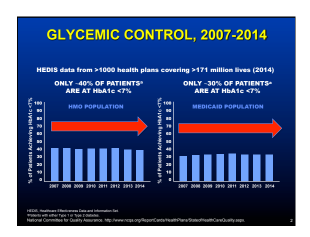
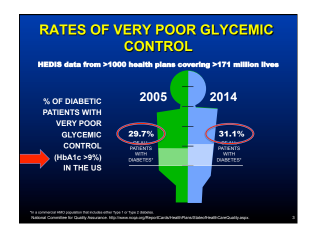
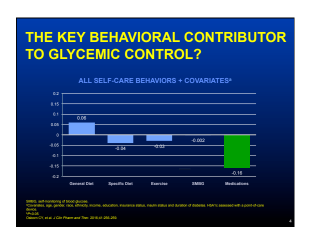
Lecture 1: 8:15 – 9:10 a.m.

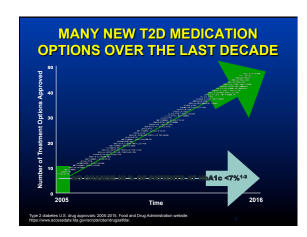
William Polonsky, PhD, CDE, Presents:

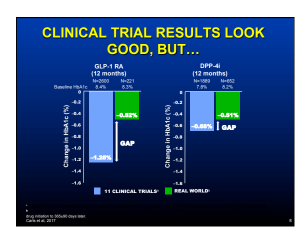
The Efficacy Mirage in Type 2 Diabetes: Why Do Clinical Trial Results Disappear in Real-World Practice?

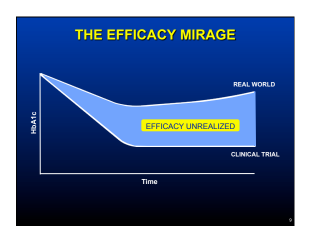


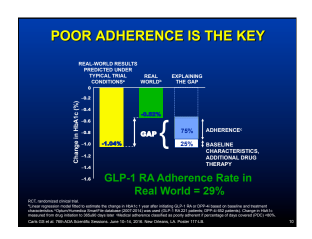


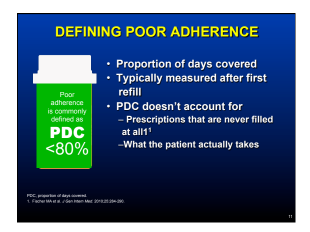


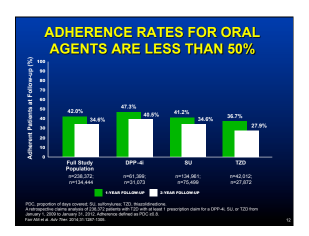


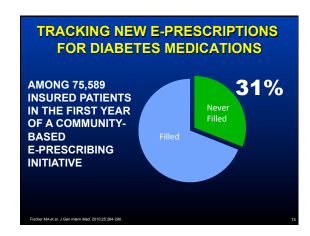


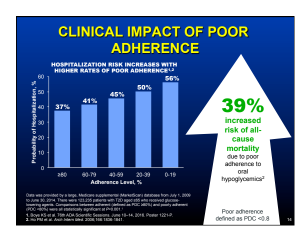












INTERVENTION STRATEGIES TO ADDRESS MEDICATION ADHERENCE

- 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: 0.29Behavioral strategies: 0.33

• Addressing habits: 0.37

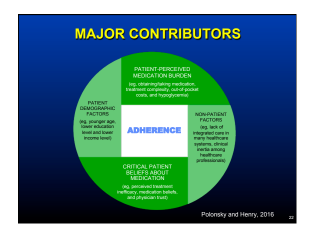
• No behavioral strategies: 0.28



Conn and Ruppar, 2017

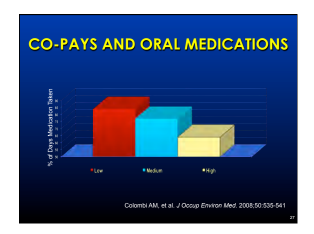
THE PRESUMED PROBLEM: FORGETFUL/DISORGANIZED

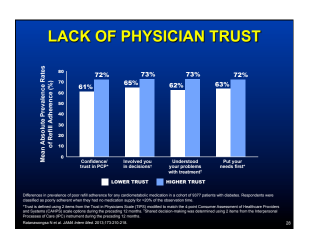


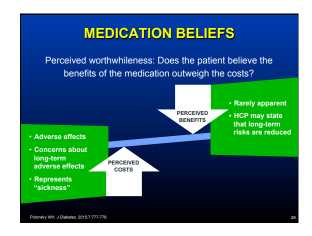












MEDI ROY	CATION BELIEFS SAM
Takes 2 oral medications for T2D and basal insulin; his last HbA1c was 6.8%	WHO IS DOING BETTER WITH HIS DIABETES? Doesn't take any medications for T2D; his last HbA1c was 9.1%
is <u>not</u> determined It is your m Even if you are r	ou are, and your risk of complications, by how much medication you take. netabolic results that matter. not taking pills or insulin, high blood ikely lead to future problems.

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? 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?"
 - o Anchoring strategies

Anchoring Medication to Daily Events *A daily event (a meal, TV show, bedtime, brushing my teeth) reminds me.* *Littenberg B, et al. BMC Fam Prac. 2006;7:1.

SO WHAT TO DO?



- 1. Ask correctly
- 2. Forgetfulness
- 3. Treatment complexity
 - o Simplify if possible
 - o Provide additional details as needed

SO WHAT TO DO?



- 1. Ask correctly
- 2. Forgetfulness
- 3. Treatment complexity
- 4. Patient-provider trust
 - · Listen, listen, listen

SO WHAT TO DO?



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

Challenging Harmful Beliefs

- Taking your medications is one of the most powerful things you can do to positively affect your health
- 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

.

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!

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www.behavioraldiabetes.org

Lecture 2: 11:30 – 12:30 p.m.

Tricia Santos Cavaiola, MD, Presents:

Which One, and When? Oral Medications for the Treatment of Type 2 Diabetes and Their Cardiovascular Affects

Case 1: Edward

- 62 year old centrally obese male (BMI 42) with with a 15-year history of type 2 diabetes also with dyslipidemia, HTN, ED, OSA, bladder cancer and CAD
- Family Hx: 3 brothers with type 2 diabetes (1 deceased/CAD)
- Notes: No home glucose monitoring data (He does not bring his meter to clinic as he "forgets" it every time)
 - Diabetes Meds: Metformin 500mg BID, glipizide 20mg BID, sitagliptin 50 mg BID, empagliflozin 10 mg QD, and glargine 100 units QHS started 6 months ago
 - Current A1c 10.5% (9.6% 1 year ago, 10.1% 2 years ago)
 - · Creatinine 1.4 mg/dl, eGFR 50
 - · LDL 92 mg/dl, Triglycerides 356 mg/dl, HDL 22 mg/dl

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

A He needs prandial insulin

B He needs a GLP-1 RA

C He is very ignorant about what to eat regarding his diabetes

D His diabetes regimen is too complicated

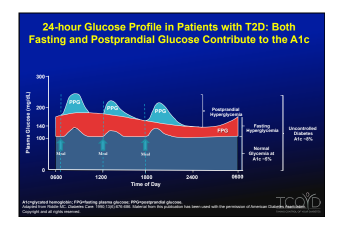
E He is most likely poorly adherent with his medications

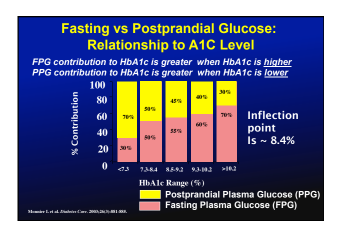


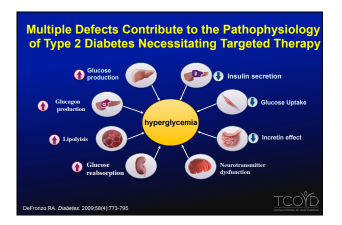
Glycemic Target Goals for Patients with Type 2 Diabetes

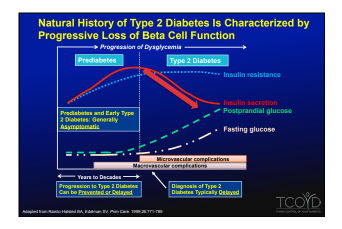
Treatment Goal	ADA	AACE
HbA _{1C} (%)	< 7	≤ 6.5
FPG (mg/dL)	80-130	<110
Preprandial glucose (mg/dL)	80–130	< 110
Postprandial glucose (mg/dL)	< 180*	< 140**

* Peak PPG; ** 2 Hr PPG American Diabetes Association. *Diabetes Care*. 2015; 38(suppl 1):S33-S40. Handelsman, Y., et al. (2015). Endocr Pract 21(0): 1-87.









