

Bangladesh Endocrine Society INSULIN GUIDELINE



Bangladesh Endocrine Society (BES) Insulin Guideline

First Edition, 2018

All rights reserved by Bangladesh Endocrine Society (BES)

Published by
Bangladesh Endocrine Society (BES)
www.bes-org.net
E-mail: endobd2012@gmail.com

Bangladesh Endocrine Society (BES) Insulin Guideline

First Edition, 2018.

Published

02 November 2018, Dhaka, Bangladesh

BES Insulin Guideline Task Force

Convener: Dr Md Fariduddin

Member Secretary: Dr Shahjada Selim

Members:

Dr Tareen Ahmed

Dr FariaAfsana

Dr Nazmul Kabir Qureshi

Dr Ahmed Salam Mir

Dr Md Faruque Pathan

Dr Md Hafizur Rahman

All rights reserved by: Bangladesh Endocrine Society (BES)

Published by

Bangladesh Endocrine Society (BES)

www.bes-org.net

E-mail: endobd2012@gmail.com

Printing

Congressia

Contents

Section 1: Insulin is a life-saving hormone	7
Section 2: Insulin Delivery Devices, Technique and Storage	10
Section 3: Insulin Initiation and Intensification in DM	12
Section 4: Insulin Therapy at Hospital Settings	18
Section 5: Insulin Therapy in Hyperglycemic Crises	21
Section 6: Using Insulin During Ramadan Fasting	26
Section 7: Insulin Use in Pregnancy	30
Section 8: Insulin in Chronic Kidney Disease (CKD)	35
Section 9: Insulin in Chronic Liver Disease (CLD)	36
Section 10: Common barriers to insulin initiation and strategies to overcome them	37

Preface

Inception of creation of “**Bangladesh Endocrine Society (BES) Insulin Guideline**” was made by Executive Committee (2016-2018) of Bangladesh Endocrine Society (BES), on its 2nd EC Meeting on 03 March 2017, through a newly formed Scientific Sub-committee.

Subsequently, the Scientific Sub-committee of BES appointed **BES Insulin Guideline Task Force** to make the dream come true. The endeavor became evident on 13th BES EC meeting, held on 18th August 2018, where the first edition of “**Bangladesh Endocrine Society (BES) Insulin Guideline**” was submitted to EC.

The purpose of creating this manual is to serve clinicians regarding insulin uses in different clinical scenarios considering resource availability and circumstances prevailing in Bangladesh.

Heartfelt gratitude goes to all members of the **BES Insulin Guideline Task Force** who worked more than a year-long to bring the vision into daylight. We express thanks to the Scientific Sub-committee of BES [2016-2018] for their efforts and guidance all through its journey.

Dr Md Faruque Pathan
President
Bangladesh Endocrine Society

Dr Md Hafizur Rahman
General Secretary
Bangladesh Endocrine Society

Dr Md Fariduddin
Convener
BES Insulin Guideline Task Force

Bangladesh Endocrine Society (BES) Insulin Guideline
First Edition, 2018.

Abbreviations

BES	Bangladesh Endocrine Society
CV	Cardiovascular
CGM	Continuous Glucose Monitoring
CSII	Continuous Subcutaneous Insulin Infusion
DM	Diabetes Mellitus
DKA	Diabetic ketoacidosis
eGFR	Estimated Glomerular Filtration Rate
FPG	Fasting Plasma Glucose
GDM	Gestational Diabetes Mellitus
HHS	hyperosmolar hyperglycemic state
MNT	Medical Nutrition Therapy
OAD	Oral Antidiabetic Drugs
PPG	Post Prandial Plasma Glucose
SMBG	Self Monitoring of Blood Glucose
T1DM	Type 1 Diabetes Mellitus
T2DM	Type 2 Diabetes Mellitus
U	Units

Section 1: Insulin is a life-saving hormone.

Introduction

Key Points:

- *Insulin still remains the oldest and arguably the best treatment option for diabetes.*
- *Early initiation of insulin in the course of diabetes has many beneficial effects.*
- *Discovery of insulin in 1921 by Banting, Best, James Collip and Macleod changed the horizon of diabetes management.*
- *Currently available insulin preparations can be divided into two types- conventional insulin and insulin analogues.*
- *There are some distinct advantages of insulin analogues over the conventional insulins: mealtime flexibility, less absorption variability, peak-less profile of basal insulin, and less chance of early post-meal hyper and late post meal hypoglycemia by the rapid acting bolus insulins.*

Diabetes mellitus is a metabolic disorder characterized by chronic or persistent hyperglycemia, due to defect in insulin secretion and/or insulin action. In the pre-insulin era it was a rapidly fatal polyuric disorder; the discovery of insulin has transformed its natural history into a chronic and complex condition resulting in significant morbidity and mortality. In 2015 the estimated number of people with diabetes worldwide was 415 million, which is projected to become 642 million by the year 2040. About 5 million adults died from diabetes in 2015[1]. In Bangladesh, the overall age adjusted prevalence of diabetes was 7.4% in 2015[2].

Lifestyle modification (including medical nutrition therapy and physical exercise) and pharmacologic agents remain the cornerstones of management of diabetes. Currently we have a huge number of oral antidiabetic agents, starting with the sulfonylureas and biguanides since 1950s, followed by the alpha glucosidase inhibitors, thiazolidinediones, DPP-IV inhibitors and the most recently added SGLT-2 inhibitors. GLP-1 agonists and amylin analogues are also available as non-insulin injectable antidiabetic agents. However, insulin still remains the oldest (and arguably the best) treatment option for diabetes.

Many studies suggest that insulin initiation early in the course of diabetes has many beneficial effects, including preservation of β -cell function. It has been found that people who experienced the greatest improvements in β -cell function were able to maintain normoglycemia for longer with lifestyle management alone. Early intensive insulin therapy was observed to increase serum adiponectin and nitric oxide concentrations, decrease TNF- α levels and significantly improved endothelial injury/dysfunction, thereby exerting beneficial effects on the vasculature. Meta-analyses of several prospective studies investigating the effect of intensive glycemic control on CV outcomes suggest that intensive glycemic control reduces the risk of CV outcomes without increasing the risk of mortality. Therefore, it is beneficial for people with type 2 diabetes having high HbA1c levels to start insulin early for normalization of their glycemic status, after which they can be moved onto standard care[3].

Discovery of insulin in 1921 by Banting, Best, Collip and Macleod changed the horizon of diabetes management. The frequency of acute complications greatly diminished; but due to the short duration of action it was difficult to achieve good glycemic control round the clock. The picture improved with the discovery of Protamine insulin in 1936 by Hans Christian Hagedorn, followed by NPH insulin in 1946 and Lente insulins in 1952. The commercially available insulins were mostly bovine or porcine in origin till 1980, when rDNA origin human insulin became available. Currently available insulin preparations can be divided into two types- conventional insulin (having the same amino acid sequence as human insulin) and insulin analogues (where a specific change has been made in the amino acid sequence to favorably alter its pharmacokinetics). Conventional insulins may be short acting or intermediate acting according to their duration of action; for the insulin analogues they are rapid acting or long acting[4]. Insulin analogues mimic the physiologic insulin secretion pattern (i.e. maintenance of a peak-less basal insulin level throughout the day, and peaks of bolus insulin during mealtime that perfectly match the post meal glucose excursions).

Table-1: Types of insulin [5].

Insulin Type	Onset	Peak	Duration	Appearance
Bolus (prandial) insulins				
Rapid-acting insulin analogues				
• Insulin Aspart	10-15 min	1-1.5 hour	3-5 hours	Clear
• Insulin Glulisine	10-15 min	1-1.5 hour	3-5 hours	Clear
• Insulin Lispro	10-15 min	1-2 hours	3.5-4.75 hours	Clear
Short-acting (Regular) insulins	30 min	2-3 hours	6.5 hours	Clear
Basal insulins				
Intermediate-acting (NPH)	1-3 hours	5-8 hours	Upto 18 hours	Cloudy
Long-acting insulin analogues				
• Insulin Detemir	90 min	N/A	24 hours	Clear
• Insulin Glargine	90 min	N/A	24 hours	Clear
• Insulin Degludec			42 hours	Clear

There are some distinct advantages of insulin analogues over the conventional insulins, like greater meal-time flexibility, less absorption variability, peak-less profile of basal insulin minimizing the risk of hypoglycemia, and less chance of early post-meal hyper and late post meal hypoglycemia by the rapid acting bolus insulins. Indeed, the insulin analogues mimic the normal physiology most precisely [6].

There are different regimes for insulin administration (Table-2), which are applicable for subcutaneous insulin administration only. In some specific situations, intravenous administration of regular short acting or rapid acting analogue insulin is required; similar is also applicable for during insulin pump use.

Table-2: Different types of Insulin Regimen

Regimen	Description
Once daily	NPH or Basal analogue at bedtime
Twice daily	
• Premixed	Less mealtime flexibility
• Co-formulation	
• Split-mixed	
Multiple daily injections	Offers more mealtime flexibility
• Basal plus	One long acting analogue at bedtime, plus one injection of rapid acting analogue with the largest meal
• Basal bolus	One long acting analogue at bedtime, plus two or three injections of rapid acting analogue with meal
Continuous subcutaneous insulin infusion	Insulin pump

Table-3: Comparison of different insulin regimens.

Types of Insulin Regime	Compliance	Cost	Self management education	Monitoring	Need for Care giver facilities	Chance of hypoglycemia	Flexibility
Conventional							
Premixed	(+)	Average	Easy	(++)	(+++)	(+++)	(-)
Split-mixed	(-)	Average	Complicated	(++)	(++)	(+)	(+)
Analogue							
Premixed	(+)	High	Easy	(+)	(++)	(++)	(-)
Co-formulations	(+)	High	Easy	(+)	(+)	(+)	(+)
Basal only	(++)	High	Easy	(+)	(+)	(-)	(+)
Basal-bolus	(-)	High	Easy	(++)	(+)	(-)	(++)

The selection of type and regime of insulin is individualized depending upon particular clinical scenario (type of diabetes, glycemic status, hypoglycemia, glycemic variability, presence of complications and comorbidities) and patient's perspective (compliance, diabetes self-management education, socioeconomic condition, availability). It is the responsibility of the physician to consider all the relevant factors before choosing the appropriate regime for a particular individual. In this context, we can recapitulate the ever memorable quote by Elliott Joslin, "Insulin is a remedy primarily for the wise and not for the foolish. Everyone knows it requires brains to live long with diabetes, but to use insulin successfully requires more than brains" [8].

References:

1. IDF diabetes atlas 2015. Available at www.diabetesatlas.org/resources/2015-atlas.html. Accessed on May 17, 2017.
2. IDF diabetes atlas summary table 2015. Available at http://diabetesatlas.org/index.php?option=com_attachments&task=download&id=106:2020-11-05_Summary_table. Accessed on May 17, 2017.
3. Hanefeld M. Use of insulin in type 2 diabetes: What we learned from recent clinical trials on the benefits of early insulin initiation. *Diabetes & Metabolism* 2014; 40(6): 391-399.
4. Gough S, Narendran P. Insulin and Insulin Treatment. In: Holt RIG, Cockram C, Flyvbjerg A, Goldstein BJ (eds). *Textbook of Diabetes*. Oxford: Wiley-Blackwell, 2010: 427-429.
5. Canadian Diabetes Association Clinical Practice Guidelines Expert Committee. *Clinical Practice Guidelines: Pharmacotherapy in Type 1 Diabetes*. *Can J Diabetes* 2013; 37: S56-S60.
6. Rolla A. Pharmacokinetic and Pharmacodynamic Advantages of Insulin Analogues and Premixed Insulin Analogues Over Human Insulins: Impact on Efficacy and Safety. *The American Journal of Medicine* 2008; 121: S9 –S19.
7. Donner T. Insulin – Pharmacology, Therapeutic Regimens and Principles of Intensive Insulin Therapy. [Updated 2015 Oct 12]. In: De Groot LJ, Chrousos G, Dungan K, et al., editors. *Endotext* [Internet]. South Dartmouth (MA): MDText.com, Inc.; 2000-. Available from: <https://www.ncbi.nlm.nih.gov/books/NBK278938>. Accessed on May 28, 2017.
8. Joslin EP. Insulin, Section II. In: Joslin EP. *The Treatment of Diabetes Mellitus*. Philadelphia: Lea &Febiger, 1928: 69.

