hypos NOT a problem
- Spike after lunch

Top Patterns

This graph shows your data averaged over 14 days



D

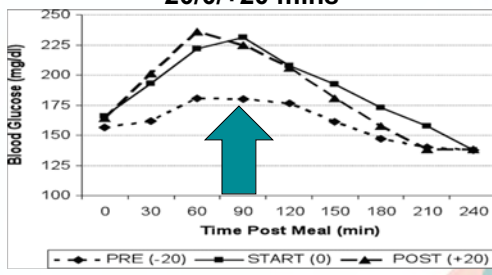
Shark Attack



- Lows after eating are VERY common
- Can result in a "rage bolus"
- Results in lows after and getting on the rollercoaster

TCOYD

Postprandial Glucose bolus at - 20/0/+20 mins



Cobby et al. Diab Tech Therap 2010;12: 173-177

TCOYD

What About the Low Carb Thing?



- It works
- Reduces margin of error
- Not easy to adhere to but given "Atkins craze", lots of tips on low carb snacks/meals/etc.
- TRY it for one week to see the effect of carbs on your BG

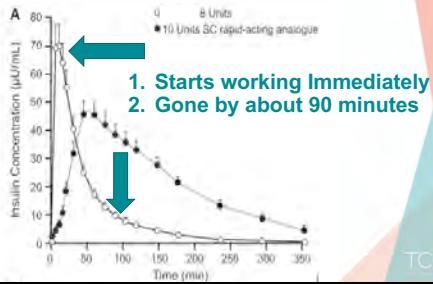
TCOYD

What is Inhaled Insulin?



TCOYD

Why is it Cool?



TCOYD

New, "Faster Acting" Insulins

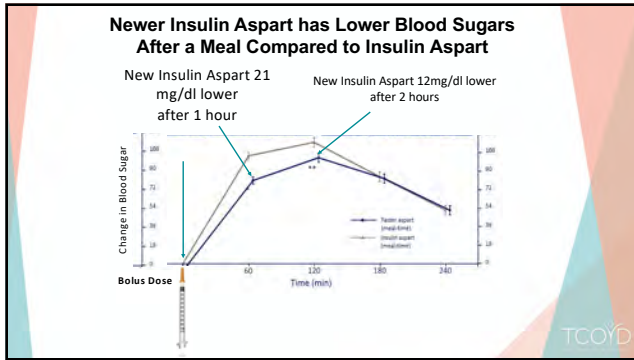


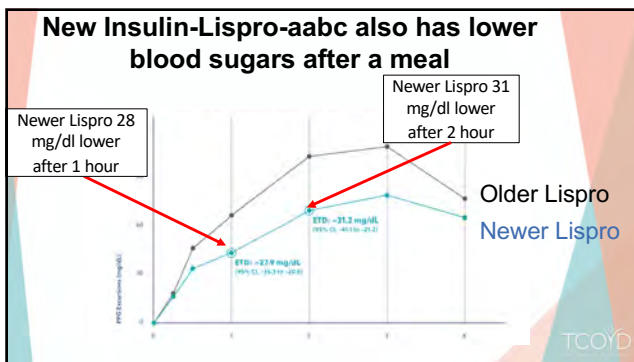
**Insulin Aspart
(Fiasp)**



**Insulin Lispro-aabc
(Lyumjev)**

TCOYD





- ### Case 3 Learning Points
- Bolus 15-30 minutes BEFORE you eat
 - Break up meal into two parts
 - Try low carb
 - Try inhaled insulin or newer, rapid-acting insulins
- TCOYD

To Be Discussed...

- Incidence and pathophysiology
- Demographics of T1D in the U.S.
- A1c and time in range (TIR)
- Overview of pumps and CGM devices
- Interpreting CGM downloads in ~ 30 secs.
- Identifying and addressing common problems
- New insulin and glucagon formulations
- Advances in hybrid and closed AP

TCOYD

Lecture 3: 12:15 – 1:45 p.m. PST

Schafer Boeder, MD, Presents:

Effective Use of Oral Medications for Type 2 Diabetes:
Lowering Cardiovascular Risk While Improving Glycemic Control

Effective Use of Oral Medications for Type 2 Diabetes: Lowering Cardiovascular Risk While Improving Glycemic Control

Schafer Boeder, MD
Assistant Professor of Medicine
Division of Endocrinology, Diabetes and Metabolism
University of California San Diego School of Medicine

WWW.TCOYD.ORG
Taking Control Of Your Diabetes, 501(c)3 is a not-for-profit educational organization.



Treatment Should be Individualized and Defects Addressed by Agents with Complementary MOAs

Progressive decline in beta-cell function

Impaired insulin secretion



Decreased incretin effect



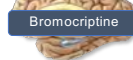
Increased lipolysis



Optimal Pharmacotherapy for Hyperglycemia in Type 2 Diabetes:

- Usually requires combinations of multiple agents with complementary mechanisms of action
- Should aim to achieve the best possible glycemic control with the least possible side effects
- Should help reduce ASCVD in patients at high risk or with pre-existing CVD

Increased hepatic glucose production



Neurotransmitter dysfunction

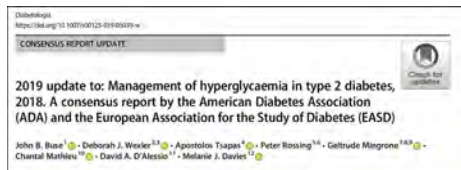


Decreased glucose uptake

Adapted from DeFronzo RA. *Diabetes*. 2009;58(4):773-795.



Key Updates to the 2018 ADA/EASD Consensus Recommendations



General Recommendations

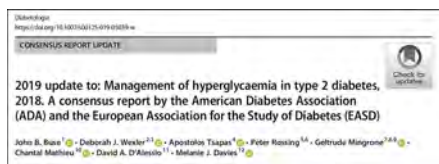
- In appropriate, high-risk individuals with T2D, decision to treat with GLP-1 RA or SGLT-2 inhibitor to reduce MACE, hHF, CV death or CKD progression should be considered independently of baseline A1c or A1c target
- Providers should engage in shared decision making around initial combination therapy in new onset cases of T2D

GLP-1 RA Recommendations

- For patients with T2D and established ASCVD, where MACE is the gravest threat, the level of evidence for MACE benefit is greatest for GLP-1 RAs
- To reduce risk of MACE, GLP-1 RA can also be considered in patients with T2D without established CVD with indicators of high risk (>55 y/o with coronary, carotid, or LE artery sclerosis >50%, LVH, eGFR <60 ml/min/1.73, or albuminuria)



Key Updates to the 2018 ADA/EASD Consensus Recommendations

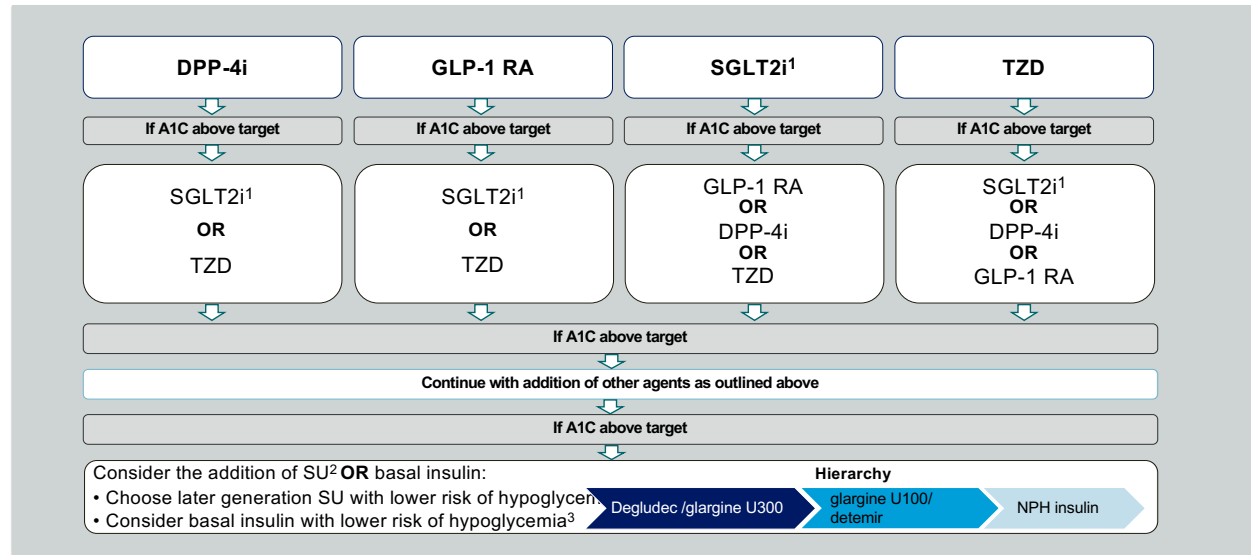


SGLT-2 Inhibitor Recommendations

- For patients with or without established ASCVD, but with HFrEF (EF <45%) or CKD (eGFR 30 to 60 ml/min/1.73 m² or UACR >30mg/g, particularly UACR >300mg/g), the level of evidence for benefit is greatest for SGLT2 inhibitors
- SGLT2 inh. are recommended in patients with T2D and HF, particularly those with HFrEF, to reduce hHF, MACE and CV death
- SGLT2 inh. are recommended to prevent the progression of CKD, hHF, MACE and CV death in patients with T2D and CKD
- Patients with foot ulcers or at risk of amputations should only be treated with SGLT2 inh. after careful shared decision making around risks and benefits with comprehensive education on foot care and amputation prevention

