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Glycosylated Hemoglobin (HbA1c): An Indispensible Tool in the Management of Diabetes Mellitus

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6 Abstract

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Diabetes mellitus was known to mankind since antiquity although the name diabetes mellitus 7 is introduced much later. The disease is characterized by elevated blood sugar and estimation 8 of Glycosylated hemoglobin (HbA1c) remains one of the most important and indispensable 9 tools of investigation besides fasting and postprandial blood sugar levels. The glycosylation is 10 the nonenzymetic conjugation of a sugar to the amino group of proteins. The authenticity and 11 use of HbA1c in the management of diabetes are established by many landmark trials and 12 studies like the Diabetes Control and Complication Trial, (DCCT), UK Prospective Diabetes 13 Study (UKPDS), Standardization of Glycated Hemoglobin testing, National Glycohemoglobin 14 Standardization Programme, etc. These studies have developed the methods of HbA1c 15 estimation, the reference range for identification of pre-diabetics and diabetics, and range at 16 which micro and macrovascular complications occur. If the pitfall of HbA1c is kept in mind, it 17 is a proven tool in the management and follow-up of Diabetes Mellitus. 18

Index terms— diabetes mellitus, hba1c, DCCT, UKPDS, NGSP, IFCC, master equation, macro and micro vascular complications

²² 1 I. Introduction

iabetes mellitus was known to mankind since antiquity. Its mention is there as early as 1500 BC in Egyptian 23 manuscripts, although the name diabetes mellitus was added later on. The term diabetes means to siphon (Greek) 24 25 since the patient passes a large amount of urine and Mellitus means sweet like honey (Latin). [1] The disease is 26 characterized by elevated blood sugar and estimation of glycated Hb (HbA1c) remains one of the most important and indispensable investigations, besides estimation of blood sugar, fasting and postprandial. The blood glucose 27 concentration depends upon the many factors like food intake, exercise, stress, etc, but the concentration of 28 HbA1c in the blood reflects the average glucose over preceding 8-12 weeks. Thus estimation of HbA1c provides 29 additional criteria to assess the control of Diabetes and is free from diurnal fluctuations which occur with blood 30 glucose. The adaptation of HbA1c estimation has become an integral exercise in the management of diabetic 31 patients because it offers 32

Author: e-mail: drsujanagrawal@gmail.com Several advantages like patient need not be fasting, the sample can be collected at any time, it is stable and has got little biological variability. It also predicts the development of micro and macro vascular complications. [2], [3], [4] a) The HbA1c

In normal adults, haemoglobin usually contains HbA (?97% of the total) (Table ??), HbA2 (?2.5%), and HbF (?0.5%).

HbA is made up of four polypeptide chains, two ?-and two ?-chains. The glycosylation is the nonenzymatic attachment of a sugar to the amino group of proteins. Analysis has revealed that HbA1c has a hexose attached covalently to the NH2-terminal value residue of the ?-chains of HbA. [5] The International Union of Pure and Applied Chemistry have defined HbA1c as the fraction of the ?-chains of hemoglobin that has a stable hexose adduct on the NH2-terminal amino acid value. [6] b) The landmark trials/studies [3] This study was done to ensure that the results obtained by DCCT trials are comparable to with UKPDS trials. The estimation of HbA1c is done by ion-exchange, high performance liquid chromatography (HPLC) in DCCT Trials. Ten year follow-up

demonstrated that the risk of myocardial infarction was significantly lower in patients who had lower HbA1c 45 at the end of UKPDS trial. [7] 3. Standardization of glycated haemoglobin testing American Association for 46 Clinical Chemistry (AACC) established a committee in 1993 to standardize Glycosylated haemoglobin testing. 47 [8] The NGSP (National Glycohemoglobin Standardization Programme) was created three years later to execute 48 the protocol developed by AACC committee. The DCCT and UKPDS trials have established beyond doubt the 49 direct relationship between HbA1c concentration and outcome risks in patients with diabetes. The concept is 50 that all clinical laboratories should report an HbA1c value equivalent to that reported in the DCCT and UKPDS. 51 [9] The goal of the NGSP is to standardize glycated hemoglobin test results to those of the DCCT and UKPDS. 52

⁵³ 2 International Federation of Clinical Chemistry and

Laboratory Medicine (IFCC) in 1995. The primary objective of this committee was to develop a true reference 54 method for HbA1c. [10] Instead of standardizing to a comparison method, the primary objective of the IFCC 55 committee was to develop a true reference method for HbA1c. The NGSP and IFCC networks have complementary 56 roles in the HbA1c standardization process. The two networks work together to form a solid basis for the 57 establishment of a reliable and accurate HbA1c measurement platform. 5. The master equation a linear 58 relationship can be derived between HbA1c results of IFCC reference method and NGSP network. [11] The 59 calculated regression equation is: NGSP=0.09148(IFCC) + 2.152. This equation is called "master equation" and 60 permits conversion between the two sets of values. But, the HbA1c values measured by the IFCC method are 61 significantly lower than the NGSP values, and moreover, the difference is not constant. 62