Section 2:

Insulin Delivery Devices, Technique and Storage

Key Points:

- *Plastic fixed-needle syringes designed for single use are available in different sizes (U100, U40).*
- *Pen injector devices have been designed to make injections easier and more flexible.*
- *An insulin pump for CSII is an alternative to treatment with MDI. Sensor-augmented pumps are provided with CGM system.*
- *Insulin injections are usually given into the deep subcutaneous tissue at 45-90° angle by two-finger pinch of skin. A wait of 15 seconds after pushing in the plunger helps to ensure complete expulsion of insulin.*
- *Sites of insulin injections are: Abdomen (the preferred site for faster and uniform absorption and less affected by muscle activity or exercise), Front/lateral thigh, Lateral aspect of arm and lateral upper quadrant of the buttocks.*
- *Insulin must never be frozen, direct sunlight or warming should be avoided.*
- *Insulin should not be used if there is change in appearance. Unused insulin should be stored in a refrigerator (4-8°C).*
- *When in use the insulin may be kept in room temperature but it retains its potency much better if kept in refrigerator.*
- *In hot climates where refrigeration is not available, cooling jars, earthenware pitcher or a cool wet cloth around the insulin will help to preserve insulin activity*

1. Insulin delivery devices

1.1.: Insulin syringe

Plastic fixed-needle syringes are available in different sizes (U100, U40). These syringes are designed for single use. However, many individuals reuse them without significant risk of infection [1]. Patient can change it every 3-4 days interval if adequate hygiene can be maintained.

1.2.: Pen injector device

Pen injector devices containing insulin in pre-filled cartridges have been designed to make injections easier and more flexible. The dose is dialed up and then pushed. These are available in reusable (cartridges can be replaced when required) or one-time use devices. Needle to be changed 3-4 days intervals or when tip becomes blunt.

1.3.: Continuous subcutaneous insulin infusion (CSII)

An insulin pump is an alternative to treatment with MDI. The pump is worn at waist or other convenient places, and insulin is delivered through tube into subcutaneous needle placed over abdomen. Basal insulin is delivered continuously, and the bolus dose is person-activated. Rapid acting insulin analogues are usually used in these devices, both as basal as well as bolus.

Sensor-augmented pumps are provided with CGM system. Newer generation pumps automatically shut-off to prevent hypoglycemia when the sensor has fallen below a preset threshold. The newer smart pumps can automatically calculate meal or correction boluses based on insulin-to-carbohydrate ratios and insulin sensitivity factors.

2. Injection technique

2.1.: Injections are given into the deep subcutaneous tissue at 45-90° angle by two-finger pinch of skin. The pinch is recommended to ensure a strict subcutaneous injection, avoiding intramuscular injection. Injections can be given perpendicularly without lifting a skin fold when needles are smaller and there is enough

subcutaneous fat. The needles should be inserted fully, otherwise there is a risk of intradermal injections. A wait of 15 seconds after pushing in the plunger helps to ensure complete expulsion of insulin through the needle, especially in pens [2]. Cleaning or disinfection of skin is advisable, but may not necessary unless hygiene is a real problem [3,4].

2.2.: Vials (also the pen devices) of cloudy insulin must always be gently rolled (not shaken) 10-20 times, to mix the insulin suspension. When a mixture of two insulins is drawn up (e.g. regular mixed with NPH), the regular insulin is to be drawn up into the syringe before the intermediate acting one. The mixture must be administered immediately.

3. Injection sites [5,6]:

3.1.: Abdomen (the preferred site when faster and uniform absorption is required and it may be less affected by muscle activity or exercise).

3.2.: Front/lateral thigh (the preferred site for slower absorption of longer acting insulin).

3.3.: Lateral aspect of arm (assistance is required for injection).

3.4.: The lateral upper quadrant of the buttocks (used less often).

3.5.: Rotation of injection sites are important within the same area of injection.

4. Storage of insulin

4.1.: Insulin must never be frozen.

4.2.: Direct sunlight or warming (e.g. in hot climates) damages insulin.

4.3.: Insulin should not be used if there is change in appearance (clumping, frosting, precipitation, or discoloration).

4.4.: Unused insulin should be stored in a refrigerator (4-8 °C) to retain its potency up to expiry date.

4.5.: When in use the insulin may be kept in room temperature (if not too hot) without much loss of efficacy. But it retains its potency much better if kept in refrigerator.

4.6.: In hot climates where refrigeration is not available, cooling jars, earthenware pitcher [7] or a cool wet cloth around the insulin will help to preserve insulin activity.

4.7.: After first usage, an insulin vial should be discarded after a certain period as instructed by manufacturer.

References

1. Schuler G, Pelz K, Kerp L. Is the reuse of needles for insulin injection systems associated with a higher risk of cutaneous complications? *Diabetes Res Clin Pract* 1992;16:209–212.
2. Ginsberg BH, Parkes JL, Sparacino C. The kinetics of insulin administration by insulin pens. *Horm-Metab Res* 1994;26:584–587.
3. Fleming DR, Jacober SJ, Vandenberg MA, et al. The safety of injecting insulin through clothing. *Diabetes Care* 1997;20:244.
4. McCarthy JA, Covarrubias B, Sink P. Is the traditional alcohol wipe necessary before an insulin injection? Dogma disputed. *Diabetes Care* 1993;16:402.
5. Koivisto VA, Felig P. Alterations in insulin absorption and in blood glucose control associated with varying insulin injection sites in diabetic patients. *Ann Intern Med* 1980; 92:59.
6. Sindelka G, Heinemann L, Berger M, et al. Effect of insulin concentration, subcutaneous fat thickness and skin temperature on subcutaneous insulin absorption in healthy subjects. *Diabetologia* 1994; 37:377.
7. Rangawala S, Shah P, Hussain S, Goenka S, Pillai K. Insulin stored in matka (earthen pitcher) with water for 60 days does not reduce in bio-activity. *J Pediatr Endocrinol Metab* 1997;10 (Suppl. 2):Abstract 347.

Section 3:

Insulin Initiation and Intensification in DM

Key points:

- *Insulin should be initiated in T2DM patients with $HbA1c \geq 10\%$, symptomatic hyperglycemia, $PPG > 19.4$ mmol/L, $FPG > 16.6$ mmol/L, If the glycemic target is not achieved by using three non-insulin agents by at least 3 months, during acute illness, surgery, stress, emergencies, pregnancy and lactation, as initial therapy in T2DM with severe hyperglycemia, severe metabolic decompensation (eg. DKA, HHS), T1DM.*
- *Insulin regimens are: Basal alone, Basal plus, basal bolus, split-mixed, pre-mixed, co-formulation, combination with injectible agents.*
- *Initiation can be done by basal insulin with the dose of 10 units or 0.1-0.2 U/kgbw. While with split-mixed or pre-mixed regimen, insulin may be initiated at dose of 0.2-0.3 U/kgbw. Intensification should be done with increase of 10-20% of dose or 2-4 units of insulin as per SMBG records once or twice weekly until glycemic targets are achieved.*
- *Switching should be done from one regimen to other one when appropriate.*
- *Basal bolus regimen (MDI) is preferred than pre-mixed regimen in T1DM.*
- *In children and adolescent MDI, CSII or 1-3 dose of bolus insulin regimen can be chosen.*

A.: Insulin Initiation and Intensification in Type 2 DM

1.: Indication of using insulin in T2DM: [1]

Insulin should be initiated in Type 2 DM in following conditions:

- *$HbA1c \geq 10\%$*
- *Symptomatic hyperglycemia*
- *$PPG > 19.4$ mmol/L, $FPG > 16.6$ mmol/L*
- *If the glycemic target is not achieved by using three non-insulin agents (metformin/pioglitazone, secretagogue, α Gi/DPP4i/SGLT2i) by at least 3 months*
- *In some specific situations: Short term use of insulin therapy in patients with T2DM may also be considered in the following conditions: acute illness, surgery, stress and emergencies, pregnancy and lactation, as initial therapy in T2DM with severe hyperglycemia*
- *Severe metabolic decompensation (eg. DKA, HHS)*

2.: Insulin Therapy

Many patients with type 2 diabetes eventually require and benefit from insulin therapy. The progressive nature of type 2 diabetes should be explained to patients frequently. Equipping patients with an algorithm for self-titration of insulin doses based on self-monitoring of blood glucose (SMBG) improves glycemic control in patients with type 2 diabetes. Comprehensive education regarding SMBG, diet, and the avoidance of and appropriate treatment of hypoglycemia are critically important in any patient using insulin[1].

2.1.: Basal Insulin

2.1.a.: Initiation

Basal insulin is preferably better to start if A1C is $> 9.0\%$ in newly detected diabetes or in an old patient of type 2 diabetes. Basal insulin alone is the most convenient initial insulin regimen, beginning at 10 units per day or 0.1–0.2 units/kg/day, depending on the degree of hyperglycemia. Basal insulin is usually prescribed in conjunction with metformin and sometimes one additional noninsulin agent[1].

2.1.b.:Adjustment

Adjust basal insulin by 10-15% or 2-4 U once-twice weekly to reach FBG target (at least 3 days apart). In cases hypoglycemia (symptomatic or absolute): Determine and address cause (s); reduce dose by 4 U or 10-20% of the current basal insulin[1].

2.1.c.:Follow up

After 3 month A1C should be done to evaluate the attainment of glycemic target [1].

2.2.: Basal Plus Insulin

2.2.a.:Initiation

When FBG target is reached or if the dose of basal insulin > 0.5 U/kg/day treat the PPG excursion with mealtime insulin. The ideal meal time insulin is the rapid acting one (aspart, lispro and glulisine). In Bangladesh, the largest meal is usually lunch. So, start: 4 U [or 0.1 U/kg or 10% basal dose] before lunch[1]. The short acting human insulin might be used if rapid acting insulins are not available (with increased chances of hypoglycemia).

2.2.b.:Adjustment:

The premeal insulin (rapid acting analogue/ short acting human insulin) dose will be increased by 1-2 U or 10-15% once –twice weekly until SMBG targets are reached.

2.2.c.:Follow up:

If hypoglycemia occurs, determine and address cause(s) and if there is no clear reason(s) causing hypoglycemia, the corresponding dose should be reduced by 2-4 U or 10-20%[1].

2.3.:Basal Bolus Insulin

Many individuals with type 2 diabetes may require mealtime bolus insulin (before 2 or 3 major meals) in addition to basal insulin. Rapidacting analogs are preferred due to their prompt onset of action. If the current A1C of the patient (newly detected patient or already on OADs/ other drugs) is $> 10\%$, basal bolus insulin should be initiated[1].

2.3.a.:Initiation

The recommended starting dose of mealtime insulin is 4 units or 0.1 U/kg, or 10% of the basal dose in patients who are on basal insulin. When starting mealtime bolus insulin, consideration should be given to decreasing the basal insulin dose.

In insulin naïve patients, the starting dose of basal insulin and bolus insulin typically be 50: 50. The daily dose may be 0.1 to 0.2 U/Kg of body weight. The $\frac{1}{2}$ of the total daily dose should be allocated at fixed time, preferably at night as basal insulin. The premeal bolus insulin will be divided in to 3 nearly equal doses and be taken before each major meals.

2.3.b.:Adjustment:

Increase dose(s) by 1-2 units or 10-15% once or twice weekly to achieve SMBG target.

Two types of insulin should not be titrated at a time. Firstly FPG may be targeted then PPG.

2.4.: Split Mixed Insulin

Two times NPH and 2 or 3 times premeal short acting insulin may be given as an alternative to basal bolus/ premixed/co-formulation insulin. This regimen may be suitable when frequent titration is intended such as while switching from intravenous insulin to subcutaneous insulin (in DKA, HHS and other parenteral feeding situations). In split mixed regimen, $\frac{2}{3}$ of dose comprises intermediate acting and $\frac{1}{3}$ comprises as short- acting insulin.

