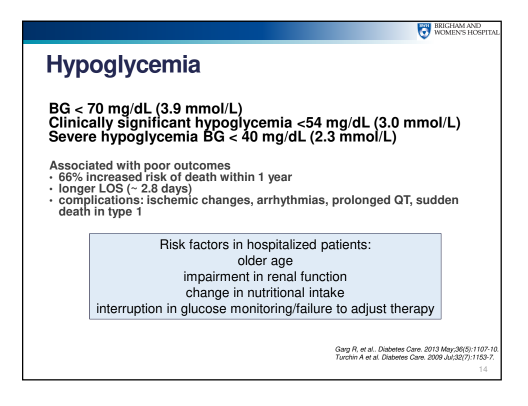
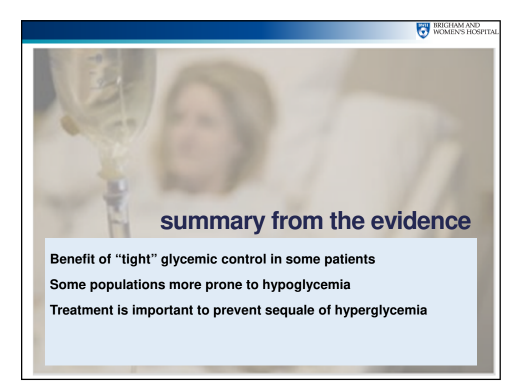
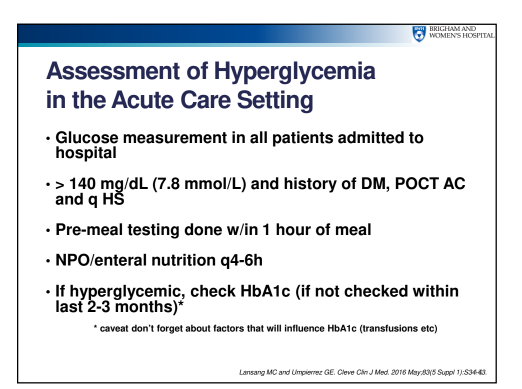


| <i>lortality</i> | | | otal no. patien | | Favours IIT Favours control | r |
|------------------|------------------------------------|-----------|-----------------|---------------------|-----------------------------|---------------|
| | Study Mixed ICU | IIT | Control | Risk ratio (95% CI) | | |
| | 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) | • | |
| | 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) | | \rightarrow |
| | 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) | | \rightarrow |
| | 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) | | |
| | Grey et al. ²⁸ | 4/34 | 6/27 | 0.53 (0.17-1.69) | | |
| | - | e 140 | - | - | | |
| Tiaht" alv | cemic c | ontr | ol d | loes na | ot benefit a | all natien |
| ingin giy | | onu | | | | an putien |
| | | | | | wiels of hours | |
| | tnose w | itn i | ncre | easea | risk of hyp | ogivcen |
| | | - | | | 71 | - 3 7 |
| | | | | | Risk ratio (95% CI) | |

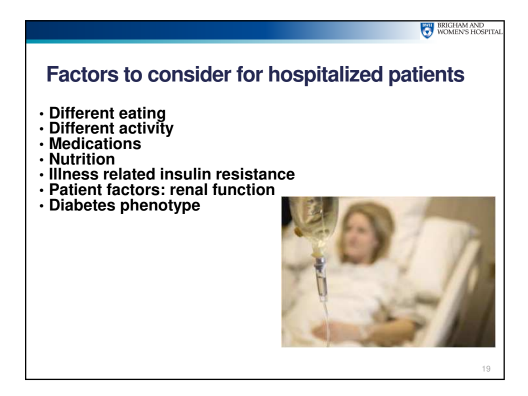






| Organization | Critically III | Non-critically III Patient |
|--------------------------------|---|---|
| ADA/AACE | < 140-180 mg/dL | Pre-meal <140 mg/dL |
| | Initiate insulin >180 mg/dL | Random < 180 mg/dL* |
| ACP | 140-200 mg/dL Recommends against IIT | |
| Critical Care Society | 140-180 mg/dL 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 | |
| Joint British Diabetes Society | | 6-10 mmol/L (108-180 mg/dL) acceptable range 4-12 mm/L (72-216 mg/dL) |

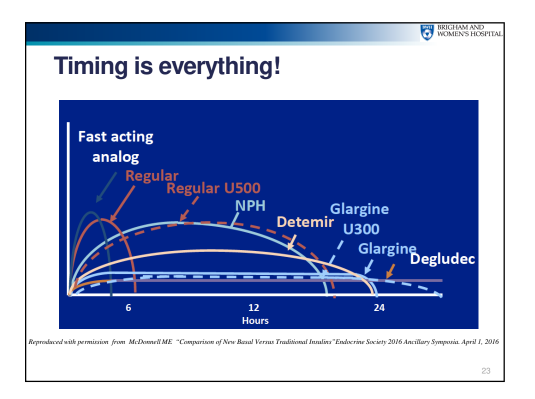
| Critically III Patient < 180 mg/dL (< 10.0 mmol/L) | Non-critically III Patient Pre-meal <140 mg/dL (< 7.8 mmol/L) |
|---|--|
| | Random < 180 mg/dL (< 10.0 mmol/L Higher glucose levels < 200 mg/dL (< 11.1 mmol/L) may be acceptable in some patients (terminally ill, multiple medical comorbidities) |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

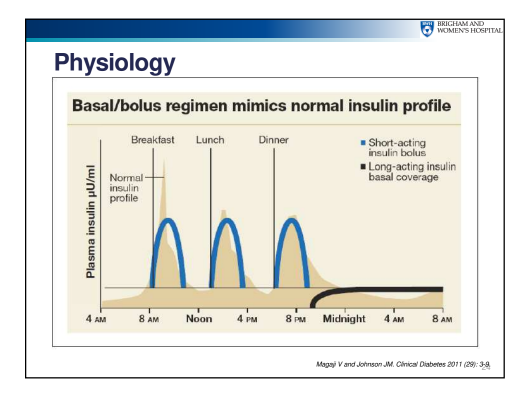


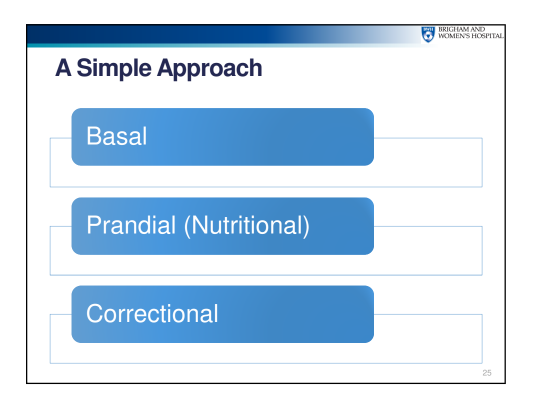
| | Advantages | Disadvantages |
|-------------------|---------------------------|---|
| Metformin | Low risk for hypoglycemia | MALA risk in patients with hypoperfusion (RI, cirrhosis, HF) |
| Sulfonylureas | | Risk of hypoglycemia (RI, reduced po intake) |
| TZDs | Low risk of hypoglycemia | Slow onset,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

