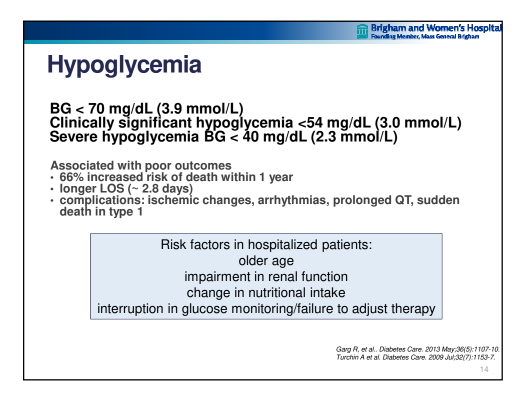
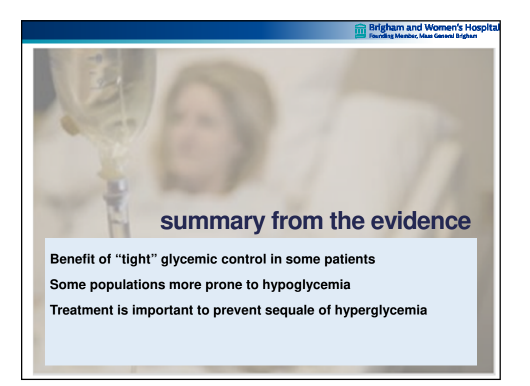
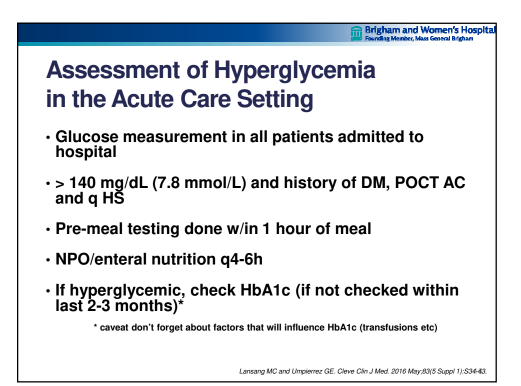


Iorauty	'		otal no. patient	ts	Favours IIT	Favours control	
lortality	Study	IIT	Control	Risk ratio (95% CI)	←	\rightarrow	_
	Mixed ICU						
	Yu et al. ³⁹	4/28	4/27	0.96 (0.27-3.47)			
	Henderson et al. ³¹	5/32	7/35	0.78 (0.28-2.22)			
	Mitchell et al. ³⁵	9/35	3/35	3.00 (0.89–10.16)	-	•••••	
	Wang et al. ³⁸	7/58	26/58	0.27 (0.13-0.57)			
	Azevedo et al. ²²	38/168	42/169	0.91 (0.62-1.34)		_	
	McMullin et al. ³⁴	6/11	4/9	1.23 (0.49-3.04)			
	Devos et al. ¹³	107/550	89/551	1.20 (0.93-1.55)			
	Brunkhorst et al. ¹¹	98/247	102/288	1.12 (0.90-1.39)	-	-	
	lapichino et al. ³²	15/45	12/45	1.25 (0.66-2.36)	_	-	
	He et al. ³⁰	16/58	29/64	0.61 (0.37-1.00)			
	Zhang et al.40	4/168	6/170	0.67 (0.19-2.35)			
	De La Rosa Gdel et al.12	102/254	96/250	1.05 (0.84-1.30)		-	
	Arabi et al. ¹⁰	72/266	83/257	0.84 (0.64-1.09)	-		
	Mackenzie et al. ³³	39/121	47/119	0.82 (0.58-1.15)		-	
	NICE-SUGAR ¹⁸	829/3010	751/3012	1.10 (1.01-1.20)			
	All mixed ICU patients	1351/5051	1301/5089	0.99 (0.87-1.12)		<u>></u>	
	Medical ICU						
	Bland et al. ²⁵	1/5	2/5	0.50 (0.06-3.91)			
	Van den Berghe et al.9	214/595	228/605	0.95 (0.82-1.11)			
	Walters et al.37	1/13	0/12	2.79 (0.12-62.48)		,	•
	Farah et al.27	22/41	22/48	1.17 (0.77-1.78)	-	-	
	Oksanen et al.36	13/39	18/51	0.94 (0.53-1.68)	_	-	
	Bruno et al. ²⁶	2/31	0/15	2.50 (0.13-49.05)		· · · · >	
	All medical ICU patients	253/724	270/736	1.00 (0.78-1.28)			
	Surgical ICU						
	Van den Berghe et al.8	55/765	85/783	0.66 (0.48-0.92)			
	Grev et al.28	4/34	6/27	0.53 (0.17-1.69)			
		~~~~	3.00				
تight" gly	cemic c	ontr	ol d	loes no	ot ben	efit all	patien
• • •							-
specially	those wi	ith i	ncre	eased I	risk ni	f hyno	alvcem
specially				casca	130.01	πιγρο	grycen
					Risk ratio		

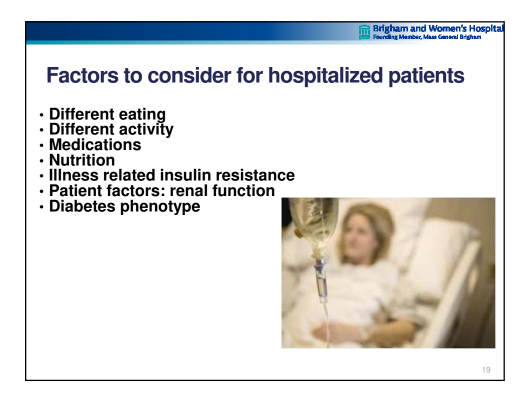






Organization	Critically III	Non-critically III Patient
DA/AACE	< 140-180 mg/dL Initiate insulin >180 mg/dL	Pre-meal <140 mg/dL Random < 180 mg/dL*
NCP	140-200 mg/dL Recommends against IIT	
Critical Care Society	140-180 mg/dL Initiate insulin >150 mg/dL	
Endocrine Society		Pre-meal < 140 mg/dL Random < 180 mg/dL* Adjust regimen < 100 mg/dL
Society of Thoracic Surgeons	Cardiac surgery: IV insulin <180 mg/dL peri-op ≤ 110 mg/dL fasting or premeal	
loint British Diabetes Society		6-10 mmol/L (108-180 mg/dL) acceptable range 4-12 mm/L (72-216 mg/dL)

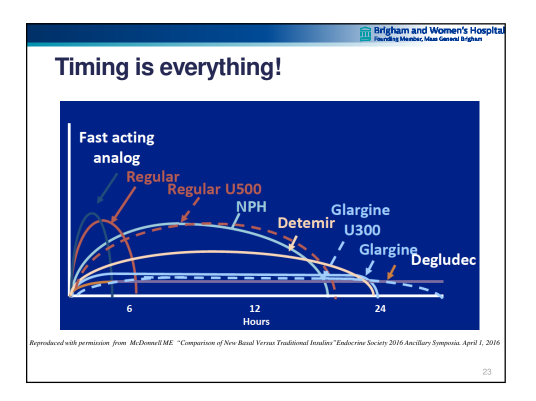
< 180 mg/dL (< 10.0 mmol/L)	Pre-meal <140 mg/dL (< 7.8 mmol/L)
	<ul> <li>Random &lt; 180 mg/dL (&lt; 10.0 mmol/L)</li> <li>Higher glucose levels &lt; 200 mg/dL (&lt; 11.1 mmol/L) may be acceptable in some patients (terminally ill, multiple medical acceptable)</li> </ul>