The three preeminent clinical diabetic organizations, the American Diabetic Association (ADA), the European Association for the Study of Diabetes (EASD), and the International Diabetes Federation (IDF), attempted to resolve the dispute by agreeing to consider reporting HbA1c with estimated average glucose (ear). [12], [13] a linear relationship was established between HbA1c and mean blood glucose. [14] c) Reporting of HbA1c

Another significant development was how to report HbA1c.In 2007 IFCC adapted reporting by International 67 System of Units (SI units) rather than the percentage. This type of reporting avoids the confusion of using 68 percentage (as the unit), which is having different reference interval. Thus the IFCC values are now expressed as 69 millimoles of HbA1c per mole of HbA1c. [15], [16] In 2004, a working group was constituted with representatives 70 from ADA, ESAD, and IDF with a view to harmonizing HbA1c reporting. This group was termed as 71 ADA/ESAD/IDF working group of the HbA1c assay. The consensus statement published in 2007 reiterates that 72 the HbA1c should be reported worldwide in IFCC units (mmol/mol) and derived NGSP units (%) using IFCC-73 NGSP master equation. [17] Reporting of HbA1c HbA1c estimation is recommended by International guidelines as 74 a preferred measure when evaluating the overall control of diabetes and patient's risk for complications. [18], [19] 75 HbA1c is recognized as a reliable marker for the overall glucose exposure and its direct consequences. In setting 76 a target level of HbA1c in an individual patient, due consideration should be given to patient's health, the risk 77 of hypoglycemia, comorbidities, and his/her specific health issues. The IDF (International Diabetes Federation) 78

and the American College of Endocrinology (ACE) recommend HbA1c values below 6.5% while ADA (American Diabetes Association) recommends control below 7.0% for most patients. [20] Thus the estimation of HbA1c has
become a key measure for diagnosing, screening and monitoring diabetes, and is an indispensable tool.

i. Criteria for Identification of Prediabetic A Prediabetic is defined as, a person, having impaired fasting
glucose and impaired glucose tolerance, based on 2-hour OGTT (oral glucose tolerance test) and FPG (Fasting
Plasma Glucose). The patients having HbA1c values 6.00% to 6.49% are considered at high risk for developing
diabetes as advised by ADA and WHO. [21], [22], [23] Table ??: Criteria for identification of Prediabetic. [23]
Measurement ii.

⁸⁷ 3 Criteria for diagnosis of Diabetes

The 1997 Expert Committee on the Diagnosis and Classification of Diabetes Mellitus encouraged that that it 88 is based on the glycaemic level at which microvascular complications develops. [24] Evaluating HbA1c in 0.5% 89 increments, investigators found that the incidence of diabetic retinopathy rose above baseline at HbA1c of 6.5%, 90 the now accepted diagnostic value. The committee established that there is increased risk of diabetic retinopathy at 91 fasting plasma glucose level greater than or equal to 126mg/dl (7.0mmol/L). It was done to define a cut-off point 92 of HbA1c for diagnostic purposes. In a DETECT-2 trial, the investigators found microvascular complications 93 above 6.5% hbA1c. [25] Diabetes can be diagnosed by using venous plasma sample for HbA1c and fasting plasma 94 glucose, 2-hour oral glucose tolerance test (OGTT) adds to the diagnostic accuracy. OGTT also identifies the 95 patients with impaired glucose tolerance; this along with impaired fasting glucose is a marker for impaired beta 96 97 cell function with future progression to frank diabetes. 98 Random plasma glucose >200mg/dL (11.1 mmol/L) and symptoms of hyperglycemia like polyurea, polydipsia,

⁹⁹ blurred vision and weight loss also confirm the diagnosis (of Diabetes Mellitus). At any given time fulfillment
¹⁰⁰ of any two of the three criteria clinches to the diagnosis. [23] Table ??: Criteria for the diagnosis of Diabetes.
¹⁰¹ [23] Measurement The condition that shortens the life-span of RBCs or causes a rapid turnover of it shortens the
¹⁰² exposure of cells to as follows:

Acute or chronic blood loss due to any causes, hemolytic anemia, spherocytosis, splenomegaly, Hemoglobinopathies and associated anemia, chronic malaria, glucose-6-phosphate-dehydrogenase (G-6-