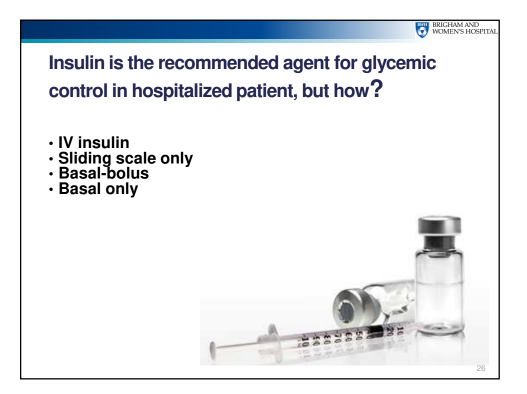


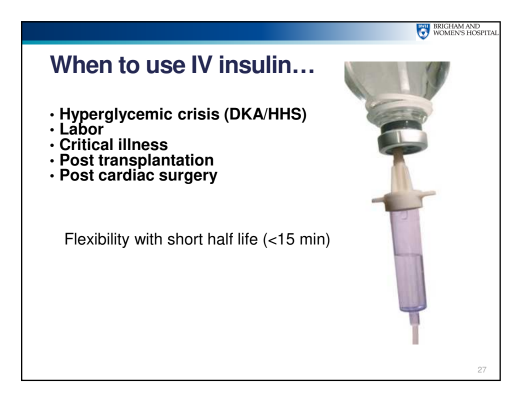
| Type of Insulin | Name | Onset | Peak | Duration |
|------------------------|--|-----------|--------------|--------------------|
| Rapid Acting | Aspart (Novolog) Lispro (Humalog) Glulisine (Apidra) | 5-15 min | 1-2 h | 4-6 h |
| Short Acting | Regular (Humulin R, Novolin R) | 30-60 min | 2-4 h | 6-10 h |
| Intermediate Acting | NPH (Humulin N, Novolin N) | 2-4 h | 6-12 h | 12-18 h |
| Long Acting | Glargine (Lantus, Basaglar) Determir (Levemir) | 2-4 h | None | 22-24 h 17-24 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 |

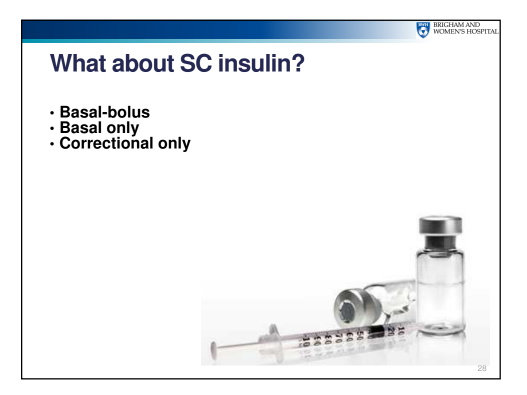


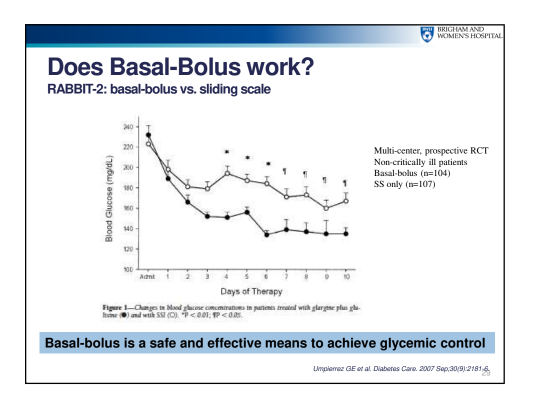


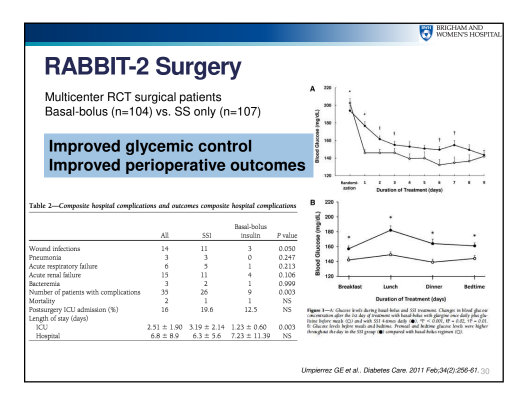


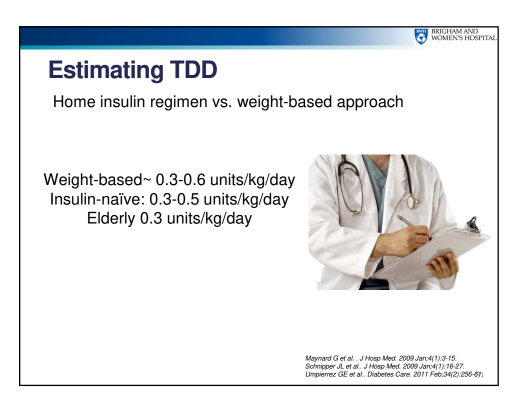


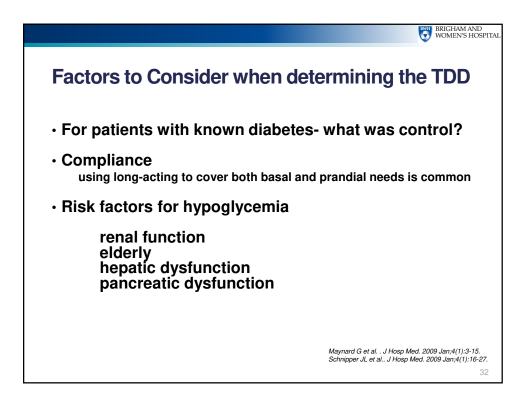




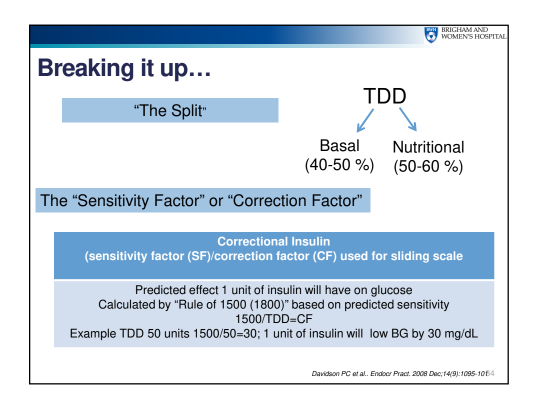


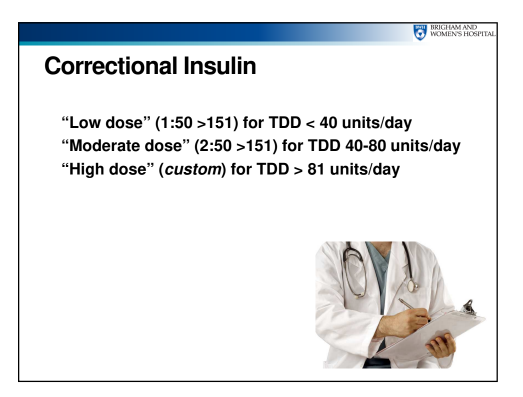






| | | BRIGHAM AND WOMEN'S HOSPITAL |
|---|--|--|
| Estimating TDD Remember this is a place to start | Baseline weight-based TDD estimate | 0.5 units/kg/day, adjust by factors listed below |
| | Age > 70 years | -0.1 units/kg/day |
| | Renal insufficiency (eGFR < 45) | -0.1 units/kg/day |
| | Hepatic 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 | = |