	medical comorbidities)

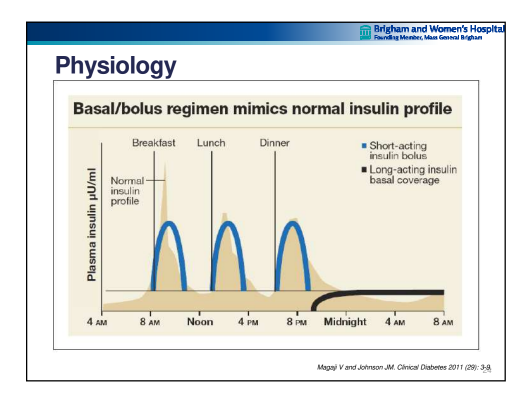


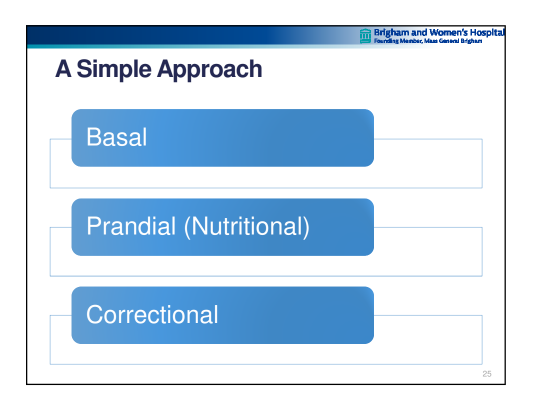
Medication	Advantages	Disadvantages
Metformin	Low risk for hypoglycemia	MALA risk in patients with hypoperfusion (RI, cirrhosis, HF)
Sulfonylureas		Risk of hypoglycemia (RI, reduced po intake)
TZDs	Low risk of hypoglycemia	Slow onset, fluid retention C/I HF or hepatic dysfunction
DPP4-inhibitors	Low risk of hypoglycemia	
GLP-1 agonists	Low risk of hypoglycemia	GI effects Perioperative planning
SGLT-2 inhibitors	Low risk of hypoglycemia	Limited data Increased risk GU infections Risk of dehydration, hypotension, euglycemic DKA Perioperative planning

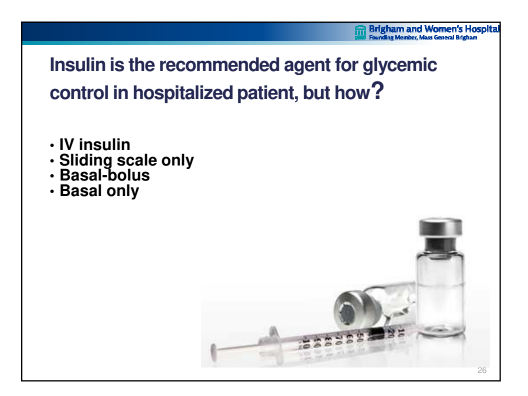


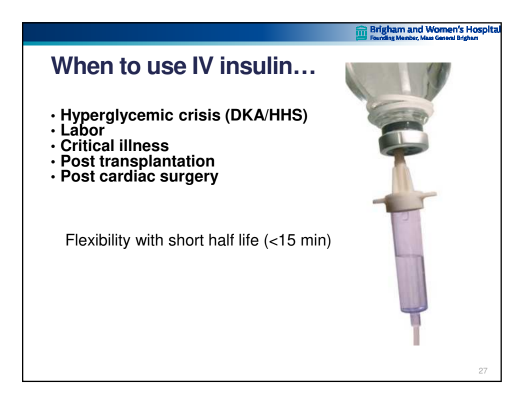
Type of Insulin	Name	Onset	Peak	Duration
Rapid Acting	Aspart (Novolog) Lispro (Humalog) Glulisine (Apidra)	5-15 min	1-2 h	4-6 h
Short Acting	Regular (Humulin R, Novolin R)	30-60 min	2-4 h	6-10 h
ntermediate Acting	NPH (Humulin N, Novolin N)	2-4 h	6-12 h	12-18 h
ong Acting	Glargine (Lantus, Basaglar)	2-4 h	None	22-24 h
	Glargine U-300 (Toujeo) Degludec U-100, U-200 (Tresiba)	6 h 1h	none none	22-36 h 42 h
Pre-Mixed Insulin	NPH/regular (Humulin 70/30,Novolin 70/30)	30-60 min	2-12 h	12-18 h
	Lispro protamine/lispro (Humalog 75/25, Humalog 50/50)	5-15 min	1-2 h	12-18 h
	Aspart Protamine/Aspart (Novolog 70/30)	5-15 min	1-2 h	12-18 h

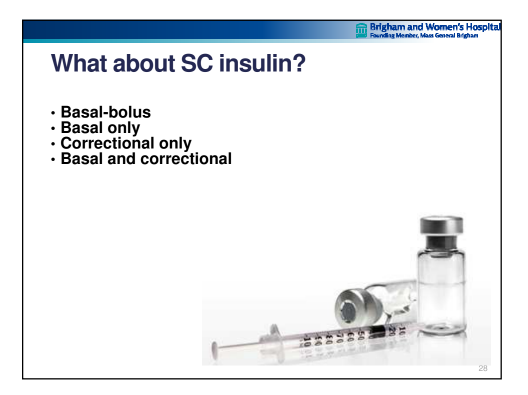


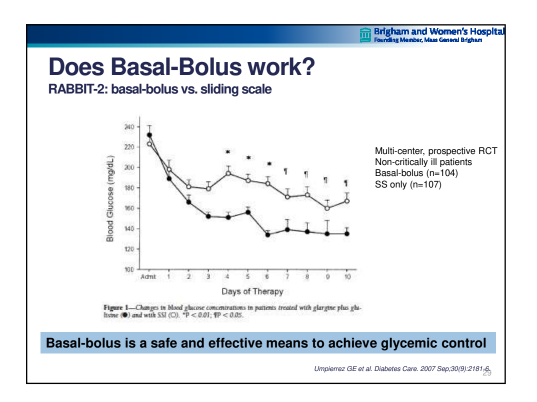


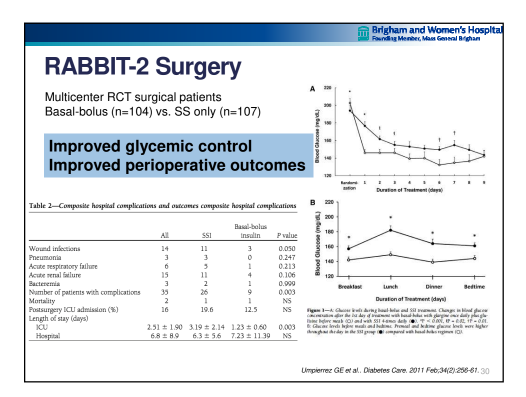


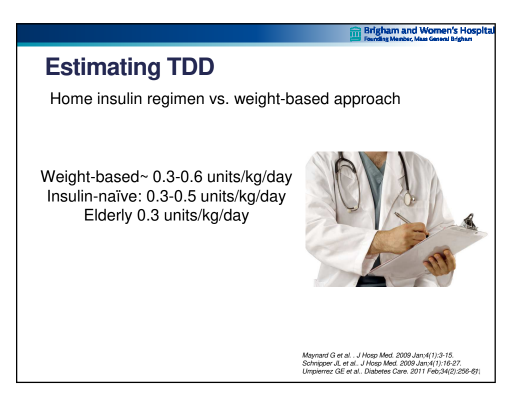


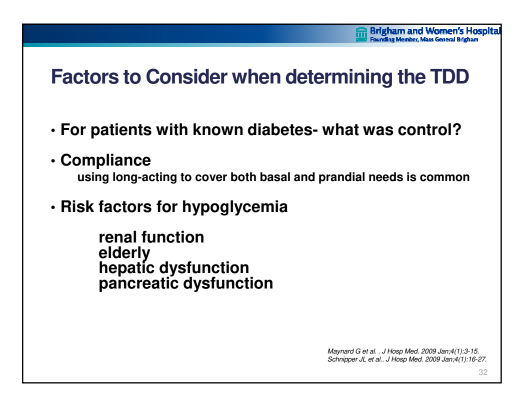




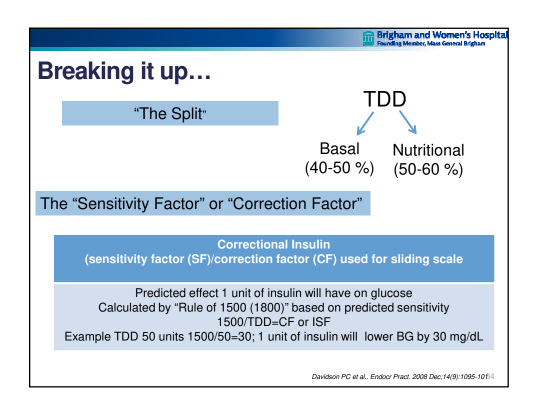


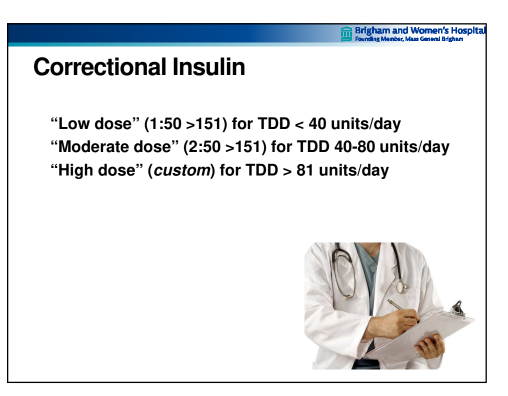






		gham and Women's Hospita ding Member, Mass General Brigham
Estimating TDD Remember this is a place to start	Baseline weight-based TDD estimate	0.5 units/kg/day, adjust by factors listed below
	Age > 70 years	-0.1 units/kg/day
	Renal insufficiency (eGFR < 45)	-0.1 units/kg/day
	Hepatic insufficiency (advanced cirrhosis)	-0.1 units/kg/day
	Pancreatic deficiency (chronic pancreatitis, CF, s/p pancreatectomy)	-0.1 units/kg/day
	HbA1c >10%	+0.1 units/kg/day
	Currently on glucocorticoids with the equivalent of prednisone 40 mg/day or greater	+0.1 units/kg/day
	FINAL TDD estimate	=



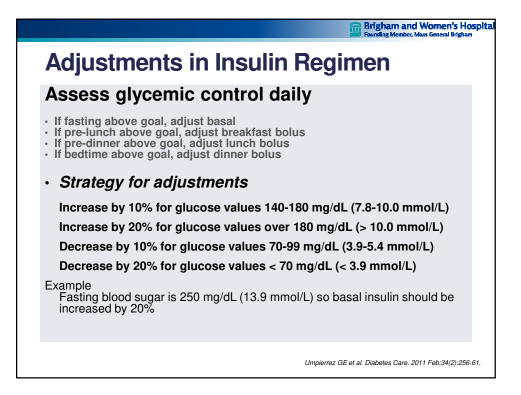


Brigham and Women's Hospita Founding Member, Mass General Brigham
Example Calculation
60 kg patient Normal renal function
Step 1: Estimate TDD (0.5 units/kg x wt) 60 x 0.5= 30 units
Step 2: Determine "the split" (usually 50% basal, 50% prandial) 50% of 30 units= 15 15 units basal insulin 15 units total for prandial/3 (b/l/d)= 5 units AC
Step 3: Determine the "correction" (AKA sliding scale) 1500/TDD=CF 1500/30=50 (for every 1 unit of insulin, expect decrease by ~50 mg/dL)