9 FDA-Approved Classes of Oral Medications for Type 2 Diabetes

- > 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)
- Bile acid sequestrant (colesevelam)*
- > Dopamine receptor agonists (bromocriptine meslate)*
- Alpha glucosidase inhibitors (acarbose, miglitol)*
- * not discussed in detail in this presentation

http://www.fda.gov/drug

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
- Adding diabetes medication instead of switching is the rule rather than the exception
- Always address the ABCs (A1c and Aspirin {81mg if over 50 y/o}, BP {<140/90 mm/Hg} and Cholesterol {LDL<100mg/dl or <70 if CAD present})
- Spending time with the patient and his support person to explain why you are starting a new medication and what benefits it will have over the long term, as well as answering any concerns will improve adherence

telman SV Henry RR. Diagnosis and management of type 2 diabetes 12th Edelman SV (TCOYDIV), 3 September 2015, Get Type 2 Diabetes and type

Case 2: Collin

- > 52 year old centrally obese male
- > 1-year history of type 2 diabetes, diagnosed with dyslipidemia and HTN
- Family History: Both Parents had type 2 diabetes, HTN and CAD
- > Notes: BMI 37 (1yr ago it was 34, 2 yrs ago it was 31)
 - Diabetes therapy included only Metformin 1000 mg BID
 - Current A1c 8.5% (7.6% 6 months ago, 7.1% at diagnosis)
 - · Creatinine 1.3 mg/dl, eGFR 65
 - · LDL 112 mg/dl, Triglycerides 256 mg/dl, HDL 29 mg/dl

What class of agent would you add to Collin's current regimen (no one right or wrong answer)?

- A Sulfonylurea
- B DPP-4 inhibitor (sita-, saxa-, lina- or 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)
- F Thiazolidinedione (pioglitizone)

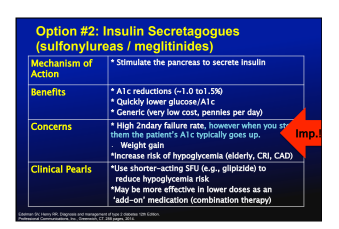


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Summary Of ADA Algorithm Step 1: start with metformin unless contraindicated Step 2: Use any other option for diabetes available in the entire universe Step 3: Use any other option for diabetes available in the entire universe except what you used in steps 1 and 2 Step 4: Use any other option for diabetes available in the entire universe except what you used in steps 1, 2 and 3 Is this helpful?

Must Individualize Therapy TO TO TO THE TRANSPORT OF THE

MOA	* Reduces hepatic glucose output
Benefits	* Significant A1c reductions (~1 to 1.5%) * Favorable to neutral effects on body weight * No hypoglycemia * Generic (low cost)
Concerns	*GI side effects (often dose-related), sustained release formulations may help * Contraindicated in chronic renal insufficiency see below * Potential for lactic acidosis (rare)
Clinical Pearls	*Start with low dose and up-titrate dose to improve GI tolerance or use long acting release formulation *eGFR <60 to ≥45 OK to use/monitor kidneys *eGFR <45 to ≥30 OK to use 50% maximum dose/ monitor kidney function every 3 months If you stop metformin, substitute with a different agent *Check B-12 levels



	Generic Name	Trade Name
Glinides	Nateglinide	Starlix
	Repaglinide	Prandin
Sulfonylureas	Glimepiride	Amaryl
	Glipizide	Glucotrol
	Glipizide (extended release)	Glucotrol XL
	Glyburide	DiaBeta, Micronase
		Glynase PressTab

Mechanism of Action	* Reduce insulin resistance
Benefits	No hypoglycemia Durable glycemic control Positive effect on lipids († HDL-C, converts small dense to large buoyant LDL-C)
Concerns	* Weight gain * Edema (precipitating CHF) Bone fractures primarily in caucasion women Risk of bladder cancer has been disproven
Clinical Pearls	* Effective in prediabetes, best used early in the natural history (balance with potential side effects) * Be cautious in combo with insulin (fluid retention)

Case 3: Jamie

42 year old African American obese male

Type 2 diabetes diagnosed at age 35

PMH: HTN, dyslipidemia

FH: T2DM, early CAD

Alc 8.3% on maximum doses of metformin and SFU
No home glucose monitoring data; "forgets" his meter and log book when he comes to clinic

Creatinine 1.4 mg/dl, eGFR 55, BMI 36

BP normally above 140/90 mmHg; on no HTN meds

BP normally above 140/90 mmHg; on no HTN meds

What therapeutic intervention would you change/initiate if you were evaluating Jamie once you have confirmed he is adherent with his medications? A Initiate basal insulin therapy B Add a DPP4 inhibitor C Add a SGLT2 inhibitor D Add a GLP1-RA E Intensify lifestyle modification and education

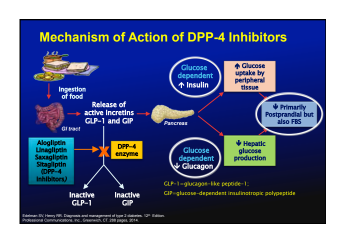
Case 3: Jamie (continued)

- ▶ Treatment History
 - A DPP-4 inhibitor was added to his regimen
 - He was sent to a CDE with his wife
- Follow up was arranged for one month instead of the usual
 3 to 4 months
- Jamie did well without weight gain or hypoglycemia
- The A1c fell to 7.4%
- His PCP eventually started an ACE inhibitor to get his BP below 140/90 mm/Hg and a statin to get his LDL <100 mg/dl
- It took almost 12 months to get his A1c, BP and lipids at goal as he was resistant to starting new medications.