2.4.a.:Initiation:

In insulin naïve patients, the starting dose of NPH and short acting insulin typically be 2/3: 1/3. The daily dose may be 0.2 to 0.3 U/Kg of body weight. The 2/3of the total daily dose [which may be for a 70 kg adult patient of type 2 diabetes ($70 \times 0.3 = 21$, 2/3 of that = 14 U)] should be allocated as NPH; 2/3rd of the NPH dose should taken 30 minutes before breakfast and 1/3rd 30 minutes before dinner. The premeal insulin will be divided in to 2 or 3 nearly equal doses and be taken before each major meals [for the above mentioned patient it may 7U+ 7U+7U] [2,3].

2.4.b.:Adjustment:

Dose tritartion will be done by monitoring the SMBG.

2.5.: Premixed Insulin

Premixed insulin products contain both a basal and prandial component, allowing coverage of both basal and prandial needs with a single injection. NPH/regular 70/30 insulin, for example, is composed of 70% NPH insulin and 30% regular insulin. These may also contain 75/25, 50/50 and other ratios of NPH/regular insulins.

There are analog premixed insulins containing different proportions of intermediate acting and rapid acting insulin e.g.; 70/30 aspart mix, 75/25 or 50/50 lispro mix).Co-formulation insulin analogue insulin contains 70/30 ratio of degludec/aspart. Each approach has its advantages and disadvantages.

2.5.a.: Initiation and Titration of Premix insulin

Premix insulin can be started once daily with 10 U either in the morning (AM), if nighttime / pre-dinner glucose is high or in the night (PM), if the morning glucose (FBG) is high. If the total insulin dose exceeds 20 U, then premix insulin can be given twice daily, before breakfast and before dinner (AM & PM), distributed as two third in morning and one third in evening.

Premix analogues should be distributed in two equal halves in morning and evening when single dose exceeds 30 units. Also premix insulin may be started twice daily in case of patients with higher HbA1c, or if blood glucose FBG and PPG both are suboptimal.

The usual initiating dose is about 10 units of premix insulin either morning (AM) or night (PM). The blood sugar levels need to be monitored every week and the dose titrated. The simple principle of “starting low and scaling slow” must be applied. Second dose to be given ifdaily insulin requirement > 20 units orpre dinner blood glucose persists > 150 mg/dl (8.3 mmol/L).

2.5.b.: Monitoring procedure while on Premix insulin

Premix insulin can be given once daily before breakfast or dinner. Then it can be intensified to twice or thrice daily. Dinner (PM) or morning (AM) dose needs to be titrated based upon prebreakfast or predinner blood glucose respectively. Titration should be done at regular interval (at least weekly) until glyce-mic goals are achieved.

2.5.c.: Adjustment:

Suggested titration is 1 to 2 units added to pre-breakfast dose and/or pre-dinner dose every 3 days interval until target BG values are reached.Prebreakfast premixed insulin achieves pre-dinner target BG value.Pre-dinner premixed insulin achieves target fasting BG value.

Premixed conventional human insulin should be given 30 minutes before meals.Biphasic analogue and coformulation insulin should be given immediately before meal.

If BG targets are not reached, continue to increase the relevant dose until both targets achieved.The individual may need to self-monitor BG 2 times a day to safely titrate insulin.

2.6.: Combination Injectable Therapy

If basal insulin has been titrated to an acceptable fasting blood glucose level (or if the dose is >0.5 units/kg/day) and A1C remains above target, consider advancing to combination injectable therapy [Figure 1].

When initiating combination injectable therapy, metformin therapy should be maintained while SU may be discontinued on an individual basis to avoid unnecessarily complex or costly regimens (i.e., adding a fourth antihyperglycemic agent). In general, GLP-1 receptor agonists should not be discontinued with the initiation of basal insulin. Sulfonylureas, DPP4 are typically stopped once more complex insulin regimens beyond basal are used. In patients with suboptimal blood glucose control, especially those requiring large insulin doses, adjunctive use of a DPP4i, thiazolidinedione or SGLT2 inhibitor may help to improve control and reduce the amount of insulin needed, though potential side effects should be considered. Once an insulin regimen is initiated, dose titration is important with adjustments made in both mealtime and basal insulins based on the blood glucose levels and an understanding of the pharmacodynamic profile of each formulation (pattern control) [4,5].

2.7.: Switching Between Insulin Regimen

2.7.a.: Regular human insulin and human NPH/Regular premixed formulations (70/30) are less costly alternatives to rapid-acting insulin analogues and premixed insulin analogs, respectively. But their pharmacodynamic profiles may make them less optimal and frequent hypoglycemic events may hamper treatment adherence.

2.7.b.: Consider switching patients from one regimen to another (i.e., premixed analog insulin three times daily to basal-bolus regimen or vice-versa) if A1C targets are not being met and/or depending on other patient considerations (willing to reduce injection frequency, care giver dependency, when frequent monitoring of BG not possible).

2.7.c.: Self-monitoring of blood glucose (SMBG) can be helpful in determining appropriate targets for therapy, when to initiate insulin therapy, and the most useful insulin regimen for a particular patient. It is useful to monitor responses to therapy and to identify and treat glycemic excursions above or below target levels. Clinical experience suggests that SMBG is an important component of effective therapy.

2.7.d.: Switching from Basal to Premixed regimen

For those patients on combination OADs and basal insulin not achieving HbA1c targets despite optimal FBG, with post-prandial hyperglycemia, another option for intensification would be to switch to a premixed regimen [6,7].

This option is usually appropriate for patients who prefer a simpler regimen and are unable to accept 3–4 injections per day. This regimen is more suitable for those with a rigid lifestyle. Sulphonylureas should be stopped but Metformin should be continued.

Dose for dose transfer can be used where the total daily dose of basal insulin is used to determine total daily dose (TDD) of premixed insulin. TDD is then administered in two divided doses, usually equal in amount i.e. split dose 50:50 at pre-breakfast and pre-dinner [6].

2.7.e.: Switch to premixed twice daily

- *Total dose transfer should be done.*
- *Split dose 50:50 ratio [pre-breakfast : pre-dinner]*
- *Titrate dose once or twice a week to reach pre-breakfast or pre-dinner BG targets.*
- *Stop SU, continue metformin*
- *Consider premixed analogue insulin.*

Premixed analogues may be considered in patients experiencing hypoglycaemia with conventional premixed insulin and in those who desire greater flexibility as administration of premixed analogue does not require specific timing prior to meals and may be injected just prior to, during, or immediately after a meal.

2.7.f.: Switching from Premixed to Basal bolus regimen:

For those patients on premixed regimen (twice or three times daily) and not achieving HbA1c targets despite optimised dose, another option for intensification would be to switch to basal-bolus regimen.

This option is appropriate for patients who require greater flexibility in dose adjustment as it potentially allows pre-meal rapid / short-acting insulin to be adjusted individually according to blood glucose level (correctional bolus) along with the carbohydrate meal content of the meal.

The initial total daily dose following the switch may be guided by using a simple dose calculation of 0.5units/kg or by a total dose for dose transfer from the prior total daily dose on the previous regimen. Following determination of total daily- dose requirement, proportion of basal to prandial insulin requirement may be estimated using a ratio of 50:50. A smaller proportion of basal insulin may also be used such as between 25 – 40% of total daily dose in certain circumstances. The basal dose is usually administered at bedtime (conventional insulin or analogue) and the prandial portion is divided into three to cover the three main meals. Estimation of the pre-meal dose should take into consideration the size of the meal, in terms of the carbohydrate content. Subsequently the basal and pre-meal insulin should be titrated or optimised accordingly towards attaining glycemic targets[6].

2.7.g.: Switching from single to multiple premixed regimen

For those patients already on a single premixed insulin regimen, usually in combination with single or multiple OADs and not achieving blood glucose and HbA1c targets despite optimising insulin and OAD doses, an option for intensification would be to initiate additional pre-meal doses of premixed insulin.

For those on single dose conventional premixed insulin, usually prior to evening meals, one additional dose may be initiated prior to the morning meal. In those receiving premixed analogue insulin, additional doses may be initiated at both morning and midday meals, either sequentially or simultaneously. It is not usual to administer conventional premixed insulin more than twice daily in view of concern for between-meal hypoglycaemia [8-11].

B.: Insulin Initiation and Intensification in Type 1 Diabetes

3.: Adults with type 1 diabetes

3.1.: Offer basalbolus insulin regimens (MDI), rather than twicedaily mixed insulin regimens, as the insulin injection regimen of choice for all adults with type 1 diabetes[11].

3.2.: Long acting insulin

Twicedaily insulin detemir, glargine or once daily degludec may be considered as basal insulin therapy for adults with type 1 diabetes. Conventional insulin regimen may be an alternative either once or twice daily.

3.3.: Rapid acting insulin

Offer rapidacting insulin analogues injected before meals, rather than rapidacting soluble human or animalinsulins, for mealtime insulin replacement for adults with type 1 diabetes. Do not advise routine use of rapidacting insulin analogues after meals for adults with type 1 diabetes.

3.4.: Split Mixed Insulin

Thricedaily human insulin regimen (two times NPH and 3 times premeal short acting insulin) for adults with type 1 diabetes is recommended if an MDI basalbolus insulin regimen is not possible. Consider a trial of a twicedaily analogue mixed insulin regimen if an adult using a twicedaily human mixed insulin regimen has hypoglycemia that affects their quality of life.

3.5.:Optimizing insulin therapy

Consider adding metformin to insulin therapy if an adult with type 1 diabetes and a BMI of 23 kg/m² or above wants to improve their blood glucose control while minimizing their effective insulin dose [11].

4. Children and young people with diabetes

4.1.: While the insulin regimen should be individualized for each patient, there are three basic types of insulin regimen: 1. MDI basalbolus insulin regimens: one daily injection of long acting insulin analogue injections and three rapidacting insulin analogue before meals which is the preferred insulin regimen. 2. CSII therapy: a programmable pump and insulin storage device that gives a regular or continuous amount of insulin (usually a rapidacting insulin analogue or shortacting insulin) by a subcutaneous needle or cannula. 3. One, two or three insulin injections per day: these are usually injections of shortacting insulin mixed with intermediateacting insulin.

4.2.: For children and young people with type 1 diabetes, MDI basalbolus insulin regimens should be considered from diagnosis. If an MDI regimen is not appropriate for a child or young person with type 1 diabetes, consider CSII therapy as recommended in CSII therapy for the treatment of diabetes mellitus [11]. For children of less than 1 year of age, degludec is not recommended.

References:

1. Pharmacologic Approaches to Glycemic Treatment: Standards of Medical Care in Diabetes - 2018. *Diabetes Care* 2018; 41 (Suppl. 1): S73-S85.
2. Heller SR. Insulin treatment in type 2 diabetes: Medical Education Partnership. As reported in *Insulin Made Easy* edited by A H Barnett: P 18. 2001
3. Wallace TM, Mathews DR. Intensive management of glycaemia. Medical Education Partnership. As reported in *Insulin Made Easy* edited by A H Barnett: P 41-42. 2001.
4. Paparella S. Avoiding errors with insulin therapy. *J Emerg Nurs* 2006;32:325-328.
5. Triplitt C, Wright A, Chiquette E. Incretin mimetics and dipeptidyl peptidase-IV inhibitors: potential new therapies for type 2 diabetes mellitus. *Pharmacotherapy* 2006;26:360-374.
6. Hirsch IB, Bergenstal RM, et al. A real-world approach to insulin therapy in primary care practice. *Clin Diabetes* 2005;23:78-86.
7. Texas Department of State Health Services. Insulin algorithm for type 2 diabetes mellitus in children and adults. Publication #45-11647. Available at: <http://www.dshs.state.tx.us/diabetes/PDF/algorithms/INST2.pdf>. Accessed March 30, 2007.
8. Inzucchi SE, et al. *Diabetes Care* 2015;38:140-149.
9. IDF Clinical Guidelines Task Force. Global guideline for type 2 diabetes. Brussels: IDF, 2012.
10. AACE/ACE Comprehensive Type 2 Diabetes Management Algorithm 2016 Published in *Endocr-Pract* 2016;22:84-113.
11. NICE short clinical guidelines. 87: London 2009.