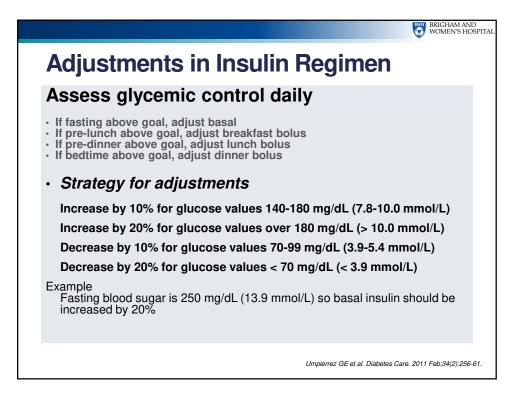


| BRIGHAM AND |
|---|
| Example Calculation |
| 60 kg patient Normal renal function |
| Step 1: Estimate TDD (0.5 units/kg x wt) 60 x 0.5= 30 units |
| Step 2: Determine "the split" (usually 50% basal, 50% prandial) 50% of 30 units= 15 15 units basal insulin 15 units total for prandial/3 (b/l/d)= 5 units 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 comorbidities) |
| | |
| | |
| | |

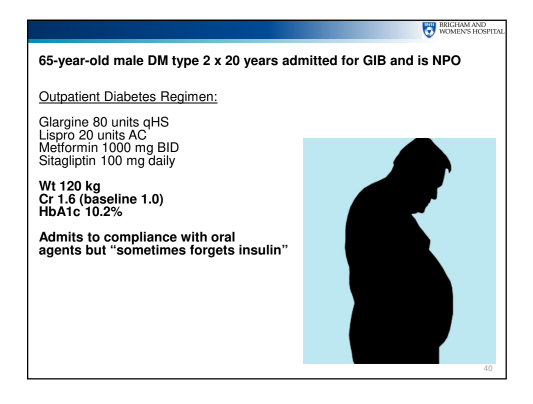
BRIGHAM AND WOMEN'S HOSPITAL



BRIGHAM AND WOMEN'S HOSPITAL

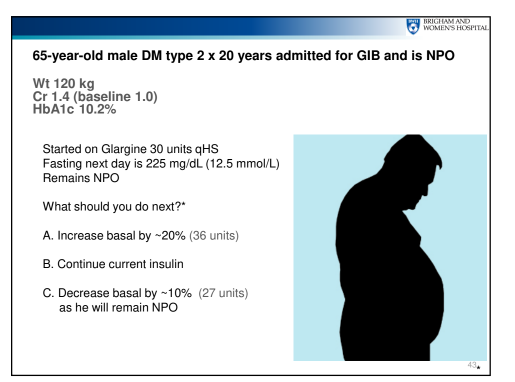
Tailor to Clinical Scenario

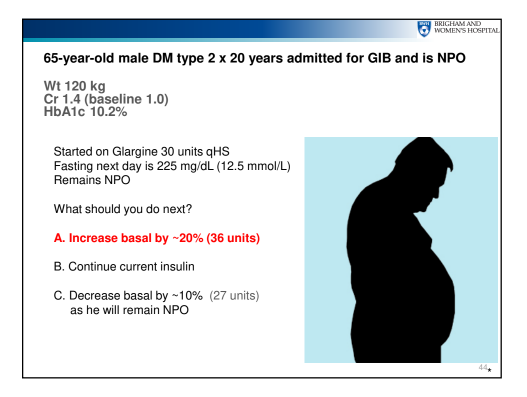
| | Example insulin regimen | |
|---|--|----|
| NPO | Basal insulin (long or intermediate acting insulin if basal requirement) Regular insulin correction scale q6h | |
| Unreliable po intake | Basal insulin (long or intermediate acting insulin if basal requirement) RAI with dose reduction for decreased 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" | vith safety " hold if TF/TPN held" vill require frequent glucose monitoring and may require dextrose support for duration of pharmacologic activity extrose at rate of TF if needed to "ride out" insulin action | 39 |

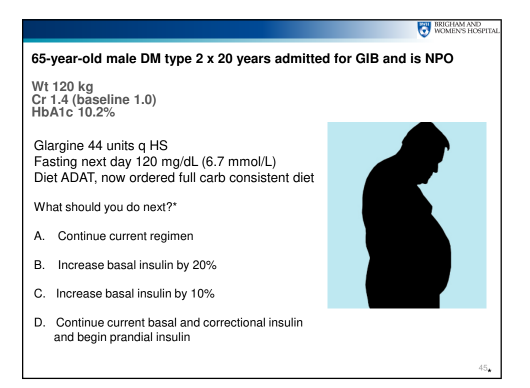


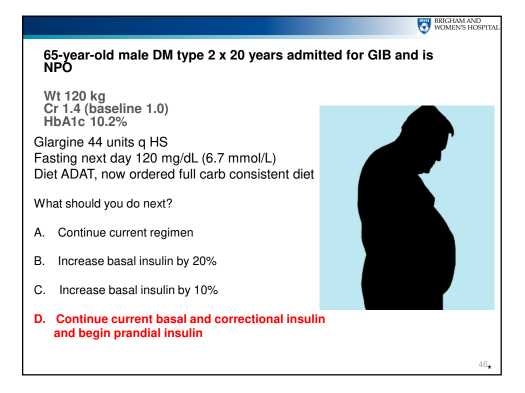
| | | WOMEN'S HOSPITAL |
|---|--|--|
| Estimating TDD Remember this is a place to start | Baseline weight-based TDD estimate | 0.5 units/kg/day, adjust by factors listed below |
| | Age > 70 years | -0.1 units/kg/day |
| Wt 120 kg Cr 1.6 (baseline 1.0) HbA1c 10.2% | Renal insufficiency (eGFR < 45) | -0.1 units/kg/day |
| NDATC 10.2% | 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 |