36

## **Target Glucose Levels**

Critically III Patient	Non-critically III Patient
< 180 mg/dL(< 10.0 mmol/L)	Pre-meal <140 mg/dL (< 7.8 mmol/L) Random < 180 mg/dL (< 10.0 mmol/L)
	•Higher glucose levels < 200 mg/dL (< 11.1 mmol/L) may be acceptable in some patients (terminally ill, multiple medical comorbidities)
	medical comolodities)

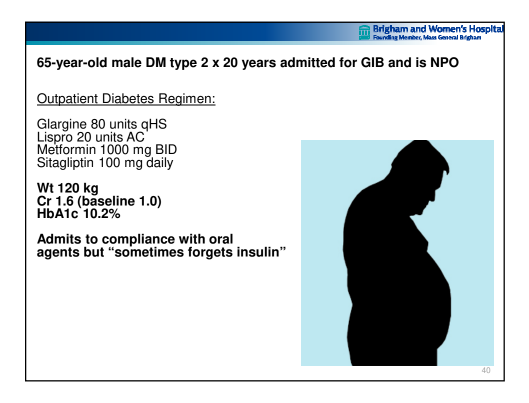
🔐 Brigham and Women's Hospital



Brigham and Women's Hospital Founding Member, Mass General Brigham

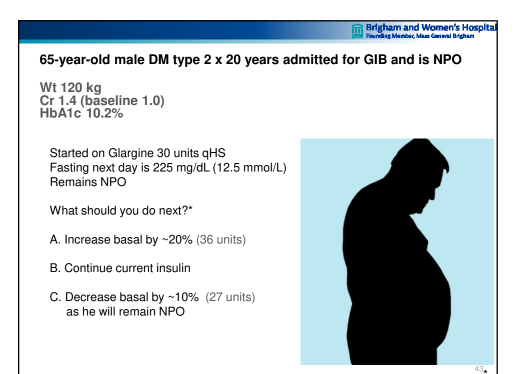
## **Tailor to Clinical Scenario**

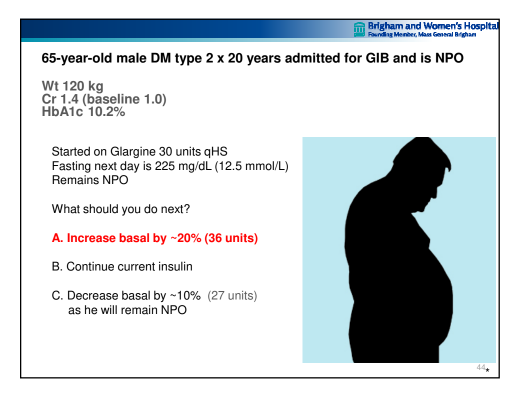
	Example insulin regimen			
NPO	Basal insulin (long or intermediate acting insulin if basal requirement) Regular insulin correction scale q6h			
Unreliable po intake	Basal insulin (long or intermediate acting insulin if basal requirement) RAI with dose reduction for decreased po intake and correction scale (or correction only)			
Reliable po intake	Basal insulin (long or intermediate acting insulin if basal requirement) RAI with meals, correction scale with RAI to be given with nutritional dose			
Parenteral nutrition	Basal insulin (long or intermediate acting insulin if basal requirement) Nutritional insulin given as regular insulin added to TPN bag			
Enteral nutrition	Continuous EN: nutritional dose/4 given as regular insulin q6h ^ Cycled EN: NPH^ at onset (12h cycle), RAI or short acting insulin pending cycle length^ Bolus EN: RAI with bolus ^			
Steroids	Basal insulin (long or intermediate acting insulin if basal requirement)-consider NPH RAI with "stacked doses" "NPH on top of" program			
"If TF/TPN interrupted patient v of last SC insulin given"	^ recommend using order set with safety " hold if TF/TPN held" ''If TF/TPN interrupted patient will require frequent glucose monitoring and may require dextrose support for duration of pharmacologic activity of last SC insulin given" If hypoglycemia, may give IV dextrose at rate of TF if needed to "ride out" insulin action 3			

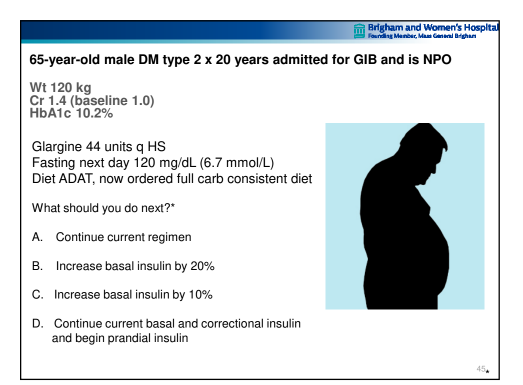


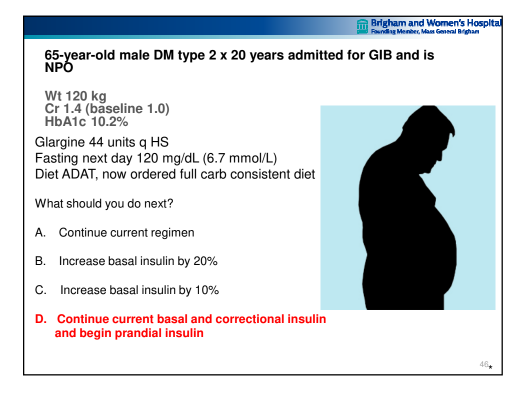
		igham and Women's Hospita Inding Member, Mass General Brigham
Estimating TDD Remember this is a place to start	Baseline weight-based TDD estimate	0.5 units/kg/day, adjust by factors listed below
	Age > 70 years	-0.1 units/kg/day
Wt 120 kg Cr 1.6 (baseline 1.0) HbA1c 10.2%	Renal insufficiency (eGFR < 45)	-0.1 units/kg/day
	Hepatic insufficiency (advanced cirrhosis)	-0.1 units/kg/day
	Pancreatic deficiency (chronic pancreatitis, CF, s/p pancreatectomy)	-0.1 units/kg/day
	HbA1c >10%	+0.1 units/kg/day
	Currently on glucocorticoids with the equivalent of prednisone 40 mg/day or greater	+0.1 units/kg/day
	FINAL TDD estimate	= 0.5 units/kg/day