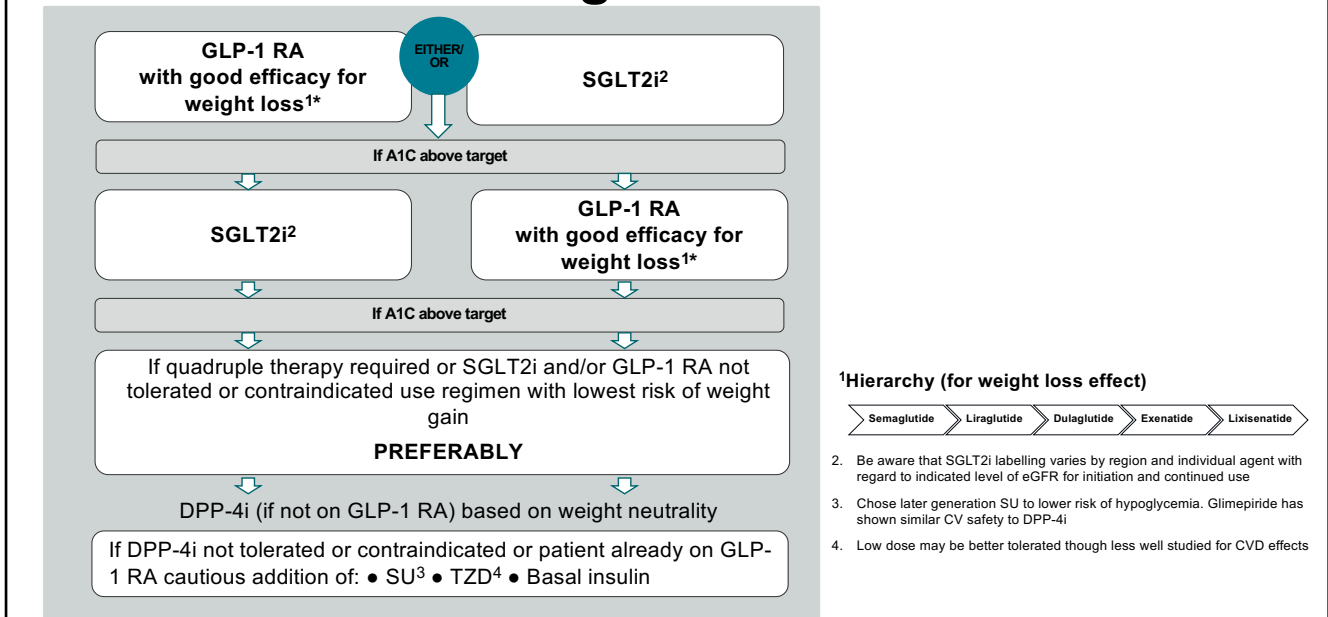


Compelling Need to Minimize Hypoglycemia

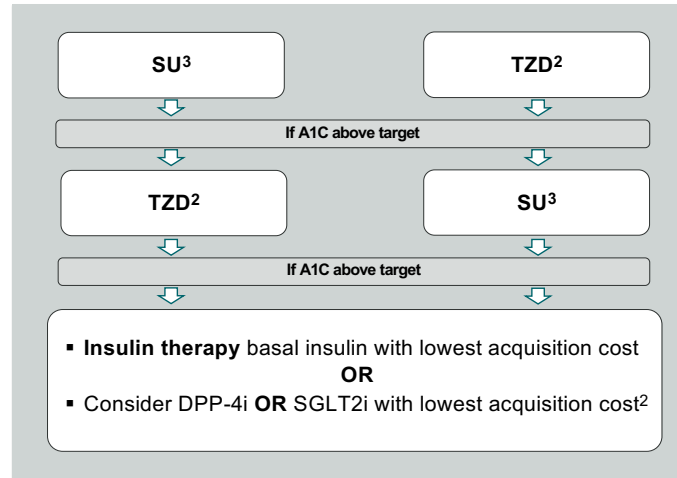


1. Be aware that SGLT2i labelling varies by region and individual agent with regard to indicated level of eGFR for initiation and continued use
2. Choose later generation SU to lower risk of hypoglycemia. Glimepiride has shown similar CV safety to DPP-4i
3. Degludec / glargine U300 < glargine U100 / detemir > NPH insulin

Compelling Need to Minimize Weight Gain or Promote Weight Loss



Second-Line Therapy for T2D if Cost is a Major Issue



1. If no specific comorbidities (i.e., no established CVD, low risk of hypoglycemia and lower priority to avoid weight gain or no weight-related comorbidities)
2. Consider country- and region-specific cost of drugs. In some countries, TZDs relatively more expensive and DPP-4i relatively cheaper
3. Chose later generation SU to lower risk of hypoglycemia. Glimepiride has shown similar CV safety to DPP-4i

Case 1: 32-year-old male with T2D for two years

- Medical history: central obesity, dyslipidemia, HTN, and CAD s/p MI
- Family Hx: Strongly positive for T2D, obesity, and CAD
- Notes: Very few home glucose monitoring results
 - Diabetes meds: metformin, SFU, DPP-4 inh., SGLT-2 inh., and basal insulin
 - Current A1c: 11.4% (10.6% one year ago, 10.1% two years ago)
 - Creatinine: 1.4 mg/dL, eGFR 65, mL/min/1.73 m²

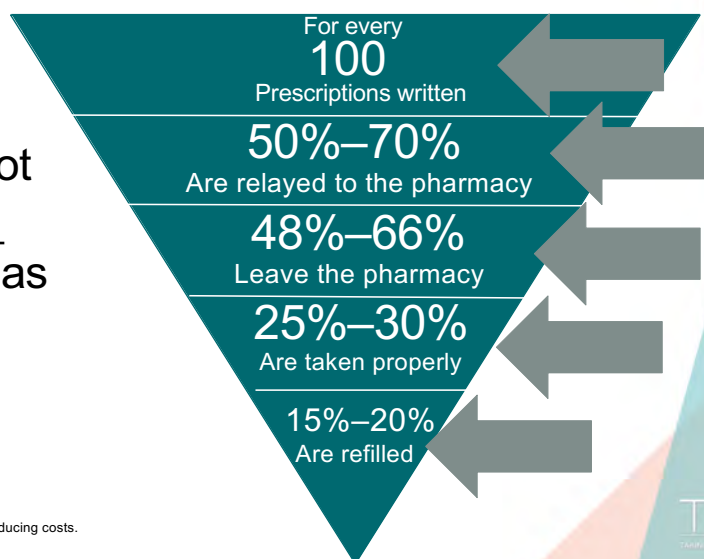
What is the most likely reason why this patient has not achieved his A1c goal?

A	He needs prandial insulin
B	He needs a GLP-1RA
C	Poor adherence with his medication
D	His diabetes regimen is too complicated

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TAKING CONTROL OF YOUR DIABETES

“Poor Adherence” with Type 2 Medications in the Real World

Prescriptions are not always filled, taken properly, or refilled as directed



National Association of Chain Drug Stores. Pharmacies: improving health, reducing costs. PrinciplesOfHealthcare.pdf. Accessed September 28, 2017.

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Nine FDA-Approved Classes of Oral Meds for T2D

- Metformin (first line therapy unless contraindicated)
- Sulfonylureas, meglitinides
- Glitazones (pioglitazone, rosiglitazone)
- DPP-4 inhibitors (sitagliptin, saxagliptin, linagliptin, alogliptin)
- SGLT-2 inhibitors (canagliflozin, dapagliflozin, empagliflozin, ertugliflozin)
- NEW ORAL GLP-1 Receptor Agonist (oral semaglutide)
- Bile acid sequestrant (colesevelam)*
- Dopamine receptor agonists (bromocriptine mesylate)*
- Alpha glucosidase inhibitors (acarbose, miglitol)*

* not discussed in detail in this presentation

<http://www.fda.gov/drugs>



Clinical Treatment Pearls

- Always confirm as best you can if the patient is adherent with his/her medications (check refill history)
- The higher the baseline A1C, the greater the fall in A1C with any therapeutic intervention
- Always address the modifiable risk factors (hypertension, dyslipidemia, smoking)
- Spending time with the patient and his/her support person(s) in meaningful shared decision-making addressing their health care priorities and concerns will improve adherence

Edelman SV, Henry RR. Diagnosis and management of type 2 diabetes. 12th Edition. Professional Communications, Inc., Greenwich, CT. 288 pages, 2014.

Edelman SV (TCOYDtv). 3 September 2015. Get Type 2 Diabetes and Live Longer Because of it (video) <https://www.youtube.com/watch?v=x24AbWnjVa8>



Case 2: 69-year-old centrally obese female with T2D for nine years

- Medical history: Obesity (BMI 34 kg/m²), dyslipidemia, OSA, breast cancer s/p lumpectomy and hormonal therapy in remission
- Family Hx: Both parents had type 2 diabetes
- Notes:
 - eGFR 75 mL/min/m², UACR normal (<30mg/g creatinine)
 - A1C 8.5%
 - Diabetes therapy is metformin and a SFU
 - LDL 121 mg/dL, triglycerides 266 mg/dL, HDL 39 mg/dL

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What class of agent would you add to this patient's current regimen of metformin and a SFU

A	Thiazolidinedione (pioglitazone)
B	DPP-4 inhibitor (sita-, lina-, saxa- and alogliptin)
C	SGLT-2 inhibitor (cana-, empa-, ertu- or dapagliflozin)
D	Basal insulin given once a day
E	GLP-1 RA (liraglutide, exenatide, dulaglutide, semaglutide)

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Update on Metformin, SFUs, and TZDs (all generic)

Metformin

- eGFR <60 to ≥ 45 OK to use full dose/monitor kidneys
- eGFR <45 to ≥ 30 OK to use 50% maximum dose/monitor renal function every 3-6 months (PI says yearly)
- Check B-12 levels

SFU

- High secondary failure rate; **however, when you stop them, the patient's A1c typically goes up**
- Increase risk of hypoglycemia (elderly, CKD, CAD), weight gain

TZD (pioglitazone)

- Effective in prediabetes, best used early in the natural history (balance with potential side effects)
- Be cautious in combo with insulin (fluid retention)
- Contraindicated in the setting of heart failure
- Weight gain
- Fracture risk is increased
- Risk of bladder cancer questionable, and the risk is low (~1/5000 in the general population)



Case 3: 56-year-old AA female diagnosed with type 2 diabetes at age 46

- PMH: HTN, dyslipidemia, obesity and NAFLD (non alcoholic fatty liver disease)
- A1C 9.2% on maximum doses of metformin and SFU
- Occasional mild hypoglycemia
- No home glucose monitoring data
- eGFR 50 mL/min/m², BMI 51 kg/m²
- BP normally above 140/90 mmHg; on no HTN meds



What therapeutic intervention would you change/initiate if you were evaluating this patient, once you have confirmed she is adherent with her medications?

A	Add pioglitazone
B	Add a DPP-4 inh.
C	Add a SGLT-2 inh.
D	Add a GLP-1 RA
E	Combination of a DPP-4 inh & SGLT-2 inh.



High CV Risk or Established ASCVD, CKD, and/or HF

Consider independently of baseline A1C of individualized A1C target

ASCVD PREDOMINATES

- Established ASCVD
- Indicators of high ASCVD risk (age ≥ 55 years with coronary, carotid or lower extremity artery stenosis $>50\%$, or LVH)

PREFERABLY
GLP-1 RA with proven CV benefit¹
OR
SGLT2i with proven CVD benefit if eGFR adequate

If A1C above target

If further intensification is required or patients is no unable to tolerate GLP-1 RA and /or SGLT2i, choose agents demonstrating CV safety:

- For patients on a GLP-1 RA, consider adding SGLT2i with proven CVD benefit¹
- DPP4i if not on GLP-1 RA
- Basal insulin⁴
- TZD⁵
- SU⁶

HF OR CKD PREDOMINATES

- Particularly HFrEF (LVEF $<45\%$)
- CKD: Specifically eGFR 30-60 mL/min/2.73 m² or UACR >30 mg/g, particularly UACR >300 mg/g

PREFERABLY
SGLT2i with evidence of reducing HF and/or CKD progression in CVOTs if eGFR adequate³
OR
If SGLT2i is not tolerated or contraindicated or if eGFR less than adequate² add GLP-1RA with proven CVD benefit

If A1C above target

- Avoid TZD in the setting of HF
- Choose agents demonstrating CV safety
- For patients on a SGLT2i, consider adding GLP-1 RA with proven CVD benefit¹
- DPP4i (not saxagliptin) in the setting of HF (if not on GLP-1 RA)
- Basal insulin⁴
- SU⁶

Case 3 Continued: Treatment History

- A DPP-4/SGLT2 inhibitor combination pill was added to her regimen (once a day and one co-pay)
- Follow up was arranged for one month instead of the usual 3 to 4 months to confirm adherence and engage patient
- She did well with a 10-pound weight loss and no hypoglycemia after the SFU dose was cut in half
- The A1C fell from 9.5% to 7.4%
- SBP decreased from 150 to 141 mmHg
- After 6 months she was started an ARB and a statin to get her BP below 140/90 mmHg and her LDL <100 mg/dl



