Factory-Calibrated Continuous Glucose Sensors: 
The Science Behind the Technology

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Abstract

The use of commercially available continuous glucose monitors for diabetes management requires sensor calibrations, which until recently are exclusively performed by the patient. A new development is the implementation of factory calibration for subcutaneous glucose sensors, which eliminates the need for user calibrations and the associated blood glucose tests. Factory calibration means that the calibration process is part of the sensor manufacturing process and performed under controlled laboratory conditions. The ability to move from a user calibration to factory calibration is based on several technical requirements related to sensor stability and the robustness of the sensor manufacturing process. The main advantages of factory calibration over the conventional user calibration are: (a) more convenience for the user, since no more fingersticks are required for calibration and (b) elimination of use errors related to the execution of the calibration process, which can lead to sensor inaccuracies. The FreeStyle Libre™ and FreeStyle Libre Pro™ flash continuous glucose monitoring systems are the first commercially available sensor systems using factory-calibrated sensors. For these sensor systems, no user calibrations are required throughout the sensor wear duration.

Keywords: Continuous Glucose Monitoring, Glucose sensor, Calibration, Factory calibration, Subcutaneous.

Introduction

Since the introduction of commercially available continuous glucose monitoring systems in 2000, significant progress in terms of system performance and convenience of use has been achieved, garnering positive expectation on the clinical utility and adoption of this technology. The first system introduced by MiniMed was a retrospective system, with data being available to the user or healthcare professional at the end of the sensor wear time. These early sensors could only be used for up to 3 days and needed a minimum of four calibrations per day. Over the following years the systems became easier to use, the accuracy of the systems improved, and the allowed wear duration was extended. However, until recently all systems still required daily blood glucose (BG) tests for recalibration to maintain accurate sensor glucose readings throughout sensor wear.

Most currently available continuous glucose monitoring systems employ enzymatic amperometric sensors measuring glucose in the interstitial subcutaneous tissue. The measurement signal is an electrical current. That current is proportional to the glucose concentration at the measurement site, with a small background current, which can be accounted for as a signal offset if necessary. To display glucose information to the user of the system, the sensor signal will have to be converted from an electrical current to a glucose value. This conversion is called calibration, and involves a BG test by the user. Assuming a linear sensor response to glucose and a negligible or known background signal, the sensor sensitivity to glucose can be calculated from one sensor current value and its corresponding time-matched BG reading. The sensor sensitivity represents the calibration factor, which can be used to convert the sensor electrical response into a glucose value moving forward from the calibration time point.

The user calibration process has several disadvantages. First, it is a burden to the user of the sensor system, since each calibration process requires a painful and time-consuming BG test. More importantly, the accuracy of the BG test directly determines the accuracy of the sensor system. Certain user mistakes like not washing hands before a BG test can...
lead to wrong glucose numbers. Some sensor systems require
the user to enter the BG value manually for calibration, where
transcription error and delayed BG entry can affect sensor ac-
curacy. On the other side, assuming the BG test was performed
correctly, if at the time of calibration the sensor signal has a
temporarily falsely reduced or elevated value, for example,
caused by interfering substances, the calculated sensor sensi-
tivity will not be correct and the following sensor data will
cause by interfering substances, the calculated sensor sensi-
tivity will not be correct and the following sensor data will
caused by interfering substances. The term factory calibration
refers by itself only to the process of determining the initial sensor sensitivity during the
manufacturing process. However, it is widely understood and
expected that factory-calibrated sensors do not require any
 recalibrations by the user, including no recalibrations during
sensor wear.

To be able to provide factory-calibrated sensors to the user
there is a set of requirements beyond the general require-
ments shared among glucose sensors, as outlined in Table 1.

The first three requirements are related to the design and
manufacturing of the sensor and the chemistry involved,
whereas the last requirement depends on the biology of the
interstitial tissue.

With respect to consistency of the sensor manufacturing
process, it is important to identify the sensor components
which do affect its sensitivity. For an amperometric sensor,
the sensing area located on the working electrode containing
the enzyme and the membrane covering the enzyme and
limiting the flux of glucose from the tissue to the enzyme are
the critical components. Therefore, it is essential to develop
processes to reproducibly deposit the enzyme on the working
 electrode and to create a uniform coating of the glucose-
 limiting membrane. Variations in sensing layer area and mem-
brane thickness between sensors have to be kept small, which
requires a high-precision manufacturing equipment given that
the areas involved are in the range of less than 1 mm² and the
membrane thickness is typically less than 100 μm. Sensor
design and architecture determine the options for manufactur-
ing methods. Therefore, if factory calibration is the goal, it is
crucial that these limitations are taken into consideration early in
the development process, so that the sensor architecture will
allow the use of robust manufacturing processes.