		41

120 kg patient
Impaired renal function HbA1c >10 %
Step 1: Estimate TDD (0.5 units/kg x wt) 120 x 0.5= 60 units
Step 2: Determine "the split" (usually 50% basal, 50% prandial) 50% of 60 units= 30 30 units basal insulin NPO, no standing prandial insulin
Step 3: Determine the "correction" (AKA sliding scale) 1500/TDD=CF 1500/60=25 (for every 1 unit of insulin, expect decrease by ~25 mg/dL)

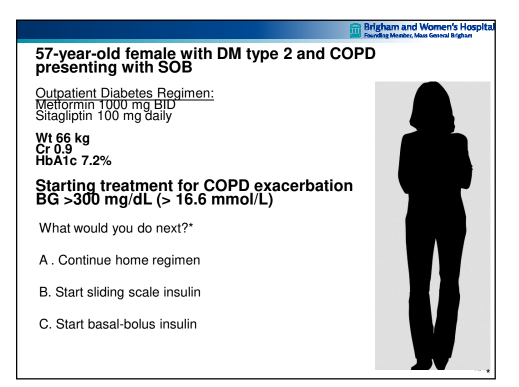


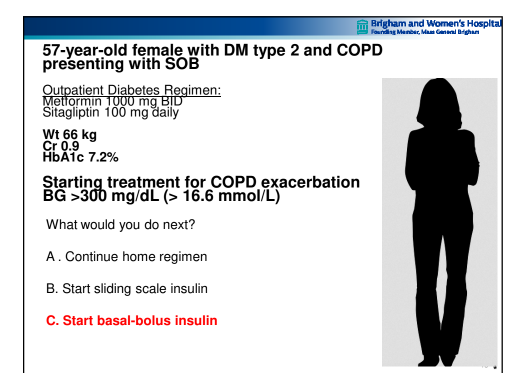


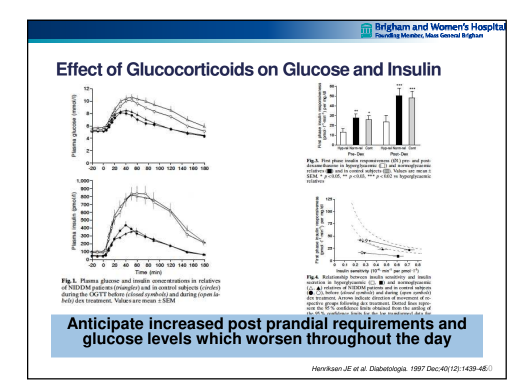


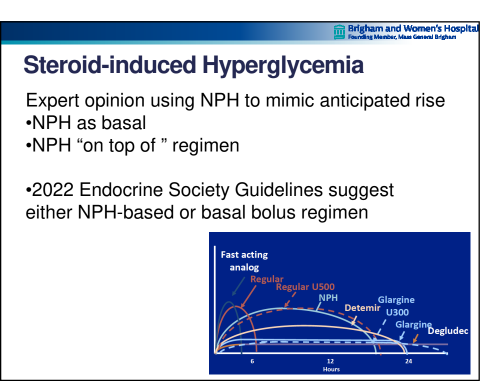


	Brigham and Women's Hospital Founding Member, Mass General Brigham			
Calculation	Diet advanced, anticipate need for prandial insulin, may consider reduced dosed until eating reliably			
<ul> <li>120 kg patient</li> <li>impaired renal function</li> <li>HbA1c&gt;10 %</li> </ul>				
Step 1: Estimate TDD (0.5 units/kg x wt) 120 x 0.5= 60 units-~ 30 units basal -now titrated to 44 units				
Step 2: Determine "the split" (usually 50% basal, 50% prandial) 50% of 60 units= 30 30 units basal insulin NPO, no standing prandial insulin 10 units AC with weight-based, if using new basal 15 units AC if worried about po intake may give 8-10 initially				
Step 3: Determine the "correction" (AKA sliding scale) 1500/TDD=CF 1500/60=25 (for every 1 unit of insulin, expect decrease by ~25 mg/dL)				
Helpful to have carb cons	istent diet for safety of insulin dosing			





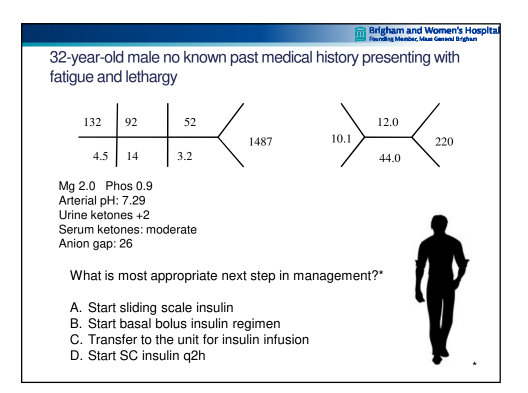


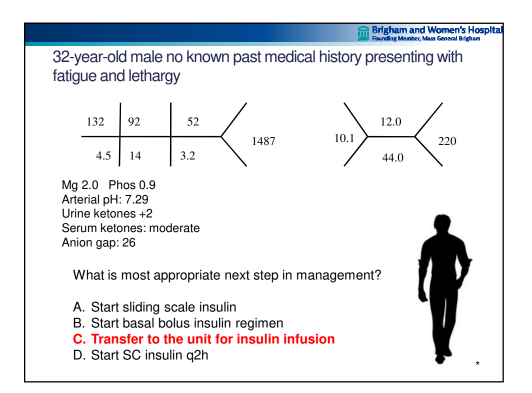


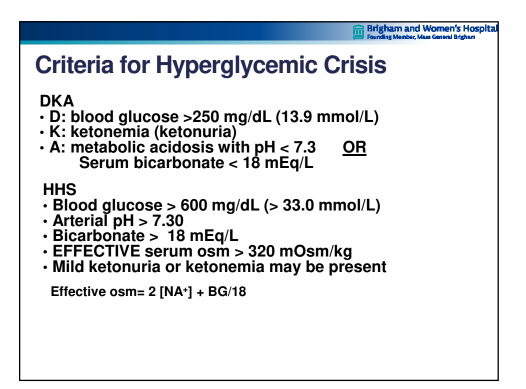
Steroid-induced Hyperglycemia				
Diabetes UK Position Statements Management of hyperglycaemia and ste (glucocorticoid) therapy: a guideline fro British Diabetes Societies (JBDS) for Inpa	m the Joint			
A. Roberts ¹ , J. James ² and K. Dhatariya ³ , on behalf of the Join (JBDS) for Inpatient Care*	t British Diabetes Societies			
Candiff and Vale UniversityLocal Health Board, Candiff, UK, ² University HospitalisLeicenter NHE Trust, Leicenter, UN NHS Roundation Trust, Nervelch, UK	Card Thanks and Norwesh University Hespitals			
Accepted 12 May 2018				
Stress Hyperglycemia	Consider SU or basal insulin (in AM)			
DM type 2 (not on insulin)	SU ± basal insulin (given in AM)			
DM type 2 (on insulin)	Basal insulin: (consider switch to AM and increase dose)			
	Premixed insulin: increase morning dose			
	MDI: increase lunch and dinner RAI			
DM type 1	Increase basal, increase lunch and dinner RAI			

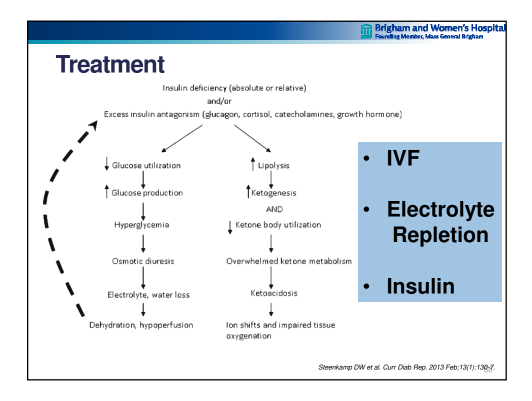
		gham and Women's Hospita ning Member, Mass General Brigham
Estimating TDD Remember this is a place to start	Baseline weight-based TDD estimate	0.5 units/kg/day, adjust by factors listed below
Wt 66 kg Cr 0.9 HbA1c 7.2%	Age > 70 years	-0.1 units/kg/day