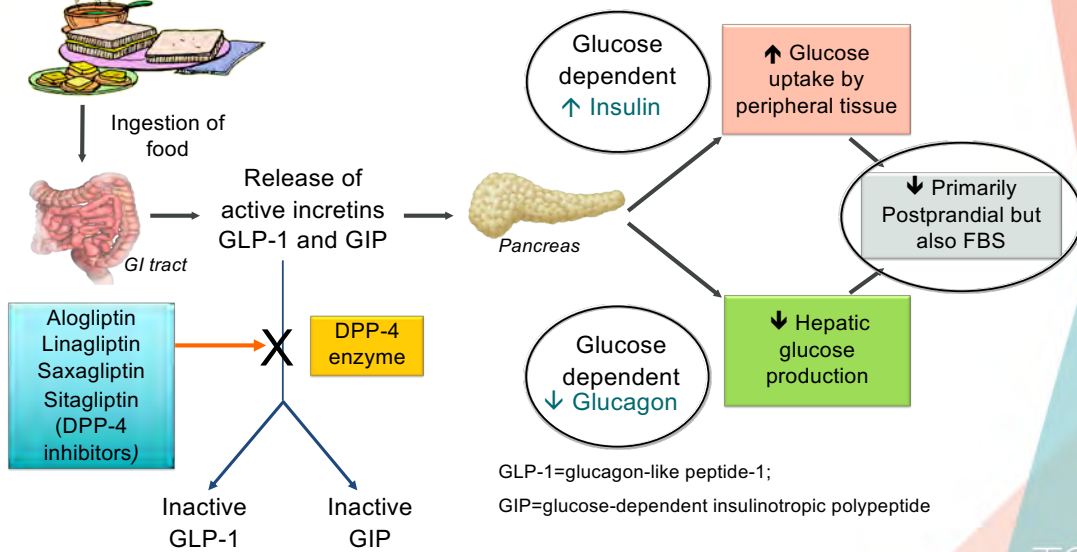
DPP-4 Inhibitors

Mechanism of Action	Inhibit the enzyme, DPP-4, that normally inactivates GLP-1 and other incretins within minutes
Benefits	<ul style="list-style-type: none"> • Once daily oral administration • Virtually no side effects • Can be added to any diabetes drug except GLP-1 RAs • A1c reduction ~ 0.5-1% range (depends on baseline A1c)
Concerns	<ul style="list-style-type: none"> • Dose adjustment with renal insufficiency (only for sita-, saxa- and alogliptin), not for linagliptin • Warnings and precautions: pancreatitis, heart failure (saxa- and alo-), acute renal failure, angioedema, Stevens-Johnson, severe arthralgia, bullous pemphigoid
Clinical Pearls	<ul style="list-style-type: none"> • Efficacy of the DPP-4 inhibitors is similar • All DPP-4 inhibitors come in combination pill with metformin (and some are available in combination with pioglitazone or an SGLT2i)

Edelman SV, Henry RR. Diagnosis and management of type 2 diabetes. 12th Edition. Professional Communications, Inc., Greenwich, CT. 288 pages. 2014.



Mechanism of Action: DPP-4 Inhibitors



Edelman SV, Henry RR. Diagnosis and management of type 2 diabetes. 12th Edition. Professional Communications, Inc., Greenwich, CT. 288 pages, 2014.

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Generic and Trade Names: DPP-4 Inhibitors

	Generic Name	Trade Name
DPP-4 Inh.	Alogliptin	Nesina
	Linagliptin	Tradjenta
	Saxagliptin	Onglyza
	Sitagliptin	Januvia

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Combination Pills with a DPP-4 Inhibitor

Generic Name	Trade Name	Daily Dose Range (mg)	Recommended Frequency
Sitagliptin/metformin	Janumet	50/500, 50/1000	Twice with meals
Saxagliptin/metformin ER	Kombiglyze XR	5/500, 2.5/1000, 5/1000	Once daily with evening meal
Linagliptin/metformin	Jentadueto	2.5/500, 2.5/850, 2.5/1000	Twice with meals
Linagliptin/empagliflozin	Glyxambi	5/10, 5/25	Once daily
Dapagliflozin/saxagliptin	Qtern	10 mg/5mg	Once daily
Alogliptin/pioglitazone	Oseni	25/15, 25/30, 25/45, 12.5/15, 12.5/30, 12.5/45	Once daily
Alogliptin/metformin	Kazano	12.5/500, 12.5 mg/1000	Twice with meals
Ertugliflozin/sitagliptin	Steglujan	5/100, 15, 100	Once daily
Saxagliptin/dapagliflozin/metformin XR	Qternmet XR	2.5/2.5/1000, 2.5/5/1000, 5/5/1000, 5/10/1000	Once daily

Newest triple combination: Empagliflozin/linagliptin/metformin (Trijardy XR)

Case 4: 70-year-old obese female with T2D for 25 years

- A1C 8.4% on maximum doses of metformin, a SFU, and a DPP-4 inh.
- Medical Hx: HTN, arthritis, recent admission for CHF
- Family Hx: Type 2 diabetes and obesity (both parents)
- Notes:
 - Fearful of injections and gaining weight BMI 31 kg/m²
 - Osteoporosis and CKD 3A (eGFR 58 mL/min/m²)
 - Home glucose monitoring shows FBS (147-219 mg/dL) with a few post dinner values (188 to 275 mg/dL)

How would you treat this patient to lower her A1c?

A	Add a TZD
B	Add a SGLT-2 inh. (cana-, dapa-, empa-, ertugliflozin)
C	Try to convince her to add a GLP-1 RA (exena-, liraglu-, dulaglu-, semaglutide)
D	Try to convince her to add a basal insulin at bedtime

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Case 4 Continued

- Low dose SGLT-2i was added to her regimen and then titrated to the maximum dose after one month
- A1C dropped to 7.3% (baseline 8.4%) and she lost 15 lbs
- She experienced a yeast infection which was easily treated with oral fluconazole and she did not want to stop the SGLT2i
- LDL-C increased from 100 to 108 mg/dL (8% rise), HDL-C increased 10%, and her TGs decreased by 25%

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SGLT-2 Inhibitors

Mechanism of Action	Reduce renal glucose reabsorption and increases urinary glucose excretion
Benefits	<ul style="list-style-type: none"> • No hypoglycemia (except when being used with SFU or insulin) • Mean A1c reduction ~1% (starting from a baseline A1c of ~8.0%) • Weight loss (2-5% of body weight) and systolic BP reduction (2-6mmHg)
Concerns	<ul style="list-style-type: none"> • Genital mycotic infections. In women (6 to 12% higher than comparator) and in uncircumcised males (2 to 6% higher than comparator) • Hypotension secondary to volume contraction especially in the elderly, those on loop diuretic use and in patients with reduced renal function. • 4 to 8% elevation in LDL cholesterol (TGs goes down and HDL goes up) • Assess renal function (discussed later) • New label warnings: DKA (discussed later), Fournier's Gangrene, acute kidney injury, UTI, risk of amputation (discussed later), bone fractures
Clinical Pearls	<ul style="list-style-type: none"> • Cana now approved for renal protection and can be used with an eGFR down to 30 • Empa- Dapa- and canagliflozin showed positive CVD outcome trials (discussed later) • Can be added to any other oral agent or injectable • Inform women to practice good hygiene and look out for yeast infections (may want to suggest to have some anti yeast infection medication at home such as miconazole)

Edelman SV, Henry RR. Diagnosis and management of type 2 diabetes. 12th Edition. Professional Communications, Inc., Greenwich, CT. 288 pages, 2014.



Generic and Trade Names: SGLT-2 Inhibitors

	Generic Name	Trade Name
SGLT-2 Inhibitor	Canagliflozin Dapagliflozin Empagliflozin Ertugliflozin	Invokana Farxiga Jardiance Steglatro

Canagliflozin:

- Suggested starting dose: 100 mg daily before first meal of day (eGFR >45 mL/min)/with CKD can use to an eGFR of 30ml/min
- Increase to 300 mg daily if tolerating 100 mg daily and eGFR > 60 mL/min

Dapagliflozin:

- Starting dose: 5mg daily in morning with or without food (eGFR for both doses > 60 mL/min)
- Increase to 10 mg daily if tolerating and need additional glycemic control

Empagliflozin:

- Starting dose: 10 mg daily in morning with or without food (eGFR>45 mL/min)
- Increase to 25 mg daily if tolerating and need additional glycemic control (eGFR>45 mL/min)

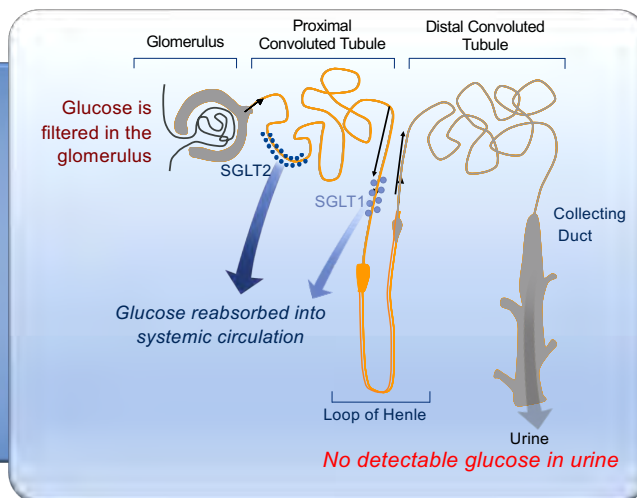
Ertugliflozin:

- Starting dose: 5 mg daily in morning with or without food (eGFR for both doses >60 mL/min)
- Increase to 15 mg daily if tolerating and need additional glycemic control



Renal Handling of Glucose in a Non-Diabetic Patient

- 180 g/day/1.73 m² is filtered glucose load¹
- SGLT-2 transports 90% of filtered glucose out of the tubular lumen¹⁻⁴

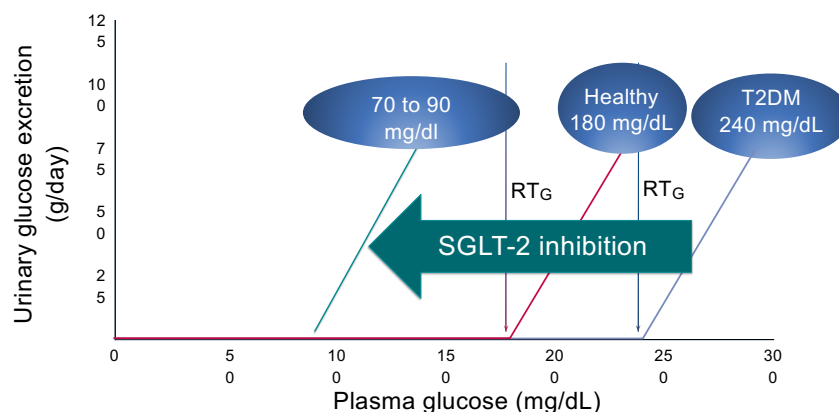


SGLT = sodium-glucose co-transporter.

1. Wright EM et al. *J Intern Med.* 2007;261(1):32-43. 2. Kanai Y et al. *J Clin Invest.* 1994;93(1):397-404. 3. You G et al. *J Biol Chem.* 1995;270(49):29365-29371. 4. Wright EM. *Am J Physiol Renal Physiol.* 2001;280(1):F10-F18.



Renal Glucose Reabsorption in Normal, T2D, and with SGLT-2 Inhibition



Adapted with permission from Abdul-Ghani, DeFronzo RA.

T2DM = type 2 diabetes mellitus.

1. Cowart SL, Stachura ME. In: Walker HK et al, eds. *Clinical Methods: The History, Physical, and Laboratory Examinations.* 3rd ed. Boston, MA: Butterworths; 1990:653-657. 2. Abdul-Ghani MA, DeFronzo RA. *Endocr Pract.* 2008;14(6):782-790. 3. Nair S, Wilding JP. *J Clin Endocrinol Metab.* 2010;95(1):34-42. 4.: Janssen Research & Development LLC. FDA Briefing Document. Endocrinologic and Metabolic Drugs Advisory Committee, 2013.



