

Blood Glucose Monitoring Test Systems for Prescription Point-of-Care Use

Draft Guidance for Industry and Food and Drug Administration Staff

DRAFT GUIDANCE

This guidance document is being distributed for comment purposes only. Document issued on: November 30, 2018.

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U.S. Department of Health and Human Services
Food and Drug Administration
Center for Devices and Radiological Health

Preface

Public Comment

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Table of Contents

I.	INTRODUCTION	5
II.	BACKGROUND	6
	CLIA WAIVER OF PROFESSIONAL USE METERS.....	8
III.	SCOPE	9
IV.	REDUCING THE RISK OF BLOODBORNE PATHOGEN TRANSMISSION	9
	A. VALIDATED CLEANING AND DISINFECTION PROCEDURES	11
	B. DEMONSTRATION THAT THE DEVICE IS ROBUST TO CLEANING AND DISINFECTION PROCEDURES.....	12
V.	DEVICE DESCRIPTION	13
VI.	PERFORMANCE EVALUATION FOR PRESCRIPTION-USE BGMS	14
	A. PRECISION EVALUATION STUDY	14
	B. LINEARITY EVALUATION STUDY.....	16
	C. METHOD COMPARISON/USER EVALUATION	16
	1. <i>General Study Design</i>	16
	2. <i>Data Analysis</i>	21
	D. INTERFERENCE EVALUATION	22
	1. <i>Endogenous/Exogenous Substances</i>	23
	2. <i>Hematocrit</i>	25
	3. <i>Oxygen</i>	28
	E. FLEX STUDIES.....	28
	1. <i>Test Strip Stability Testing</i>	30
	2. <i>System Operating Conditions Testing</i>	32
	3. <i>Altitude Effects</i>	32
	4. <i>Error Codes for Samples Outside the Measuring Range</i>	32
	<i>You should perform adequate analyses to demonstrate that your meter provides the appropriate error codes when measured glucose concentrations are outside of the BGMSs claimed measuring range, and include these results in your 510(k) submission</i>	32
	5. <i>Short Sample Detection</i>	33
	6. <i>Sample Perturbation Study</i>	33
	7. <i>Intermittent Sampling</i>	33
	8. <i>Testing with Used Test Strips</i>	34
	F. METER CALIBRATION AND QUALITY CONTROL MATERIAL	34
VII.	TEST STRIP LOT RELEASE CRITERIA	35
VIII.	THIRD PARTY TEST STRIPS	35
IX.	SOFTWARE	35
X.	LABELING	36

Contains Nonbinding Recommendations
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APPENDIX 1.SOURCES OF ERROR TO CONSIDER FOR BGMSS41
APPENDIX 2. SPECIAL 510(K)S AND BGMSS.....44

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Guidance for Industry and Food and Drug Administration Staff

1 *This draft guidance, when finalized, will represent the current thinking of the Food and*
2 *Drug Administration (FDA or Agency) on this topic. It does not establish any rights for*
3 *any person and is not binding on FDA or the public. You can use an alternative*
4 *approach if it satisfies the requirements of the applicable statutes and regulations. To*
5 *discuss an alternative approach, contact the FDA staff or Office responsible for this*
6 *guidance as listed on the title page.*

I. Introduction

This draft guidance document describes studies and information that FDA recommends be used when submitting premarket notifications (510(k)s) for blood glucose monitoring systems (BGMSs) which are for prescription point-of-care use. When finalized, this guidance document is intended to guide manufacturers in conducting appropriate performance studies and preparing 510(k) submissions for these device types, and will replace the final guidance entitled “Blood Glucose Monitoring Test Systems for Prescription Point-of-Care Use” issued on October 11, 2016.