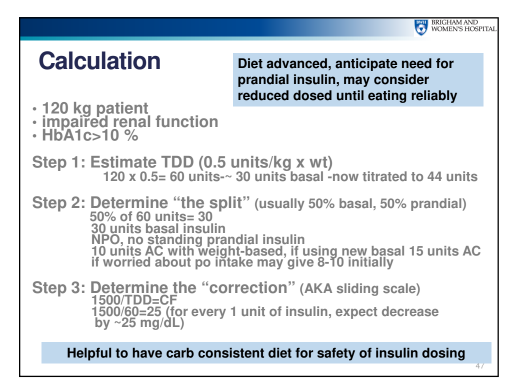
| | BRIGHAM AND WOMEN'S HOSPITAL |
|---|---------------------------------|
| Calculation | |
| 120 kg patient Impaired renal function HbA1c >10 % | |
| Step 1: Estimate TDD (0.5 units/kg x wt) 120 x 0.5= 60 units | |
| Step 2: Determine "the split" (usually 50% basal, 5 50% of 60 units= 30 30 units basal insulin NPO, no standing prandial insulin | 50% prandial) |
| Step 3: Determine the "correction" (AKA sliding = 1500/TDD=CF 1500/60=25 (for every 1 unit of insulin, expect of by ~25 mg/dL) | • |
| | 42 |

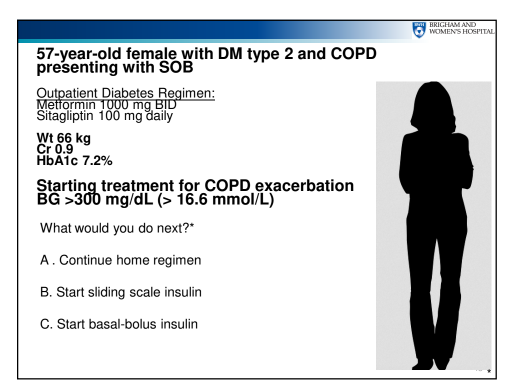


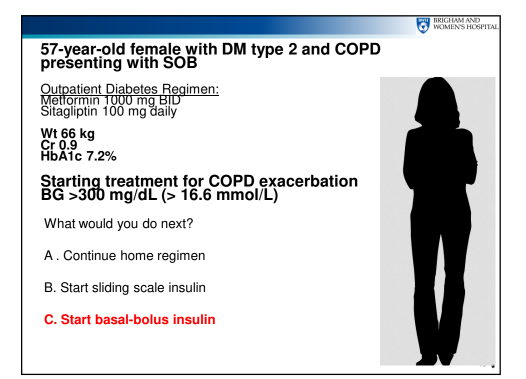


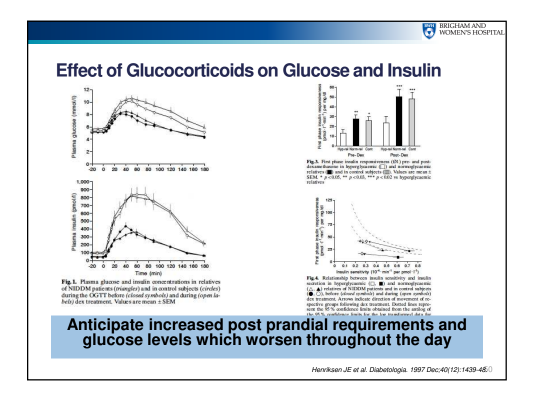


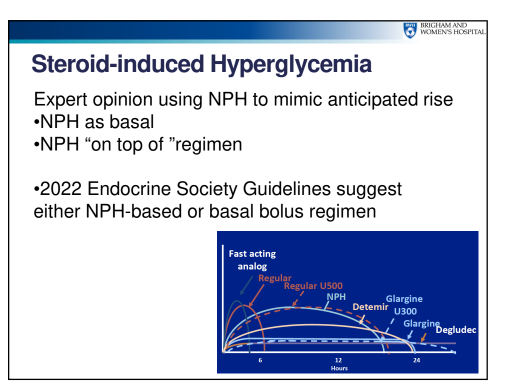








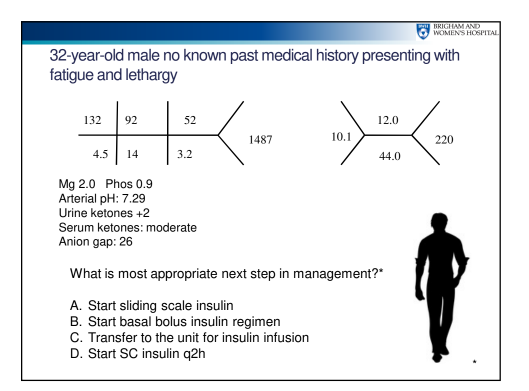


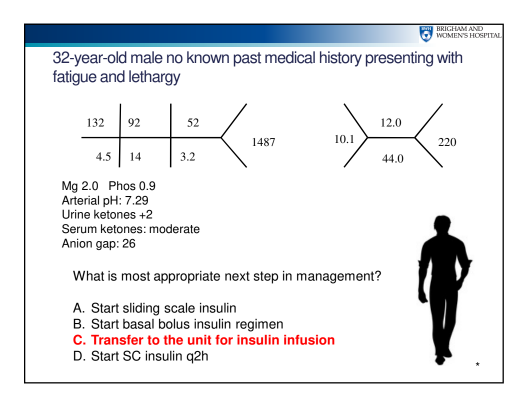


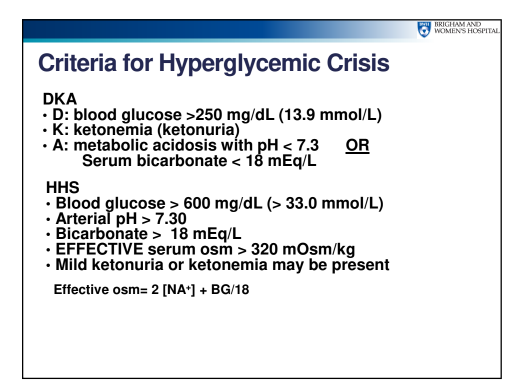
| Diabetes UK Position Statements Management of hyperglycaemia and sto (glucocorticoid) therapy: a guideline fro British Diabetes Societies (JBDS) for Inpa | m the Joint |
|--|--|
| A. Roberts ¹ , J. James ² and K. Dhatariya ³ , on behalf of the Join (JBDS) for Inpatient Care* | t British Diabetes Societies |
| "Cadiff and Vale University, cool Health Board, Cardiff, UK, "University HospitalsLeikenter NMG That, Leikenter, D NMG Roundation Truct, Normelch, UK Accepted 72 May 2018 | Card Transit Let Novel Conseq (Hights) |
| | |
| 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 |