Option #4:	DPP-4 Inhibitors
Mechanism of Action	* Inhibit the enzyme, DPP-4, that normally inactivates GLP-1 and other incretins within minutes
Benefits	* 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	* Dose adjustment with renal insufficiency (only for sita-,saxa-and alogliptin), not for linagliptin * Rare reports of hypersensitivity skin reactions * No association or signal for pancreatitis and pancreatic cancer (2013 FDA hearing on pancreatic cancer and incretins)
Clinical Pearls	* Efficacy of the DPP-4 inhibitors is similar * All DPP-4 inhibitors come in combination pill with metformin (Alo- is combined with Pio- and Lina- is combined with empagliflozin)

	Generic Name	Trade Name
OPP4-Inhibitors	Alogliptin	Nesina
	Linagliptin	Tradjenta
	Saxagliptin	Onglyza
	Sitagliptin	Januvia



Generic Name	Trade Name	Daily Dose Range (mg)	Recommended Frequency
Sitagliptin/metformin	Janumet	50/500, 50/1000	Twice with meals
Saxagliptin/metforminER	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
Alogliptin/metformin	Kazano	12.5/500, 12.5 mg/ 1000	Twice with meals
Ertugliflozin/sitagliptin	Steglujan	5/100, 15, 100	Once daily 🕠

		MLAR		
	Alogliptin	Linagliptin	Saxagliptin	Sitagliptin
Usage and Indications	* Use with diet and exercise to improve glycemic control in type 2 diabetes * Combination studies with SFUs, MET, pioglitazone and Insulin			
Dosage Administration	Once daily, with or without food	Once daily, with or without food	Once daily, with or without food	Once daily, with or without food
	Tablets: 25mg 12.5mg (CrCl <50), & 6.25mg (CrCl <30)	Tablets: 5mg No dose adjustment needed for renal function	Tablets: 5mg & 2.5mg (CrCl <50)	Tablets: 100mg, 50mg (CrCl <50), & 25mg (CrCl <30)
Contraindications	Hypersensitivity	Hypersensitivity (i.e., urticaria, angioedema, or bronchial hyperreactivity)	Hypersensitivity	Hypersensitivity (i.e. anaphylaxis or angioedema)
Warnings and precautions	"When used with a SFU or insulin, a lower dose of SFU or insulin may be needed to reduce the risk of hypoglycemia			
	*Post-marketing rep patients with histo		D/C if suspect pancrea	titis; Use with caution

Case 4: Susan

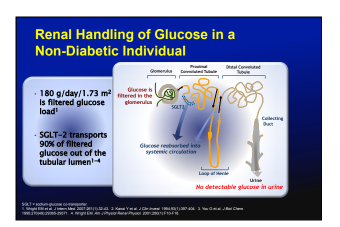
- > 58 year old obese female
- > Type 2 diabetes diagnosed 10 years ago
- > A1 c 8.7%, (one year ago it was 8.2%) and adamantly refused any injectable agent
- > On max. doses of metformin and a DPP4-inhibitor
- > Family History: Type 2 diabetes and obesity (both parents)
- Notes
 - <u>Very</u> fearful of injections and gaining weight
 - Normal renal function, BMI 31kg/m²
 - HGM shows FBS (137-221 mg/dl), and a few post dinner values (187 to 265mg/dl)

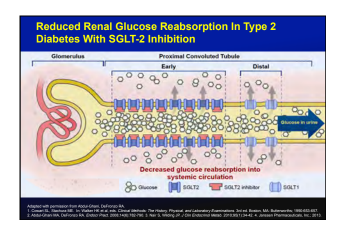
How would you treat Susan to lower her A1c?

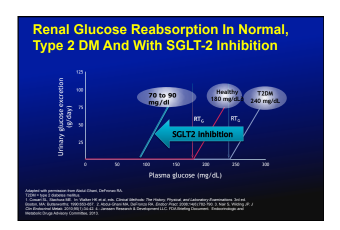
Α	Add a SFU
В	Add a TZD
С	Start a SGLT-2 inhibitor (cana-, dapa-,empa- ertugliflozin)
D	Try to convince her to add a GLP-1 receptor agonist (exenatide, liraglutide, dulaglutide, semaglutide)
E	Try to convince her to add a basal insulin at bedtime

Optio	n #5: SGLT-2 Inhibitors
Mechanism of Action	* Reduces renal glucose reabsorption and increases urinary glucose excretion
Benefits	* 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	* Genital mycotic infections. In women (6 to 12% higher than comparator) and in uncircumdised 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 IDL Cholesterol (TGS goes down and HDL goes up) * Assess renal function (discussed later) * New label warnings: DKA (discussed later)/bone fractures/risk of amputation DISCUSSED LATER WITH CVOT DATA
Clinical Pearls	1st oral medication that leads to statistically significant weight loss Empa-and canapilflozin showed positive CVD outcome trials(discussed later) Can be added to any other oral agent or injectable Tell 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 Monostat) Physician day reference (RRM ed.) (2014 Montale, NJ. Physician Desk Reference.

Generic and Trade Names (dose range) Generic Name Trade Name SGLT-2 Inhibitor Dapagliflozin Farxiga Empagliflozin Jardiance Entugliflozin Jardiance Steglatro Canagliflozin: Suggested starting dose: 100 mg daily before first meal of day (eGFR >45 mL/min) increase to 300 mg daily if tolerating 100 mg daily and eGFR > 60 mL/min Dapagliflozin: Starting dose: 5 mg daily in morning with or without food (eGFR for both doses > 60) increase to 10 mg daily if tolerating and need additional glycemic control Empagliflozin: Starting dose: 5 mg daily in morning with or without food (eGFR>45) increase to 25 mg daily in morning with or without food (eGFR for both doses > 60) increase to 25 mg daily in morning with or without food (eGFR for both doses > 60) increase to 25 mg daily in morning with or without food (eGFR for both doses > 60) increase to 15 mg daily in morning with or without food (eGFR for both doses > 60) increase to 15 mg daily in morning with or without food (eGFR for both doses > 60) increase to 15 mg daily if tolerating and need additional glycemic control







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
2. Many but not all of the reports for DKA were in patients with LADA
3. Be especially cautious in insulin using patients (since glucagon levels go up with SGLT2s and by lowering the insulin too much an imbalance of glucagon to insulin may occur, leading to DKA)
4. Be especially cautious in women with a history of UTIs, pyelonephritis and/or genital mycotic infections

ooks M. SGLT2 Inh Diabetes Drugs May Cause Ketoacidosis: FDA. Retrieved from http rondu N, et al. Diabetes Care September 2015 38:1680-1686; 2015

Case 4: Susan continued

Low dose SGLT-2 inhibitor was added to her regimen and then titrated to the maximum dose after one month



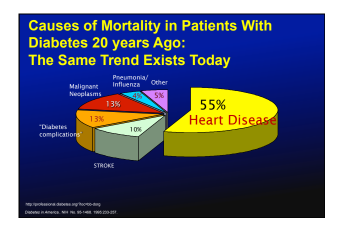
- > A1c dropped to 7.5% (baseline 8.7%) and she lost 15 lbs
- She was more motivated to improve her lifestyle habits and her A1c came down to 7.2% over the next 4 months
- She experienced a yeast infection which was easily treated with a topical antifungal and she did not want to stop the SGLT2 inhibitor
- She also said she had increased urination in the mornings for the first few weeks but that stopped
- LDL went from 100 to 108 mg/dL (8% rise) and her TGs of dropped by 25%

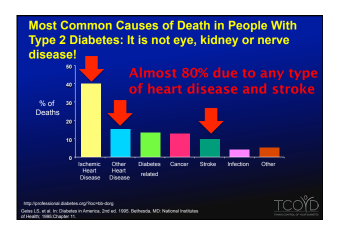
Which of the following statement is <u>true</u> regarding SGLT-2 inhibitors?