	Renal insufficiency (eGFR < 45)	-0.1 units/kg/day
	Hepatic insufficiency (advanced cirrhosis)	-0.1 units/kg/day
	Pancreatic deficiency (chronic pancreatitis, CF, s/p pancreatectomy)	-0.1 units/kg/day
	HbA1c >10%	+0.1 units/kg/day
	Currently on glucocorticoids with the equivalent of prednisone 40 mg/day or greater	+0.1 units/kg/day
	FINAL TDD estimate	= 0.6 unit/kg/day

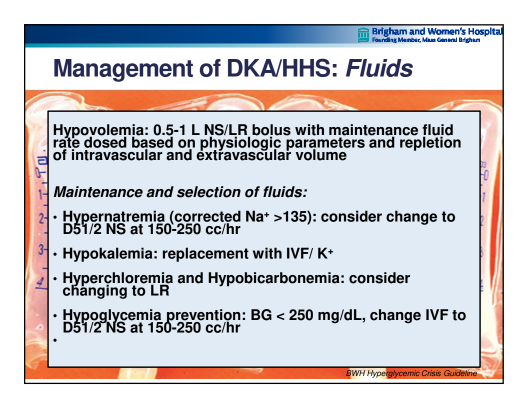
	Brigham and Women's Hospita Founding Member, Mass General Brigham			
Calculation <ul> <li>66 kg patient</li> <li>normal renal function</li> <li>HbA1c 7.2%</li> </ul>	May use 50/50 or consider 40/60 split, using NPH and/or "stacked RAI" with steroids. Anticipate decreased requirements as steroids tapered			
Step 1: Estimate TDD (0.6 units/kg x wt) 66 x 0.6= 40 units				
Step 2: Determine "the split" (usually 50% basal, 50% prandial) 50% of 40 units= 20 units 20 units basal insulin (if using NPH can split 10/10 or 13/7) 20 units prandial insulin 20/3=6 units RAI AC (another strategy 4/5/6)				
1500/TDD=CF	orrection" (AKA sliding scale) unit of insulin, expect decrease			

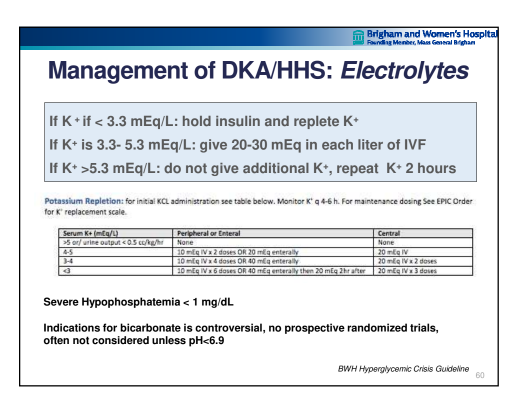


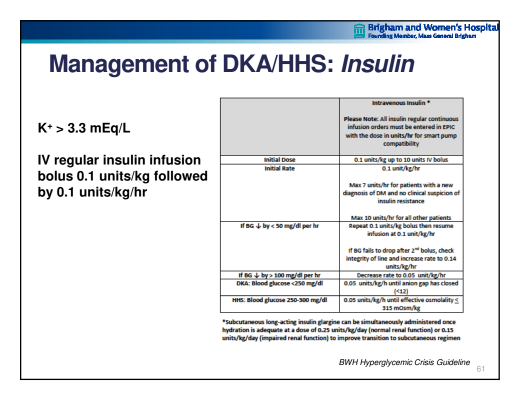




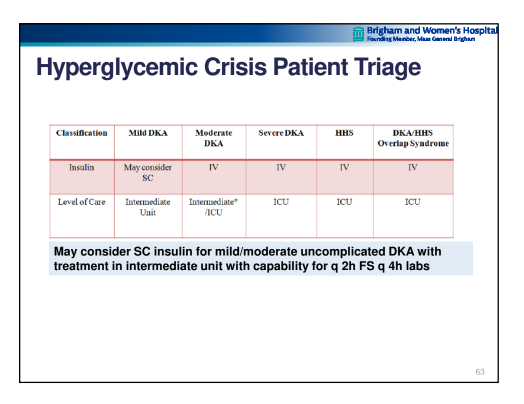


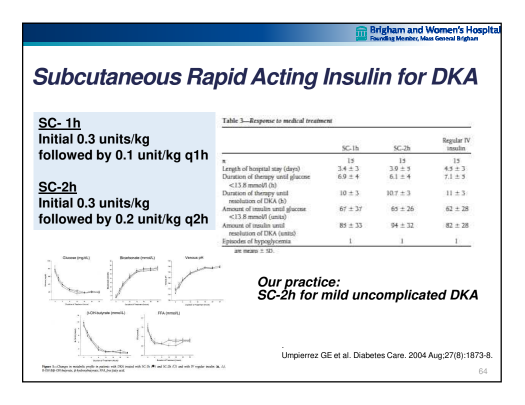






## m Brigham and Women's Hospital Hyperglycemic Crisis: Classification of Patients Mild DKA Moderate DKA Severe DKA HHS Blood glucose (mg/dL) >250 >250 >250 >600 pH < 7.30 7.12-7.24 < 7.15 >7.30 15-18 10 to < 15 <10 HCO, >18 Urine/Serum Ketones + + ÷ +/-Serum Osm (Osm_) >320 AG elevated elevated elevated variable Mental Status alert alert/drowsy stupor/coma stupor/coma Modified from Kitabchi et al. Diabetes Care 2009 32(7):1335-1343 62

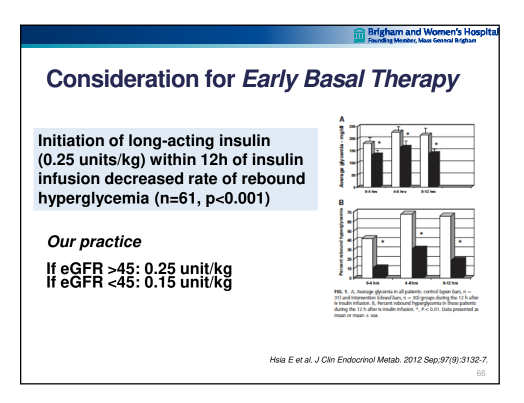


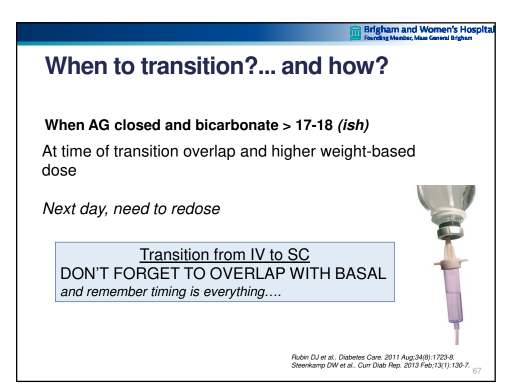


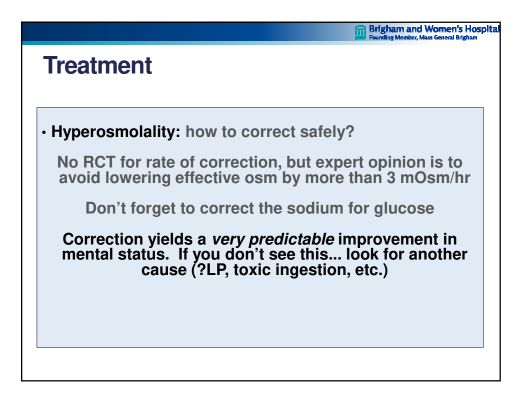
## Example Protocol for Use of Subcutaneous Insulin Protocol in Treatment of Mild Uncomplicated DKA

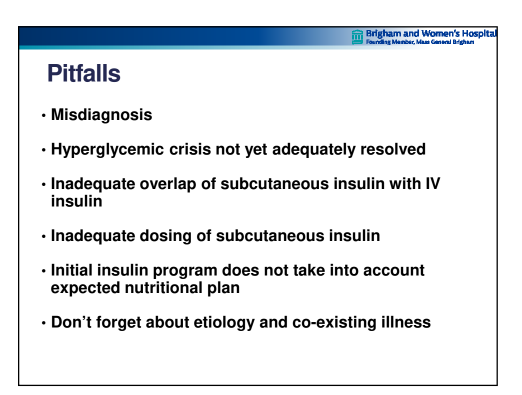
	Subcutaneous Insulin Aspart	Subcutaneous Insulin Glargine
Initial Dose	0.3 units/kg maximum 20 units	0.25 units/kg if GFR >40; 0.15 units/kg if GFR <40
Subsequent Dose	0.2 units/kg every 2 hours maximum 10 units	Redose in 24 hours based on response to initial dose
Blood glucose <250 mg/dl	0.05-0.1 units/kg every 2 hours	

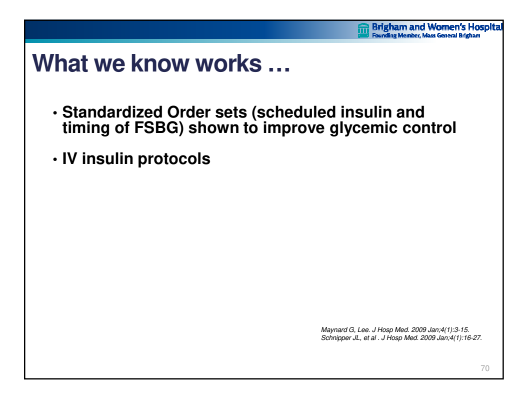


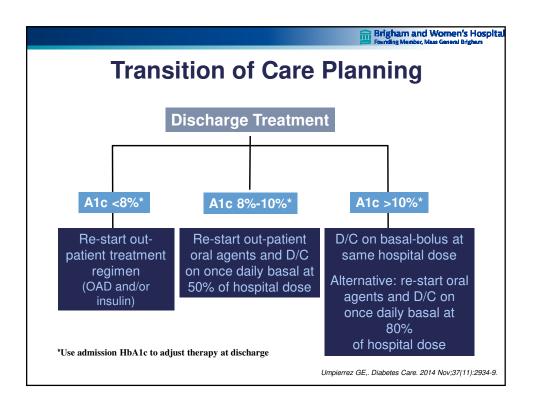




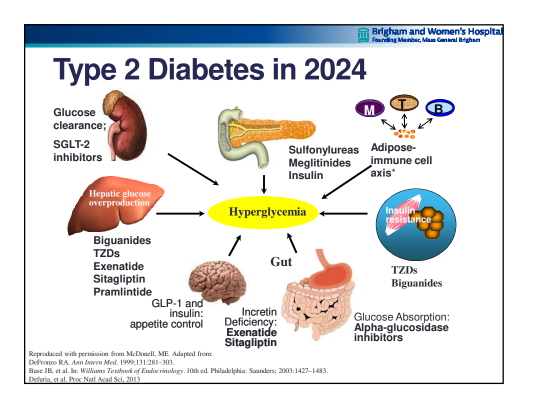


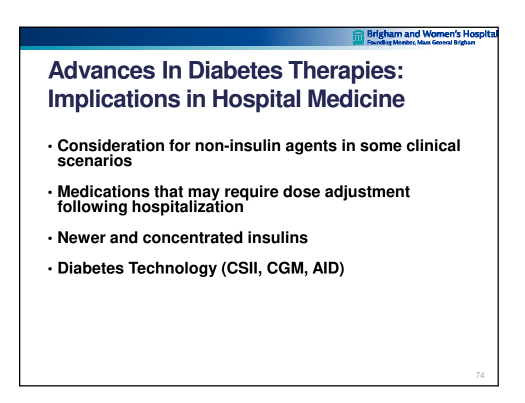






Brigham and Women's Hospital Bounding Member, Mass General Brigham				
Transition of Care Checklist				
Diabetes Education ("survival skills")				
Insulin Teaching (if applicable, should include pen and vial/syringe)				
Glucometer Teaching				