FDA Drug Safety Communication: the Prescribing Information for ALL SGLT-2 inhibitors was updated to include new Warnings and Precautions for Ketoacidosis, Urosepsis and Pyelonephritis

December 14, 2015

1. Extremely low incidence, mostly type 1's and type 2's receiving insulin
2. Complex mechanism related to paradoxical increase in glucagon promoting ketosis in the setting of glycosuria so extreme hyperglycemia is limited (watch for euglycemic DKA)
3. Be especially cautious in women with a history of UTIs, pyelonephritis and/or genital mycotic infections
4. August 2018: New warning for extremely rare but serious infection called Fournier's gangrene

Brooks M. SGLT2 Inhibitors May Cause Ketoacidosis: FDA. Retrieved from <http://www.medscape.com/viewarticle/844754>
Erondu N, et al. Diabetes Care September 2015 38:1680-1686; 2015

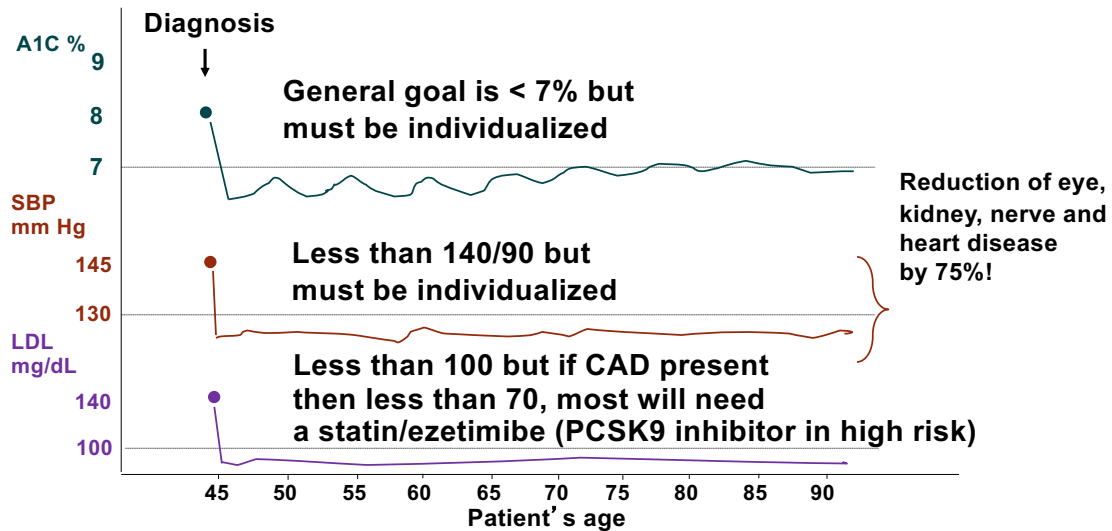


What is the most common cause of death in type 2 diabetes?

A	Nephropathy including end-stage renal disease requiring dialysis or transplantation
B	Complications from peripheral and autonomic neuropathy
C	Heart disease or stroke
D	Complications from obesity
E	Peripheral arterial disease



Primary Objectives of Effective Management: Important Basics...The 'ABCs'



American Diabetes Association. *Diabetes Care*. 2017;40(suppl 1)

Blood Pressure Management

Individualize BP Goals:

<140/90 mmHg (10-yr CV risk <15%)

<130/80 mmHg (10-yr CV risk >15%)

Dyslipidemia Management

Individualize lipid Goals:

LDL < 100mg/dl in all PWD

LDL < 70 mg/dl if ASCVD present

Triglycerides less than 200mg/dl

HDL as high as you can get it!

Diabetes Care Volume 43, Supplement 1, January 2020, S111-S134

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Table 10.2—High-intensity and moderate-intensity statin therapy*

High-intensity statin therapy (lowers LDL cholesterol by $\geq 50\%$)	Moderate-intensity statin therapy (lowers LDL cholesterol by 30–49%)
Atorvastatin 40–80 mg	Atorvastatin 10–20 mg
Rosuvastatin 20–40 mg	Rosuvastatin 5–10 mg
PCSK9 inhibitors (evolocumab and alirocumab) if LDL not at goal on maximally tolerated statin/ezetimide	Simvastatin 20–40 mg
Just approved in 2020 bempedoic acid (Nexletol), first in class LDL medication used for add on to statin therapy	Pravastatin 40–80 mg
	Lovastatin 40 mg
	Fluvastatin XL 80 mg
	Pitavastatin 1–4 mg

*Once-daily dosing. XL, extended release.

Diabetes Care Volume 43, Supplement 1,
January 2020, S111-S134

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Management Of Hypertriglyceridemia

1. Elevated triglycerides combined with low HDL levels are part of the insulin resistant state and metabolic syndrome.
2. Diet, exercise and improved glycemic control will improve but not typically normalize elevated TG levels in type 2 DM.
3. The goal is to get the TGs to below 200mg/dl, which in turn will elevate the HDL levels
4. Fibric acid derivatives such as fenofibrate are commonly used to treat high TGs.
5. Icosapent ethyl is an omega-3 fatty acid that has the formal indication from the FDA to reduce heart attacks and strokes in patient who have or are at risk for ASCVD.

N Engl J Med 2019; 380:11-22 REDUCE-IT trial

TCOYD
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Non-Insulin CVOTs in T2D: DPP-4 Inhibitors

Study	SAVOR	EXAMINE	TECOS	CAROLINA	CARMELINA
DPP4-i	saxagliptin	alogliptin	sitagliptin	linagliptin	linagliptin
Comparator	placebo ✓	placebo ✓	placebo ✓	sulfonurea ✓	placebo ✓
N	16,500	5,400	14,000	6,000	8,200
Results	NEUTRAL 2013	NEUTRAL 2013	NEUTRAL June 2015	NEUTRAL 2017	NEUTRAL 2017

TCOYD
TAKING CONTROL OF YOUR DIABETES

Non-Insulin CVOTs in T2D: SGLT-2 Inhibitors (Primarily driven by a reduction in heart failure)

Study	EMPA-REG	CANVAS	DECLARE	EMPEROR-Reduced	VERTIS-CV
SGLT-2-i	empagliflozin ✓	canagliflozin ✓	dapagliflozin ✓	empagliflozin ✓	ertugliflozin
Comparator	placebo	placebo	placebo	placebo	placebo
N	POSDITIVE 7,000	POSDITIVE 10,000	POSDITIVE 10,000	POSDITIVE 7,000	3,900
Results	Sept 2015	2017	2018	2020	Late 2020

Courtesy of Silvio Inzucchi MD, Yale University

TCOYD
TAKING CONTROL OF YOUR DIABETES

Non-Insulin CVOTs in T2D: GLP-1 RA (Primarily driven by a reduction in death due to cardiovascular disease)

Study	LEADER	ELIXA	SUSTAIN 6	EXSCEL	HARMONY	REWIND	PIONEER 6
GLP1-RA	Lira-glutide	Lixi-senatide	Semaglutide	Exe-natide LR	Albi-glutide	Dula-glutide	Oral semaglutide
Comparator	placebo	placebo	placebo	placebo	placebo	placebo	placebo*
N	16,500	14,000	6,000	5,400	9,400	8,200	3,100
Results	POSITIVE	NEUTRAL	POSITIVE	NEUTRAL	POSITIVE	POSITIVE*	POSITIVE*

*CV death less with oral sema; no difference in non-fatal MI or non-fatal stroke. Median time in study: 15.9 months NEJM 2019;381:841-851.

Adapted from a slide courtesy of Silvio Inzucchi MD, Yale University



Diabetes Medications FDA Approved for CV Risk Reduction

Empagliflozin (based on EMPA-REG trial data)

- to reduce the risk of cardiovascular death in adult patients with type 2 diabetes mellitus and established cardiovascular disease
- also FDA Fast Track designation provided to reduce the risk of cardiovascular death and hospitalization for heart failure in people with CHF (based on EMPEROR-Reduced trial)

Liraglutide (based on LEADER data)

- to reduce the risk of major adverse cardiovascular (CV) events (CV death, non-fatal myocardial infarction, or non-fatal stroke) in adults with type 2 diabetes mellitus and established CVD

Canagliflozin (based on CANVAS program data)

- to reduce the risk of major adverse cardiovascular events in adults with type 2 diabetes mellitus and established cardiovascular disease

Semaglutide (based on SUSTAIN 6)

- To reduce the risk of major adverse cardiovascular events (MACE) including cardiovascular death, non-fatal heart attack, or non-fatal stroke in adults with type 2 diabetes and established cardiovascular disease (CVD).

Dulaglutide (based on REWIND data)

- for the reduction of major adverse cardiovascular events (MACE) in adults with type 2 diabetes who have established cardiovascular (CV) disease or multiple cardiovascular risk factors.



Not All CVOTs are Created Equal

- Differences in study design: powered for safety or superiority
- Patient characteristics: age, weight, co-morbid complications, presence of CVD and risk factors
- Comparators may be different
- Weight gain and hypoglycemia differences
- Regional differences
- Outcomes differ (mortality, non-fatal and fatal MI, stroke, etc.)
- Study conduct and adherence may effect results

Gautam Das, Journal of Diabetes Research & Clinical Metabolism 2015, <http://www.hoajonline.com/journals/pdf/2050-0866-4-3.pdf>

TCOYD
TAKING CONTROL OF YOUR DIABETES

Diabetes Medications FDA Approved for Renal Disease

- Canagliflozin (CREDENCE study): Reduce the risk of end-stage kidney disease, doubling of the serum creatinine, cardiovascular death and hospitalization for CHF in patients with type 2 diabetes with nephropathy (eGFR between 30 and 90 ml/min) and albuminuria > 300mg
- Dapagliflozin (DAPA-CKD study): Reduced the risk of the composite of a sustained decline in eGFR of at least 50%, end-stage kidney disease, or death from cardiovascular or renal cause in patients with CKD (regardless of the presence or absence of diabetes)
- EMPA-KIDNEY: On-going

Jardine MF et al. Am J Nephrol. 2017;46(6):462-472; Perkovic V et al. N Engl J Med. 2019;380(24):2295-2306; Neal B et al. N Engl J Med. 2017;377(7):644-657; Zinman B et al. N Engl J Med. 2015;373(22):2117-2128

TCOYD
TAKING CONTROL OF YOUR DIABETES

Key Principles of Management of T2D

- Glycemic targets and glucose-lowering therapies should be individualized
- Diet, exercise, and diabetes self-management education and support are the foundations of therapy
- Unless contraindicated, metformin is the preferred first line drug
- After metformin, the first consideration is whether the patient has established ASCVD, HF, or CKD. If not, then whether hypoglycemia, weight, or financial status are dominant issues. Shared decision making is KEY!



Key Principles of Management of T2D

- GLP-1 RA are the preferred first injectable therapy over basal insulin except in patients with very poor glycemia control
- Many patients over time will require insulin therapy alone or in combination with other agents to maintain glycemic control
- Vascular disease is the most common cause of death and prevention strategies need to be emphasized (A1c, aspirin, blood pressure, cholesterol, smoking cessation, and diabetes drugs that reduce ASCVD/heart failure)



Thank you!