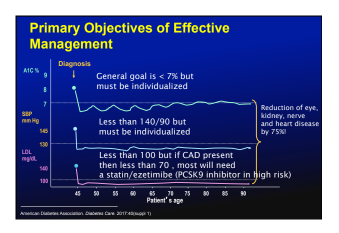
Α	They are contraindicated with loop diuretics and a history of DKA
В	They should not be used in women or men with a history of UTIs
С	They can be used safely with pioglitazone and GLP-1 RAs
D	They are approved for both type 1 and type 2 diabetes
Е	Men who are not circumcised should not use them

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

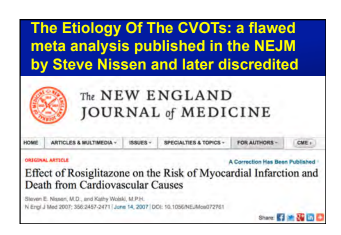
A	Nephropathy including end stage renal disease requiring dialysis or transplantation
В	Complications from peripheral and autonomic neuropathy
С	Stroke or cardiovascular disease
D	Complications from obesity
E	Peripheral arterial disease

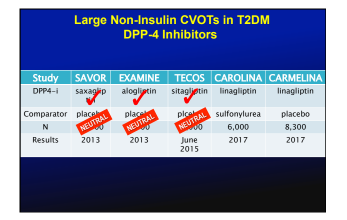






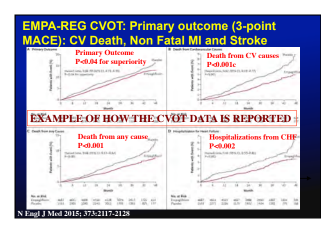
Study	Micro	vasc	cv	D	Mort	ality
UKPDS 33 (7.0 vs. 7.9%)	•	•	(=)	•	()	•
DCCT / EDIC* (7.2 vs. 9.1%)	•	•	\Leftrightarrow	•	(•
ACCORD (6.4% vs. 7.5%)	1	þ	Ġ	→	4	
ADVANCE (6.3% vs. 7.0%)	7	þ	⇔	\Leftrightarrow	①	\Leftrightarrow
VADT (6.9% vs. 8.4%)	1	þ	\Leftrightarrow	•	(\leftarrow





Large Non-Insulin CVOTs in T2DM GLP-1 Receptor Agonists						
Study	LEADER	ELIXA	SUSTAIN 6	EXSCEL	REWIND	
GLP1-RA	liraglutide	lixisepatide	semaglutide	exenatide LR	dulaglutide	
Comparator	planVE POSITIVE	NEUTRAL	POSITIVE	NEUTRAL NEUTRAL	placebo	
N	20,500	4,000	0,000	3,400	8,300	
Results	2016	2015	2016	2018	2019	
Courtey of Silvio Inzucchi MO, Yale University						

Large Non-Insulin CVOTs in T2DM SGLT-2 Inhibitors					
Study	EMPA-REG	CANVAS	DECLARE	NCT01986 881	
SGLT-2-i	empaglifozin	canaglifozin	dapagliflozin	ertugliflozin	
Comparator	placebo	placeb POSITIVE	placebo	placebo	
N	P0500	P.300	22,200	3900	
Results	Sept 2015	2017	2019	2020	
Courtesy of Silvio Inzucchi M	D, Yale University				



Real-World CV Study on SGLT-2 Inhibitors (CVD reduction may be a class effect?)

CVD-REAL study shows SGLT-2 inhibitors significantly reduced hospitalizations for heart failure and death versus other type-2 diabetes medicines

- CVD-REAL study assessed data from 300,000+ patients
 - > (87% did not have history of CV disease)
- Reduced rate of hospitalization for heart failure by 39% and allcause mortality by 51%

New FDA Indication for Diabetes Medications

- Diabetes medications FDA approved for CV risk reduction
- Empagliflozin (based on EMPA-REG data)
- Reduction in risk of CV death in patients with type 2 diabetes and established CV disease
- 2. Liraglutide (based on LEADER data)
 - Reduction in risk of major CVevents in patients with type 2 diabetes and established CV disease
 - Canagliflozin and semaglutide under review



New FDA Warning for Diabetes Medications

- FDA warning for lower limb amputation
- > 2 fold increase in amputation in the CANVAS CVOT
- Relative risk 0.63 (canagliflozin) vs 0.34 (placebo) amputations per 100 patient years
- No increased risk of amputation in the phase 3 clinical trial program (~10,000 patients)



Not All	CVOTS	: Ara (Proato	d Faula
NOL AII	01013		JI Gale	u Lyua



- > Differences in study design: powered for safety or
- Patient characteristics: age, weight, co-morbid complications, presence of CAD
- > Comparators may be different
- > Weigh gain and hypoglycemia differences
- > Time to first event
- > Regional differences
- Outcomes differ: overall mortality, non fatal and fatal MI, stroke, etc.
- > Adherence may effect results

Das, Journal of Diabetes Research & Clinical Metabolism 2015, sw.hoajonline.com/journals/pdf/2050-0866-4-3.pdf Courtesy of Mikail Kosiborodi MD, Saint Lukes

Key Principles of Management of Type 2 Diabetes

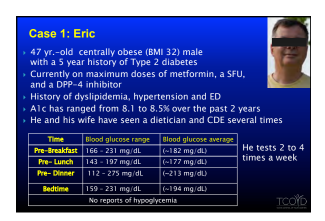
- Glycemic targets & glucose-lowering therapies should be individualized
- > Diet, exercise and education are the foundations of therapy
- > Unless contraindicated, metformin is optimal 1st line drug
- After metformin, combination therapy with 1-3 other oral and/or injectable agents; minimize side effects
- > Many patients over time will require insulin therapy alone or in combination with other agents to maintain glycemic control
- CAD is the most common cause of death and prevention strategies need to be emphasized

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

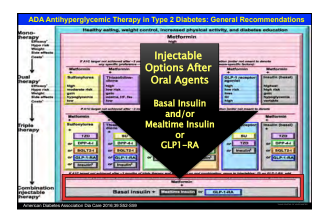
Lecture 3: 1:15 – 2:15 p.m.

Carol Wysham, MD, Presents:

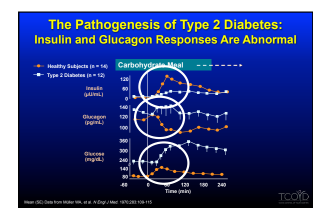
Clinical Applications of Injectable Agents: GLP-1 Receptor Agonists, Basal Insulin and More Intensive Regimens

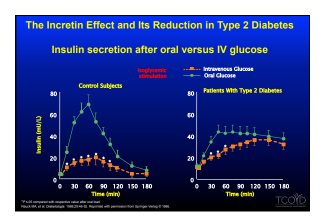


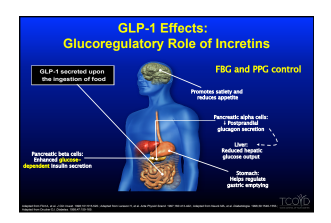
Vhich of the following would you ecommend for Eric if he were your patie					
А	Initiate basal insulin				
В	Initiate a GLP-1 Receptor Agonist (RA)				
С	Initiate a basal bolus insulin regimen				
D	Initiate a fixed combination of a basal insulin and a GLP-RA				

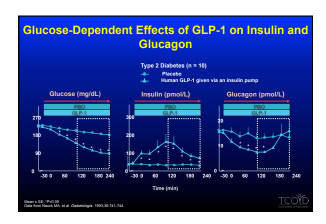


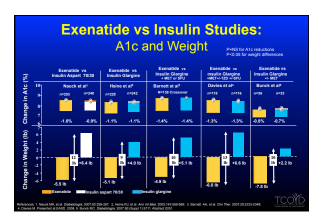
	rs GLP-1 RA
Insulin: Injectable once or twice a day	GLP-1 RA: Injectable once a day or once weekly
Need to titrate dose targeting the FBG	Reduced need to titrate dose to BG, but increase dose slowly to avoid GI side effects
Need to institute home glucose monitoring (SMBG)	"No" need for SMBG
Important 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 dabetes. 12° Editor. Professional Communications. Inc., Greenarch, CT. 255 pages, 2014.	TCOY(