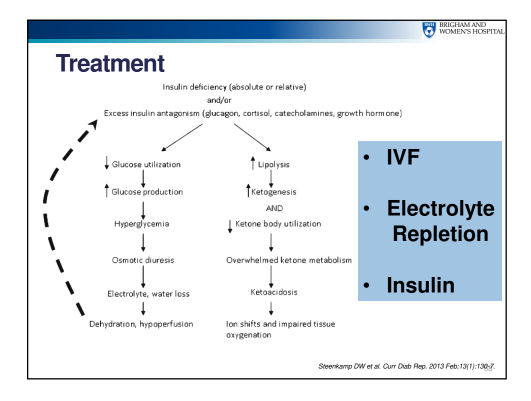
| | | WOMEN'S HOSPITAL |
|---|--|--|
| Estimating TDD Remember this is a place to start | Baseline weight-based TDD estimate | 0.5 units/kg/day, adjust by factors listed below |
| Wt 66 kg Cr 0.9 HbA1c 7.2% | Age > 70 years | -0.1 units/kg/day |
| | Renal insufficiency (eGFR < 45) | -0.1 units/kg/day |
| | 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 |

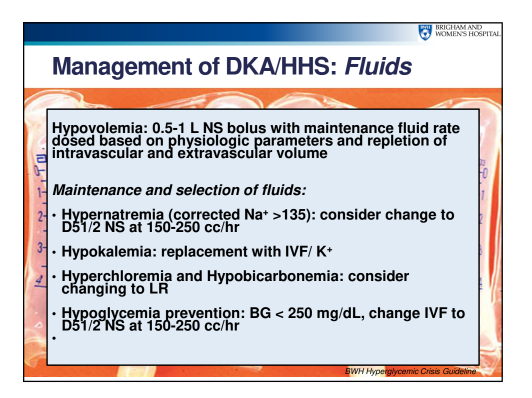
| | BRIGHAM AND WOMEN'S HOSPITA | | |
|--|--|--|--|
| Calculation 66 kg patient normal renal function HbA1c 7.2% | May use 50/50 or consider 40/60 split, using NPH and/or "stacked RAI" with steroids. Anticipate decreased requirements as steroids tapered | | |
| Step 1: Estimate TDD (0.6 units/kg x wt) 66 x 0.6= 40 units | | | |
| 50% of 40 units= 20 un 20 units basal insulin 20 units prandial insu | (if using NPH can split 10/10 or 13/7) | | |
| Step 3: Determine the "co 1500/TDD=CF 1500/40=38 (for every 1 by ~40 mg/dL) | rrection" (AKA sliding scale) unit of insulin, expect decrease | | |



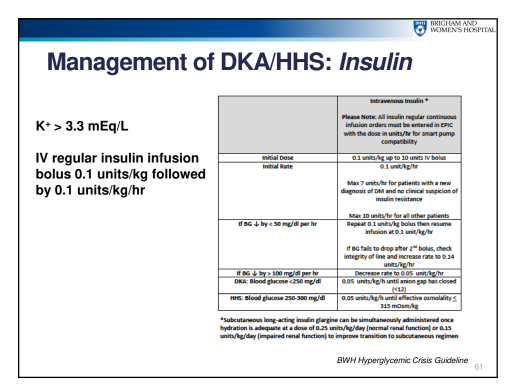




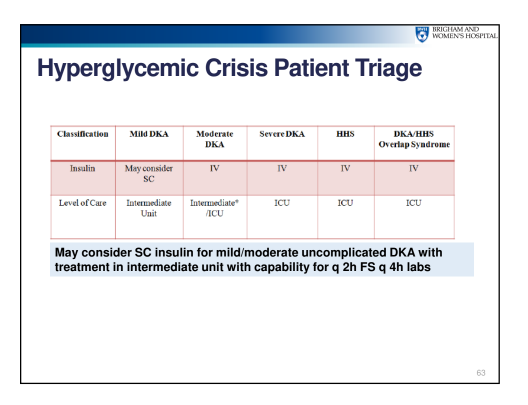


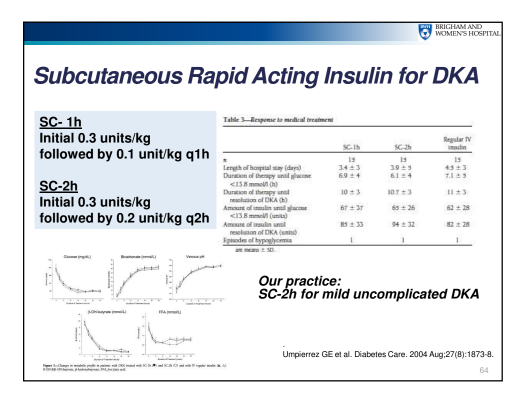


| K * if < 3.3 mEq/L | .: hold insulin and replete K* | |
|--|---|---|
| K+ is 3.3-5.3 mF | q/L: give 20-30 mEq in each lit | er of IVF |
| 10 13 0.0- 0.0 IIIL | | |
| | lo not give additional K+, repea | at K+2 hours |
| $K + \langle h \rangle$ $K + \langle h \rangle$ | | |
| K ⁺ >5.3 mEq/L: 0 | 3 | |
| - | administration see table below. Monitor K ⁺ q 4-6 h. For main | |
| assium Repletion: for initial KCL (* replacement scale. | administration see table below. Monitor K ⁺ q 4-6 h. For main | |
| assium Repletion: for initial KCL " replacement scale. Serum K+ (mEq/L) | administration see table below. Monitor K ⁺ q 4-6 h. For main Peripheral or Enteral | tenance dosing See EPIC Or |
| assium Repletion: for initial KCL (* replacement scale. Serum K+ (mEq/L) >5 ar/ urine output < 0.5 cc/kg/hr | administration see table below. Monitor K ⁺ q 4-6 h. For main Peripheral or Enteral None | itenance dosing See EPIC Of |
| assium Repletion: for initial KCL (* replacement scale. Serum K+ (mEq/L) >S or/ urine output < 0.5 cc/kg/hr 4-5 | administration see table below. Monitor K ⁺ q 4-6 h. For main Peripheral or Enteral None 10 mEq IV x 2 doses OR 20 mEq enterally | tenance dosing See EPIC Or |
| assium Repletion: for initial KCL (* replacement scale. Serum K+ (mEq/L) >S or/ urine output < 0.5 cc/kg/hr 4-5 3-4 | administration see table below. Monitor K ⁺ q 4-6 h. For main Peripheral or Enteral None 10 mEq IV x 2 doses OR 20 mEq enterally 10 mEq IV x 4 doses OR 40 mEq enterally | Central None 20 mEq IV 20 mEq IV x 2 doses |
| assium Repletion: for initial KCL (* replacement scale. Serum K+ (mEq/L) >S or/ urine output < 0.5 cc/kg/hr 4-5 | administration see table below. Monitor K ⁺ q 4-6 h. For main Peripheral or Enteral None 10 mEq IV x 2 doses OR 20 mEq enterally | Central None 20 mEq IV |



| | 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 |
| HCO, | 15-18 | 10 to < 15 | <10 | >18 |
| Urine/Serum Ketones | + | + | . + | +/- |
| Serum Osm (Osm_) | | | | >320 |
| AG | elevated | elevated | elevated | variable |
| Mental Status | alert | alert/drowsy | stupor/coma | stupor/com |