7
8 This guidance is not meant to address self-monitoring blood glucose test systems (SMBGs) for
9 over-the-counter (OTC) home use by lay-users. FDA addresses those device types in another
10 guidance entitled “Self-Monitoring Blood Glucose Test Systems for Over-the-Counter Use”
11 (SMBG guidance -
12 [http://www.fda.gov/downloads/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocu](http://www.fda.gov/downloads/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocuments/UCM380327.pdf)
13 [ments/UCM380327.pdf](http://www.fda.gov/downloads/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocuments/UCM380327.pdf)). FDA is also issuing a revised draft of the SMBG guidance
14 ([https://www.fda.gov/downloads/MedicalDevices/DeviceRegulationandGuidance/GuidanceDoc](https://www.fda.gov/downloads/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocuments/UCM626742.pdf)
15 [uments/UCM626742.pdf](https://www.fda.gov/downloads/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocuments/UCM626742.pdf)) to reflect similar clarifications to the ones proposed in this draft
16 guidance.
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18 For the current edition of FDA-recognized standards referenced in this document, see the [FDA](#)
19 [Recognized Consensus Standards Database Web site](#).¹

20
21 FDA's guidance documents, including this guidance, do not establish legally enforceable
22 responsibilities. Instead, guidances describe the Agency's current thinking on a topic and should
23 be viewed only as recommendations, unless specific regulatory or statutory requirements are
24 cited. The use of the word *should* in Agency guidances means that something is suggested or
25 recommended, but not required.
26

27 **II. Background**

28
29 Portable blood glucose meters that measure blood glucose values are used by millions of people
30 every day as an aid in diabetes self-management. These types of devices are also used by
31 healthcare professionals in a variety of clinical settings including acute and chronic care
32 facilities, general hospital wards and intensive care units, physicians' offices, assisted living
33 facilities, and nursing homes.
34

35 Historically, the FDA has not recommended different types of information in premarket
36 submissions (510(k)s) for BGMSs intended to be used by healthcare professionals as compared
37 to over-the-counter SMBGs intended for home use by lay-users. However, it has become
38 increasingly clear that these different use settings comprise distinct intended use populations
39 with unique characteristics that can impact device design specifications, and that manufacturers
40 should take these unique characteristics into account when designing their devices. In order to
41 distinguish between FDA recommendations for prescription use blood glucose meters, which
42 are intended for use in point-of-care professional healthcare settings, and those intended for
43 OTC self-monitoring by lay-users, the Agency is issuing two separate guidances for (i) BGMSs
44 intended for use in point-of-care professional healthcare settings, and (ii) SMBGs intended for
45 home use for self-monitoring by lay-users. FDA believes that by making this distinction, each of
46 the devices can be better designed to meet the needs of their intended use populations, thereby
47 providing greater safety and efficacy.
48

49 In recent years, concerns have been raised related to infection control issues involving blood
50 glucose meters and lancing devices. According to the Centers for Medicare and Medicaid
51 Services (CMS) and the Centers for Disease Control and Prevention (CDC), blood glucose
52 meters can transmit bloodborne pathogens if these devices are contaminated with blood
53 specimens and are shared between users without effective cleaning, disinfecting, and
54 appropriate infection control measures.² Because BGMSs used in professional healthcare
55 settings are more likely to be used on multiple patients, this type of use requires certain design

¹ <http://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfStandards/search.cfm>

² See information at <http://www.cdc.gov/injectionsafety/blood-glucose-monitoring.html>.

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56 features and the capacity for cleaning and disinfection to prevent the spread of bloodborne
57 pathogens.³

58

59 In addition, concerns have been raised regarding the inability of currently cleared BGMSs to
60 perform effectively in professional healthcare settings because these devices have not been
61 adequately evaluated in some of the populations in which they are being used. Patients in
62 professional healthcare settings are often fundamentally different than lay-users using these
63 devices at home. Patients in professional healthcare settings can be acutely ill and medically
64 fragile and are more likely to present physiological and pathological factors that could interfere
65 with glucose measurements relative to lay-users. Errors in BGMS device accuracy can lead to
66 incorrect insulin dosing, which, when combined with other factors, can lead to increased
67 episodes of hypoglycemia. For hospitalized patients who may be seriously ill, glucose meter
68 inaccuracies could further increase risk to health.