Q&A TO FOLLOW

WWW.TCOYD.ORG

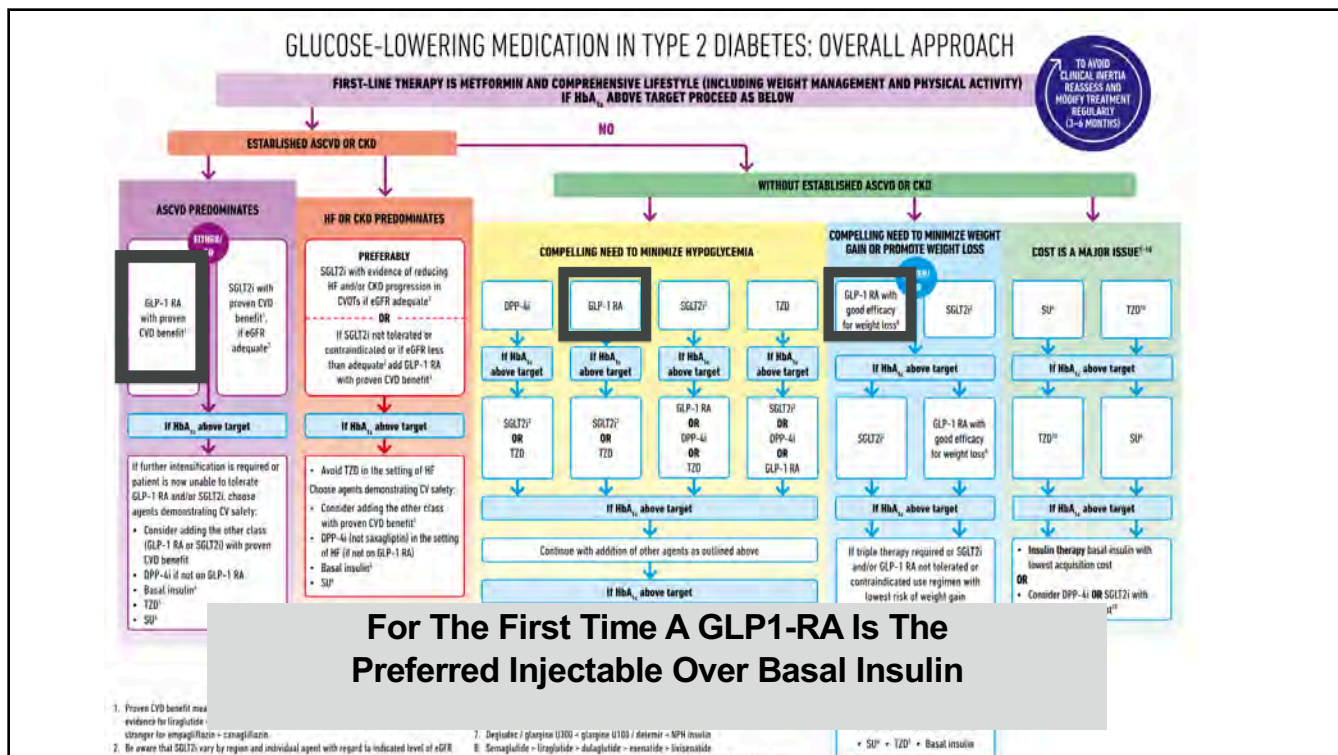
Taking Control Of Your Diabetes, 501(c)3 is a not-for-profit educational organization.



Lecture 4: 2:15 – 3:30 p.m. PST

Steven V. Edelman, MD, Presents:

Practical Application of Injectable Agents and Their Cardiovascular Effects:
Individualized Treatment Strategies



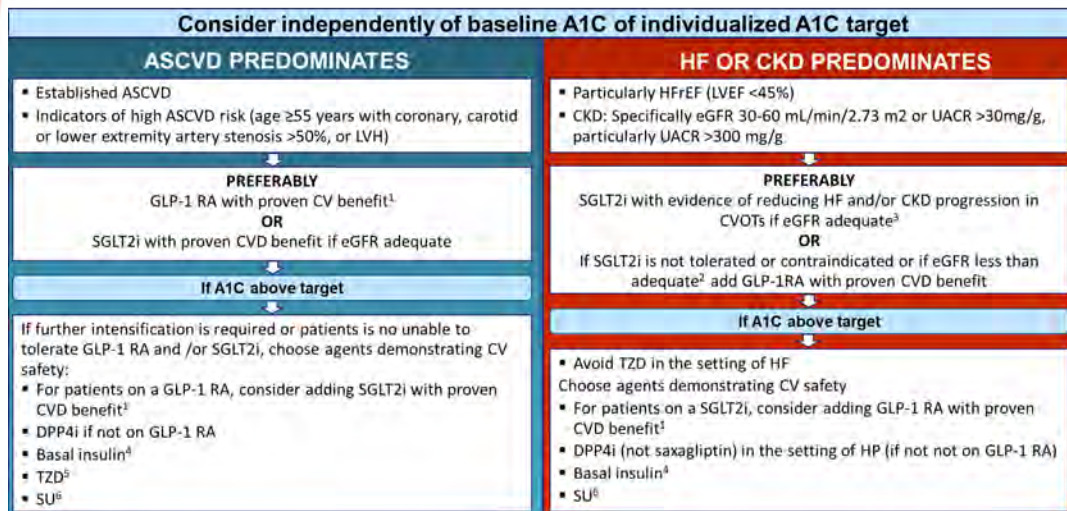
Case 1: 54 year old male with type 2 diabetes for 10 years

- History of dyslipidemia, hypertension, NAFLD
- Strong family history of type 2 diabetes
- Currently on metformin, SFU and a DPP4 inhibitor
- Recent myocardial infarction s/p 4 cardiac stent insertions
- A1c 9.3%
- Creatinine 1.3 eGFR 70
- HGM data: ranges from 82 to 379 mg/dl
- Bedtime average 210 mg/dl SD 76mg/dl
- Morning average 221 mg/dl

Which of the following would you recommend for this patient?

A	Initiate basal insulin
B	Initiate a GLP-1 Receptor Agonist (RA)
C	Initiate premixed insulin (70/30) BID
D	Initiate a fixed combination of a basal insulin and a GLP-1RA

High CV risk or established ASCVD, CKD and/or HF



1. Proven CVD benefit means it has label indication of reducing CVD events 2. Be aware that SGLT2i labelling varies by region and individual agent with regard to indicated level of eGFR for initiation and continued use 3. Empagliflozin, canagliflozin and dapagliflozin have shown reduction in HF and to reduce CKD progression in CVOTs. Canagliflozin has primary renal outcome data from CREDENCE. Dapagliflozin has primary HF outcome data from DAPA-HF 4. Degludec and U100 glargine have demonstrated CVD safety 5. Low dose may be better tolerated though less well studied for CVD effects 6. Chose later generation SU to lower risk of hypoglycemia. Glimepiride has shown similar CV safety to DPP-4i.

Basal Insulin

vs

GLP-1 RA

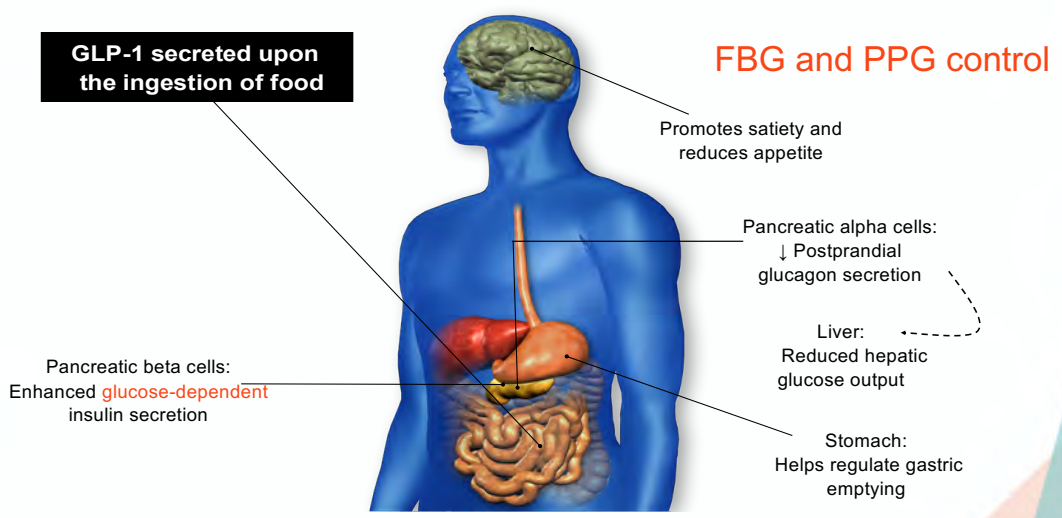
(an incretin hormone)

Insulin: Injected once or twice a day	GLP-1 RA: Injectable once or twice a day, injectable once weekly, or oral once daily
Need to titrate dose to achieve the desired FBS	Titrate to the highest acceptable dose to avoid nausea
Need to institute home glucose monitoring (SMBG)	“No” need for SMBG
Important to have frequent follow up when initiating basal insulin (days to weeks)	Follow up not as crucial
Weight gain	Weight loss
Hypoglycemia	No Hypoglycemia

Edelman SV, Henry RR. Diagnosis and management of type 2 diabetes. 12th Edition. Professional Communications, Inc., Greenwich, CT. 288 pages, 2014.



GLP-1 Effects: Glucoregulatory Role of Incretins



Adapted from Flint A, et al. *J Clin Invest*. 1998;101:515-520.; Adapted from Larsson H, et al. *Acta Physiol Scand*. 1997;160:413-422.; Adapted from Nauck MA, et al. *Diabetologia*. 1996;39:1546-1553.; Adapted from Drucker DJ. *Diabetologia*. 1998;47:159-169.



GLP-1 Receptor Agonists

Mechanism of Action	<ul style="list-style-type: none"> Mimic the effects of human GLP-1
Benefits	<ul style="list-style-type: none"> Significant A1c reductions (1.0 to 3.0% depending on baseline) Shorter acting GLP-1 RAs have greater effects on PPG Weight loss No hypoglycemia Once daily, twice daily and once weekly formulations
Concerns	<ul style="list-style-type: none"> GI side effects (typically nausea) Contraindicated in patients with a personal or family history of MTC or MEN2 Relative contraindication in patients with a history of pancreatitis (important to know the etiology)
Clinical Pearls	<ul style="list-style-type: none"> Ideal choice in obese patients with poor control, especially those on large doses of insulin “No” need to initiate or increase glucose testing Several with positive CVOT results

Edelman SV, Henry RR. Diagnosis and management of type 2 diabetes. 12th Edition. Professional Communications, Inc., Greenwich, CT. 288 pages, 2014.



Generic and Trade Names: GLP-1 RAs

	Generic Name	Trade Name
GLP-1 Receptor Agonists	Exenatide Twice-daily Once-weekly	Byetta Bydureon
	Liraglutide Once-daily	Victoza
	Dulaglutide Once-weekly	Trulicity
	Lixisenatide Once-daily	Adlyxin
	Semaglutide Once weekly	Ozempic
	Oral Semaglutide Once daily	Rybelsus



Generic and Trade Names: GLP-1 RAs, Continued

	Generic Name	Trade Name
Basal Insulin/GLP-1 Receptor Agonist Fixed Combination	Glargine/lixisenatide once daily	Soliqua
	Degludec/liraglutide once-daily	Xultophy



Summary of Completed GLP-1 receptor agonists Cardiovascular Outcome Trials (CVOTs)

MACE Outcomes

	Study Drug n/N (%)	Placebo n/N (%)	Hazard Ratio	95% CI		P-Value (superiority)
ELIXA (lixisenatide vs PBO)	406/3034 (13.4%)	399/3034 (13.2%)	1.02	0.89, 1.17		0.81
LEADER (liraglutide vs PBO)	609/4668 (13%)	694/4672 (14.9%)	0.87	0.78, 0.97		0.01*
SUSTAIN-6* (semaglutide vs PBO)	108/1648 (6.6%)	146/1649 (8.9%)	0.74	0.58, 0.95		<0.001*
EXSCCEL (exenatide vs PBO)	839/7356 (11.4%)	905/7396 (12.2%)	0.91	0.83, 1.00		0.06 <0.001 (NI)
Harmony Outcomes (albiglutide vs PBO)	338/4731 (7.1%)	428/4732 (9.1%)	0.78	0.68, 0.90		0.0006

*Superiority testing not a prespecified analysis

0 1 2
◀ Favors Treatment Favors Placebo ▶

1. Pfeffer MA, et al. N Engl J Med. 2015;373(23):2247-57. 2. Bentley-Lewis R, et al. Am Heart J. 2015;169(5):631-638.e7. 3. Marso SP, et al. Am Heart J. 2013;166(5):823-30.e5. 4. Marso SP, et al. N Engl J Med. 2016;375(4):311-22. 5. Marso SP, et al. N Engl J Med. 2016;375(19):1834-1844. 6. Holman RR, Bethel MA, et al. N Engl J Med. 2017;377(13):1228-1239. 7. Hernandez A, et al. Lancet. (online first October 2, 2018).