Section 4:

Insulin Therapy at Hospital Settings

Key Points:

- For inpatients, best tool to differentiate between hyperglycemia and pre-existing diabetes mellitus (DM) is A1C. A value of A1C $\geq 6.5\%$ suggests pre-existing DM.
- Target plasma glucose during hospital stay (including intensive care unit) is between ≥ 7.8 mmol/L to ≤ 10.0 mmol/L.
- Glycemic target is best achieved with judicious use of insulin.
- Recommended Insulin therapy protocol at non-ICU settings: Subcutaneous basal insulin \pm prandial insulin \pm Correction insulin regimen is recommended. Sliding scale insulin (SSI) regimen is not recommended. Premixed insulins can be used for stable persons or before discharge.
- Naso-gastric (NG) tube feeding/ Per-enteral tube feeding: In patients on per enteral tube feeding, two episodes of feeding should be ensured after the evening prandial S/C insulin to avoid nocturnal hypoglycemia.
- Recommended Insulin therapy protocol at ICU settings: Short acting regular insulin in IV route is recommended to use by IV infusion pump device at rate 0.5 to 12 units/ hour. Another approach is weight-based calculation of insulin dose e.g.; 0.01 to 0.02 to 0.05 units of insulin/Kg body weight/hour.
- Peri-operative insulin therapy: Perioperative PG target is between 6.0 – 10.0 mmol/L. At night before the intended procedure, regular insulin regimen should be used.

Mortality and morbidity of patients admitted into hospital are associated with hyperglycemia [1,2,3]. At inpatient settings, hyperglycemia is defined as RPG ≥ 7.8 mmol/L. Best tool to differentiate between hyperglycemia and pre-existing diabetes mellitus (DM) is A1C. A value of A1C $\geq 6.5\%$ suggests DM [4]. Target plasma glucose during hospital stay (including intensive care unit) is between ≥ 7.8 mmol/L to ≤ 10.0 mmol/L and this target is best achieved with judicious use of insulin [4].

In this regard, practical “guideline” is less formulated due to fewer researches, various clinical settings and less experience with newer agents [4,5,6]. We (Bangladesh Endocrine Society) propose following insulin therapy for treating hyperglycemia in various inpatient settings.

1. Insulin therapy protocol at non-ICU settings:

1.1: In non-ICU settings, physiological Insulin replacement protocol in subcutaneous (SC) route for management of inpatient hyperglycemia is basal insulin \pm prandial insulin \pm Correction (Multiple dose insulin) regimen when meal consumption is regular.

1.2.: While meal intake is discrete or the patient is under the settings of nothing per oral (NPO), naso-gastric feeding (NG) and or receiving total par-enteral nutrition (TPN), this regimen is defined as Basal \pm Nutritional \pm Correction insulin regimen [4,5,6].

1.3: Basal-prandial insulin regimen is superior to sliding scale insulin (SSI) because SSI regimen lacks basal component which is required to suppress and or prevent gluconeogenesis and ketogenesis [7].

1.4: Physiological Insulin replacement protocols in SC route for various inpatient settings are as following [4,5,6]:

- **Step 1:** Define nutritional status of patient, that is whether patient is eating or under settings of NPO, tube feeding, TPN, etc.
- **Step 2:** Then, estimate approximate total daily dose (TDD) of insulin. One approach is to take history of previous total dose, current A1c and adjust the dose accordingly. When records are not available or patient is insulin naïve, another approach is weight-based calculation of TDD of insulin as following; TDD of insulin: 0.3 to 0.5 units of insulin/kg body weight/Day.

- **Step 3:** Split TDD of insulin into basal and prandial/ nutritional components as 50/50 or 30/70 ratio or 30 to 50% basal and 70 to 50% prandial/ nutritional components. NPH, long acting basal analogues insulins are used as basal component and short acting regular or rapid acting analogues insulins are used as prandial/ nutritional component in SC route.

Example: A 70 kg diabetic male with random plasma glucose of 12 mmol/L and A1c 7.2% is admitted into general medicine ward. Patient can take meal routinely. Before admission he was on premixed (30/70) regular insulin at a dose 24+0+20 SC. So, his TDD of insulin is 24+20=44. Then, split TDD into 30% basal (e.g.; 15 units basal analogue given SC once or 7 units NPH twice daily) and 70% prandial component (e.g.; 70% of 44= 30 units; so 10 units regular short acting/ rapid acting analogue given SC 3 times before 3 major meals).

- **Step 4:** Correction or supplement dose is given with regular short acting or rapid acting analogue insulin in SC route which may be required if pre-meal PG is high. After 24 hour plasma glucose monitoring (pre-meals, 2 hour post-meals, q 4-6 hourly), re-adjust the dose. The regimen should be re-assessed on daily basis depending on plasma glucose level and condition of patient. Detailed protocol in various settings are mentioned in Table 1.

1.5.: Naso-gastric (NG) tube feeding/ Per-enteral tube feeding: In patients on per enteral tube feeding, two episodes of feeding should be ensured after the evening prandial S/C insulin to avoid nocturnal hypoglycemia.

2. Insulin therapy protocol critical at ICU settings:

2.1: Insulin in intravenous (IV) route is used during these settings, e.g.; MICU, SICU, major Surgery, cardiovascular procedures, MI, NPO, DKA, high dose steroids, gastroparesis, dose finding strategy etc [5].

2.2: Short acting regular insulin in IV route is used by infusion pump device at these settings. As per PG values, infusion rate may vary between 0.5 to 12 units of insulin/ hour. Another approach is weight based calculation of insulin dose e.g.; 0.01 to 0.02 to 0.05 units of insulin/Kg body weight/hour [5,8].

2.3: Switching from IV insulin to SC insulin regimen protocol [4,5,6]:

- **Step 1:** first find out TDD of insulin which may be required by patient in next 24 hours. This is best calculated by following formula: TDD of insulin for next 24 hours= Stabilized hourly rate of insulin x 20. For example, if patient and his PG were stable with 2 units/ hour of regular short acting insulin during switching the route, then TDD requirement for next 24 hours would be approximately 2x 20= 40 units of insulin.
- **Step 2:** Then, divide TDD of insulin into 30-50 % basal insulin and 70 to 50% into 3 prandial/nutritional insulin components. SC insulin should be started 1-2 hour before discontinuation of IV insulin. Correction dose with short acting of insulin may be required. Re-assess the regimen and dose after 24 hours and re-adjust accordingly if required.

3. Peri-operative insulin therapy [4,5,6]:

3.1: At perioperative period, PG target is between 6.0 – 10.0 mmol/L.

3.2: Surgical procedure is encouraged to be started at early morning. At night before the intended procedure, regular insulin regimen should be used.

3.3: At morning during short procedures, ½ of NPH or long acting basal insulin in SC route is given and PG is monitored every 30-60 minutes interval and if PG is above target then correction dose with regular short acting insulin or rapid acting analogue insulin in SC route is administered Q 4-6 hourly.

3.4: During prolonged procedures, IV infusion of 5% dextrose + IV short acting regular insulin in drip is used with SC correction dose of regular short acting insulin or rapid acting analogue insulin Q 4-6 hourly if needed. Fluid balanced should be maintained.

4. Use of Pre-mixed insulin:

Premixed insulins can be used for stable persons or before discharge [4,6].

Table 1: Physiological Insulin replacement protocols for various inpatient settings [5].

Settings	TDD of insulin	Basal insulin S/Cq 24 or 12 h Basal -I or NPH	Prandial/ Nutritional insulin S/C q 4-6 h Reg-I, Rapid-I	Correction/ Supplement dose S/C insulin with Reg-I, Rapid-I	IV insulin
On insulin therapy- Eating	0.4-0.6 U/Kg/day	30%-50 %	70-50%	over 10 mmol/L. 1 mmol/L =0.5- 1 unit I	
On insulin therapy- NPO	0.4-0.6 U/Kg/day	30%-50 %	70-50%	Same	Insulin drip 5% DNS at 75-100 ml/hr (+ Reg I 1-4 U/100ml)
Oral agents- Eating	0.3 u/kg/day	30%-50 %	70-50%	Same	
Oral agents- NPO	0.3 u/kg/day	20%-50 %	80-50%	Same	5% DNS at 75-100 ml/hr (+ Reg I 1-4 U/100ml)
Newly diagnosed hyperglycemia - eating	0.3u/kg/day	30%-50 %	70-50%	Same	
Newly diagnosed hyperglycemia - NPO	0.2u/kg/day	Basal +/-		Same	5% DNS at 75-100 ml/hr (+ Reg I 1-4 U/100ml)
Enteral tube feeding	0.3u/kg/day	<40%:	>60%		
On Steroid	0.4-0.6 U/Kg/day	30%-50 %	70-50%	Same	Insulin pump

N.B.; TDD: total daily dose of insulin; U: unit; Kg: kilogram; I: insulin; Reg I: regular short acting insulin; Rapid I: rapid acting analogue insulin; NPH: Neutral Protamine Hagedorn; .

References:

- McAlister FA, Majumdar SR, Blitz S, Rowe BH, Romney J, Marrie TJ. The relation between hyperglycemia and outcomes in 2,471 patients admitted to the hospital with community-acquired pneumonia. *Diabetes Care* 2005;28:810–815.
- Falguera M, Pifarre R, Martin A, Sheikh A, Moreno A. Etiology and outcome of community-acquired pneumonia in patients with diabetes mellitus. *Chest*. 2005 Nov;128(5):3233-9.
- Frisch A, Chandra P, Smiley D, Peng L, Rizzo M, Gatcliffe C, Hudson M, Mendoza J, Johnson R, Lin E, Umpierrez GE. Prevalence and clinical outcome of hyperglycemia in the perioperative period in noncardiac surgery. *Diabetes Care* 2010;33:1783–1788
- American Diabetes Association. Standards of Medical Care in Diabetes, 2017. *Diabetes Care* 2017;40(supp 1):s120—s127.
- Clement S, Baithwaite SS, Magee MF, et al. Management of diabetes and hyperglycemia in hospitals. *Diabetes Care* 2004;27:553–591.
- Wesorick D, O'Malley C, Rushakoff R, Larsen K, Magee M. Management of Diabetes and Hyperglycemia in the Hospital: A Practical Guide to Subcutaneous Insulin Use in the Non-Critically Ill, Adult Patient. *J of Hosp Med* 2008; 3(5) : S17-S28.
- Umpierrez GE, Andres P, Smiley D, et al. Randomized study of basal-bolus insulin therapy in the inpatient management of patients with type 2 diabetes (Rabbit 2 trial). *Diabetes Care* 2007;30:2181–2186.
- van den Berghe G, Wouters P, Weekers F, Verwaest C, Bruyninckx F, Schetz M, Vlasselaers D, Ferdinande P, Lauwers P, Bouillon R. Intensive insulin therapy in the critically ill patients. *N Engl J Med* 2001;345:1359–1367.

Section 5:

Insulin Therapy in Hyperglycemic Crises

Key points:

- *Serious acute metabolic complications of diabetes are Diabetic ketoacidosis (DKA) and the hyperosmolar hyperglycemic state (HHS).*
- *The combination of uncontrolled hyperglycemia, metabolic acidosis, and ketonemia/ketonuria characterizes DKA.*
- *HHS is characterized by severe hyperglycemia, hyperosmolality, and dehydration in the absence of significant ketoacidosis.*
- *The most common precipitating factor in the development of DKA and HHS is intercurrent infection; other precipitating factors include discontinuation of insulin therapy, inadequate insulin therapy, pancreatitis, myocardial infarction, cerebrovascular accident, new-onset type 1 diabetes, psychological problem etc.*
- *The principles of treatment of DKA and HHS includes- correction of dehydration, hyperglycemia, electrolyte imbalances and identification of comorbid precipitating events along with frequent patient monitoring.*
- *Continuous administration of short acting regular insulin via intravenous infusion is the mainstay and preferred route because of its short half-life and easy titration.*

Serious acute metabolic complications of diabetes are Diabetic ketoacidosis (DKA) and the hyperosmolar hyperglycemic state (HHS). In the USA, DKA is responsible for more than 500,000 hospital days per year [1,2]. The combination of uncontrolled hyperglycemia, metabolic acidosis, and ketonemia/ketonuria characterizes DKA. HHS is characterized by severe hyperglycemia, hyperosmolality, and dehydration in the absence of significant ketoacidosis. Table 1 outlines the diagnostic criteria for DKA and HHS. These metabolic derangements result from the combination of absolute or relative insulin deficiency and an increase in counter-regulatory hormones (glucagon, catecholamines, cortisol, and growth hormone). Majority of the patients with DKA belong to type 1 diabetes. However, patients with type 2 diabetes can also develop DKA during stress and acute illness such as trauma, surgery, or infections.