Confirm patient has diabetes supplies:				
<ul> <li>Medications* (if using insulin vial-syringe; if insulin pen-pen needles)</li> <li>Test strips (must match glucometer)</li> <li>Lancets</li> </ul>				
Clear communication with patient regarding discharge regimen*				
Follow-up appointment scheduled				
PCP aware of any dose adjustments				
*Medications and supplies will vary depending on insurance coverage- often human insulin cheaper than analogs; helpful to know coverage for pen vs. vial/syringe prior to discharge ¹²				





📻 Brigham and Women's Hospital **Metformin and Risk of Acidosis** 

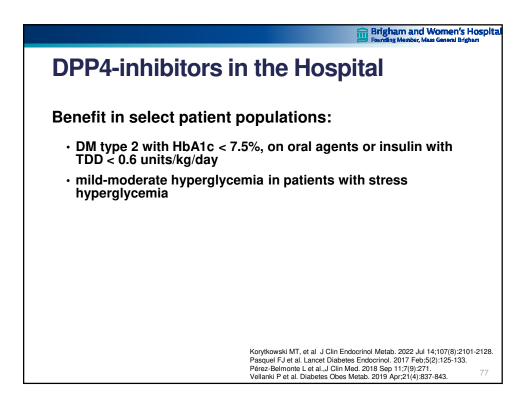
Association of Time-Dependent Metformin Use With Acidosis Hospitalization by Time-Dependent Estimated Glomerular Filtration Rate (eGFR) Category in Geisinger Health System

Table 3.

Parameter	$\rm HR^{8}$ (95% CI) for Acidosis Associated With Metformin Use by Time-Dependent eGFR Category, mL/min/1.73 m^2								
	Overall ^b	≥90	60-89	45-59	30-44	<30			
Person-time (on metformin/off metformin)	188 578/281 536	80 653/98 905	79 788/102 110	21 232/40 861	6358/29 834	548/9827			
Acidosis events (on metformin/off metformin)	737/1598	206/323	288/446	157/286	64/314	22/229			
Unadjusted (n = 75 413)	0.89 (0.81-0.97)	0.77 (0.65-0.92)	0.82 (0.71-0.95)	1.05 (0.87-1.28)	0.95 (0.73-1.25)	1.71 (1.10-2.64)			
Demographic adjusted ^C (n = 75 413)	0.89 (0.81-0.97)	0.75 (0.63-0.90)	0.82 (0.71-0.96)	1.07 (0.88-1.30)	0.98 (0.75-1.28)	1.76 (1.14-2.73)			
Fully adjusted ^d (n = 72 232)	0.98 (0.89-1.08)	0.88 (0.73-1.05)	0.87 (0.75-1.02)	1.16 (0.95-1.41)	1.09 (0.83-1.44)	2.07 (1.33-3.22)			
Fully adjusted with time-dependent medication use ^e (n = 72 232)	0.94 (0.83-1.05)	0.80 (0.66-0.97)	0.81 (0.68-0.95)	1.14 (0.93-1.40)	1.13 (0.85-1.49)	2.21 (1.42-3.44)			
Sensitivity analyses									
Fully adjusted ^d excluding baseline insulin users (n = 60 112)	1.02 (0.91-1.13)	0.88 (0.71-1.09)	0.89 (0.75-1.06)	1.21 (0.97-1.50)	1.16 (0.87-1.57)	2.22 (1.41-3.51)			
Fully adjusted ^d including adjustment for baseline hemoglobin $A_{1c}$ (n = 58 093)	1.01 (0.90-1.14)	0.84 (0.67-1.04)	0.93 (0.78-1.12)	1.23 (0.98-1.55)	1.07 (0.78-1.46)	2.22 (1.37-3.59)			
Fully adjusted ^d in incident diabetes mellitus cohort (n = 49 839)	0.91 (0.79-1.04)	0.85 (0.68-1.06)	0.82 (0.66-1.01)	1.15 (0.86-1.53)	0.88 (0.55-1.39)	2.37 (1.20-4.71)			
Fully adjusted ^d with early censoring of metformin ( $n = 72232$ )	1.04 (0.95-1.15)	0.93 (0.78-1.12)	0.93 (0.80-1.09)	1.23 (1.01-1.50)	1.17 (0.89-1.54)	2.26 (1.45-3.51)			

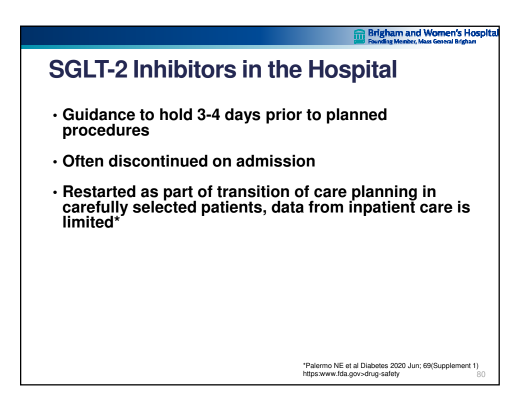
Lazarus B1,2, Wu A1, Shin et al. Association of Metformin Use With Risk of Lactic Acidosis Across the Range of Kidney Function: A Community-Based Cohort Study. JAMA Intern Med. 2018 Jul 1;178(7):903-910.