CVOTs of GLP-1 RAs (SGLT2 Inhibitors Indicated for CHF/CKD)

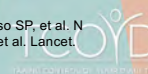
Hospitalization for Heart Failure

	Study Drug n/N (%)	Placebo n/N (%)	Hazard Ratio	95% CI		P-Value
ELIXA (lixisenatide vs PBO)	122/3034 (4.0%)	127/3034 (4.2%)	0.96	0.75, 1.23		0.75
LEADER (liraglutide vs PBO)	218/4668 (4.7%)	248/4672 (5.3%)	0.87	0.73, 1.05		0.14
SUSTAIN-6 (semaglutide vs PBO)	62/1648 (3.6%)	54/1649 (3.3%)	1.11	0.77, 1.61		0.57
EXSCCEL (exenatide vs PBO)	219/7356 (3.0%)	231/7396 (3.1%)	0.94	0.78, 1.13		

Harmony Outcomes
(albiglutide vs PBO)
HR 0.85 (0.70, 1.04); p=0.113
Composite of CV death or HHF

0 1 2
◀ Favors Treatment Favors Placebo ▶

1. Pfeffer MA, et al. N Engl J Med. 2015;373(23):2247-57. 2. Bentley-Lewis R, et al. Am Heart J. 2015;169(5):631-638.e7. 3. Marso SP, et al. Am Heart J. 2013;166(5):823-30.e5. 4. Marso SP, et al. N Engl J Med. 2016;375(4):311-22. 5. Marso SP, et al. N Engl J Med. 2016;375(19):1834-1844. 6. Holman RR, Bethel MA, et al. N Engl J Med. 2017;377(13):1228-1239. 7. Hernandez A, et al. Lancet (online first October 2, 2018).



ITCA 650—Medical Device To Deliver a GLP-1RA (exenatide)

TECHNOLOGY



MEDICATION: EXENATIDE

- Subcutaneous delivery system; short office procedure
- Small micropump
- maintains stability at temps ≈37°C
- secretes medication for ≥ 12 months

- Previously-approved GLP-1 therapeutic which demonstrates:
 - glycemic control
 - weight loss
 - safety



Not yet approved by the FDA



Case 2: 72 year old Caucasian woman with type 2 diabetes for 23 years

- On maximal doses of metformin, SU, and a SGLT-2 inhibitor
- She adamantly does not want to take insulin for fear of weight gain
- PMH: dyslipidemia, hypertension, papillary thyroid cancer and obesity (BMI=31)
- Both parents and two siblings have type 2 diabetes and early CVD
- eGFR 65 ml/min
- Her A1c is 8.8 % (goal for this patient at least less than 8%)
- Average FBS is in the 180s (does not test at other times)

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THERAPEUTIC COORDINATION OF YOUR DIABETES

What would you recommend now for this patient?

A	Start a DPP4 inhibitor
B	Try to convince her to start basal insulin and titrate the dose to get her FBS less than 140mg/dl
C	Start a GLP1-RA
D	Initiate a fixed combination of a basal insulin and a GLP-1RA

TCOYD
THERAPEUTIC COORDINATION OF YOUR DIABETES

Case 2 continued

- She agreed to start a once weekly GLP-1RA (exenatide, dulaglutide or semaglutide)
- When prescribing once-weekly GLP-1 RA, inform patient that it may take several weeks to reach equilibration and, with once-weekly exenatide, skin nodules may occur (self limited and resolve in a few days to weeks).
- She experienced no nausea or hypoglycemia. Over the next three months she lost 13 pounds and her A1c fell from 8.8% to 7.2%.

* Increased frequency of SMBG testing not a requirement with GLP-1 receptor agonists

Before GLP-1*

FBS (mg/dl)	PPG (mg/dl)
-------------	-------------

Average 188	
-------------	--

After GLP-1*

FBS (mg/dl)	PPG (mg/dl)
-------------	-------------

Average 139	
-------------	--

	Average 167
--	-------------

TCOYD
TRANSFORMING YOUR DIABETES

Fixed Combinations Of Basal Insulin and GLP- Receptor Agonist

Insulin degludec/liraglutide
Insulin glargine/lixisenatide

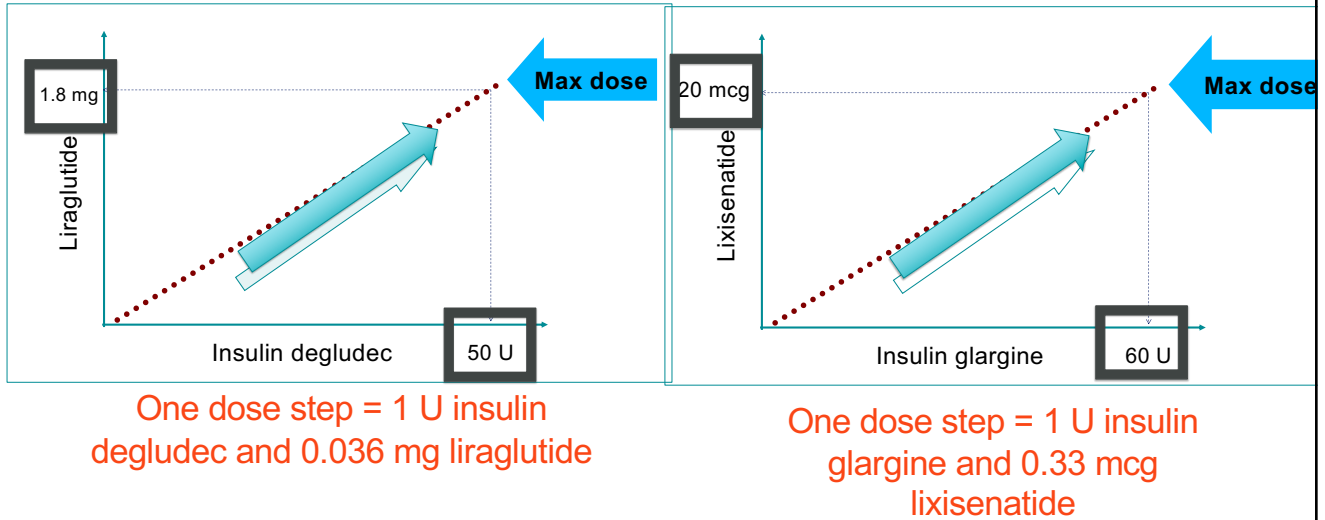


- | | |
|---|---|
| <ul style="list-style-type: none"> ▪ 1 dose step (unit) has 1 unit insulin degludec and 0.036 mg of liraglutide (max. dose is 50 insulin degludec/1.8mg liraglutide) ▪ Injected once daily at same time each day with or without food | <ul style="list-style-type: none"> ▪ 1 dose step (unit) has 1 unit insulin glargine and 0.33 mcg lixisenatide (max. dose is 60 insulin glargine/20 mcg lixisenatide) ▪ Injected once daily within one hour prior to the first meal of the day |
|---|---|

Lancet Diabetes Endocrinol. 2014 Nov;2(11):856-8, 2017 PDR Pls

TCOYD
TRANSFORMING YOUR DIABETES

Fixed-Ratio Combination of Insulin Degludec And Liraglutide And U-100 Glargine and Lixisenatide



Buse JB, et al. *Diabetes Care*. 2014; 37:2926-33.

Insulin Degludec/Liraglutide vs. Insulin Glargine/Lixisenatide

Pen dose steps (units): insulin degludec + liraglutide	Pen dose steps (units): insulin glargine + lixisenatide
10 dose steps=10 units insulin degludec +0.36 mgs of liraglutide 50 dose steps=50 units insulin degludec +1.8 mgs of liraglutide	15 dose steps=15 units insulin glargine + 5 mcg of lixisenatide 30 dose steps=30 units insulin glargine + 10 mcg of lixisenatide 60 dose steps=60 units insulin glargine + 20 mcg of lixisenatide
Starting dose: 16 dose steps which has 16 units insulin degludec + 0.58 mgs of liraglutide	Starting dose: If glargine U-100 dose is <30, start at 15 dose steps which has 15u glargine + 5mcg lixi If glargine U-100 dose is >30, start at 30 dose steps which has 30u glargine + 10 mcg lixi
Titrate according to FBG, as if you were using basal insulin alone, generally 2 dose steps at a time, usually every 3-4 days	Titrate according to FBG, as if you were using basal insulin alone, generally 2-4 dose steps at a time, usually weekly
Maximum dose is 50 units of insulin degludec and 1.8 mgs of liraglutide	Maximum dose is 60 units of insulin glargine and 20 mcgs of lixisenatide

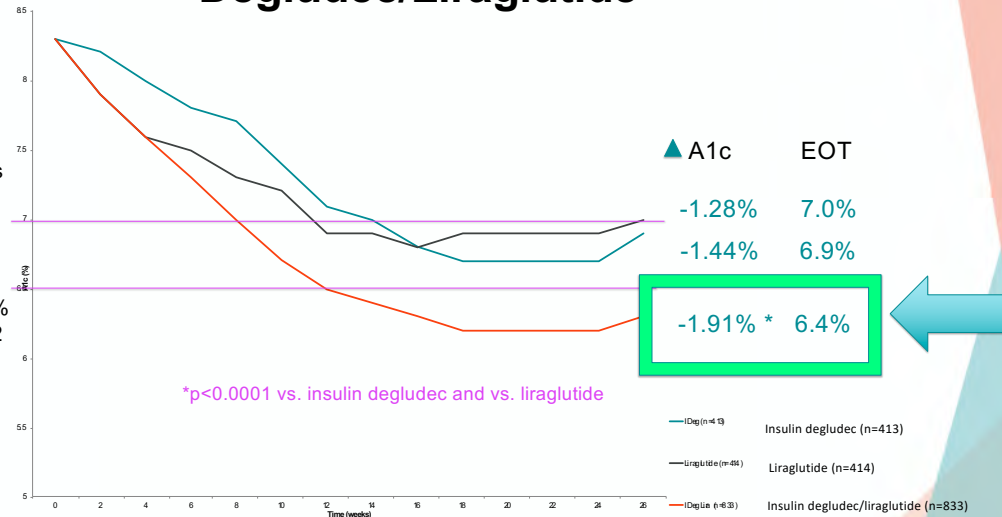
PNP (PL for both) 2017



A1c of 8.3% at baseline drops to 6.4% with Insulin Degludec/Liraglutide

Inclusion Criteria:

- Type 2 diabetes
- Insulin naïve treated with metformin ± pioglitazone
- A1c 7.0 – 10.0%
- BMI < 40 kg/m²
- Age ≥ 18 years

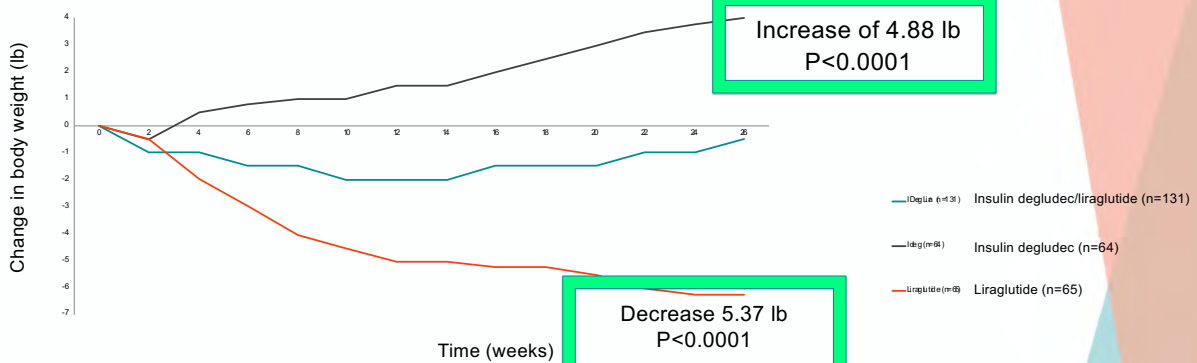


Mean values (+SEM) based on FAS and LOCF imputed data; EOT = end of trial; p-values are from an ANCOVA ADA/EASD A1c target < 7.0%; AACE A1c target < 6.5%