69

70 Previously, most blood glucose meters, even those intended to be used by healthcare
71 professionals, were submitted to FDA with claims for OTC home use by lay-users. Sponsors
72 evaluated these devices for self-use by healthy people with diabetes or by healthcare
73 professionals on healthy people with diabetes. However, they were actually being used by
74 healthcare professionals as point-of-care (POC) devices to monitor blood glucose levels in
75 diabetic and non-diabetic patients in various states of health. Scientific and clinical issues
76 specific to the professional healthcare setting, which could affect glucose meter performance,
77 were never evaluated for these devices. Use of BGMSs in professional healthcare settings on
78 patients in various states of health and receiving intensive medical intervention and therapy,
79 when they were evaluated and cleared based on studies performed in healthy subjects, can put
80 patients at risk. Therefore, when devices are intended for use in professional healthcare
81 settings, the intended use population should be accurately defined, distinct performance
82 parameters should be met, and sponsors should demonstrate substantial equivalence of the
83 device for that particular use.

84

85 The intent of this guidance is to describe the studies that should be conducted to demonstrate
86 BGMS performance for devices which are intended to be used in diverse professional
87 healthcare settings on subjects in various states of health. Intended use populations for a BGMS
88 may include patients in all professional healthcare settings, patients in specific healthcare
89 settings (e.g., in emergency response vehicles), patients in long-term care facilities, or patients
90 at a physician's office. The Agency expects that not all sponsors will seek clearance for their
91 device to be used across all professional healthcare settings. BGMSs intended for POC use in
92 specific professional healthcare settings should be studied in those specific populations in
93 accordance with the recommendations in this guidance, and labeled appropriately. For BGMSs
94 intended for use in many or all professional healthcare settings, it may be necessary to identify
95 sub-populations in which the BGMS may function differently than in the broader intended use

³ Thompson, N.D. and Perez, J.F. (2009) Eliminating the blood: Ongoing outbreaks of hepatitis B virus infection and the need for innovative glucose monitoring technologies. *Journal of Diabetes Science and Technology*. 3(2), 283-288.

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96 population. The identification and study of patient subpopulations is described in greater detail in
97 Section VI.C below. In all cases, performance studies should account for factors such as
98 disease state, patient condition, physiological state, and medications that might affect device
99 performance in the intended use population for that BGMS.

100 *CLIA waiver of professional use meters*

101

102 FDA’s clearance of a 510(k) submission for SMBGs intended for OTC home use allows
103 automatic CLIA waived categorization (see 42 U.S.C. 263a(d)(3)). As described above, most
104 blood glucose meters on the market today, even those used in healthcare professional settings,
105 were previously submitted to FDA with claims for OTC use by lay-users and were therefore
106 given CLIA waived categorization pursuant to regulation (see 42 CFR 493.15). The use of
107 blood glucose meters cleared for OTC use in professional healthcare settings poses a number of
108 additional risks to patients, as described above. By contrast, clearance of BGMSs as
109 prescription devices intended for point-of-care use in professional healthcare settings, as
110 described in this guidance, means that FDA expects that clearance of BGMSs for prescription
111 point-of-care use will be categorized upon clearance as moderate complexity. However, FDA
112 recognizes the importance of having CLIA-waived BGMSs in point-of-care professional
113 healthcare settings and intends, through the studies described in this guidance, to facilitate CLIA
114 waiver for these devices by recommending that the information described below be submitted in
115 a dual 510(k)/CLIA waiver submission or an associated application for CLIA Waiver, and
116 enabling BGMSs to be CLIA waived concurrently with their 510(k) clearance.⁴

117

118 FDA has proposed several studies in this guidance that can be performed in a way that will
119 allow sponsors to request FDA review of both their 510(k) submission and CLIA waiver for
120 their BGMSs concurrently. For example, the recommended number of samples (for each
121 sample type: arterial, venous, capillary blood) requested in the Method Comparison/User
122 Evaluation (Section VI-C below) is specifically recommended to allow results from this study to
123 be used to support a CLIA waiver application. The sponsor should plan to conduct these studies
124 using untrained intended users in a CLIA waived setting. Based on feedback from the clinical
125 community, we understand that because of the settings in which these devices are used, and the
126 types of users who use them in clinical practice, it is beneficial to patients and the healthcare
127 community that BGMSs be CLIA waived; therefore, sponsors should design their studies with
128 CLIA waiver in mind. We recommend that sponsors refer to FDA’s guidance entitled
129 [“Recommendations: Clinical Laboratory Improvement Amendments of 1988 \(CLIA\) Waiver](#)
130 [Applications for Manufacturers of In Vitro Diagnostic Devices”](#)⁵ to further understand how the
131 studies described and recommended in this guidance can be performed to support CLIA waived
132 status. We also encourage sponsors to contact the Agency with questions prior to starting their