The sensor sensitivity is determined as part of the factory
calibration process at the end of the sensor manufacturing pro-
cess. This information is assigned to every individual sensor
usually in the form of a code. However, sensors are not being
used immediately after they are produced, and there will be a
period of time between the production and the use date. During
that time the sensor sensitivity cannot change. Otherwise, the
initially assigned sensitivity is no longer valid and the sensor
will provide false data once inserted and used by the patient.

| Requirement                        | Objective                                      | Testing environment |
|------------------------------------|-----------------------------------------------|---------------------|
| Consistent sensor manufacturing    | Reduce sensitivity variation between sensors.  | In vitro            |
| Shelf life stability               | Maintain sensor sensitivity over the assigned shelf life. | In vitro            |
| Wear stability                     | Maintain sensor sensitivity over the wear duration. | In vitro/ in vivo   |
| Consistent blood/tissue relationship| Demonstrate consistent BG-to-ISF-glucose gradient between subjects. | In vivo             |

BG, blood glucose.
Similar to the requirement that the sensor needs to be stable over its assigned shelf life, it also needs to be stable over its use period. If the sensitivity of the sensor to glucose does not change over the wear time, then recalibrations are not necessary. Alternatively, if the sensor response does change, recalibrations by the user can compensate for that sensor drift. Sensor drift is the reason why all currently available sensor systems, except FreeStyle Libre, require BG-based fingerstick calibrations by the user, typically twice a day.

The sensor stability over its use period is determined by two fundamentally different sensor properties. The first is the ability of the sensor to detect glucose with a stable sensor response. This property is governed by the underlying sensor chemistry and the enzyme involved, and can be demonstrated through in vitro tests. The second property is related to the biocompatibility of the sensor. The foreign body response to the sensor inserted into the subcutaneous tissue may lead to a change in sensor response. Therefore, to keep the signal stable over the wear period, the sensor design and the membrane chemistry either have to minimize the foreign body response or be able to prevent that response from interfering with the sensor signal. The sensor stability in vivo has to be determined through clinical trials.

The last requirement for the feasibility of factory calibration is the only requirement that is not related to the sensor itself. Since the currently available sensor systems are measuring glucose in the interstitial fluid, but are expected to predict the BG concentration, a consistent ratio between blood and tissue glucose is required. Many studies have been performed to estimate the absolute value of the interstitial glucose concentration and its relationship to BG. No clear consensus has been achieved to date, but most recent publications tend to estimate the tissue glucose to be around 90% of that of the blood under steady-state conditions. However, most studies only report an average value and attribute any variations to the experimental conditions and errors. Therefore, data need to be generated with respect to the variation of the blood to tissue glucose ratio variation both within a subject at different body sites or between different subjects.

The accuracy of a factory-calibrated sensor system will depend on the variations of the parameters associated with the requirements in Table 1. For example, minimizing the variation in sensitivity from sensor to sensor through manufacturing controls will minimize the sensor performance variance from individual to individual. The overall accuracy of the factory-calibrated sensor is achieved by applying appropriate specifications for the requirements in Table 1. Each specification will impact the accuracy independently, and therefore, it is up to the manufacturer to choose a set of specifications which will guarantee a desired accuracy level.

Implementation of Factory Calibration for FreeStyle Libre

The FreeStyle Libre and FreeStyle Libre Pro flash continuous glucose monitoring systems are the first commercially available factory-calibrated sensor systems. To our knowledge, no scientific studies have been published previously evaluating the feasibility of factory calibration besides the ones leading to FreeStyle Libre. All calibration-related studies and publications were focused on understanding and improving the standard BG-based fingerstick calibration or overcoming transient effects, such as lag and signal artifacts that can impact calibration. This demonstrates the novelty of this alternative approach and also possibly the superiority of the chemistry used in FreeStyle Libre over other sensor systems.

The development of the FreeStyle Libre sensor was guided by the requirements as outlined in Table 1. The chemistry as well as the architecture of the sensor was optimized to provide the necessary stability and robustness.

The FreeStyle Libre sensor is an enzymatic amperometric 3-electrode sensor system. The chemistry is based on the Wired Enzyme technology, which has been utilized in the FreeStyle Navigator continuous glucose monitoring system. This technique uses mediator molecules which are crosslinked together with the enzyme into a polymer matrix. Glucose molecules diffuse from the interstitial tissue through the outer membrane into the enzyme matrix and are oxidized by the enzyme glucose oxidase. The resulting electrons are transferred from the enzyme to mediator molecules (an osmium complex) and then shuttled to the working electrode using neighboring mediator molecules. The required electrical potential at the working electrode is only 40 mV versus a Ag/AgCl reference electrode. A low electrical potential minimizes the oxidation of electroactive species at the working electrode and thereby minimizes susceptibility to interferents.