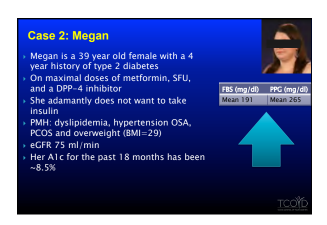




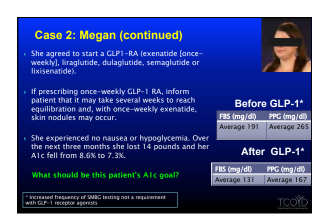
GLP-1 Receptor Agonists				
Mechanism of Action	* Mimic the effects of human GLP-1			
Benefits	Significant A1c reductions (1.0 to 2.0%) Shorter acting GLP-1 RAs have greater effects on PPG Statistically significant weight loss No hypoglycemia (due to GLP-1 RA directly) Once daily and once weekly formulations			
Concerns	Cliside 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	Ideal choice in obese patients with poor control, especially those on large doses of insulin "No" need to initiate or increase glucose testing One of the most powerful agents for type 2 diabetes			
man SV, Henry RR. Diagnosis and man Greenwich, CT. 288 pages, 2011.	agement of type 2 diabetes. Eleventh Edition. Professional Communications,			

Generic and Trade Names: GLP-1 RAs					
	Generic Name	Trade Name			
GLP-1 Receptor	Exenatide				
Agonists	Twice-daily	Byetta			
	Once-weekly	Bydureon			
	Liraglutide				
	Once-daily	Victoza			
	Dulaglutide '				
	Once-weekly	Trulicity			
	Lixisenatide				
	Once-daily	Adlyxin			
	Semaglutide				
	Once weekly	Ozempic			
Basal Insulin/					
GLP-1Receptor	Glargine/lixisenatide	Soliqua			
Agonist Fixed	Degludec/liraglutide	Xultophy			
Combination	both once-daily	TCOYL			

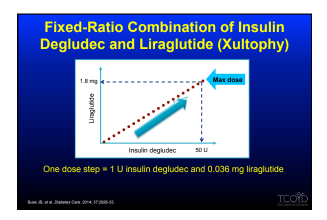
ITCA 650—Medi Type 2	cal Device	
TECHNOLOGY	MEDICATION: EXENATIDE	
proceedure	• Previously– approved GLP–1 therapeutic which demonstrates: –glycemic control	
• Small micropump -maintains stability at temps ≈37°C -maintains stability for ≥ 12 months	-weight loss -safety	
Not yet approved by the FDA		

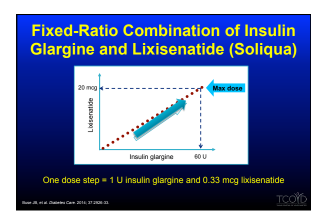


Wha	t would you recommend now for Megan?
Α	Start a SGLT2 inhibitor
В	Try to convince her to start basal insulin
С	Start a GLP-1 RA and discontinue the DPP-4 inhibitor
D	Start a fixed combination of a basal insulin and a GLP-RA
	TCOYD Sent act or or or or or

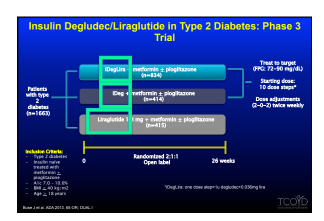


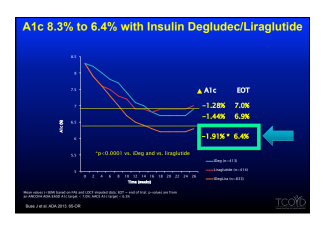


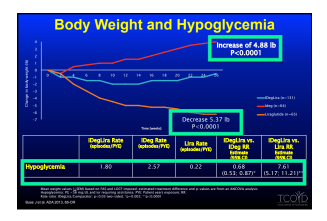




Insulin Degludec/Liraglutide v	s. Insulin Glargine/Lixisenatide
Pen dose steps (units): insulin degludec + liraglutide (Xultophy)	Pen dose steps (units): insulin glargine + lixisenatide (Soliqua)
10 dose steps=10 units insulin degludec +0.36 mgs of liraglutide 50 dose steps=50 units insulin degludec +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 liragiutide	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



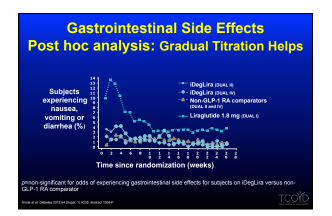


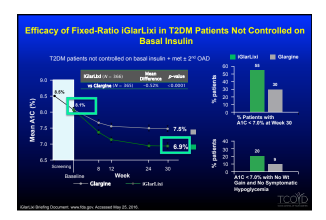


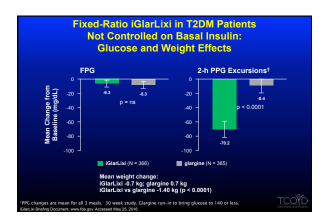
DUAL VII – Open-label trial comparing iDeg/Lira to basal-bolus insulin therapy (glargine + aspart) for 26 weeks

iDeg/lira was non-inferior to basal-bolus for glycemic control
Mean A1c reduction from 8.2 to 6.7% in both groups

iDeg/lira was associated with:
Lower insulin doses (40.1 units for iDeg/Lira group compared to 84.6 units in basal-bolus)
Less hypoglycemia: 89% less severe or symptomatic confirmed hypoglycemia compared to basal-bolus
Mean weight loss (0.9kg) versus weight gain (2.6kg) with basal-bolus

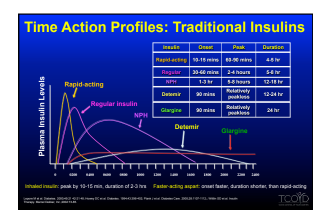




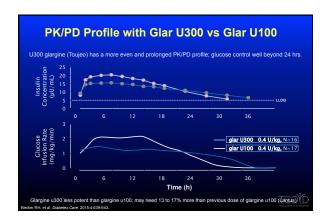


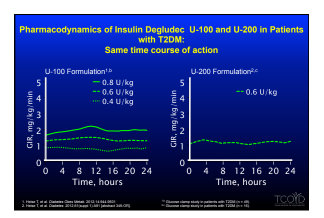
Summary: Benefits for Combining GLP-1 Receptor Agonists and Basal Insulin Analogs Combined glycemic effects of GLP-1RA and basal insulin provides greater glycemic efficacy than either of its component parts. Dose related adverse effects of each component (nausea and weight gain) are minimized. No increased 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.

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) inhaled insulin	Humulin R, Novolin R Humulin R U-500 NovoLog Fiasp Apidra Humalog Afrezza	
Basal Insulin	intermediate-acting: NPH long-acting: detemir glargine (U-100) glargine (U-300) degludec (U-100/200)	Humulin N Novolin NPH Levemir Lantus Toujeo Tresiba	
	follow-on biologic glargine (U-300)	Basaglar TCC	



Shortcomings of Traditional Basal Insulins Include: Hypoglycemia resulting in: Insulin under-dosing Insufficient glycemic control Weight gain Inconsistent insulin action...leading to inconsistent blood glucose levels Not enough flexibility with timing of injections Insufficient duration of action...therefore, requiring a minimum of 1 and, sometimes, 2 injections/day Large volume injections required for some patients

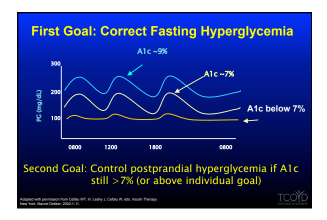




Currently on ma	emale diagnosed with ty eximum doses of 3 oral a BID and linagliptin 5 m	agents: meti			
months ago did	rt insulin for years (afrai try 10 units of glargine felt it "did not work" ar	in the mon	ning. After		
Current SMBG (
		Pre- Lunch	Pre- Dinner	Bedtime	7
	mg/dl) below:			Bedtime 185	1
Current SMBG (r	ng/dl) below: Pre-Breakfast				
Current SMBG (r	ng/dl) below: Pre-Breakfast 211		Dinner		

	taran da antara da a	
A	Patient fear of Insulin	
В	Health care provider inertia	
С	Inadequate titration of the glargine U-100	
D	Glargine U-100 should have been given at bedtime	
	y explain: A B C	A Patient fear of Insulin B Health care provider inertia C Inadequate titration of the glargine U-100 Glargine U-100 should have been given at

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. Titrating the dose is essential (self titration can work well). Use a fast-acting analog at meal time when indicated. (may only needed to be given with the largest meal) Self-monitoring of blood glucose (SMBG) is an important tool in motivating patients and in guiding dose adjustments.