⁴ For information on FDA’s CLIA administrative procedures, see FDA’s guidance entitled
“Administrative Procedures for CLIA Categorization – Guidance for Industry and Food and Drug
Administration Staff”. (<http://www.fda.gov/RegulatoryInformation/Guidances/ucm070762.htm>).

⁵<https://www.fda.gov/downloads/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocuments/ucm070890.pdf>

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133 studies to ensure that the studies they plan to perform are designed to support CLIA waived use
134 of their device.⁶
135

136 **III. Scope**

137
138 This guidance document is limited to BGMSs which are regulated under 21 CFR 862.1345.
139

140 The following product codes are within the scope of this guidance document:

- 141 CGA (glucose oxidase method)
- 142 CFR (hexokinase method)
- 143 LFR (glucose dehydrogenase method)

144

145 This document is **not** meant to address the following types of devices:

146

- 147 • SMBGs intended for home use by lay-users (product code NBW). Additional
148 considerations (labeling or other) may be needed for OTC devices.
- 149 • Devices used to screen for and/or to diagnose diabetes (such as clinical chemistry
150 analyzers).
- 151 • Continuous glucose sensors, implanted or external (e.g., continuous glucose monitoring
152 systems (CGMs) or sensors within catheters).
- 153 • Non-invasive glucose measurement devices (i.e., devices that do not require removal of
154 a blood sample from a finger or other anatomical site).
- 155 • Blood glucose test technologies labeled for specialized use (e.g., for automated
156 monitoring to aid in glycemic control protocols).

157

158 While FDA recommends that the information described in this guidance be included in
159 premarket submissions for BGMSs, submissions containing alternative information may be
160 sufficient if able to demonstrate substantial equivalence to a legally marketed predicate device.

161

162 We recommend that you contact the Division of Chemistry and Toxicology Devices in the
163 Office of In Vitro Diagnostics and Radiological Health (OIR) if you have questions regarding
164 alternative intended uses or similar technologies.

165

166 **IV. Reducing the Risk of Bloodborne Pathogen** 167 **Transmission**

168

⁶ Requirement for a Pre-Submission for a Dual 510(k) Waiver by Application
<http://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/IVDRegulatoryAssistance/ucm393233.htm>

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169 Since BGMSs use blood specimens for glucose measurement, their design and instructions for
170 use are very important factors in reducing the risk of bloodborne pathogen transmission during
171 use. This is especially important for blood glucose meters used in professional settings which
172 may be used in the care of multiple patients. According to the Centers for Medicare and
173 Medicaid Services (CMS) and the Centers for Disease Control and Prevention (CDC), blood
174 glucose meters can transmit bloodborne pathogens, such as viral hepatitis, if these devices are
175 contaminated with blood and are shared between users without effective cleaning and
176 disinfection.⁷ To minimize the risk of bloodborne pathogen transmission you should address the
177 following in your device design and labeling:

- 178
- 179 • Meters should be designed such that all external materials can be cleaned (removal of
180 organic soil) and disinfected (microbicidal process).
- 181 • All external surfaces of the meter, including seams and the test strip port, should be
182 designed for both ease of use and ease of cleaning and disinfection.
- 183 • You should develop an effective disinfection method and provide the validated cleaning
184 and disinfection procedures for your BGMS device in your 510(k) submission as well as
185 in the labeling. Cleaning and disinfection are different processes and need separate
186 validation procedures and specifications.
- 187 • You should validate the efficacy of any disinfectant you recommend for use with your
188 device, as described below. We recommend you consult the Environmental Protection
189 Agency’s (EPA) list of disinfectants that are registered for use against infectious
190 bacteria and viruses in choosing disinfectants to validate for use with your device.⁸
- 191 • Your BGMS device should be intended for use with only auto-disabling, single use
192 lancing devices. Single use lancing devices are designed to be used only once, after
193 which the blade is retracted, capped, or otherwise made unusable. The auto-disabling,
194 single use lancing device you recommend for use with your BGMS device should be
195 specified in your labeling. You should emphasize in the labeling that lancing devices are
196 for single patient use and should NEVER be used for more than one person. Your
197 labeling should instruct users to discard lancing devices in designated sharps containers.
- 198 • Labeling concerning safe device use can reduce the risk of user error. Therefore,
199 instructions for cleaning and disinfection should be clear and detailed. The various test
200 system components should be named in such a way that they are recognized as
201 belonging to the same system or family of products, and to distinguish them from similar
202 devices intended for single-patient use only (e.g., ABC blood glucose test system, ABC
203 blood glucose meter, ABC blood glucose test strips, etc.). See Section X Labeling below
204 for detailed labeling recommendations. For additional information on labeling your
205 reusable medical device, see FDA’s guidance entitled “Reprocessing Medical Devices
206 in Health Care Settings: Validation Methods and Labeling”

⁷ “Infection Prevention during Blood Glucose Monitoring and Insulin Administration”
<http://www.cdc.gov/injectionsafety/blood-glucose-monitoring.html>.

⁸ Selected EPA-registered Disinfectants available at <http://www.epa.gov/oppad001/chemregindex.htm>.

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207 (http://www.fda.gov/downloads/medicaldevices/deviceregulationandguidance/guidanced
208 ocuments/ucm253010.pdf).

209

210 Validation of cleaning and disinfection procedures involves determining both that the cleaning
211 and disinfection products are effective against the primary viruses of concern (Human
212 Immunodeficiency Virus (HIV), Hepatitis B, Hepatitis C) and that the cleaning and disinfection
213 procedures do not deteriorate the device or alter device performance. FDA’s recommendations
214 for such validation procedures are outlined in the following Subsections.

215 ***A. Validated cleaning and disinfection procedures***

216 You should select cleaning and disinfection products that do not result in physical
217 deterioration of the device overall, or any device component such as the housing, touch pad,
218 or buttons. You should make note of any physical indicators of deterioration during your
219 validation study and provide this information for our review in your 510(k) submission. The
220 disinfectant product you choose should be effective against HIV, Hepatitis B, and Hepatitis
221 C viruses. Of these viruses, Hepatitis B virus is the most difficult to kill and prior outbreak
222 episodes associated with blood glucose meters have been due to transmission of Hepatitis B
223 viruses. Therefore, disinfection efficacy studies should be performed to demonstrate
224 effectiveness of the chosen disinfectant against Hepatitis B virus. Please note that 70%
225 ethanol solutions are not effective against viral bloodborne pathogens, and the use of 10%
226 bleach solutions may lead to physical degradation of your device.

227

228 To demonstrate that your disinfection procedure is effective against Hepatitis B virus, you
229 should perform disinfection efficacy studies to demonstrate that your procedure is effective
230 with the external meter materials (e.g. case, display, buttons, etc.). Studies have
231 demonstrated that viruses can remain infective for different time periods, depending on the
232 surface. Viral survival may increase or decrease with the number of microbes present on a
233 surface. Increasing amounts of microbes can protect viruses from disinfection, and
234 damaging effects may also result from microbial proteases and fungal enzymes. Factors that
235 influence survival on surfaces include fomite properties, initial viral titer, virus strain,
236 temperature, humidity, and suspending media. The simplest disinfection method would be the
237 use of towelettes pre-saturated with a selected disinfectant. Disinfection with a towelette
238 will reduce the risk of liquid getting into the meter, therefore minimizing the chance of
239 affecting the glucose meter function. However, you should choose a disinfectant that is
240 effective against Hepatitis B virus and is compatible with your device. If you intend to claim
241 that your disinfection protocol is effective against other pathogens, you should consider
242 submitting a pre-submission to discuss this with the Agency prior to conducting your testing.
243 For information about the pre-submission process, see FDA’s guidance entitled “Requests
244 for Feedback on Medical Device Submissions: The Pre-Submission Program and Meetings
245 with Food and Drug Administration Staff

246 ([http://www.fda.gov/downloads/MedicalDevices/DeviceRegulationandGuidance/GuidanceD
247 ocuments/UCM311176.pdf](http://www.fda.gov/downloads/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocuments/UCM311176.pdf)).