The sensor design and the related manufacturing processes for the FreeStyle Libre sensor were chosen specifically to be able to manufacture identical sensors with respect to their response to glucose (Table 1). The most critical elements are the enzyme layer containing the enzyme and the glucose limiting membrane. The manufacturing equipment applying these two components has been optimized for robustness and reproducibility. Additional inspection steps ensure that every single sensor meets the predetermined specification criteria. Sensor lot release testing provides the lot calibration code and also includes a quantitative measure of within-lot variability.

The factory calibration process is based on the assumption that the in vitro sensor sensitivity predicts the in vivo sensor response. Since the sensor measurement site is the interstitial fluid and the reported value is BG, it is required to establish the relationship between the glucose concentrations of these two compartments. This can be done analytically or empirically. The analytical path will take into account all factors which are different in vitro versus in vivo, and which do influence the sensor response, for example, absolute glucose concentration, temperature, oxygen, and interfering substances. Alternatively, the in vitro to in vivo relationship can be established empirically by performing clinical studies and comparing the in vivo response to the in vitro data.

For example, the in vitro sensitivity can be calculated by examining the signal response of a sample of sensors from a lot to a set of known glucose concentrations, and then calculating the in vitro glucose sensitivity for each sensor. The nominal in vitro sensitivity of that sensor lot is then determined by taking the mean of the per-sensor in vitro sensitivities. Similarly, the in vivo sensitivity can be calculated by examining all the paired sensor/reference BG values in each sensor from a clinical study, and calculating the in vivo sensitivity for each of the sensors. Finally, the pooled in vivo sensitivity of that sensor lot is calculated by taking the mean of the per-sensor in vivo sensitivities.
Figure 1 shows the average in vivo sensor response from multiple sensor lots compared with their in vitro sensitivity. Data shown are drawn from two separate clinical studies. One study was performed in 12 subjects with diabetes, each subject wearing three sensors from six different sensor lots simultaneously over a 5-day wear period (Fig. 1: Study 1, lot 1 through 6). The other study includes 72 subjects with diabetes, each subject wore two sensors simultaneously. A total of three sensor lots were evaluated in this study (Fig. 1: Study 2, lot A through C). Capillary BG values are used as the reference BG in this analysis. While the study population and timing of the studies may have an influence on the sensitivity values and the narrow sensitivity range of the sensor lots used in the studies is limiting the statistical significance of data, we can see a correlation between the in vitro and in vivo values and an overlap of data from the two separate clinical studies.

Measuring and monitoring shelf-life stability for sensors can be performed under standard temperature conditions or under accelerated conditions at elevated temperatures. If accelerated conditions are chosen, data need to be available to determine the required exposure duration at the selected elevated temperature. These data are usually based on an Arrhenius relationship, which needs to be established for the specific sensor system. Sensor shelf life is limited by the stability of the enzyme and it is essential that the enzyme immobilization conditions are selected carefully. For the FreeStyle Libre sensor, the enzyme is immobilized in a crosslinked polymer matrix, which provides an optimized environment for enzyme stability.

Sensor stability for the FreeStyle Libre sensor over its 14-day use period has been demonstrated earlier. The in vitro tests include an initial sensitivity test, where the sensor is exposed to glucose solutions with different glucose concentrations. From the sensor response, a sensitivity value can be calculated. After this initial test, the sensors are kept in a glucose solution for 14 days to measure that glucose level continuously. After the 14-day period, another sensitivity test equal to the test at the beginning is being performed, and the resulting sensitivity is compared with the sensitivity at the beginning of the 14-day test. The difference between the initial and the final test represents the drift the sensor is experiencing over a 14-day monitoring period.

In vivo testing of sensor stability is absolutely required in addition to in vitro testing since different processes may be limiting stability in the tissue. In vivo stability is the ultimate requirement for sensor stability, and it may not be necessary to show in vitro stability if in vivo data are available. However, due to the significantly higher effort and cost to obtain clinical data, it is efficient to optimize sensor stability in vitro and, once the desired level of stability is achieved, only then to advance to the clinical stage.

Clinical data for 14-day stability have been shown previously using a sensor based on Wired Enzyme chemistry leading to the development of FreeStyle Libre. More recently, a clinical trial has been conducted using actual FreeStyle Libre sensors to evaluate accuracy of the system over a 14-day wear period. Seventy-two subjects with diabetes wore two sensors simultaneously on the back of the upper arm. Capillary BG was measured by the subjects throughout the test using the built-in FreeStyle Precision Strip Port, and compared with the glucose value reported by the factory-calibrated sensor system. The BG readings on the built-in meter are independent of, and do not influence, sensor readings.