Diabetic ketoacidosis is the most important cause of death in children with type 1 diabetes. It accounts for half of all deaths in diabetic patients younger than 24 years of age [3,4]. The overall mortality in adult subjects with DKA may be as low as <1% [1], but the rate is higher (~5%) in the elderly and in patients with concomitant life-threatening illnesses [5,6]. Mortality due to HHS is considerably higher (5–20%) [7,8].

1. Pathophysiology of DKA and HHS:

1.1.: The principal mechanism of hyperglycemia and ketosis in DKA is reduced effective insulin concentrations and increased concentrations of counter-regulatory hormones. Due to lack of insulin, free fatty acids are released into the circulation from adipose tissue (lipolysis). There is unrestrained hepatic fatty acid oxidation in the liver to ketone bodies, resulting in ketonemia and metabolic acidosis [9].

1.2.: In HHS, there is a greater degree of dehydration, but endogenous insulin secretion is more than in DKA. In HHS insulin level is inadequate to counteract hyperglycemia but adequate to prevent lipolysis and subsequent ketogenesis [10].

2. Precipitating Factors of DKA & HHS:

2.1.: The most common precipitating factor in the development of DKA and HHS is intercurrent infection [1,7].

2.2.: Other precipitating factors include discontinuation of insulin therapy, inadequate insulin therapy, pancreatitis, myocardial infarction, cerebrovascular accident, new-onset type 1 diabetes.

2.3.: Psychological problem is an important contributing factor for recurrent ketoacidosis in young patients with diabetes.

3. The principles of treatment of DKA and HHS:

- Correction of dehydration
- Correction of hyperglycemia
- Correction of electrolyte imbalances
- Identification of comorbid precipitating events
- Frequent patient monitoring.

3.1.: Administration of regular insulin via continuous intravenous infusion is the mainstay in the treatment of DKA [11]. Insulin therapy is effective regardless of the route of administration [12]. However, continuous administration of insulin via intravenous infusion is the preferred route because of its short half-life and easy titration [12,13,14].

3.2.: At the beginning patients should receive an hourly insulin infusion of 0.14 units/kg body weight [15]. The target is to decrease plasma glucose concentration at a rate of 2.8-4.2 mmol/L/hour. If plasma glucose does not decrease by 2.8-4.2 mmol from the initial value in the first hour, the insulin infusion should be increased every hour until a steady glucose decline is achieved (Figure 1). When the plasma glucose reaches 11.1 mmol/L in DKA or 16.7 mmol/L in HHS, it may be possible to decrease the insulin infusion rate to 0.02–0.05 units/kg/hour, at which time dextrose may be added to the intravenous fluids. Thereafter, the rate of insulin administration or the concentration of dextrose may need to be adjusted to maintain glucose values between 8.3 and 11.1 mmol/L in DKA or 13.9 and 16.7 mmol/L in HHS until they are resolved [16].

3.3.: The continuous intravenous insulin infusion in patients with DKA/HHS should be continued until the hyperglycemic crisis is resolved. Criteria for resolution of ketoacidosis includes a blood glucose <200 mg/dl and two of the following criteria:

- Serum bicarbonate level >15 mEq/l
- Venous pH >7.3
- Calculated anion gap <12 mEq/l.

3.4.: Resolution of HHS is indicated by normal plasma osmolality and regaining normal mental status. When resolution occurs, subcutaneous insulin therapy can be started. To prevent recurrence of hyperglycemia or ketoacidosis during the transition period to subcutaneous insulin, it is important to allow an overlap of 1–2 hours between discontinuation of intravenous insulin and the administration of subcutaneous insulin. The intravenous insulin infusion and fluid replacement should be continued despite biochemical resolution of the crises if the patient is to remain fasting/nothing by mouth. Patients with known diabetes may be given insulin at the dosage they were receiving before the onset of DKA if it was controlling glucose properly [16]. In insulin-naïve patients, a multidose insulin regimen should be started at a dose of 0.5–0.8 units/kg/day [19]. Human insulin (NPH and regular) may be given in two or three doses per day. However, basal-bolus regimen is the best option as it efficiently mimics physiologic insulin secretion. Table-2 shows a proposed method for estimation of insulin dose while switching from intravenous to subcutaneous insulin [20].

Table-1: Diagnostic criteria for DKA and HHS [10].

	DKA			HHS
	Mild	Moderate	Severe	
Arterial PH	7.25-7.30	7.00 to <7.24	<7.00	>7.30
Serum Bicarbonate (mEq/L)	15-18	10 to <15	<10	>18
Urine Ketone	Positive	Positive	Positive	Small
Effective serum osmolality	Variable	Variable	Variable	>320 mOsm/Kg
Anion gap	>10	>12	>12	Variable
Mental status	Alert	Alert/Drowsy	Stupor/ Coma	Stupor/ Coma

Effective serum osmolality= 2 [measured Na⁺ (mEq/L)] + Glucose (mmol/L)

Anion gap= (Na⁺) – [Cl⁻ + HCO₃⁻ (mEq/L)]

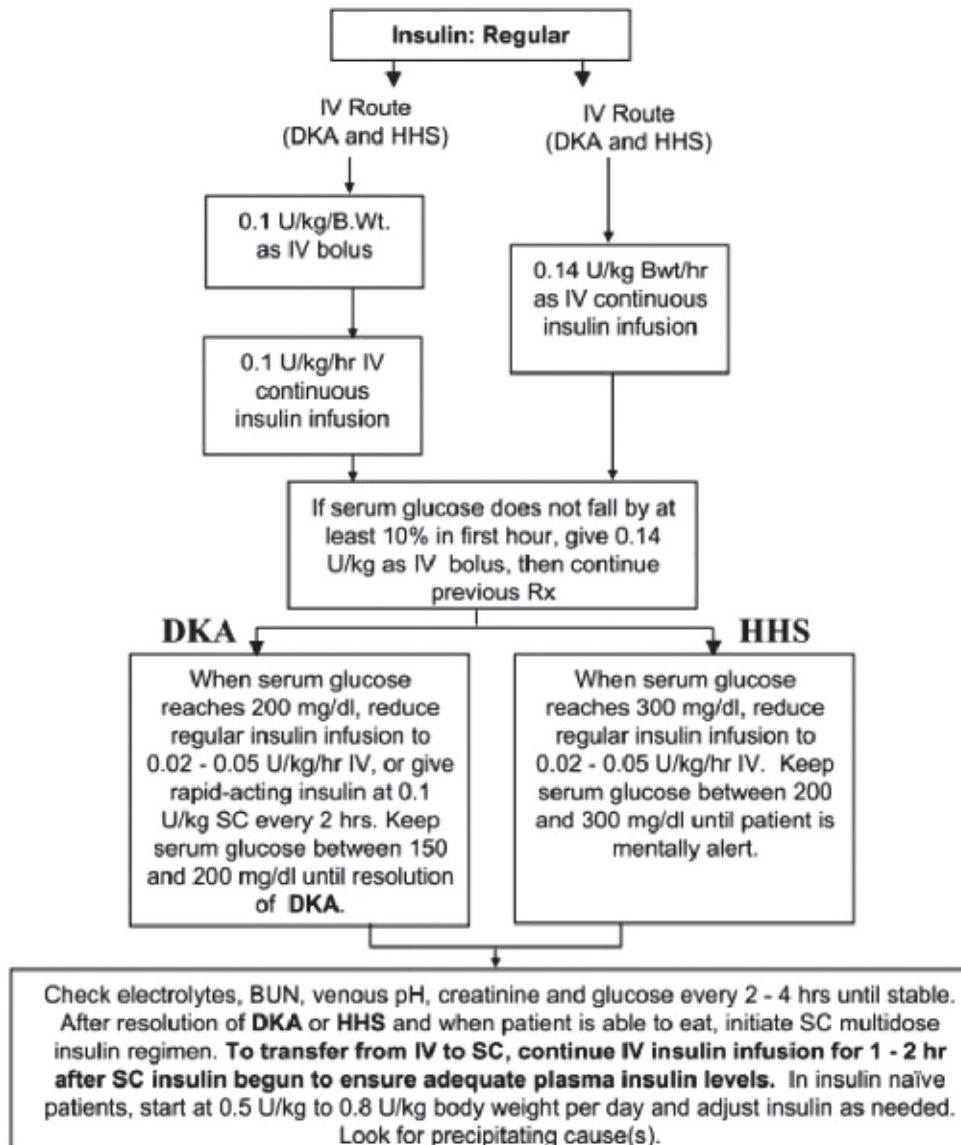


Figure-1: Insulin treatment in DKA and HHS [16].

Table 2: Estimation of insulin dose (S/C) for a patient being converted from an intravenous insulin infusion (Adapted from Hammersley et al) [20].

- Calculate average hourly insulin dose by totaling the last 6 hours' doses on the chart and dividing by 6 (e.g. 15 units divide by 6 = 2.5 units/hour. Multiply by factor of 20 to get the total daily dose (TDD) insulin (e.g. 50 units)
- For a basal : bolus insulin regimen, divide 50 : 50 basal : bolus (e.g. 25 units as basal; 25 units as bolus. Give 25 units as a basal insulin. Divide total bolus dose by three to get bolus for each meal (e.g. 8 units short - acting insulin)
- For a twice daily fixed mixture regime divide 60 : 40 morning : evening.

References:

1. National Center for Health Statistics. National hospital discharge and ambulatory surgery data [article online]. Available from <http://www.cdc.gov/nchs/about/major/hdasd/nhds.htm>.
2. Kim S. Burden of hospitalizations primarily due to uncontrolled diabetes: implications of inadequate primary health care in the United States. *Diabetes Care* 2007;30: 1281–1282.
3. Wolfsdorf J, Glaser N, Sperling MA. Diabetic ketoacidosis in infants, children, and adolescents: a consensus statement from the American Diabetes Association. *Diabetes Care* 2006;29:1150–2259.
4. White NH. Diabetic ketoacidosis in children. *Endocrinol Metab Clin North Am* 2000;29:657–682.
5. Graves EJ, Gillium BS, the National Center for Health Statistics. Detailed diagnoses and procedures: National Hospital Discharge Survey, 1995. *Vital Health Stat* 13 1997;(130):1–146.
6. Malone ML, Gennis V, Goodwin JS. Characteristics of diabetic ketoacidosis in older versus younger adults. *J Am Geriatr Soc* 1992;40:1100–1104.
7. Ennis ED, Stahl EJVB, Kreisberg RA. The hyperosmolar hyperglycemic syndrome. *Diabetes Rev* 1994;2:115–126.
8. Lorber D. Nonketotic hypertonicity in diabetes mellitus. *Med Clin North Am* 1995;79:39–52.
9. Miles JM, Haymond MW, Nissen S, Gerich JE. Effects of free fatty acid availability, glucagon excess and insulin deficiency on ketone body production in post-absorptive man. *J Clin Invest* 1983;71:1554–1561.
10. Kitabchi AE, Fisher JN, Murphy MB, Rumbak MJ. Diabetic ketoacidosis and the hyperglycemic hyperosmolar nonketotic state. In *Joslin's Diabetes Mellitus*. 13th ed. Kahn CR, Weir GC, Eds. Philadelphia, Lea &Febiger, 1994, p. 738–770.
11. Kitabchi AE, Umpierrez GE, Murphy MB, Barrett EJ, Kreisberg RA, Malone JI, Wall BM. Management of hyperglycemic crises in patients with diabetes. *Diabetes Care* 2001;24:131–153.
12. Fisher JN, Shahshahani MN, Kitabchi AE. Diabetic ketoacidosis: low-dose insulin therapy by various routes. *N Engl J Med* 1977;297:238–241.
13. Kitabchi AE, Fisher JN. Insulin therapy of diabetic ketoacidosis: physiologic versus pharmacologic doses of insulin and their routes of administration. In *Handbook of Diabetes Mellitus*. Brownlee M, Ed. New York, Garland ATPM Press, 1981, p. 95– 149.