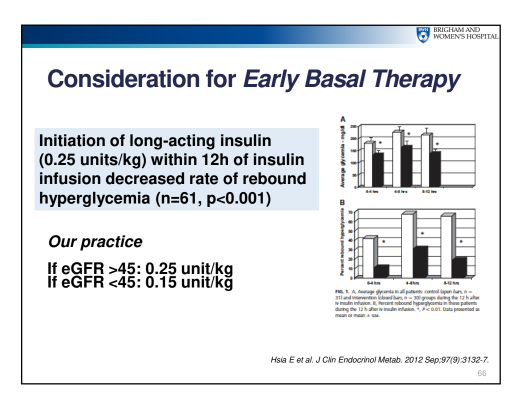


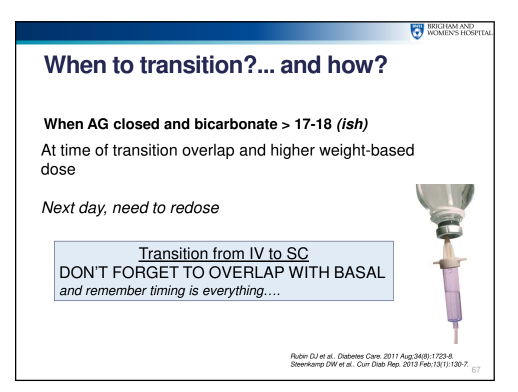


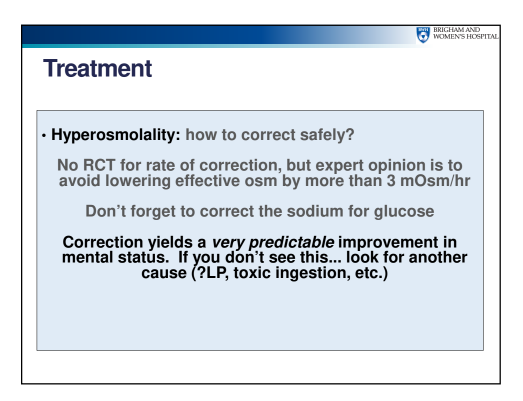
Example Protocol for Use of Subcutaneous Insulin Protocol in Treatment of Mild Uncomplicated DKA

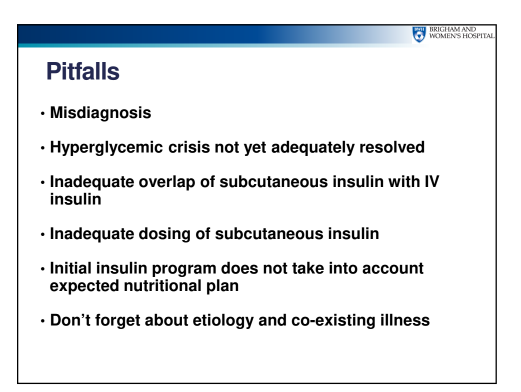
| | Subcutaneous Insulin Aspart | Subcutaneous Insulin Glargine |
|--------------------------|--|---|
| Initial Dose | 0.3 units/kg maximum 20 units | 0.25 units/kg if GFR >40; 0.15 units/kg if GFR <40 |
| Subsequent Dose | 0.2 units/kg every 2 hours maximum 10 units | Redose in 24 hours based on response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

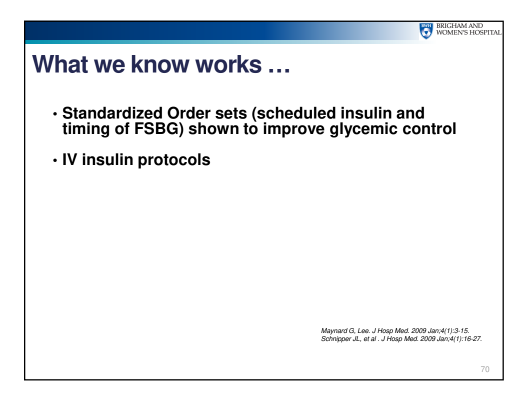
BWH Hyperglycemic Crisis Guideline

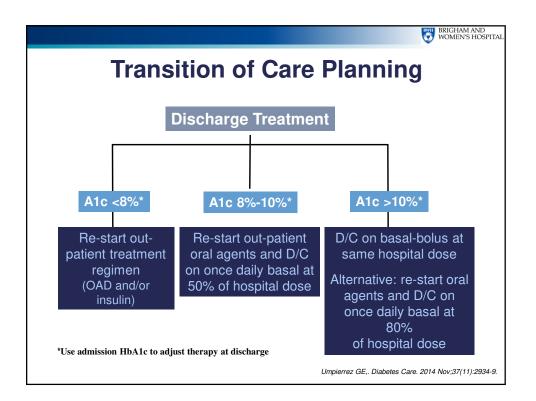




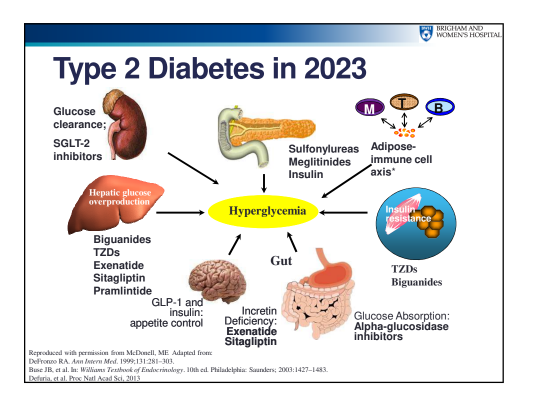


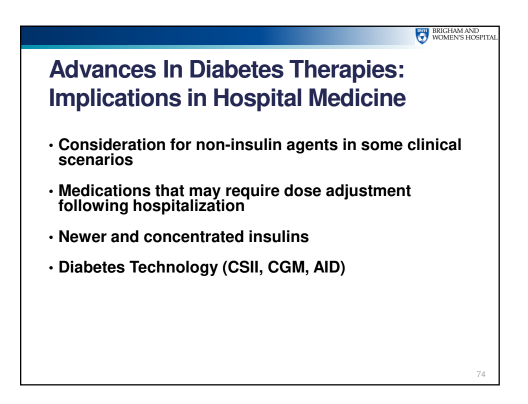






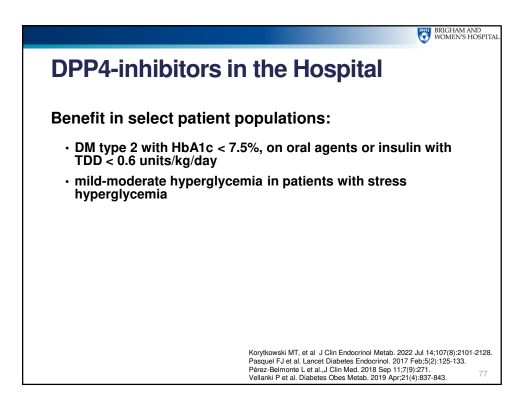
| BRIGHAM AND WOMEN'S HOSPITAL | | | | |
|--|--|--|--|--|
| Transition of Care Checklist | | | | |
| Diabetes Education ("survival skills") | | | | |
| Insulin Teaching (if applicable, should include pen and vial/syringe) | | | | |
| Glucometer Teaching | | | | |
| Confirm patient has diabetes supplies: | | | | |
| Medications* (if using insulin vial-syringe; if insulin pen-pen needles) Test strips (must match glucometer) Lancets | | | | |
| Clear communication with patient regarding discharge regimen* | | | | |
| Follow-up appointment scheduled | | | | |
| PCP aware of any dose adjustments | | | | |
| | | | | |
| *Medications and supplies will vary depending of insurance coverage- often human insulin cheaper than analogs; helpful to know coverage for pen vs. vial/syringe prior to discharg@2 | | | | |