Buse J et al. ADA 2013. 65-OR



Body Weight and Hypoglycemia



	Insulin degludec/liraglutide Rate (episodes/PYE)	Insulin degludec Rate (episodes/PYE)	Liraglutide Rate (episodes/PYE)	Insulin degludec/liraglutide vs. insulin degludec RR Estimate (95% CI)	Insulin degludec/liraglutide vs. liraglutide RR Estimate (95% CI)
Hypoglycemia	1.80	2.57	0.22	0.68 (0.53; 0.87)*	7.61 (5.17; 11.21)**

Mean weight values (±SEM) based on FAS and LOCF imputed; estimated treatment difference and p-values are from an ANCOVA analysis
 Hypoglycemia: PG <56 mg/dL and/or requiring assistance, PYE: Patient years exposure; RR: Rate ratio: I/Degl./lira/Comparator; ns=0.05 two-sided; *p<0.002; **p<0.0001

Buse J et al. ADA 2013, 65-OR

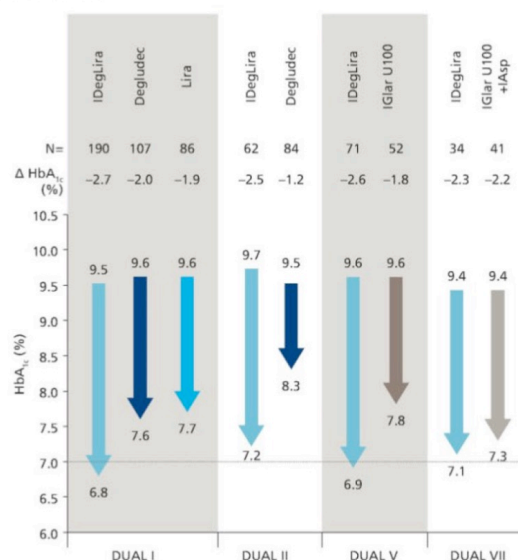


Effects of insulin degludec/liraglutide in patients with poorly controlled type 2 diabetes with HbA1c >9%: analyses from the DUAL program

Frias JP et al. Diabetes 2018 Jul; 67(Supplement 1): -
<https://doi.org/10.2337/db18-1092-P>

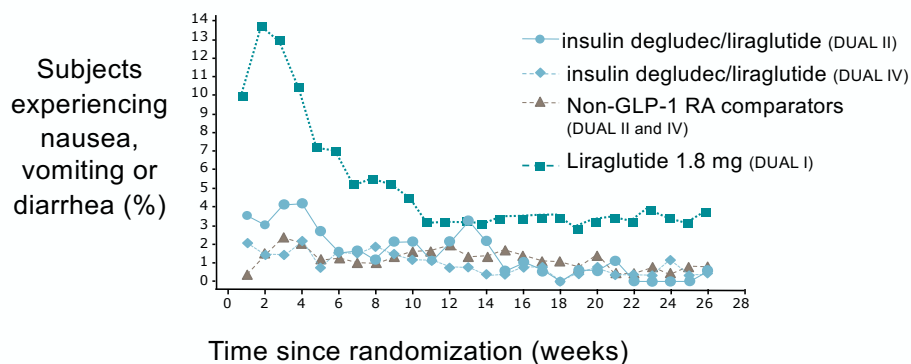
Figure 2: HbA_{1c} at baseline and end of trial

Key result



End of trial data based on full analysis set for all trials except DUAL VII (observed data; N-numbers for week 26). Arrows in graph indicate direction of change in HbA_{1c}, from baseline to end of trial. Δ HbA_{1c}, change in HbA_{1c}, from baseline to end of trial; IAsp, insulin aspart; IDegLira, insulin degludec/liraglutide; IGlar U100, insulin glargine 100 units/mL; Lira, liraglutide.

Gastrointestinal Side Effects: Gradual Titration Helps

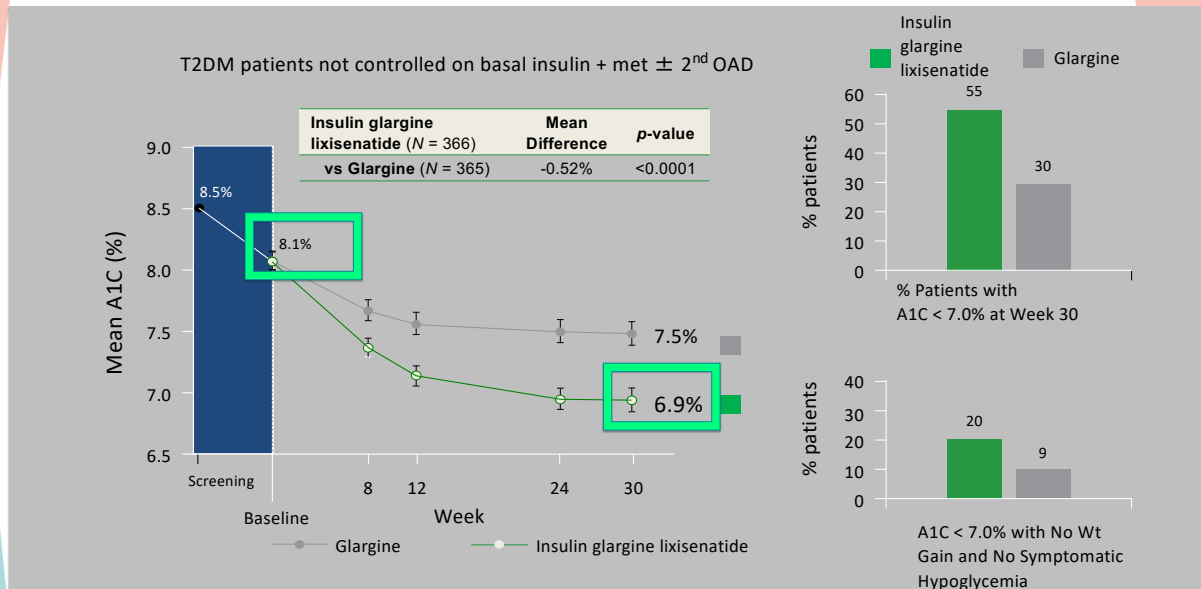


p =non-significant for odds of experiencing gastrointestinal side effects for subjects on insulin degludec/liraglutide versus non-GLP-1 RA comparator

Aroda et al. Diabetes 2015;64 (Suppl. 1):A235; abstract 1009-P

TCOYD
 TAKING CONTROL OF YOUR DIABETES

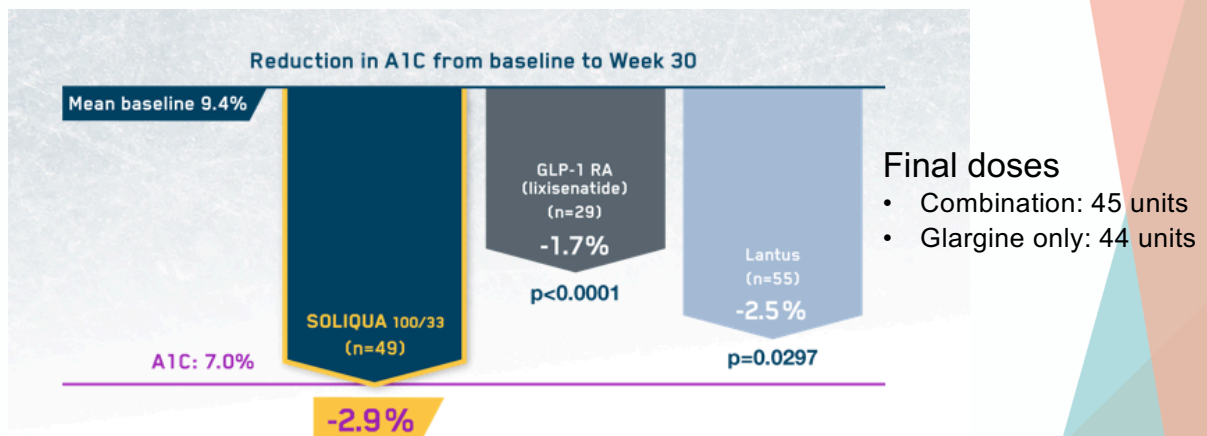
Efficacy of Fixed-Ratio insulin glargine lixisenatide in T2DM Patients Not Controlled on Basal Insulin



IGlarLixi Briefing Document. www.fda.gov. Accessed May 25, 2016.

TCOYD
TAKING CONTROL OF YOUR DIABETES

Post Hoc Analysis insulin glargine/lixisenatide In Patients With Very Poor Glycemic Control (A1c >9%). LixiLan O Study



Davis MF, et al. Diab Obes Metab. 2019. <https://doi.org/10.1111/dom.13791>.

TCOYD
TAKING CONTROL OF YOUR DIABETES

Summary: Benefits for Combining GLP-1 Receptor Agonists and Basal Insulin Analogs

- Combined glycemic effects of GLP-1RA and basal insulin provides greater glycemic efficacy than either of its component parts.
- Dose related adverse effects of each component (nausea and weight gain) are minimized.
- No increased risk of hypoglycemia in the setting of improved glycemic control as compared to basal insulin alone.
- In the setting of inadequate control on basal insulin, adding a GLP-1RA is associated with greater benefits (weight loss and minimal hypo) than adding prandial insulin.

TCOYD
THE COVERED OPIC COMPANY

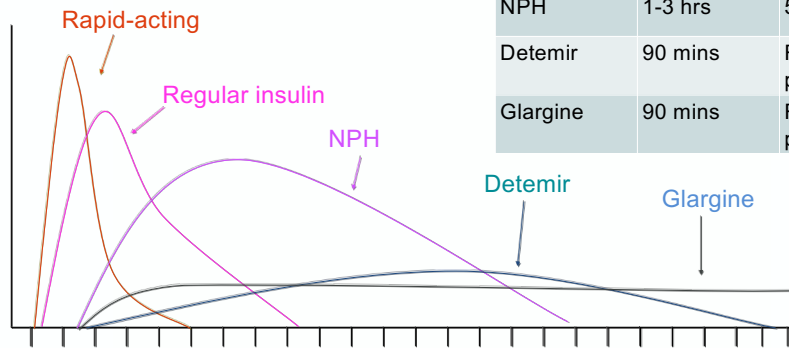
Generic and Trade Names: Insulin

	Generic Name	Trade Name
Fast-Acting Insulin	regular U-500 regular aspart faster acting aspart glulisine lispro (U-100 and U-200) Follow on biologic lispro inhaled insulin	Humulin R, Novolin R Humulin R U-500 NovoLog Fiasp Apidra Humalog Admelog Afrezza
Basal Insulin	intermediate-acting: NPH long-acting: detemir glargine (U-100) glargine (U-300) degludec (U-100/200) follow-on biologic glargine (U-100)	Humulin N Novolin NPH Levemir Lantus Toujeo Tresiba Basaglar

TCOYD
THE COVERED OPIC COMPANY

Time Action Profiles: Traditional Insulins

Insulin	Onset	Peak	Duration
Rapid-acting	10-15 mins	60-90 mins	4-5 hrs
Regular	30-60 mins	2-4 hrs	5-8 hrs
NPH	1-3 hrs	5-8 hrs	12-18 hrs
Detemir	90 mins	Relatively peakless	12-24 hrs
Glargine	90 mins	Relatively peakless	24 hrs



Inhaled insulin: peak by 10-15 min, duration of 2-3 hrs Faster-acting aspart: onset faster, duration shorter, than rapid-acting

Lepore M et al. *Diabetes*. 2000;49:21 42-21 48; Howey DC et al. *Diabetes*. 1994;43:396-402; Plank J et al. *Diabetes Care*. 2005;28:1107-1112; Wittlin SD et al. *Insulin Therapy*. Marcel Dekker, Inc.;2002:73-85.