Adding Basal Insulin to Oral Agents An Effective Strategy to Initiate Insulin Therapy

- Only 1 injection per day is typically required
- ▶ No need for mixing different types of insulin
- Convenience (usually given at night or first thing in the morning)
- > Slow, safe, and simple titration
- Low dosage compared to a full insulin regimen
- Limited weight gain especially compared to insulin only regimens
- Effective improvement in glycemic control by suppressing hepatic glucose production

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

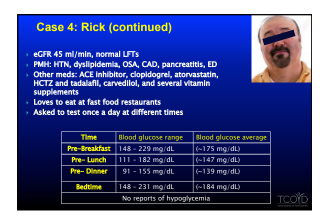
TCOY

Case 4: Rick

- 61 yr.-old overweight (BMI 30, 220lbs) male
- Type 2 diabetes diagnosed 9 years ago
- → History of CAD s/p MI 2 years ago
- Treated for 2 years with diet and exercise alone even though his A1c was above 9.5% ("did not want to take medications")
- Eventually started on metformin, sequentially followed by a sulfonylurea and a DPP-4 inhibitor (100mg sitagliptin), and his A1c fell from 9.9% to 7.9%
- It took two years (6 clinic visits) to initiate these 3 meds and get his A1c down

What should be this patient's A1c goal?

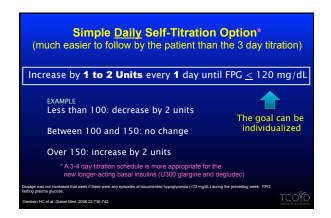


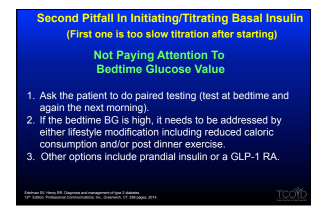


	f the following would you suggest Rick if he were your patient?	
А	Work on lifestyle and no medication addition	
В	Initiate basal insulin	
С	Start a GLP-1 RA and stop his DPP-4 inhibitor	
D	Start a SGLT-2 Inhibitor	
D	Start a SGL1-2 Inhibitor	

Case 4: Rick (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- Unnch -- Pre- Dinner -- Bedtime 128-183 mg/dL (-155 mg/dL) Alc 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







		Pearls: tion 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 12 th Edition.	Henry RR. Diagnosis i Professional Commun	Sure the pt. does not need pre dinner fast acting insulin. and management flyp 2 database. TCC

Case 5: Angela • 65 year old female on triple oral agent therapy (SFU ,met, DPP-4 inhibitor) was started on 10 units of insulin glargine (U-100) qAM in July 2011 • FPG ~ 220 mg/dL, A1c 8.5 %, wt = 176 lb • Insulin glargine (U-100) was titrated to 45u qAM from July 2011 to November 2011 • FPG 78-132 mg/dL, A1c = 7.4%, wt = 181 lbs, eEGR 62 • Patient was asked to test more frequently than usual for 3 to 4 days before meals and bedtime (pattern testing) | July 2011 | November 2011 | A1c 60 | 8.5 | 7.4 | | FPG (mg/dL) | -220 | 78-132

Case 4: Angela (cont) 65 year old woman on glargine (U-100) and 3 oral agents: SMBG data Bedtime ---- = did not test TCOYE

Which of the following would you recommend for Angela at this point?

	Pre-Breakfast	Pre- Lunch	Pre- Dinner	Bedtime
Monday	101	124		185
Tuesday	132	146	109	214
Wednesday	98	111	89	229
Thursday	78		121	201

Α	Increase basal insulin
В	Switch to premix insulin at dinner
С	Intensify regimen by adding rapid acting insulin at dinner
D	SGLT-2 inhibitor

Case	5: An	gela ((cont)

- Dinnertime bolus added:
 - Patient was started on 5 units of rapid-acting insulin analog at dinnertime and titrated up to 15 units over a few weeks based on the bedtime blood glucose levels (initial dose can be ~10% of the total basal dose). Options include lispro, aspart, glulisine, and inhaled Insulin
 - The basal insulin dose (glargine [U-100] 45 units) was titrated downward to 40 units on initiation of rapid-acting insulin based on the patient's near normal fasting blood glucose levels in order to avoid nocturnal/fasting hypoglycemia

Nathan DM et al. Diabetologia 2008;51:8-11. Raccah D et al. Diabetes Metab Res Rev. 2007;23:257-26TCOY

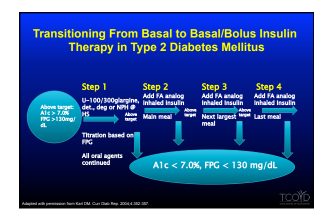
Case 5: Angela (cont)

 SMBG values on glargine (U-100) 40 units at bedtime; lispro 15 units pre-dinner

	Pre- Breakfast	Pre- Lunch	Pre- Dinner	Bedtime
Wednesday	88			136
Thursday	131		143	188
Friday	98	122		121
Saturday	112		134	169

- A1c fell from 7.4% to 6.8%.
- Angela experienced occasional mild hypoglycemia.

TCOYD





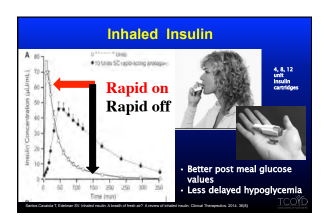


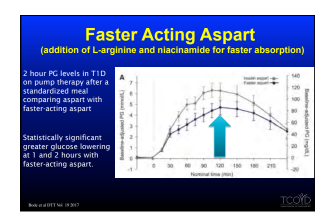






Shortcomings of Existing Bolus Insulins Include Not rapid enough: Leading to mismatch between peak postprandial glucose and peak insulin action Need to take up to ½ hour before eating Lasts too long...leading to delayed hypoglycemia Inconsistent action leading to inconsistent blood glucose levels





Case 1: Eric (Follow up!)

- 47 yr.-old centrally obese (BMI 32) male with a 5 year history of Type 2 diabetes
- Currently on maximum doses of metformin, a SFU, and a DPP4 inhibitor
- History of dyslipidemia, hypertension and ED
- Alc has ranged from 8.1 to 8.5% over the past 2 years
- He and his wife have seen a dietician and CDE several times

Time	Blood glucose range	Blood glucose average	
Pre-Breakfast	166 - 231 mg/dL	(~182 mg/dL)	
Pre- Lunch	143 - 197 mg/dL	(~177 mg/dL)	
Pre- Dinner	112 - 275 mg/dL	(~213 mg/dL)	
Bedtime 159 - 231 mg/dL (~194 mg/dL)		(~194 mg/dL)	
	No reports of hypoglycemia		

He tests 2 to 4 times a week

Which of the following would you recommend for Eric if he were your patient?