| Metformin and | l Ris | k of | Acic | losis | S | | | |
|--|---|------------------|------------------|------------------|------------------|------------------|--|--|
| | | | | | | | | |
| Table 3. | | | | | | | | |
| Association of Time-Dependent Metformin Use With Aci Category in Geisinger Health System | dosis Hospitaliza | tion by Time-D | ependent Estim | ated Glomerula | r Filtration Rat | e (eGFR) | | |
| Lategory in Geisinger Health System | | | | | | | | |
| Parameter | HR ^a (95% Cl) 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 = 72 232) | 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) | | |

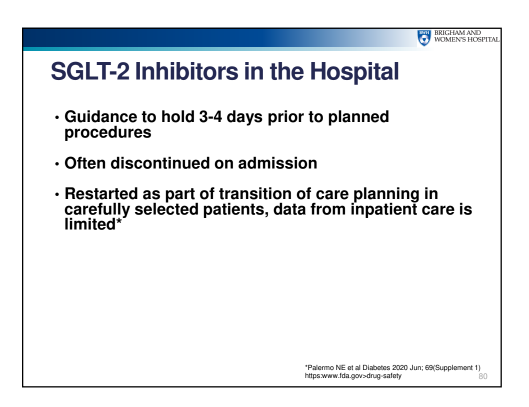
BRIGHAM AND WOMEN'S HOSPITAL Incretin-based therapy in hospitalized patients Lina-Real-World Study Basal-bolus vs basal-linagliptin Observational, multicenter 18 17 Non-critically ill patients with DM type 2 160 on oral agents (n=953) 150 DPP4i effective in patients with 3 mild-moderate hyperglycemia Minimizing injection burden 18 170 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

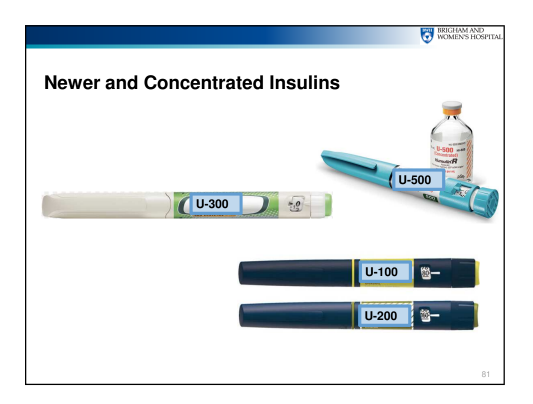


| | | | | BRIGHA WOMEN | M AND VS HOSPITAL |
|-----|----------------------|----------------|---------|-----------------|----------------------|
| Dos | se Adjustments | Based o | on Rena | al Funct | ion: |
| | agliptin agliptin | | | | |
| | GFR (ml/min) | <u>></u> 50 | 30-49 | <30 | |
| | Sitagliptin | 100 mg | 50 mg | 25 mg | |
| | Saxagliptin | 5 mg | 2.5 mg | 2.5 mg | |
| | Linagliptin | 5 mg | 5 mg | 5 mg | |
| | | | | | |
| | | | | | |

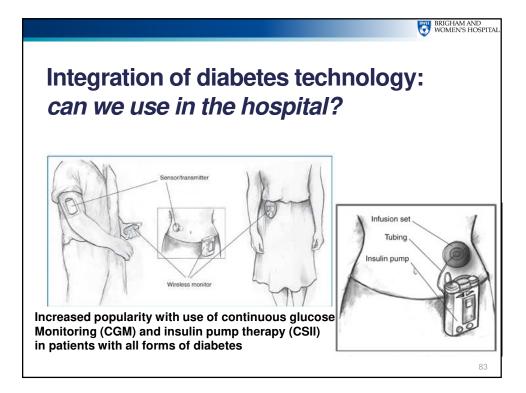
| Case patient | eristics of eul | DKA cases | 3 | | | | 5 | 6 | | 7 | | 8 | 9 |
|--|--------------------------|-------------------------|--------------|----------|------------|----------------------|-------------|------------|------------|------------|--------|------------|---------------------------------|
| uase patient Age (years) | 40 | 58 | 27 | | 2 | | 31 | 55 | | 25 | | 39 | 64 |
| eger (years) lex | Female | 30 Male | Fema | la. | Ferr | * | Female | Female | | Female | | Female | Female |
| 11/12 | T1 | T2 | 71 | ~ | T | | T1 | T1 | | T1 | | T1 | T2 |
| ADI/CSII | MDI | N/A | MD | | | | CSII | CSII | | CSII | | CSII | N/A |
| Juration (years) | 17 | 2 | 25 | | 6 | | 15 | 18 | | 13 | | 26 | 6 |
| MI (kg/m ²) | 26.5 | 26.5 | 24.3 | | 25 | 9 | 33.2 | 22.0 | | 22.0 | | 26.1 | 32.8 |
| rior A1C (% (mmol/mol)) | 11.4 (101.1) | 9.8 (83.6) | 7.8 (61 | .71 | 8.0 (8 | 53.9) | 7.0 (53.0) | 7.2 (55.2) | | 5.6 (48.6) | | 7.0 (53.0) | 7.8 (62.0) |
| anaglifiorin 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) |
| | 0.9 | 7.44 | 0.07 | | | | | | 1.45 | | | | |
| oo ₂ (mmHg) | 10 | | | | | | | | 26 | | | | 13 and then 5 |
| icarbonate (mEq/L) nion gap (mEq/L) | 6 25 | 10 | 6 35 | | 11 22 | 18 | | 15 26 | 21 | | | 24 | 15 and then 5 16 and then 19 |
| etones" | Yes (serum and urine) | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes (serum and urine) |
| Where treated | ICU | ICU | ICU | Outpt. | ICU | | Outpt | ICU. | ICU | Outpt. | Outot. | ICU | ICU |