Figure 2 shows an analysis of the 14-day stability of the sensor signal. A sensitivity value is calculated from each sensor/reference BG paired data point. For each sensor, the median of these individual sensitivity values are used to normalize data. Per-sensor normalized sensitivity values were then calculated for each day. Figure 2 shows the daily medians, interquartile ranges, and the 5th and 95th percentiles. That analysis illustrates any significant trends in the sensor sensitivity over the 14 days. We see a lower value on the first day, which is presumably related to the insertion process of the sensor and the associated trauma. From day 2 throughout day 14, the median sensor sensitivity remains...
constant, reflecting stable sensor chemistry, as well as negligible interference from the foreign body response.

The last requirement for the feasibility of factory calibration as laid out in Table 1 is the need for a constant blood to tissue glucose relationship. This requirement can be tested by using glucose sensors with identical in vitro response to glucose in different subjects and comparing the resulting sensor sensitivities from the in vivo testing. If there was a wide distribution of the ratio of tissue to BG concentration, there would be a wide distribution of the resulting sensor sensitivities. We have previously published data supporting the hypotheses that there is no difference in the tissue to BG ratio within a person at different body sites (arm vs. abdomen) as well as between subjects.9,11 We also used data from the clinical study described earlier12 and analyzed the sensor data for their sensitivity variation. Sensors with a minimum wear duration of 10 days were included in the analysis. Figure 3 shows the results in a cumulative distribution function plot, separated by the three lots used in the study. We can see that the three lots have 80%–92% of their values within 10% of their respective median and 100% of the values are within 20%. There are many factors that influence the calculation of each sensor’s in vivo sensitivity. Errors related to BG measurements,40,41 transient sensor effects,31,32,42,43 intersensor sensitivity variation used in the study, and variations in each study subject’s BG range and BG rate of change range44 can contribute to the variability observed in Figure 3. The narrow distribution indicates that the tissue to BG ratio is similar between subjects, which is required for factory calibration of sensors measuring glucose in the interstitial tissue.

**Alternative Approaches to Sensor Calibration**

If factory calibration is not feasible, there are other options to reduce the number of BG tests required for sensor calibration. Commercially available nonfactory calibrated continuous glucose monitoring systems require a minimum of two recalibrations per day and several studies suggest that accuracy can be impacted by increasing or decreasing this frequency.14,20 As previously outlined, the frequency of recalibrations is determined by the stability of the sensor over the wear period. Increasing sensor stability can, therefore, allow for a reduction in recalibration frequency for example, once a day instead of twice a day.

If sensor stability can be guaranteed throughout the sensor wear time no recalibrations may be necessary, and calibration is only needed at the beginning of sensor wear. This approach has significant risk since the calibration factor applied to the sensor throughout its wear time will be determined through only one calibration event. Some sensor systems take a hybrid approach with a robust initial calibration (multiple...
fingerstick requests) and a reduced frequency of recalibrations (e.g., once every 2 days)\textsuperscript{29,45} to minimize the overall number of BG tests required. However, many factors can impact the reliability of fingerstick calibration, resulting in calibration being one of the more dominant sources of sensor error.\textsuperscript{22,46} In daily use, recalibration requests may be skipped or not promptly entered,\textsuperscript{47,48} and there are many practical and technical factors\textsuperscript{40,41,49} limiting BG accuracy.\textsuperscript{50–53} A true factory calibration, where the sensor sensitivity is determined under laboratory conditions, is not susceptible to these use-dependent factors.

**Conclusions**

The availability of factory-calibrated glucose sensors has been predicted several years ago: “...I can see the day when accuracy will be sufficient that regulators will accept that CGM values can be used for clinical decision making, that factory calibration will be possible, that reimbursement will be a foregone conclusion, and usage will be routine so that all patients and providers will need to know how to accomplish it” (Skyler\textsuperscript{2}). With the introduction of the FreeStyle Libre flash continuous glucose monitoring system, part of this vision has become a reality. There is no need for the user to perform BG tests for sensor calibration. Calibrations performed by the user are not only a hassle and painful, but they introduce additional cost and can also lead to inaccurate sensor readings if done incorrectly. Factory calibration is performed under laboratory conditions and is part of the sensor manufacturing process. However, to be able to implement factory calibration several requirements related to sensor stability and reproducibility have to be demonstrated and maintained over the product life.

**Author Disclosures Statement**

U.H. and E.B. are employees of Abbott Diabetes Care.

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