А	Initiate basal insulin
В	Initiate a GLP-1 RA; stop DPP-4 inhibitor
С	Initiate a basal bolus insulin regimen
D	Initiate a fixed combination of a basal insulin and a GLP-RA; stop DPP-4 inhibitor

TCOÝÍ

Summary

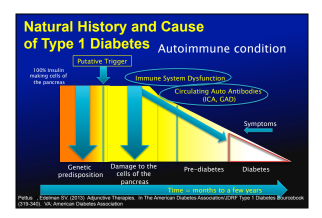
- GLP-1 agonists represent a tremendous advance in the treatment of type 2 because of glucose lowering in addition to weight loss and reducing the risk of hypoglycemia
- Combination therapy (adding basal insulin to daytime OHAs) is safe, effective and easy to implement
- The fixed combination of basal 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
- Patient and clinical inertia are serious problems
- Adherence and persistence needs to be addressed at every visit $_{\text{TCO}}$

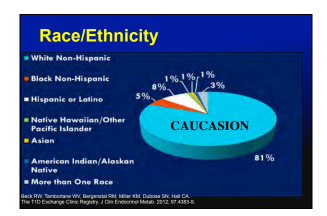
Lecture 4: 2:15 – 3:15 p.m.

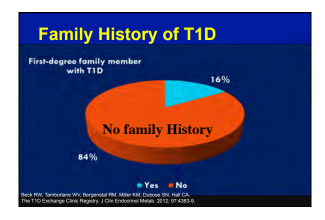
Steven V. Edelman, MD, Presents:

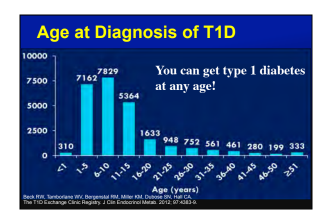
Cutting-Edge Strategies for the Treatment of People with Type 1 Diabetes

It is all about "Time In Range": Keeping the glucose levels between 70 and 180 mg/dl 1. 1st priority is getting a CGM and educate your patients to respond to the trend arrows. 2. Bolus calculations are more than just the carbohydrates and static glucose readings 3. In addition to getting the A1c below 7%, try to reduce the daily glucose fluctuations in your patients (hyper- and hypoglycemia) 4. The insulin regimen should mimic what happens in a non-diabetic state Edeman SV Taking control of your diabetes: a patient oriented book on diabetes. Fifth Edition Froblessional Communications inc. General Control of Your diabetes. Control of Your diabetes. Control of Your diabetes a patient oriented book on diabetes. Fifth Edition Froblessional Communications inc. General Control of Your diabetes. Fifth Edition

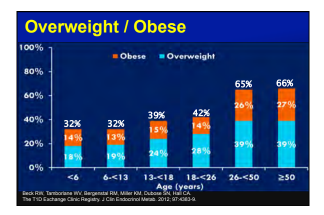










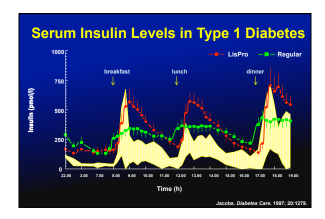


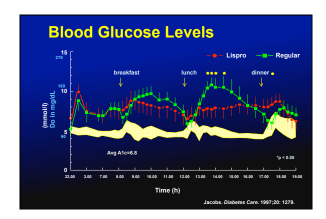
Case 1: Phil

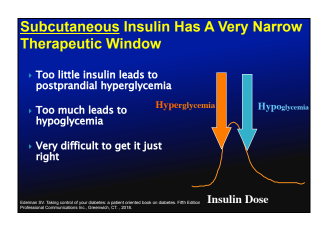
- 46 year old male with the diagnosis of type 1 diabetes at age 6 (Classic presentation of DKA)
- He has been on an insulin pump for many years
- Over the last 8 years he has developed central obesity and his insulin requirements doubled
- He also developed high blood pressure and dyslipidemia (triglycerides went up and his HDL when down).
- Family history is that his father and both paternal uncles have type 2 diabetes.

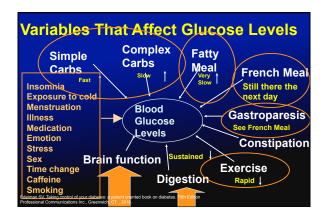
What is the most likely explanation of why Phil's insulin requirements doubled later in life?

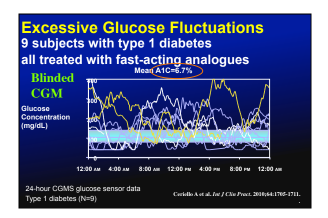
A He developed central obesity	
B He has both type 1 and type 2 diabetes	
C His A1c kept rising	
The insulin he was receiving by mail was denatured and lost potency	

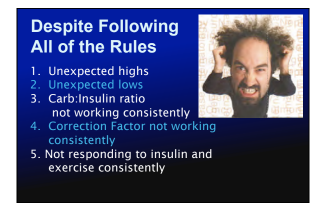


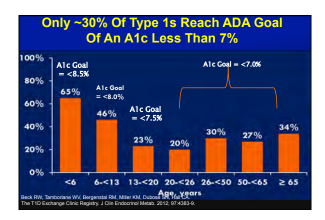


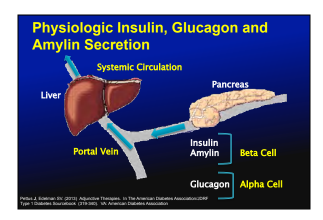


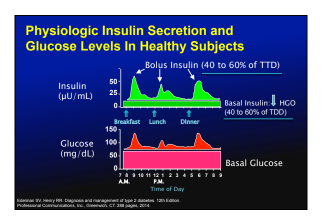












Case 2: Tom

- > 36 year old male with type 1 diabetes for 20 years
- He is on a basal bolus regimen (20 units of insulin glargine at bedtime and 16 to 22 units of fast acting meal and correction boluses throughout the day.
- His correction factor is 1:40 (goal of 125) and his insulin to carbohydrate ratio is 1:12
- Alc is 7.1%,, however his glucose values bounce from high to low and he is very frustrated.
- ▶ He tests his glucose value 6 to 8 times a day
- He tried to be as consistent as possible with his diet and exercise
- His wife is very supportive and he is motivated to do well

What therapeutic intervention do you think is the most important to help Tom with his glucose control?

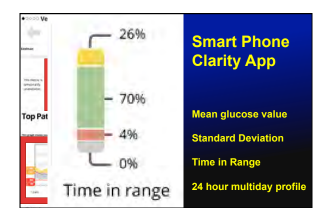
- A Put Tom on an insulin pump
- Put Tom on a continuous glucose monitor
- C Split his dose of insulin glargine so that he takes 10 units BID
- D Send Tom to a diabetes education class

Severe Hypoglycemia and A1C: DCCT¹⁵ (1993), JDRF² (2008), and STAR 3¹⁶ (2010) Studies DCCT (intensive therapy): 62 per 100 pt.yrs, A1C(6.5 yr): 9.0% → 7.2% JDRF CGM: 20.0 per 100 pt.yrs, A1C(6.5 yr): 9.0% → 7.1% STAR 3 MDI (all ages): 13.5 per 100 pt.yrs, A1C (1 yr): 8.3% → 3.1% STAR 3 SAP (all ages): 13.3 per 100 pt.yrs, A1C (1 yr): 8.3% → 3.1%