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249 We recommend you refer to the following standards when developing a disinfection protocol
250 for your device:

- 251 • ASTM standard ASTM E1053-11: *Standard Test Method for Efficacy of*
252 *Virucidal Agents Intended for Inanimate Environmental Surfaces*
- 253 • ASTM standard ASTM E2362-09: *Standard Practice for Evaluation of Pre-*
254 *saturated or Impregnated Towelettes for Hard Surface Disinfection.*

255 ***B. Demonstration that the device is robust to cleaning and disinfection***
256 ***procedures***

257 You should demonstrate through bench studies that your BGMS device is robust to cleaning
258 and disinfection procedures after multiple cleaning and disinfection cycles. You should
259 describe in your 510(k) submission the study design and results demonstrating that the
260 analytical performance of the BGMS is not impacted by the cleaning and disinfection
261 procedures.

262
263 You should address the following in your study design:

- 264
265 • Worst case scenarios with regards to cleaning and disinfection frequency and end
266 user environment should be used to determine the number of cleaning and
267 disinfection cycles that should be tested. For example, the number of times you
268 clean and disinfect the meter should be representative of the cleaning and
269 disinfection that the meter will be exposed to during its use life (typically 3-5 years).
270 A cleaning step should precede the disinfection step for each cleaning and
271 disinfection cycle.
- 272 • The disinfection contact time used in the robustness study should be identical to the
273 contact time used in the disinfection efficacy testing and described in your cleaning
274 and disinfection instructions in the labeling.
- 275 • We recommend using the same disinfectant product for both cleaning and
276 disinfection. The effects of using multiple cleaning products on the efficacy of
277 disinfectant products are not well understood.
- 278 • You should demonstrate that the test strip port and all other openings which are
279 susceptible to blood contamination and could either directly or indirectly be contacted
280 during use are able to withstand your cleaning and disinfection procedures. You
281 should ensure that you test parts of the meter that are particularly susceptible to
282 blood contamination, such as the test strip port and material seams. It is important to
283 be able to clean and disinfect all parts of your meter to reduce the risk of bloodborne
284 pathogen transmission.
- 285 • When evaluating your device after the cleaning and disinfection phase, you should
286 ensure that the procedure does not cloud or deface the display of the meter and
287 does not corrode or erode the plastic housing or buttons. All these physical indicators
288 of deterioration should be noted throughout your study and included in your 510(k)
289 submission. You should evaluate the accuracy of the meter using blood samples

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290 compared to results obtained by a comparator method (please refer to Section VI
291 below for the definition of comparator method) to ensure that accuracy is not
292 affected by repeated cleaning and disinfection. The study should also evaluate the
293 functionality of meter features (as appropriate), for example, touch screen function,
294 USB port function, speaking functions, etc., to ensure they are not affected by
295 repeated cleaning and disinfection.

- 296 • You should include infection control in your risk analyses and incorporate your
297 validated cleaning and disinfecting procedures into your risk assessment.

298

299 A description of the protocols and acceptance criteria for all studies should be included in
300 your 510(k) submission.