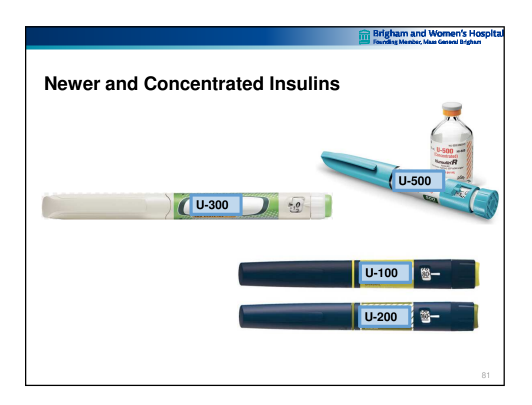
m Brigham and Women's Hospital Incretin-based therapy in hospitalized patients Lina-Real-World Study Basal-bolus vs basal-linagliptin 180 Observational, multicenter 170 Non-critically ill patients with DM type 2 160 on oral agents (n=953) 150 140 DPP4i effective in patients with 3 mild-moderate hyperglycemia Minimizing injection burden 180 170 Lower risk of hypoglycemia 16 150 p=0.401 Time of the day Pérez-Belmonte L et al., J Clin Med. 2018 Sep 11;7(9):271. 76



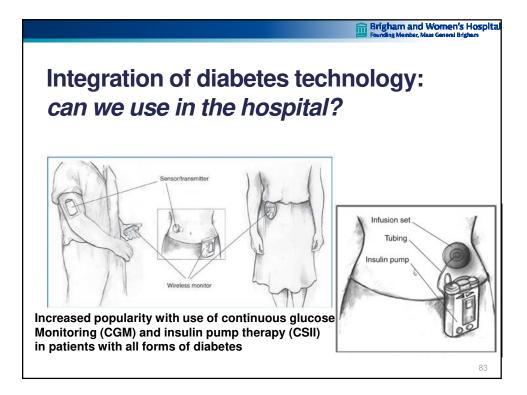
			(ham and Women' Ing Member, Mass General B	
e Adjustments	Based o	on Rena	al Funct	ion:
gliptin agliptin				
GFR (ml/min)	<u>&gt;</u> 50	30-49	<30	
Sitagliptin	100 mg	50 mg	25 mg	
Saxagliptin	5 mg	2.5 mg	2.5 mg	
Linagliptin	5 mg	5 mg	5 mg	
	gliptin agliptin GFR (ml/min) Sitagliptin Saxagliptin	gliptin agliptinGFR (ml/min)≥ 50Sitagliptin100 mgSaxagliptin5 mg	e Adjustments Based on Renagliptin agliptinGFR (ml/min)≥ 5030-49Sitagliptin100 mg50 mgSaxagliptin5 mg2.5 mg	e Adjustments Based on Renal Functgliptin agliptinGFR (ml/min)≥ 5030-49<30

able 1-Clinical character		OKA cases											
ase patient	1	2	3		4		5	6		7		8	9
ge (years) ex	40 Female	58 Maie	27 5ema		2 Ferr	*	31 Female	55		26 Female		39 Female	64 Female
ex 1/T2	Female T1	Male T2	Fema T1	Ne .	Ferr		Female T1	Female T1		Female T1		Female T1	Female T2
1/12 ADI/CSII	MDI	N/A	MD		C	-	CSII	CSII		CSII		CSII	N/A
uration (years)	17	2	25		6		15	18		13		26	6
Mi (kg/m²)	26.5	26.5	24.3		25	9	33.2	22.0		22.0		26.1	32.8
rior A1C 1% (mmol/moliil	11.4 (101.1)	9.8 (83.6)	7.8 (6)		8.0 8		7.0 (53.0)	7.2 (55.2)		5.6 (48.6)		7.0 (53.0)	7.8 (62.0)
anagliflozin dose (mg)	300	300	300	100	300	100	300	300		150		300	300
otential contributors	URI	Surgery 1 week prior	URI, alcohol	Akohol	Alcohol	Exercise, alcohol	Exercise	GI		None		URI	Surgery 12 h prior
just prior to euDKA	Yes	N/A	Yes	No	Yes	Yes	Yes	Unknown	No	No	No	Yes	N/A
resenting plasma glucose (mg/dL (mmol/L))	2 20 (12.2)	150 (8.3)	150 (8.3)	96 (5.3)	224 (12.4)	158 (8.8)	~125 (~6.9)	203 (11.3)	190 (10.6)	150 (8.3)		233 (12.9)	169 (9.4)
4		7.44	0.07										
xo ₂ (m mHg) carbonate (mEg/L)	10	10			11	18		15	26				13 and then 5
carbonate (mEq/L) nion gap (mEq/L)	25	10	35		22	18		26	21			24	15 and then 5
stones*	Yes (serum and urine)	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes (serum and urine)
/here treated	ICU	ICU	ICU	Outpt.	ICU	Inpt	Outpt	ICU	ICU	Outpt.	Outot.	ICU	ICU.