BRICHAM AND

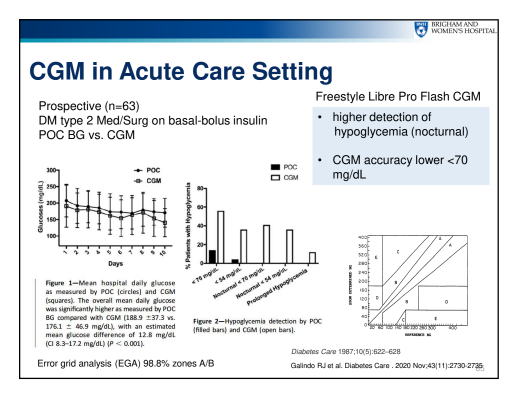




| | delivery of HUMULIN R U-500 using these devices Delivery Using a U-100 insulin syringe | Delivery Using a Tuberculin syringe |
|------------------------------------|---|--|
| HUMULIN R U-500 | Amount of HUMULIN R U-500 to draw up in the syringe in | |
| dose prescribed (units of insulin) | "unit 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 |
| | | U-souther U-souther U-souther Nonucir R |



| GM Use in | the Hospita | l: special | |
|--|--|---|-----------------------------|
| | - | | |
| onsideratio | ns | | |
| | | | |
| • | patient: imaging (MI | | |
| s of ongoing intere | st: surgery, pressors | , periods of rapid gl | ucose fluctuatic |
| act of certain medic | ations | | |
| | | | |
| Table 1 List of FDA-approved CGN CGM system | systems with features, limitations, and Key features | d interfering substances 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] | b). 1-h warm-up | preserve data b). No threshold or predictive alerts | Salicylic acid |
| | c). 14-day sensor wear d). Range 40–500 mg/dl | b). No threshold or predictive alerts | |
| Abbott Diabetes Care Freestyle Libre 2 [12] | a). No calibration required b). 1-h warm-up | a). Requires scanning every 8 h to preserve data | Ascorbic acid |
| Eloie 2 [12] | c). 14-day sensor wear | b). No predictive alarms | |
| | d). Range 40–400 mg/dl e). Optional alarms for hypoglyce- | c). Limited ability to transmit data | |
| | mia, hyperglycemia, and signal | | |
| Dexcom G6 [14] | a). No calibration required | a). 2-h warm-up | Hydroxyurea |
| | b). 10-day sensor wear c). 40–400 mg/dl range | | |
| | d). Predictive alerts for hypogly- | | |
| Medtronic MiniMed Guardian Sen- | a). 7-day sensor wear | a). 2-4 calibrations/day required | Acetaminophen |
| sor [15] | b). Predictive alerts c). Range 40–400 mg/dl | b). 2-h warm up c). 7-day sensor wear | |
| Senseonics Eversense [16] | a). 90-180 day sensor wear | a). Implantable | Mannitol, tetracycline |
| | b). Predictive hypo- and hyperglyce- mia alerts | b). 2 calibrations/day required c). 24-h warm-up | |
| | | | |



| | | RT-CO | - GM (De: | xcom | ı) vs PC | C |
|--|--|--|--|------------|------------|-------------------------------|
| Prospective RCT Non-critically ill patien Insulin treated DM typ | · / | for hypoglycer | nia | | | |
| Table 2–Glycemic outcomes | | | | 2 | | |
| | RT-CGM/GTS group ($n = 36$) | POC group ($n = 36$) | P value | | 1.69 | |
| | | | | 1.6 | | |
| lypoglycemic events/patient | | | | 1.0 | | |
| <70 mg/dL | 0.67 (0.34-1.30) | 1.69 (1.11-2.58) | 0.024 | | | |
| <70 mg/dL <54 mg/dL | 0.67 (0.34–1.30) 0.08 (0.03–0.26) | 1.69 (1.11-2.58) 0.75 (0.51-1.09) | 0.024 0.003 | 1.0 | | |
| <70 mg/dL <54 mg/dL locturnal hypoglycemic events/patient | 0.08 (0.03-0.26) | 0.75 (0.51-1.09) | 0.003 | | | 0.75 |
| <70 mg/dL <54 mg/dL | | | | | 0.67 | 0.75 |
| <70 mg/dL <54 mg/dL locturnal hypoglycemic events/patient <70 mg/dL | 0.08 (0.03–0.26) | 0.75 (0.51-1.09) | 0.003 | er Patient | 0.67 | 0.75 |
| <70 mg/dL <54 mg/dL locturnal hypoglycemic events/patient <70 mg/dL <54 mg/dL | 0.08 (0.03–0.26) 0.19 (0.09–0.41) 0.03 (0.01–0.24) | 0.75 (0.51–1.09) 0.33 (0.19–0.59) 0.11 (0.04–0.33) | 0.003 0.26 0.26 | er Patient | 0.67 | 0.75 |
| 70 mg/dt. 54 mg/dt. octurnal hypoglycemic events/patient 70 mg/dt. 54 mg/dt. ypoglycemic events (<70 mg/dt.)/patient/day | 0.08 (0.03-0.26) 0.19 (0.09-0.41) 0.03 (0.01-0.24) 0.12 (0.06-0.24) | 0.75 (0.51–1.09) 0.33 (0.19–0.59) 0.11 (0.04–0.33) 0.35 (0.23–0.54) | 0.003 0.26 0.26 0.011 | 1.2 | 0.67 | 0.75 |
| <pre><70 mg/dL <54 mg/dL coturnal hypoglycemic events/patient <70 mg/dL <54 mg/dL yogdycemic events (<70 mg/dL)/patient/day BR <70 mg/dL (%)</pre> | 0.08 (0.03-0.26) 0.19 (0.09-0.41) 0.03 (0.01-0.24) 0.12 (0.06-0.24) 0.40 (0.18-0.92) | 0.75 (0.51–1.09) 0.33 (0.19–0.59) 0.11 (0.04–0.33) 0.35 (0.23–0.54) 1.88 (1.26–2.81) | 0.003 0.26 0.26 0.011 0.002 | 1.2 | | 0.08 |
| | 0.08 (0.03-0.26) 0.19 (0.09-0.41) 0.03 (0.01-0.24) 0.42 (0.06-0.24) 0.40 (0.18-0.92) 0.05 (0.01-0.43) | 0.75 (0.51–1.09) 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) | 0.003 0.26 0.011 0.002 0.017 | 1.2 | < 70 mg/dL | 0.08 < 54 mg/dL |
| 70 mg/dL <54 mg/dL locturnal hypoglycemic events/patient <70 mg/dL <54 mg/dL <54 mg/dL MBR <70 mg/dL (%) BR <74 mg/dL (%) | 0.08 (0.03-0.26) 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) 5.91.2 (52.47-66.61) | 0.75 (0.51-1.09) 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) | 0.003 0.26 0.26 0.011 0.002 0.017 0.39 | 1.2 | | 0.08 |
| | 0.08 (0.03-0.26) 0.19 (0.09-0.41) 0.03 (0.01-0.24) 0.42 (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) | 0.75 (0.51-1.09) 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) | 0.003 0.26 0.26 0.011 0.002 0.017 0.39 0.94 | 1.2 | < 70 mg/dL | 0.08 < 54 mg/dL p=0.003 |

BRIGHAM AND WOMEN'S HOSPITAL

Glucose Monitoring in 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? |
| | | |