Pd) deficiency, sickle cell anemia etc, since they change turnover of RBCs. Erythrocyte survival is also shortened 105 in chronic hyperglycemic states. This decreased survival may underestimate the severity of hyperglycemia at 106 higher HbA1c levels. [27], [28] The end-stage renal disease generally has falsely low HbA1c levels. It is due 107 to chronic anemia and reduced red cell survival. So in end-stage renal disease, the clinician should take into 108 consideration this fact while evaluating glycaemic status. [29], [30] Pregnancy: A1C may not be a true reflection 109 of glycaemia during pregnancy primarily because of both the decreased life span of the red blood cell from 110 about 120 days to about 90 days as well as increased erythropoietin production. So, it is advised not to use the 111 criteria of HbA1c levels for the diagnosis of gestational diabetes. [31] Miscellaneous: the vitamin supplements 112 like Vitamin-E, Ribaverin, and interferon Alfa are associated with falsely lowered HbA1c levels. 113

¹¹⁴ 4 ii. Falsely elevated HbA1c levels

Any condition that prolongs the lifespan of red blood cells or decreases its turnover may cause a false elevation of HbA1c levels. These conditions include Iron deficiency anemia, vitamin B-12, and folate deficiency anemia. [32] Uremia, hypertriglyceridemia (concentration > 1750mg/dL), severe hyperbilirubinemia can also show falsely elevated HbA1c levels [33][34] So in the presence one has to be cautious in its interpretation. Salicylates, opiates and lead poisoning may be responsible for the false elevation of HbA1c levels. [35] While making changes in therapy based on elevated HbA1c levels these conditions must be taken into consideration to avoid, inappropriate therapeutic regimen and hypoglycemia.

In situations where the HbA1c results do not give a true picture due to false elevation or false low results, it is desirable to use alternative indices. These include fructosamine, glycated albumin, 1, 5-Anhdroglucitol (1, 5-AG), and continuous glucose monitoring (CGM).

125 5 II. Conclusion

The measurement of plasma glucose is the most reliable mean to diagnose diabetes mellitus. HbA1c measurement 126 remains the gold standard to monitor glycaemic control and also an important diagnostic criterion. The results 127 should be interpreted keeping in mind that, the other clinical conditions and co-morbidities, may give falsely 128 elevated or lowered levels of hbA1c. In such situations, the alternative measure of glycaemic control should be 129 considered like fructosamine, glycated albumin, 1, 5-AG or continuous glucose monitoring. The importance of 130 the estimation of HbA1c is that it is validated as a predictor of diabetes related outcomes. The diagnostic HbA1c 131 cut-off point of 6.5% is associated with retinopathy prevalence and is the diagnostic thresholds for Fasting plasma 132 glucose and 2-hr post-prandial-glucose. The HbA1c range of 5.7-6.4% in addition to impaired fasting glucose 133 (IFG), and IGT (impaired glucose tolerance) have been included in the "categories of increased risk for diabetes". 134 The study of HbA1c has answered a very basic question of glycaemic control after which complications are 135 prone to occur. It has got certain inherent limitation, despite that estimation of HbA1c remains the gold standard 136 in the management of Diabetes mellitus. 137

The key learning points 1. An HbA1c level ? 6.5% is the cut -off point, used for the diagnosis of diabetes mellitus. 2. The values of HbA1c in the range of 5.7% to 6.4%. are included in the "categories of increased risk for diabetes". 3. HbA1c is formed by the glycosylation of Haemoglobin, so anything which interferes with a lifespan of RBCs or process of glycosylation may give a false result. 4. The use of Hba1c levels for monitoring of diabetes control should take into consideration the other co-morbidities. 5. The combination of FGT and HbA1c

¹⁴³ significantly enhances the diagnostic accuracy of these tests. ¹

¹Glycosylated Hemoglobin (HbA1c): An Indispensible Tool in the Management of Diabetes Mellitus

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HbA1c %	Mean plasma g	clucose (mg/dL) IFCC Units (mmol/mol
6	126	42
6.5	141	48
7	154	53
7.5	169	59
8	183	64
8.5	198	69
9	212	75
9.5	226	80
10	240	86
11	269	97
12	298	108
IFCC= International Federation	of Clinical Chemistry and Laborat	tory Medicine
d) Pitfalls in HbA1c Measureme	e e	~

i. Falsely lowered HbA1c levels

Figure 1: Table 3 :

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[Note: $C \otimes 2018$ Global Journals. 21. World Health Organization. Use of glycated haemoglobin (HbA1c) in the diagnosis of diabetes mellitus. Abbreviated report of a WHO consultation. World Health Organization website. http://www.who.int/diabetes/publications/report-hba1c_2011. pdf. Published 2011. Accessed May 27, 2018.]

Figure 2:

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