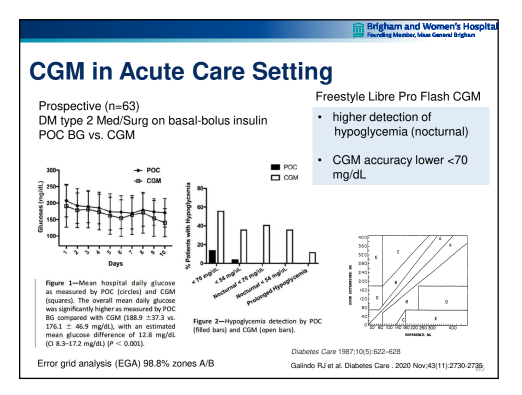




	delivery of HUMULIN R U-500 using these devices Delivery Using a U-100 insulin syringe	Delivery Using a Tuberculin syringe
HUMULIN R U-500	Amount of HUMULIN R U-500 to draw up in the syringe in	
dose prescribed (units of insulin)	"unit marking"	"volume marking"
	Conversion: Divide prescribed dose by 5	Conversion: Divide prescribed dose by 500
25 Units	Draw to the 5 unit mark on syringe	Draw to the 0.05 mL mark on syringe
50 Units	Draw to the 10 unit mark on syringe	Draw to the 0.1 mL mark on syringe
75 Units	Draw to the 15 unit mark on syringe	Draw to the 0.15 mL mark on syringe
100 Units	Draw to the 20 unit mark on syringe	Draw to the 0.2 mL mark on syringe
125 Units	Draw to the 25 unit mark on syringe	Draw to the 0.25 mL mark on syringe
150 Units	Draw to the 30 unit mark on syringe	Draw to the 0.3 mL mark on syringe
175 Units	Draw to the 35 unit mark on syringe	Draw to the 0.35 mL mark on syringe
200 Units	Draw to the 40 unit mark on syringe	Draw to the 0.4 mL mark on syringe
225 Units	Draw to the 45 unit mark on syringe	Draw to the 0.45 mL mark on syringe
250 Units	Draw to the 50 unit mark on syringe	Draw to the 0.5 mL mark on syringe
500 Units	Draw to the 100 unit mark on syringe	Draw to the 1.0 mL mark on syringe
		Source and the source of the s



			righam and Wornen's unding Member, Mass General B
GM Use in [•]	the Hospita	I: special	
	-	-	
onsideratio	ns		
	patient: imaging (MI	, · · ·	
	st: surgery, pressors	, periods of rapid gli	ucose fluctuation
ct of certain medica	ations		
Table 1 List of FDA-approved CGM	systems with features, limitations, and	interfering substances	
CGM system	Key features	Limitations	Known interfering substance
Abbott Diabetes Care FreeStyle	a). No calibration required	a). Requires scanning every 8 h to	Ascorbic acid
Libre 14 day System [13]	<ul> <li>b). 1-h warm-up</li> <li>c). 14-day sensor wear</li> <li>d). Range 40–500 mg/dl</li> </ul>	preserve data b). No threshold or predictive alerts	Salicylic acid
Abbott Diabetes Care Freestyle Libre 2 [12]	<ul> <li>a). No calibration required</li> <li>b). 1-h warm-up</li> </ul>	<ul> <li>a). Requires scanning every 8 h to preserve data</li> </ul>	Ascorbic acid
Line 2 [12]	c). 14-day sensor wear d). Range 40-400 mg/dl	b). No predictive alarms c). Limited ability to transmit data	
	<ul> <li>e). Optional alarms for hypoglyce- mia, hyperglycemia, and signal</li> </ul>	c). Limited ability to transmit data	
	loss		A MARKAN STRANDOR STRANDOR ST
Dexcom G6 [14]	<ul> <li>a). No calibration required</li> <li>b). 10-day sensor wear</li> </ul>	a). 2-h warm-up	Hydroxyurea
	<ul> <li>c). 40–400 mg/dl range</li> <li>d). Predictive alerts for hypogly- cemia</li> </ul>		
Medtronic MiniMed Guardian Sen- sor [15]	<ul> <li>a). 7-day sensor wear</li> <li>b). Predictive alerts</li> </ul>	<ul> <li>a). 2–4 calibrations/day required</li> <li>b). 2-h warm up</li> </ul>	Acetaminophen
201 [10]	c). Range 40–400 mg/dl	c). 7-day sensor wear	
	a). 90-180 day sensor wear	<ul> <li>a). Implantable</li> <li>b). 2 calibrations/day required</li> </ul>	Mannitol, tetracycline
Senseonics Eversense [16]	<ul> <li>b). Predictive hypo- and hyperglyce- mia alerts</li> <li>c). Conditional MRI compatibility</li> </ul>	c). 24-h warm-up	



		· · · · · ·	J			
Prospective RCT Non-critically ill patien Insulin treated DM typ	( )		GM (De) mia	xcom	ı) vs PC	)C
Table 2—Glycemic outcomes	RT-CGM/GTS group $(n = 36)$	POC group $(n = 36)$	P value	2	1.69	
Hypoglycemic events/patient	K1-Colw/G15 group (II - 56)	POC group (n - 56)	P Value	1.6		
	0.67 (0.34-1.30)	1.69 (1.11-2.58)	0.024			
<70 mg/dL						
<54 mg/dL	0.08 (0.03-0.26)	0.75 (0.51-1.09)	0.003	tig 1.2		
<54 mg/dL locturnal hypoglycemic events/patient				1.2		
<54 mg/dL locturnal hypoglycemic events/patient <70 mg/dL	0.19 (0.09-0.41)	0.33 (0.19-0.59)	0.26	uts ber Patient	0.67	0.75
<54 mg/dL locturnal hypoglycemic events/patient <70 mg/dL <54 mg/dL	0.19 (0.09–0.41) 0.03 (0.01–0.24)	0.33 (0.19–0.59) 0.11 (0.04–0.33)		er Patier	0.67	0.75
<54 mg/dL locturnal hypoglycemic events/patient <70 mg/dL <54 mg/dL lypoglycemic events (<70 mg/dL)/patient/day	0.19 (0.09-0.41)	0.33 (0.19-0.59)	0.26 0.26	er Patier	0.67	0.75
<pre>&lt;54 mg/dL locturnal lypoglycemic events/patient &lt;70 mg/dL &lt;54 mg/dL lypoglycemic events (&lt;70 mg/dL)/patient/day BR &lt;70 mg/dL (%)</pre>	0.19 (0.09–0.41) 0.03 (0.01–0.24) 0.12 (0.06–0.24)	0.33 (0.19–0.59) 0.11 (0.04–0.33) 0.35 (0.23–0.54)	0.26 0.26 0.011	0. Events Per Patier 8	0.67	0.75
<54 mg/dL locturnal hypoglycemic events/patient <70 mg/dL <54 mg/dL ypoglycemic events (<70 mg/dL)/patient/day	0.19 (0.09-0.41) 0.03 (0.01-0.24) 0.12 (0.06-0.24) 0.40 (0.18-0.92)	0.33 (0.19–0.59) 0.11 (0.04–0.33) 0.35 (0.23–0.54) 1.88 (1.26–2.81)	0.26 0.26 0.011 0.002	0. Events Per Patier 8		0.08
<pre>&lt;54 mg/dL Octurnal hypoglycemic events/patient &lt;70 mg/dL &lt;54 mg/dL ypoglycemic events (&lt;70 mg/dL)/patient/day BR &lt;70 mg/dL (%) BR &lt;54 mg/dL (%)</pre>	0.19 (0.09-0.4) 0.03 (0.01-0.24) 0.12 (0.06-0.24) 0.40 (0.18-0.92) 0.05 (0.01-0.43)	0.33 (0.19-0.59) 0.11 (0.04-0.33) 0.35 (0.23-0.54) 1.88 (1.26-2.81) 0.82 (0.47-1.43)	0.26 0.26 0.011 0.002 0.017	No. Events Per Patie	< 70 mg/dL	0.08 < 54 mg/dL
<54 mg/dL octumal hypoglycemic events/patient <70 mg/dL <54 mg/dL ypoglycemic events (<70 mg/dL)/patient/day BR <70 mg/dL (%) BR <54 mg/dL (%)	0.19 (0.09-0.41) 0.03 (0.01-0.24) 0.12 (0.06-0.24) 0.04 (0.18-0.92) 0.05 (0.01-0.43) 59.12 (52.47-66.61)	0.33 (0.19-0.59) 0.11 (0.04-0.33) 0.35 (0.23-0.54) 1.88 (1.26-2.81) 0.82 (0.47-1.43) 54.69 (47.96-62.37)	0.26 0.26 0.011 0.002 0.017 0.39	No. Events Per Patie		0.08
<pre>&lt;54 mg/dL octurnal hypoglycemic events/patient &lt;70 mg/dL &lt;54 mg/dL ypoglycemic events (&lt;70 mg/dL)/patient/day BR &lt;54 mg/dL (%) BR &lt;54 mg/dL (%) BR 70-180 mg/dL (%) R &gt;100-250 mg/dL (%)</pre>	0.19 (0.09-0.41) 0.03 (0.01-0.24) 0.12 (0.06-0.24) 0.40 (0.18-0.92) 0.05 (0.01-0.43) 59.12 (52.47-66.61) 29.88 (26.11-34.19)	0.33 (0.19-0.59) 0.11 (0.04-0.33) 0.35 (0.23-0.54) 1.88 (1.26-2.81) 0.82 (0.47-1.43) 54.69 (47.96-62.37) 30.10 (26.11-34.70)	0.26 0.26 0.011 0.002 0.017 0.39 0.94	No. Events Per Patie	< 70 mg/dL	0.08 < 54 mg/dL p=0.003

		Brigham and Women's I Founding Member, Mass General Brig	
Glucos	e Monitoring ir	n Hospitalized Patients	
	Advantages	Disadvantages	
POC testing	Readily available	Labor intensive (IV q1-2h) Patient preference Does not provide full 24h glycemic profile	
CGM	Provides 24h glycemic profile Potential prediction of hypoglycemic event Alarm for asymptomatic hypoglycemia	Cost?	
			87