TCOYD
TRANSFORMING YOUR DIABETES

Benefits Of U-300 Glargine And Degludec In Type 1 and Type 2 Diabetes

- Less intra-subject variability
- Less hypoglycemia
- Less weight gain
- Flat, stable and prolonged action greater than 24 hours
- Tell patients it takes 4 to 5 days to reach equilibration and they may need correction doses
- 1 to 1 conversion from prior basal dose (patients switching from U-100 to U-300 glargine may need ~15% more)
- Both insulins come in easy to use pens

Riddle MC et al. *Diabetes Care*. 2014;37:2755-2762; Yki-Järvinen H et al. *Diabetes Care*. 2014; Published ahead of print: doi: 10.2337/dc14-0990
 Bolli GB et al. Poster presented at EASD 2014: P947; Bajaj H. Oral presentation at CDA 2014: #14; Home P et al. Abstract presented at EASD 2014: 0148
 Bajaj H et al. Poster presented at CDA 2014: P112; Matsuhisa M et al. Poster presented at EASD2014: P975; Terauchi Y et al. Poster presented EASD 2014: P976

TCOYD
TRANSFORMING YOUR DIABETES

Case 3: 66 year old obese female diagnosed with type 2 diabetes 9 years ago

- Currently on maximum doses of 3 oral agents: metformin 1000 mg BID, SFU and a SGLT2 inhibitor. She was intolerant to GLP-1RAs.
- Her PCP started 10 units of insulin glargine in the morning. After 3 months on 10 units she felt it “did not work” and she stopped it.
- A1c > 8.5% for the past 2 years, eGFR 89, LFTs normal
- Current SMBG (mg/dl) below:

	Pre-Breakfast	Pre-Lunch	Pre-Dinner	Bedtime
Monday	211	---	---	185
Tuesday	247	---	174	---
Wednesday	181	---	---	196
Thursday	226	---	179	---



Which of the following is the single most likely explanation for her failure with basal insulin:

A	Poor adherence
B	Initial dose was too little
C	Inadequate titration of the glargine U-100
D	Glargine U-100 should have been given at bedtime



Simple Daily Self-Titration Option*

(much easier to follow by the patient than the 3 day titration)

Increase by 1 to 2 Units every 1 day until FPG \leq 120 mg/dL

EXAMPLE

Less than 100: decrease by 2 units

Between 100 and 150: no change

Over 150: increase by 2 units


The goal can be
individualized

* Daily titration works well with all old and new basal insulins

Dosage was not increased that week if there were any episodes of documented hypoglycemia (<72 mg/dL) during the preceding week. FPG, fasting plasma glucose.

Gerstein HC et al. Diabet Med. 2006;23:736-742.


TALKING COVERED. YOUR CARE.

Self Titration Clinic Form

Starting/Adjusting Long-Acting Basal Insulin

1. Give Basal insulin once a day at Morning
2. Starting dose: 20 units
3. Every 1 day(s), adjust your dose based on your fasting blood sugar that morning before eating or drinking:
 - a. If fasting blood sugar is over 140, then increase your dose by 2
 - b. If fasting blood sugar is under 90, then decrease your dose by 2
 - c. If fasting blood sugar is between 90 and 140, then keep the same Lantus dose

Important:

The purpose of long active basal is to provide a background amount of insulin throughout the day and at night while you sleep. It is not meant to treat high blood sugars caused by eating food, so you should not change your dose based on blood sugar numbers during the day when you are eating.


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Case 4: 55 year old obese Latino male with a 22 year history of type 2 diabetes

- CKD stage 3b (eGFR 37 ml/min)
- History of ASCVD s/p MI and CHF
- HTN, dyslipidemia, OSA , NAFLD and h/o pancreatitis
- Currently treated with low dose metformin, SFU, DPP4 inhibitor and canaflioflozin (initiated by nephrology)
- A1c 8.9%

Time	Blood glucose range	Blood glucose average
Pre-Breakfast	148 – 229 mg/dL	(175 mg/dL)
Pre- Lunch	111 – 182 mg/dL	(147 mg/dL)
Pre- Dinner	91 – 155 mg/dL	(139 mg/dL)
Bedtime	148 – 231 mg/dL	(184 mg/dL)
No reports of hypoglycemia		

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Which of the following would you suggest for this patient?

A	Initiate pioglitazone
B	Initiate basal insulin
C	Start a GLP-1 RA and stop his DPP-4 inhibitor
D	Change to a different SGLT-2 Inhibitor

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Case 4: continued

- Insulin degludec U-200 was added at night (20 units) and titrated up to 120 units over the next 10 weeks
- He was asked to test 2x/day (pre-breakfast and bedtime)
- It is important to make sure the patient is not going to bed high

Pre-Breakfast	82 – 155 mg/dL	(~122 mg/dL)
Pre- Lunch	----	----
Pre- Dinner	----	----
Bedtime	128 – 183 mg/dL	(~155 mg/dL)

- A1c dropped to 7.1%, no hypoglycemia. Gained 2 lbs in 3 months
- Oral agents can be continued unless hypoglycemia occurs during the day, in which case the sulfonylurea should be reduced or withdrawn

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Clinical Pearls: Combination Therapy with Basal Insulin

1	Start with 10 to 20 units (based on FBS, weight)
2	The key to success is frequent follow up after initiation to avoid “failure” (most patients will need 40 to 70 units/day)
3	Have the patient follow a self-titration regimen and return to clinic or follow up in some other manner (phone, fax, email, telehealth, etc.) <u>relatively soon</u>
4	You can usually limit SMBG to only once a day in the morning but check at bedtime once in awhile to make sure the pt. does not need pre dinner fast acting insulin.

Edelman SV, Henry RR. Diagnosis and management of type 2 diabetes.
12th Edition. Professional Communications, Inc., Greenwich, CT. 288 pages, 2014.

TCOYD
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Second Pitfall In Initiating/Titrating Basal Insulin (First one is too slow titration after starting)

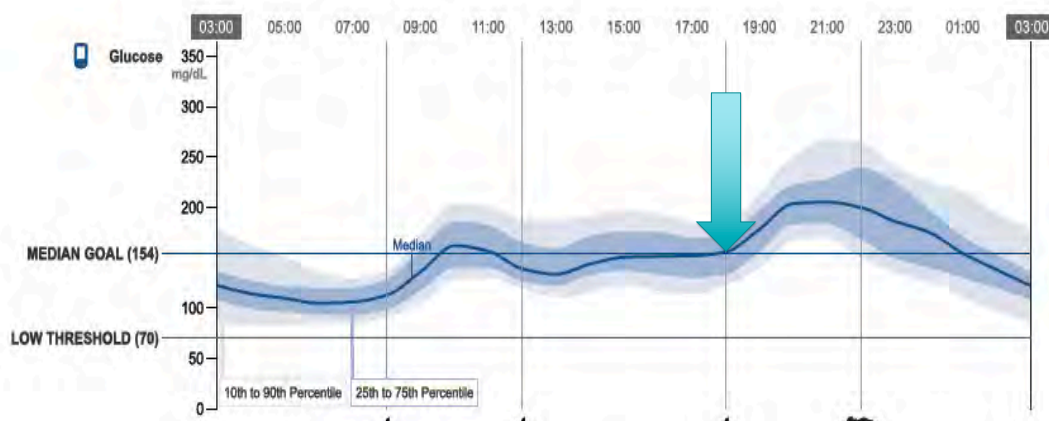
Not Paying Attention To Bedtime Glucose Value So You Avoid Overbasalinization

1. Ask the patient to do paired testing (test at bedtime and again the next morning).
2. If the bedtime BG is high, it needs to be addressed by either lifestyle modification including reduced caloric consumption and/or post dinner exercise.
3. Other options include prandial insulin or a GLP-1 RA.

Edelman SV, Henry RR. Diagnosis and management of type 2 diabetes.
12th Edition. Professional Communications, Inc., Greenwich, CT. 288 pages, 2014.

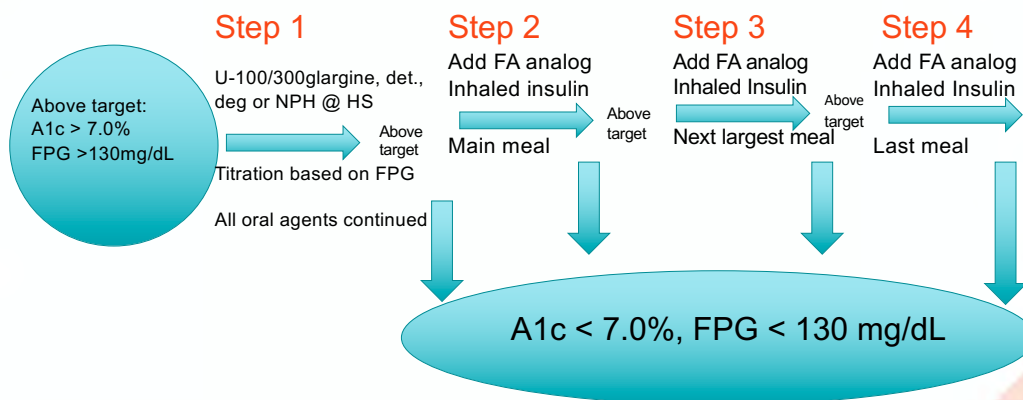
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68 Year Old Male On Oral Agents and Basal Insulin: Need For Prandial Insulin Only At Dinner



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Transitioning From Basal to Basal/Bolus Insulin Therapy in Type 2 Diabetes Mellitus



Adapted with permission from Karl DM. Curr Diab Rep. 2004;4:352-357.

TCOYD
TRANSITIONING FROM DIABETES

Initiating Insulin Therapy in Type 2 Diabetes: General Concepts

- Don't wait forever
- Address patient concerns/fears
- Consider combination therapy with oral agents
- Start with basal insulin if very poor glycemic control (A1c>9%) or in addition to a GLP-1RA
- Titration of the dose is essential (self titration can work well)
- Use a fast-acting analog as an add on to basal dose when indicated (may only be needed to be given with the largest meal)
- Self-monitoring of blood glucose (SMBG) and CGM are important tools in motivating patients and in guiding dose adjustments

Edelman SV, Henry RR. Diagnosis and management of type 2 diabetes. 12th Edition. Professional Communications, Inc., Greenwich, CT. 288 pages, 2014.

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TRANSITIONING FROM DIABETES

Summary

- GLP-1 RAs represent a tremendous advance in the treatment of type 2 because of significant glucose lowering in addition to weight loss and reducing the risk of hypoglycemia
- Combination therapy (adding basal insulin to daytime OHAs/GLP1-RAs) is safe, effective and easy to implement
- The fixed combination of basal insulin and a GLP-1 RA has clinical advantages in terms of efficacy, reduced side effects and ease of use
- The Basal Bolus approach in type 2 diabetes does not need to be four injections per day (pens, patch pumps and inhaled insulin to improve adherence)
- Adherence and persistence needs to be addressed at every visit
- Protection for ASCVD