14. Kitabchi AE, Umpierrez GE, Fisher JN, Murphy MB, Stentz FB. Thirty years of personal experience in hyperglycemic crises: diabetic ketoacidosis and hyperglycemic hyperosmolar state. *J Clin Endocrinol Metab* 2008;93:1541–1552.
15. Kitabchi AE, Murphy MB, Spencer J, Matteri R, Karas J. Is a priming dose of insulin necessary in a low-dose insulin protocol for the treatment of diabetic ketoacidosis? *Diabetes Care* 2008;31:2081–2085.
16. Kitabchi AE, Umpierrez GE, Miles JM, Fisher JN. Hyperglycemic Crisis In Adult Patients with Diabetes. *Diabetes Care* 2009; 32 (7): 1335- 1343.
17. Umpierrez GE, Latif K, Stoeber J, Cuervo R, Park L, Freire AX, Kitabchi AE. Efficacy of subcutaneous insulin lispro versus continuous intravenous regular insulin for the treatment of diabetic ketoacidosis. *Am J Med* 2004;117:291–296.
18. Umpierrez GE, Latif KA, Cuervo R, Karabell A, Freire AX, Kitabchi AE. Treatment of diabetic ketoacidosis with subcutaneous insulin aspart. *Diabetes Care* 2004; 27:1873–1878.
19. Kitabchi AE, Umpierrez GE, Murphy MB, Kreisberg RA. Hyperglycemic crises in adult patients with diabetes. *Diabetes Care* 2006;29:2739–2748.
20. Hammersley MS, James J. In-Hospital Treatment and Surgery in Patients with Diabetes. In: Holt RIG, Cockram C, Flyvbjerg A, Goldstein BJ (eds). *Textbook of Diabetes*. Oxford: Wiley-Blackwell, 2010: 521.

Section 6:

Using Insulin During Ramadan Fasting

Key points:

- *During Ramadan fast, person with diabetes can be categorized having very high, high or moderate/low risk.*
- *Pregnant women with diabetes are stratified as very high risk and are advised not to fast.*
- *During Ramadan, use of insulin analogues are better choice over regular human insulin due to a number of advantages.*
- *Structured Ramadan focused education program, Pre-Ramadan assessment, appropriate adjustment of treatment, education on meal and exercise should ideally be done 6–8 weeks before the start of Ramadan.*
- *Insulin dose should be reduced at Suhoor to avoid hypoglycemia.*
- *SMBG should be practiced as prescribed by physician to avoid hypo or hyperglycemia.*

Appropriate dose adjustment and choosing tailored insulin regimen is crucial for people having diabetes who wish to fast during holy month of Ramadan [1, 2]. During Ramadan fast, person with diabetes can be categorized having very high, high or moderate/low risk [3]. Very high-risk group is advised not to fast. Pregnant women with diabetes are stratified as very high risk and are advised not to fast. [3].

1. Insulin treatment for T2DM during Ramadan:

- 1.1. Insulin regimen for T2DM may include the use of premixed insulin (human & analogue), co-formulation of basal analogue, rapid acting analogue, basal analogue, split-mixed regimen (human), Intermediate-acting (human insulin), basal analogue without or with rapid acting analogue (basal plus, basal bolus regimen) and may be used in conjunction with OADs [4, 5, 6].
- 1.2. Limited data are available regarding the optimal insulin type or regimen for people with T2DM during Ramadan but results from several studies indicate that appropriate modification and individualization of insulin regimens are required [7, 8, 9, 10, 11, 12].
- 1.3. During Ramadan, use of insulin analogues is recommended over human insulin due to a number of advantages e.g.; less hypoglycemia, less weight gain, flexibility with meal time [13].
- 1.4. Pre-Ramadan assessment, appropriate adjustment of treatment, education on meal, exercise and monitoring of blood glucose monitoring should ideally be done 6–8 weeks before the start of Ramadan [3].
- 1.5. Patients with T2DM with poor glycaemic control despite of using multiple daily injections (MDI) of insulin can possibly be benefitted from CSII [14].
- 1.6. The “South Asian Consensus Guideline: Use of insulin in diabetes during Ramadan” states that ‘Once- or twice-daily injections of intermediate or long-acting insulin along with pre-meal rapid-acting insulin is the management of choice’ [15].
- 1.7. Recommended dose adjustments and SMBG-guided dose titrations can be found in Tables 1, 2 and 3 [3, 16]. Dose adjustment while switching from premixed human to premixed analogue insulin and while using insulin pump can be found in Tables 4 and 5. [3, 17, 18, 19].

2. Insulin treatment for T1DM during Ramadan:

- 2.1. Religious leaders, in alliance with diabetes experts, do not recommend fasting in individuals with T1DM [3, 20]. However, many persons with T1DM choose to fast.

- 2.2. The decision to fast during Ramadan must be respected and If the person is stable and healthy, he or she can do so with strict medical supervision and focused education [3].
- 2.3. Recommended insulin dose adjustment for person with T1DM during Ramadan fast can be found in Tables 1, 3, 5 and 6[3].
- 2.4. Adolescents with T1DM must be aware of all potential risks associated with Ramadan fasting. Insulin dose adjustment can be found in Table 6[3].
- 2.5. Capillary blood glucose should be monitored several times during the day [3].

3. Frequency of SMBG during Ramadan:

Recommended schedule as following[3]:

- 3.1. Moderate/Low risk: 1–2 times a day e.g.; 1 or 2 hour before Iftar / 4-6 hour after Suhoor, 2 hour after Iftar/ dinner.
- 3.2. Very high/ High Risk: several times a day e.g.; 1 or 2 hour before Iftar, 2 hour after Iftar/ dinner, before Suhoor, 4-6 hour after Suhoor.
- 3.3. Dose titration should be done according to SMBG results every 3rd day or as frequently as required.

4. When to break Fast:

Fasting should be broken if any of following conditions arises [3].

- 4.1. Blood glucose <3.9 mmol/L (<70 mg/dL).
- 4.2. Re-check within 1 h if blood glucose is between 3.9–5.0 mmol/L (70–90 mg/dL).
- 4.3. Blood glucose >16.7 mmol/L (>300 mg/dL) or symptomatic.
- 4.4. Symptoms of hypoglycaemia or acute metabolic complications.

Tables:

Table 1: Pre-mixed (human/ analogue) insulin dose adjustment during Ramadan both T2DM & T1DM.		
Once-daily dosing	Usual dose at iftar	
Twice-daily dosing	Usual morning/ higher dose at iftar.	Reduce evening/lower dose 50% if BG controlled or 0-25% if BG is uncontrolled and prescribe at suhoor.
Thrice-Daily dosing (analogue)	Usual morning/ higher dose at iftar.	Reduce evening/lower dose 50% if BG controlled or 0-25% if BG is uncontrolled and prescribe at suhoor. Omit lunch-time dose.
SMBG guided Dose titration: Should be carried out every 3 rd days.		

Table 2: Co-formulation (analogue) insulin dose adjustment during Ramadan.		
Once-daily dosing	Usual dose at iftar	
Twice-daily dosing	Usual breakfast/ lunch dose at iftar.	Reduce evening dose 30-50% and prescribe at suhoor.
SMBG guided Dose titration: Should be carried out every 3 rd days.		

Table 3: Intermediate (NPH)/Long acting (basal analogue) and Short/ rapid (analogue) acting insulin dose adjustment during Ramadan both T2DM & T1DM.		
NPH/Basal analogue: Once-daily dosing	Reduce dose by 15-30% and prescribe at iftar	
NPH/ Basal analogue: Twice-daily dosing	Usual morning dose at iftar.	Reduce evening dose 50% and prescribe at suhoor.
Short acting insulin/ rapid acting analogue	Usual morning dose at iftar.	Reduce evening dose 50% and prescribe at suhoor, omit lunch-time dose if dinner is not taken.
SMBG guided Dose titration: Should be carried out every 3 rd days.		

Table 4: Switching human Pre-mixed to analogue premixed insulin & dose adjustment during Ramadan		
Once-daily dosing	Reduce 20-30% of morning dose and prescribe at iftar	
Twice-daily dosing	Reduce 20-30% of morning dose and prescribe at iftar	Reduce evening/lower dose 60% and prescribe at suhoor.
SMBG guided Dose titration: Should be carried out every 3 rd days.		

Table 5: Insulin pump dose adjustment during Ramadan both T2DM & T1DM.		
Basal rate	Increase dose by 0-30% during early hours after iftar	Reduce dose by 20-40% during last 3-4 hours of fasting
Bolus rate	As per carbohydrate counting and insulin sensitivity principles	

Table 6: Intermediate (NPH)/Long acting (basal analogue) and Short / rapid (analogue) acting insulin dose adjustment during Ramadan in Adolescent T1DM.		
NPH/Basal analogue: Once-daily dosing	Reduce dose by 30-40% and given at iftar	
NPH/ Basal analogue: Twice-daily dosing	Usual morning dose at iftar.	Reduce evening dose 50% and prescribe at suhoor.
Short acting insulin/ rapid acting analogue	Usual morning dose at iftar.	Reduce evening dose 25-50% and prescribe at suhoor, omit lunch-time dose if dinner is not taken.
SMBG guided Dose titration: Should be carried out every 3 rd days.		

References:

1. Al-Arouj M, Bouguerra R, Buse J, et al. Recommendations for management of diabetes during Ramadan. *Diabetes Care* 2005;28:2305-11.
2. Al-Arouj M. Risk stratification of Ramadan fasting in person with diabetes. *J Pak Med Assoc* 2015;65:S18-21.
3. International Diabetes Federation and the DAR International Alliance. *Diabetes and Ramadan: Practical Guidelines*. Brussels, Belgium: International Diabetes Federation, 2016. www.idf.org/guidelines/diabetes-in-ramadan and www.daralliance.org
4. Joshi S and Joshi P. A review of insulin and insulin regimens in type 2 diabetes. *S Afr Fam Pract* 2009;51:97-102.
5. Bellido V, Suarez L, Rodriguez MG, et al. Comparison of basal-bolus and premixed insulin regimens in hospitalized patients with type 2 diabetes. *Diabetes Care* 2015;38:2211-6.
6. Mosenzon O and Raz I. Intensification of insulin therapy for type 2 diabetic patients in primary care: Basal-bolus regimen versus premix insulin analogs: When and for whom? *Diabetes Care* 2013;36:S212-8.
7. Salti I. Efficacy and safety of insulin glargine and glimepiride in subjects with type 2 diabetes before, during and after the period of fasting in Ramadan. *Diabet Med* 2009;26:1255-61.
8. Akram J and De Verga V. Insulin lispro (Lys(B28), Pro(B29) in the treatment of diabetes during the fasting month of Ramadan. Ramadan Study Group. *Diabet Med* 1999;16:861-6.
9. Hui E, Bravis V, Salih S, et al. Comparison of Humalog Mix 50 with human insulin Mix 30 in type 2 diabetes patients during Ramadan. *Int J Clin Pract* 2010;64:1095-9.
10. Mattoo V, Milicevic Z, Malone JK, et al. A comparison of insulin lispro Mix25 and human insulin 30/70 in the treatment of type 2 diabetes during Ramadan. *Diabetes Res Clin Pract* 2003;59:137-43.
11. Shehadeh N and Maor Y. Effect of a new insulin treatment regimen on glycaemic control and quality of life of Muslim patients with type 2 diabetes mellitus during Ramadan fast - an open label, controlled, multicentre, cluster randomised study. *Int J Clin Pract* 2015;69:1281-8.
12. Soewondo P, Adam JM, Sanusi H, et al. A multicenter, prospective, non-interventional evaluation of efficacy and safety of using biphasic insulin aspart as monotherapy, or in combination with oral hypoglycemic agent, in the treatment of type 2 diabetic patients before, during, & after Ramadan. *J Indones Med Assoc* 2009;59:574-9.
13. Grunberger G. Insulin analogs—Are they worth it? Yes! *Diabetes Care* 2014;37:1767-70.
14. Reznik Y, Cohen O, Aronson R, et al. Insulin pump treatment compared with multiple daily injections for treatment of type 2 diabetes (Opt2mise): A randomised open-label controlled trial. *Lancet* 2014;384:1265-72.
15. Pathan MF, Sahay RK, Zargar AH, et al. South Asian Consensus Guideline: Use of insulin in diabetes during Ramadan. *Indian J Endocrinol Metab* 2012;16:499-502.
16. Hassanein M, et al. Efficacy and Safety Analysis of Insulin Degludec/Insulin Aspart (IDegAsp) Compared with Biphasic Insulin Aspart 30 (BIAsp 30 [NovoLog® Mix 70/30]): A Phase 3, Multicenter, International, Open-Label, Randomized, Treat-to-Target Trial in Patients with Type 2 Diabetes Fasting during Ramadan. Abstract presented at ENDO 2017, April 1-4 2017, Orlando FL.
17. Hassanein M, Belhadj M, Abdallah K, Bhattacharya AD, Singh AK, Tayeb K, et al. Management of Type 2 diabetes in Ramadan: Low ratio premix insulin working group practical advice. *Indian J Endocrinol Metab* 2014;18:794-9.
18. Benbarka MM, Khalil AB, Beshyah SA, et al. Insulin pump therapy in Moslem patients with type 1 diabetes during Ramadan fasting: an observational report. *Diabetes Technol Ther* 2010;12:287-90.
19. Khalil AB, Beshyah SA, Abu Awad SM, et al. Ramadan fasting in diabetes patients on insulin pump therapy augmented by continuous glucose monitoring: an observational real-life study. *Diabetes Technol Ther* 2012;14:813-8.
20. Beshyah SA. Fasting during the month of Ramadan for people with diabetes: Medicine and Fiqh united at last. *Ibnosina Journal of Medicine and Biomedical Sciences* 2009;1:58-60.