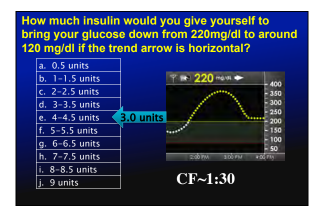


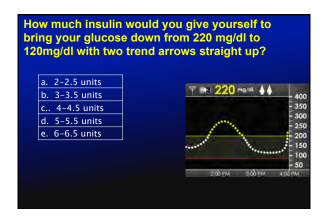


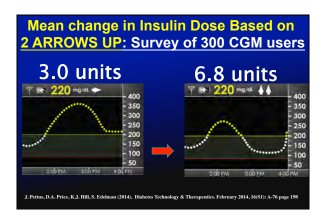


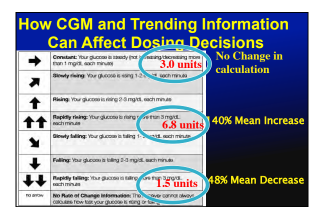


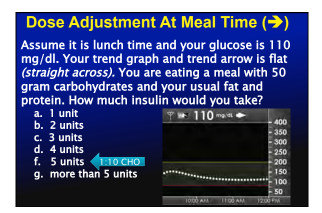
How Do Patients (n=300) with Type 1 Translate CGM Data Into Diabetes Management Decisions Mean age was 46 ± 14 years old Duration of diabetes: 22±14 years Mean A1C (self reported) was 6.9% ± 0.8% Minimal hypoglycemia Insulin delivery: 75% used CSII 25% used multiple daily injections J. Pettes, D.A. Price, K.J. Hill, S. Edelman (2014), Diabetes Technology & Therapeutics, February 2014, IG(S1): A-76 page 198

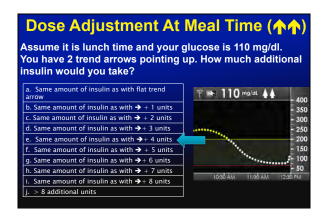


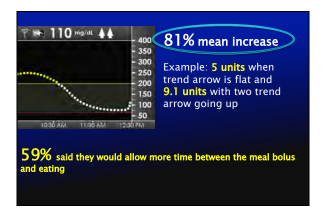


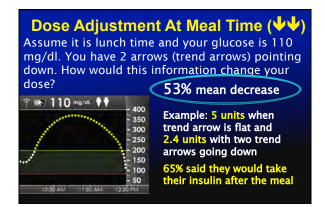


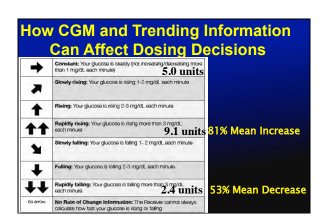


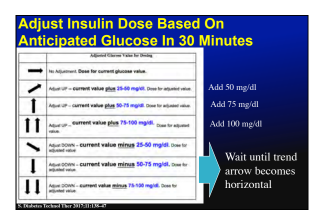


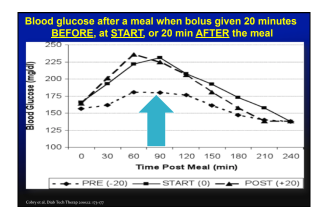




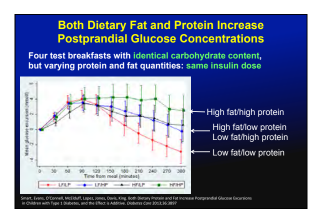




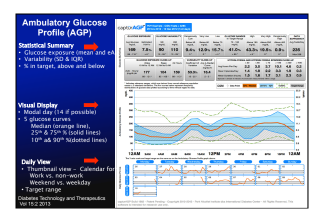


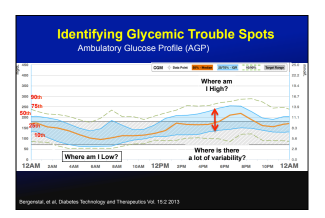


How much fast acting insulin would you recommend to a patient eating a meal with 60 grams of carbohydrates (Insulin to Carb ratio is 1 to 10), an 8 oz Filet and a salad with olives and avocados?		
Α	3 units	
В	6 units	
С	12 units	
D	More than 6 units	







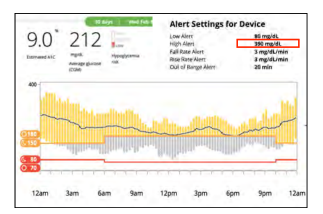


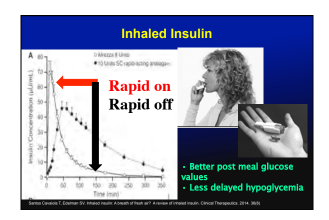


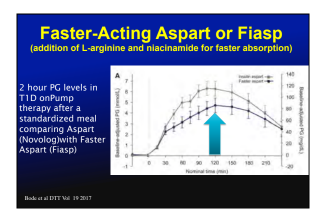
Concerns To Address With CGM

- ▶ Alarm fatigue
- High and low alert settings (80 to 180mg/dl)
- High and low snooze alarms (also known as repeat high and low alerts)
- Take advantage of the Share system
- > Stacking (taking multiple boluses too close in time

Edelman SV, Hirsch IB, Pettus JH. Practical Management of Type 1 Diabetes. Second Edition. Professional Communications, Inc., Greenwich, CT. 175-209, 201-

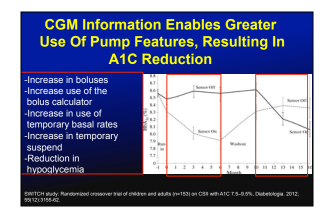


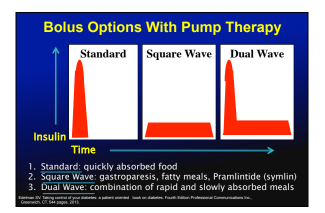


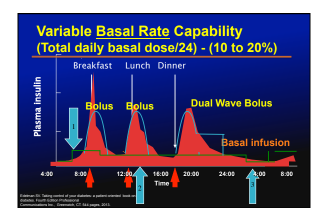




Insulin Pumps: Advantages Improved glycemic control More precise, physiologic insulin delivery Greater ability to handle dawn phenomenon, stress and other conditions that alter insulin requirements "Smart features" help to estimate insulin doses and reduce errors, i.e. stacking insulin In some situations (but not all) freedom and flexibility in lifestyle Eliminate multiple daily injections (1 stick every 3 days) very easy to respond to CGM results Reduce restrictions on eating, exercise and sleeping patterns; could have the same benefits with MDI Greater flexibility with sports, travel, work schedule and other activities (not with water sports)

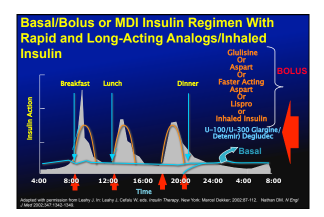


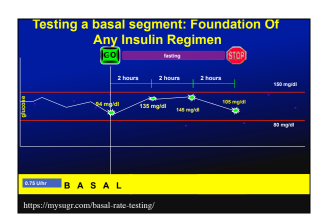




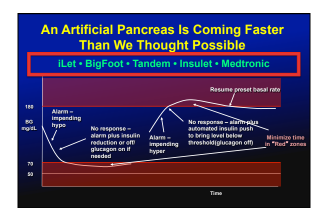


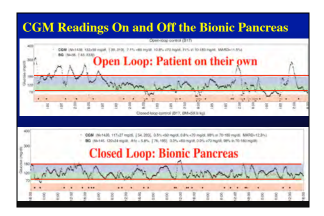
Disadvantages of Pump Therapy A disruption in short acting insulin delivery due to a dislodged catheter, blockage, or an empty reservoir can result in a fairly rapid rise in glucose concentration Severe hyperglycemia Ketoacidosis Cost of the insulin pumps Pump costs approximately \$3,500 to \$5,000 (some pumps offer pay as you go options) Monthly cost of \$30 to \$40 due to batteries, infusion lines, syringes, and adhesive tape Minor skin irritation or infections at the insulin pump catheter insertion site Very occasional abscess











Summary/Conclusions

- CGM will <u>bridge the gap</u> until a real cure for type 1 is discovered
- Numerous variables can and will affect the blood glucose levels on a daily basis
- Every day is <u>different</u> for a person with type 1 diabetes
- A glucose value at one point in time has limited value when dosing insulin
- Trend arrows can help PWD make better daily diabetes decisions