 Diabetes Management in a Point of Care Setting

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Why point of care testing?
The objective of point-of-care testing (POCT) is the rapid provision of diagnostic information to enable clinical decisions to be made at the earliest opportunity during patient care and treatment. Such rapid provision of information facilitates optimisation of the care process. The potential utility for any application of POCT can therefore be judged in terms of its contribution to decision making and to the process of care – the latter including access to care. However it is also important to be aware of the potential impact of the analytical performance of POCT measurement systems compared with laboratory analytical systems, as the majority of the core evidence on the utility of a test will have been established using laboratory based systems.

A chequered history
The role of HbA1c testing in the management of patients with diabetes has been established for several decades, whilst its role in the diagnosis has been recognised more recently. These utilities are based on the fact that the HbA1c concentration reflects the average circulating glucose concentration over the lifespan of the red cell, and the evidence that HbA1c concentration is a good predictor of the complications of diabetes e.g. cardiovascular disease (1-3). Early experience with the use of HbA1c measurement was based on laboratory-based methods including ion exchange and affinity chromatography methods, with alternative affinity and immunological methods following later. An early study of biological variation indicated that intra- and inter individual variation between non-diabetics was 1.7 and 4.0% respectively (4). Another study found an intra-individual variation of HbA1c as 1.2% in non-diabetics, with a figure of 1.75% in patients with Type 1 diabetes. Interestingly, the respective figures for fasting blood glucose were 5 and 30% (5); this illustrates one of the attractive features for using the HbA1c measurement in screening for, and management of, diabetes. A more recent evaluation of the biological variation of HbA1c in healthy individuals using an IFCC-calibrated assay found intra- and inter individual variation of 2.5 and 7.1% respectively.

These authors used this data to calculate the desirable analytical goals for imprecision, bias and total error as 1.3%, 1.9% and 3.9% respectively (6). Using similar performance criteria Lestra Winters et al (7) found that only two out of eight POCT systems for the measurement of HbA1c met the required performance criteria whilst Bruns and Boyd (8) commented on the implication of poor analytical performance on clinical decision making. In the Lenters-Westra report, it is apparent that there was variation among the laboratory reference methods, although they were all controlled and calibrated in the authors’ laboratory. Indeed, one reference is cited as a source of concern regarding the accuracy of POC instruments, yet this reference describes an accuracy drift over time that was as large in the central laboratory instrument as it was in the POC device (9). Survey results from the College of American Pathologists (10) indicate that in the field, variation within and between laboratory-based methods can be comparable to or greater than some of the POC results reported by Lenters-Westra and an analysis of these trends was given in the report by Holmes et al (9).

Digging deeper into the analysis
A recent systematic review of the use of POCT for HbA1c in the management of patients with diabetes concluded that there was “an absence of evidence in clinical trial data to date for the effectiveness of POCT for HbA1c in the management of diabetes” (11). Whilst this might be considered disappointing, it is helpful to explore how the authors came to these conclusions. Firstly, whilst there were seven studies included in the review (12-17), two approaches to the surrogate outcome measure were employed (mean change in HbA1c level, and change in proportion of patients with HbA1c ≤ 7.0%) reducing the opportunity for meta-analysis of the full cohort of patients. Secondly there was considerable heterogeneity in the patient populations studied, including both Type 1 and Type 2 diabetics in some studies, as well as the proportion of patients with HbA1c values ≤7.0% at the outset of the studies (baseline), thus limiting the opportunity for pooling of data and meta-analysis.
In addition there was very little documentation of the treatment protocols employed, and therefore no indication as to whether patients had been stratified in relation to the care they were given. Thirdly there was no indication in some of the studies as to whether the results were discussed with the patients at the time the results were generated, and in one instance it was documented that the results using POCT had not been discussed at the time of generation (14). However there was some evidence of greater treatment intensification in patients with HbA1c >7.0% in those receiving POCT.

The authors drew attention to key features in the use of POCT, which are equally applicable to routine practice as well as in research studies, namely (i) the need to stratify patients (and their treatment) according to baseline HbA1c values; (ii) define and adhere to a revised process of care using POCT; and (iii) ensure that results of POCT are discussed with patients when generated and that treatment decisions are documented and implemented. Interestingly, four observational studies, of over 5700 patients with diabetes, in which there was immediate feedback of results to patients all showed significant reductions in the HbA1c results (17-20). Indeed, one of these studies demonstrated maintenance of improved HbA1c concentrations for a period of four years (19). A recent systematic view of quality improvement (QI) strategies in the management of diabetes has shown that QIs, involving greater adherence to guidelines can help to improve HbA1c levels (20). Data mining of primary care records has shown that there is evidence of both over- and under utilisation of HbA1c tests (21). On the other hand there is good evidence to show that patient satisfaction is improved using POCT, and personal knowledge of an individual’s HbA1c levels is associated with better outcomes (as judged by HbA1c levels) (22-24).

Concerns have been expressed about the use of HbA1c in screening for diabetes including the issue of access to the test in terms of instrument and consumables costs (31). Further, in studies of opportunistic screening for Type 2 diabetes in Emergency Departments, a high prevalence has been found. However there were significant problems associated with patient follow-up for diagnostic testing and lifestyle guidance (32,33). Current guidance supports the employment of HbA1c measurement in both screening for type 2 diabetes and in the management of patients with diabetes.

Summary
There are strong arguments for the use of POCT for HbA1c where the performance characteristics of the systems are equivalent to those employed in the central laboratory. POCT offers improved access to testing, as well as enabling immediate clinical decision making, discussion with the patient and implementation of appropriate treatment and/or lifestyle advice. Furthermore POCT enables testing to be undertaken closer to the patient, affords greater convenience for the patient, thereby improving the likelihood of treatment compliance. EKF Diagnostics provides point of care HbA1c analysers certified to international standards (IFCC and NGSP) for point of care testing during screening and monitoring of diabetes.

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