Section 7:

Insulin Use in Pregnancy

Key Points:

- *Strict glycemic control is mandatory before and throughout pregnancy both in pre-gestational diabetes and GDM.*
- *Insulin is the standard of care to attain optimal glucose control in pregnancy and multiple methods and regimens are available to initiate insulin.*
- *Insulin therapy should be considered if one fails to achieve glycemic targets with non-pharmacological therapy (MNT & Physical activity) within target days and if target BGs are not achieved at any point of pregnancy after 1 to 2 weeks on MNT and exercise.*
- *Recommended insulins during pregnancy are: short-acting regular insulin, NPH, Aspart, Lispro and Detemir.*
- *Glulisine, Glargine and Degludec are pregnancy category C.*
- *Mixed use of conventional insulin with analogue insulin is not recommended.*
- *Required initial insulin dose is 0.5 to 1.0 U/kg body weight. Obese may require higher dose.*
- *Glycemic targets in pregnancy are: HbA1C ((%) <6.5%, FBG \leq 92 mg/dL (5.1 mmol/L), 1 hr PPG \leq 140 mg/dL (7.8 mmol/L) or 2-hr PPG 1: \leq 120 mg/dL (6.7 mmol/L).*

Strict glycemic control is mandatory before and throughout pregnancy both in pre-gestational diabetes and GDM, as it plays a vital role in decreasing poor maternal and fetal outcomes. Hyperglycemia itself is not teratogenic. As the prevalence of diabetes in young is increasing, the prevalence of pre-gestational diabetes is likely to increase in the pregnant population [1]. Uncontrolled diabetes during pregnancy may complicate pre-gestational diabetes with teratogenicity and GDM by poor perinatal outcome.

1. Insulin choice in Pregnancy

1.1.: Insulin is the standard of care to attain optimal glucose control in pregnancy and multiple methods and regimens are available to initiate insulin.

1.2.: Available Insulins and associated pregnancy category are given at Table 1 [2-14].

1.2.: The choice of protocol usually is based on clinician and patient's comfort and preference.

1.3.: Pregnancy does not have sufficient time as both maternal and fetal risks increases rapidly with poor glycemic control and quick control is mandatory. Accurate and timely adjustments depend on accurate blood glucose testing, type of insulin used, and consistent diet plan.

2. Insulin dosing in Pregnancy[15]:

2.1.: Required initial insulin dose is 0.5 to 1.0 U/kg body weight.

2.2.: Obese women may need higher dose.

2.3.: Treatment should be individualized and graded to reach the targets.

3. Glycemic Target [15]:

3.1.: Strict glycemic control is of utmost importance in all stages of pregnancy for women diagnosed with GDM, type 1 diabetes, or type 2 diabetes.

3.2.: Glycemic targets in pregnancy are HbA1C ((%) <6.5%, FBG \leq 92 mg/dL (5.1 mmol/L), 1 hr PPG \leq 140 mg/dL (7.8 mmol/L) or 2-hr PPG 1: \leq 120 mg/dL (6.7 mmol/L). [Table 2]

4. Insulin Protocol in pregnancy[15]:

4.1.: According to “GDM: SAFES Recommendations and action plan” insulin therapy should be considered if one fails to achieve glycemic targets with non-pharmacological therapy (MNT & Physical activity) within target days and if target BGs are not achieved at any point of pregnancy after 1 to 2 weeks on MNT and exercise

4.2.: During FIRST TRIMESTER and Third TRIMESTER [Table 3].

4.2.a.: If FPG is ≥ 92 mg/dl (≥ 5.1 mmol/L) to 109 mg/dl (6.0 mmol/L) and or 2h PPG is ≥ 120 mg/dl (≥ 6.7 mmol/L) to 139 mg/dl (7.7 mmol/L) non-Pharmacological therapy is started and continued. If BG targets are not achieved within 1 week, along with non-Pharmacological therapy, pharmacological treatment should be started.

4.2.b.: If FPG is ≥ 110 mg/dl (≥ 6.1 mmol/L) to 125 mg/dl (6.9 mmol/L) and or 2h PPG is ≥ 140 mg/dl (≥ 7.8 mmol/L) to 199 mg/dl (11.0 mmol/L) non-Pharmacological therapy is started & continued for 3 days. If good improvement after 3 days, non-Pharmacological therapy can be continued for 1 week. If BG target achieved after 1 week, then non-Pharmacological therapy is continued. If BG targets are not achieved, pharmacological treatment should be started along with non-Pharmacological therapy.

4.2.c.: If FPG is ≥ 125 mg/dl (≥ 7.0 mmol/L) and or 2h PPG is ≥ 200 mg/dl (≥ 11.1 mmol/L), along with non-Pharmacological therapy, pharmacological therapy should be started at the onset of treatment.

4.3.: During SECOND TRIMESTER [Table 4]:

4.3.a.: If FPG is ≥ 92 mg/dl (≥ 5.1 mmol/L) to 109 mg/dl (6.0 mmol/L) and or 2h PPG is ≥ 120 mg/dl (≥ 6.7 mmol/L) to 139 mg/dl (7.7 mmol/L) non-Pharmacological therapy is started & continued. If BG targets are not achieved within 2 weeks for uncomplicated cases and 1 week for complicated cases (Pre-eclampsia, ployhydramnios) pharmacological treatment should be started along with non-Pharmacological therapy.

4.3.b.: If FPG is ≥ 110 mg/dl (≥ 6.1 mmol/L) to 125 mg/dl (6.9 mmol/L) and or 2h PPG is ≥ 140 mg/dl (≥ 7.8 mmol/L) to 199 mg/dl (11.0 mmol/L) non-Pharmacological therapy can be started & continued for 1 week. If BG targets are achieved after 1 week, then non-Pharmacological therapy is continued. If BG targets are not achieved, pharmacological treatment should be started along with non-Pharmacological therapy.

4.3.c.: If FPG is ≥ 125 mg/dl (≥ 7.0 mmol/L) and or 2h PPG is ≥ 200 mg/dl (≥ 11.1 mmol/L), along with non-Pharmacological therapy, pharmacological therapy should be started at the onset of treatment.

5. Insulin therapy in diabetes with pregnancy (pre-gestational diabetes)

5.1.: Preconception care is important for all women with preexisting type 1 or type 2 diabetes. Education regarding strategies to maintain adequate nutrition and glucose control before conception, during pregnancy, and in the postpartum period is cornerstone of good pregnancy outcome.

5.2.: When women with diabetes attains glycemic control ($HbA_{1c} < 6.5\%$) before pregnancy then she will plan for conception and this glycemic target must be achieved by Insulin treatment. In that case OAD should be shifted to insulin and women getting insulin shifted to pregnancy safe one at least 3 months before planning for conception [15].

6. Premixed insulin is not a good choice of insulin during pregnancy and be considered on individual basis where patients are unwilling to or unable to take basal bolus or split mix regimen [15.]

7. Mixed use of conventional insulin with analogue insulin is not recommended [15].

Insulin	Time of onset	Peak time	Duration	Pregnancy category
Regular 40U/100U	30min	3 hr	8 hrs	B
Aspart/Lispro	10-15 min	30-90 min	3-5 hrs	B
Glulisine	10-15 min	55 min	3-5 hrs	C
NPH	1-2 hrs	4-8 hrs	10-20 hrs	B
Detemir	1-2 hrs	None	24 hrs	B
Glargine	1-2 hrs	None	24 hrs	C
Degludec	1 hrs	None	48 hrs(in steady state)	C

	mg/dl	mmol/L
FPG	<92	<5.1
1 h PPG	< 140	< 7.8
2 h PPG	< 120	< 6.7

N.B.: Adopted from GDM: SAFES Recommendations and action plan, Dhaka, SAFES, 2017 [15].

GDM PLASMA GLUCOSE TARGETS AND TREATMENT PROTOCOL BY SAFES						
	PG values			Treatment at onset	Change of treatment if Target not achieved in	Treatment reviewed & continued
FPG	≥92 mg/dl (≥5.1 mmol/L)	To	109 mg/dl (6.0 mmol/L)	NPT	1 week	NPT+PT
and/or						
2h PPG	≥120 mg/dl (≥6.7 mmol/L)	To	139 mg/dl (7.7 mmol/L)	NPT	1 week	NPT+PT
FPG	≥110 mg/dl (≥6.1 mmol/L)	To	125 mg/dl (6.9 mmol/L)	NPT	3 days	NPT+PT
and/or						
2h PPG	≥140 mg/dl (≥7.8 mmol/L)	To	199 mg/dl (11.0 mmol/L)	NPT	3 days	NPT+PT

NPT: Non-pharmacological treatment, PT: Pharmacological treatment

Table 4: TREATMENT IN 2ND TRIMESTER[15]						
GDM PLASMA GLUCOSE TARGETS AND TREATMENT PROTOCOL BY SAFES						
SECOND TRIMESTER:						
	PG values			Treatment at onset	Change of treatment if Target not achieved in	Treatment reviewed & continued
FPG	≥92 mg/dl (≥5.1 mmol/L)	To	109 mg/dl (6.0 mmol/L)	NPT	2 week/1 week Uncomplicated/complicated	NPT+PT
and/or						
2h PPG	≥120 mg/dl (≥6.7 mmol/L)	To	139 mg/dl (7.7 mmol/L)	NPT	2 week/1 week Uncomplicated/complicated	NPT+PT
FPG	≥110 mg/dl (≥6.1 mmol/L)	To	125 mg/dl (6.9 mmol/L)	NPT	1 week	NPT+PT
and/or						
2h PPG	≥140 mg/dl (≥7.8 mmol/L)	To	199 mg/dl (11.0 mmol/L)	NPT	1 week	NPT+PT
FPG	≥125 mg/dl (≥7.0 mmol/L)			NPT+PT	X	NPT+PT
and/or						
2h PPG	≥200 mg/dl (≥11.1 mmol/L)			NPT+PT	X	NPT+PT
NPT: Non-pharmacological treatment, PT: Pharmacological treatment						

References:

1. Centers for Disease Control and Prevention. National Diabetes Statistics Report: Estimates of diabetes and its burden in the United States, 2014. Atlanta, Ga., U.S. Department of Health and Human Services, 2014. Available from <http://www.cdc.gov/diabetes/data/statistics/2014StatisticsReport.html>. Accessed 3 October 2015). Humulin R U-100 [package insert] Indianapolis, Ind., Eli Lilly and Company, 2015.
2. Zuckerman LC, Werner EF, Pettker CM, McMahon-Brown EK, Thung SS, Han C. Pre-gestational diabetes with extreme insulin resistance: use of U-500 insulin in pregnancy. *ObstetGynecol* 2012;120:439–442.
3. Humulin R U-500 [package insert] Indianapolis, Ind., Eli Lilly and Company, 2014.
4. NovoLog [package insert] Bagsvaerd, Denmark, Novo Nordisk, 2015.
5. Mathiesen ER, Kinsley B, Amiel SA et al. Maternal glycemic control and hypoglycemia in type 1 diabetic pregnancy: a randomized trial of insulin aspart versus human insulin in 322 pregnant women. *Diabetes Care* 2007;30:771–776.
6. Humalog [package insert] Indianapolis, Ind., Eli Lilly and Company, 2015.
7. Apidra [package insert]. Bridgewater NJ., Sanofi-Aventis, 2015.
8. Humulin N [package insert] Indianapolis, Ind., Eli Lilly and Company, 2015.
9. Levemir [package insert] Bagsvaerd, Denmark, Novo Nordisk, 2015.
10. Herrera KM, Rosenn BM, Foroutan J, et al. Randomized controlled trial of insulin detemir versus NPH for the treatment of pregnant women with diabetes. *Am J ObstetGynecol* 2015;213:426.e1–426.e7.
11. Lantus [package insert]. Bridgewater NJ., Sanofi-Aventis, 2015.
12. Toujeo [package insert]. Bridgewater NJ., Sanofi-Aventis, 2015.
13. Tresiba [package insert] Bagsvaerd, Denmark, Novo Nordisk, 2015.
14. Afrezza [package insert] Danbury, Conn., MannKind Corporation, 2014.
15. GDM: SAFES Recommendations and action plan, Dhaka, SAFES, 2017.

Section 8:

Insulin in Chronic Kidney Disease (CKD):

Key Points:

- *In CKD, there is increased risk of hypoglycemia while using insulin due to decreased clearance.*
- *All available insulin preparations can be used in patients with CKD.*
- *Usually no dose adjustment is required for total daily dose (TDD) of insulin if the eGFR is >50 mL/min.*
- *Reduction to 75% of TDD of insulin when the eGFR is between 10 and 50 mL/min and to 50% of TDD for a eGFR of <10 mL/min may be considered which is independent of the type of insulin being used.*
- *Long acting insulin/ basal insulin analogue should be used with reduced dose during initiation and intensification.*

As kidney disease progresses there is increased risk of hypoglycemia due to decreased clearance of insulin. The kidney is responsible for about 30 to 80 % of insulin removal; reduced kidney function is associated with a prolonged insulin half-life and a decrease in insulin requirements as GFR declines. All available insulin preparations can be used in patients with CKD. The insulin type, dose and administration must be tailored to each patient to achieve goal glycemic while limiting the risk of hypoglycemia. Usually no dose adjustment is required for total daily dose (TDD) of insulin if the eGFR is >50 mL/min. Nevertheless, reduction to 75% of TDD when the eGFR is between 10 and 50 mL/min and to 50% of TDD for a eGFR of <10 mL/min may be considered which is independent of the type of insulin being used. With reduced eGFR, lower dose of long acting insulin/ basal insulin analogue should be used during initiation and intensification. Any regimen, e.g.; premixed or basal bolus may be used. Close monitoring of blood glucose and adjustment of insulin doses are required to avoid hypoglycemia [1].

Reference:

1. BetônicoCR, Silvia M O Titan, Maria Lúcia C, Giannella C, Nery M, Queiroz M. Management of diabetes mellitus in individuals with chronic kidney disease: therapeutic perspectives and glycemic control. Clinics (Sao Paulo) 2016 Jan; 71(1): 47–53.

Section 9:

Insulin in Chronic Liver Disease (CLD):

Key Points:

- *In CLD, Insulin therapy is considered as the safest and most effective therapy.*
- *In patients with DM and chronic liver disease, there is increase chance of post-prandial hyperglycemia and fasting hypoglycemia.*
- *Short-acting insulin are preferred.*
- *Insulin analogs may offer improved glycemic control compared to standard insulin with a lower risk for nocturnal and severe hypoglycemia.*
- *In CLD, short acting insulin analogues can be used just after meal if patient have nausea or reduced appetite.*

Insulin therapy is considered as the safest and most effective therapy in patients with liver dysfunction, with the limitation of increased risk of hypoglycemia. Short-acting insulin are preferred because the duration of action may vary in such situations. In patients with DM and chronic liver disease, there is increase chance of post-prandial hyperglycemia and fasting hypoglycemia. Without increasing costs, insulin analogs may offer equivalent or improved glycemic control compared to standard insulin while being associated with a lower risk for hypoglycemia, particularly nocturnal and severe hypoglycemia. The pharmacokinetics and pharmacodynamics of rapid-acting insulin analogs suggest that they can be given just after meals. This is of benefit to many patients with advanced CLD as they may have nausea and reduced appetite and hence have the option of using rapid-acting insulin analogs just after their meals depending on their intake [1].

Frequent dose adjustment and careful glucose monitoring for T2DM and CLD patients is thus very important to minimize the risk of hypoglycemia or hyperglycemia in this group of patients. Meals should be taken every three-hourly to avoid hypoglycemia.

Reference:

1. Gangopadhyay KK, Singh P. Consensus statement on dose modifications of antidiabetic agents in patients with hepatic impairment. Indian journal of endocrinology and metabolism 2017; 21 (2) : 341-354.

Section 10:**Common barriers to insulin initiation and strategies to overcome them**

Patient identified barrier	Strategies to address barrier
Fear of injection pain [1,2,3]	Demonstrate available tools and needle sizes [2]. Provide adequate practice and support of injections to overcome fear [3].
Fear of weight gain [1,2]	Dietary control and adequate exercise can minimise weight gain while also improving glycaemic control [3].
Inability to manage insulin regimen [1].	Provide education and support. Simplification of regimen and use of simple self-titration tools [1].
Hypoglycemia [1,2].	Use of long-acting analogues to reduce hypoglycemia risk. Provide education on recognition, management and avoidance of hypoglycemia. Reassure that incidence of serious hypoglycemia is rare [1].
Fear that diabetes has gotten worse or has become 'end stage' [1,2,3].	Introduce insulin as a diabetes management tool early in course of T2DM (24,25). Reassure that insulin requirement is an inevitable part of the disease course [1].
Decreased lifestyle flexibility [1,2].	Explain different insulin regimens and injection schedules [1].
Social stigma associated with injecting [1,2,3].	Introduce to all who have insulin pen stomach injecting simple and more discreet [1].
Insulin is not beneficial or can harm health [1,2,4]	Provide adequate education.

Reference:

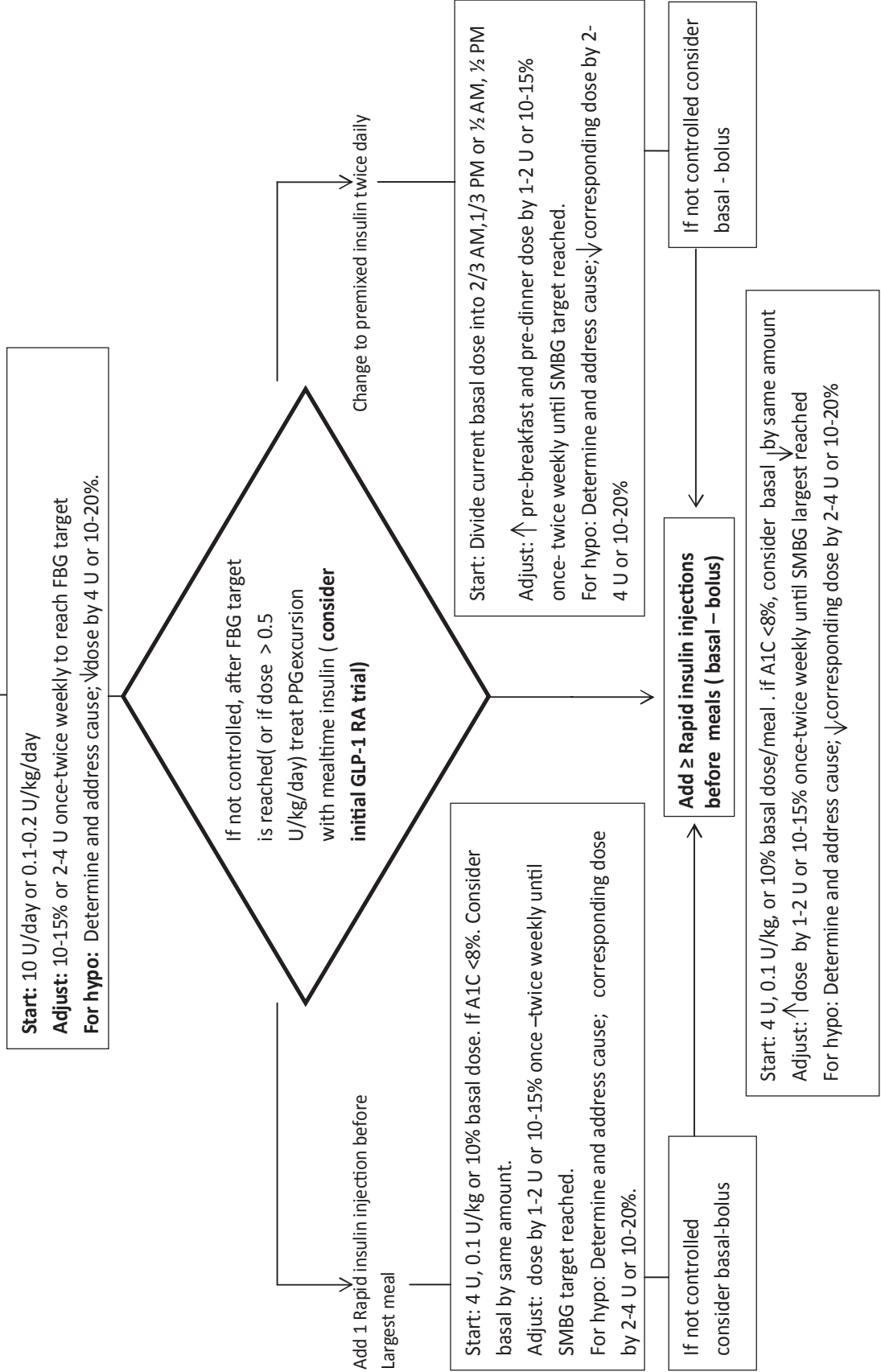
1. Peyrot M, Rubin RR, Khunti K. Addressing barriers to initiation of insulin in patients with type 2 diabetes. *Primary Care Diabetes* 2010;4:S11–S8.
2. Kunt T, Snoek FJ. Barriers to insulin initiation and intensification and how to overcome them. *International Journal of Clinical Practice* 2009;63:6–10.
3. Tan A, Muthusamy L, Ng C, Phoon K, Ow J, Tan N. Initiation of insulin for type 2 diabetes mellitus patients: what are the issues? A qualitative study. *Singapore Medical Journal* 2011;52(11):801.
4. Karter AJ, Subramanian U, Saha C, Crosson JC, Parker MM, Swain BE, et al. Barriers to insulin initiation. *The Translating Research Into Action for Diabetes Insulin Starts Project* 2010;33(4):733–5.

Appendix: Section 3

Approach to start and adjust insulin in T2DM

Basal insulin

(Usually with metformin +/- other noninsulin agent)



Notes

A series of horizontal dashed lines for writing notes.



Bangladesh Endocrine Society (BES)

Room-904, DMCH Building 2
Dhaka Medical College & Hospital
Secretariat Road, Dhaka 1000, Bangladesh
www.bes-org.net

Phone : +88 01511 552012
email : endobd2012@gmail.com

Visit www.bes-org.net/guideline to download this guideline