301

302 **V. Device Description**

303

304 You should provide the following information in the device description portion of your 510(k)
305 submission:

306

- 307 • Description of physical components of the system (including diagrams where
308 appropriate).
- 309 • Manufacturer's performance specifications.
- 310 • Description and explanation of the test principle, including chemical reactions.
- 311 • Description of the format of results, including units of measurement and whether results
312 are reported in whole blood or plasma equivalents.⁹
- 313 • Description of the composition and levels of control material recommended for use with
314 your system.
- 315 • User maintenance needs (e.g., batteries).
- 316 • Features of the device, such as data transmission capabilities or features designed to
317 enhance robustness and ease of use.
- 318 • Features designed to minimize the risk of bloodborne pathogen transmission among
319 patients.
- 320 • Description of features controlled by the software, which should describe the following:
 - 321 ○ Displays and user messages: This includes how the BGMS determines and displays
322 the glucose concentration, messages or displays that appear while a user is taking a
323 measurement, and features such as how a user can retrieve past results from
324 storage in the device.
 - 325 ○ User prompts: You should describe prompts that the BGMS provides to the user,
326 expected user responses, and timing issues (e.g., how quickly does the user need to

⁹ Note that BGMSs intended for use in the U.S. should report results in mg/dL and in plasma equivalents.

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327 respond, what happens if they respond after the allowed time). Examples of a user
328 prompt include messages to the user to add specimen to the test strip, insert the test
329 strip into the meter, calibrate the meter, or store a result, etc.

- 330 ○ Error messages and alerts: This includes any error messages or alerts that the
331 BGMS displays. You should describe how the system responds to errors in user
332 action, user inaction, or system status. Suggested examples of error messages or
333 alerts include: when a test strip is inserted incorrectly or removed prematurely; too
334 small a sample is applied to the test strip; damaged, incorrect or deteriorated test
335 strips are used; or when there is a low battery or excessively high ambient
336 temperature. This should also include the methods by which the BGMS detects and
337 alerts the user when glucose levels are outside of the linear range of the system.
338 You should describe at what point each message is triggered and describe any self-
339 diagnostic routines that the system performs.

340

341 It is important that you identify the expected responses by the user to the error messages or
342 alerts. This includes whether and how the user should input information or press certain buttons
343 to correctly set up the meter or respond to an error message or alert.

344

345 **VI. Performance Evaluation for Prescription-Use** 346 **BGMSs**

347

348 Subsections A-F below indicate the types of device performance information that you should
349 include in a 510(k) submission for a BGMS. Although many manufacturers design their BGMS
350 validation studies based on the International Standards Organizations document 15197: *In vitro*
351 *diagnostic test systems—Requirements for blood glucose monitoring systems for self-*
352 *testing in managing diabetes mellitus*, FDA believes that the criteria set forth in the ISO
353 15197 standard do not adequately protect patients using BGMSs in professional settings, and
354 does not recommend using the criteria in ISO 15197 for BGMSs.

355

356 In this guidance, the term “comparator method” refers to a laboratory-based glucose
357 measurement method that has been well-validated for precision and accuracy, and that is
358 traceable to a higher order, e.g., an internationally recognized reference material and/or method.
359 The traceability chain should include as few stages as possible to reduce bias. FDA’s current
360 thinking on the recommended study designs and device performance criteria are discussed
361 below in Subsections A-F.

362 **A. Precision Evaluation Study**

363 You should evaluate both within-run precision and intermediate precision for your BGMS
364 and include these evaluations in your 510(k) submission. The following sections outline
365 FDA’s current thinking on appropriate study design and analyses to evaluate within-run and
366 intermediate precision for BGMSs.

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Within-Run Precision Evaluation:

In this guidance, within-run precision studies are bench studies designed to evaluate imprecision under conditions of repeated measurement of the same sample with different meters and multiple test strip lots. In order to assess imprecision of the device across the claimed measuring range, you should evaluate samples containing glucose concentrations within each of the five intervals provided in Table 1 below.

Table 1. Glucose Concentrations for Precision Evaluations

Interval	Glucose Concentration Range (mg/dL)
1	30-50
2	51-110
3	111-150
4	151-250
5	251-400

376
377
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You should determine within-run precision using venous whole blood samples. Altered venous whole blood samples such as those that are spiked, diluted, or allowed to glycolyze in order to obtain the appropriate glucose concentrations are acceptable in order to facilitate coverage of the entire claimed glucose measuring range. However, you should clearly identify all altered samples (spiked, diluted, or glycolyzed) in all submitted data. A minimum of 500 test strips from at least 10 vials and 3 manufacturing lots should be used in this study. For each sample concentration, a minimum of 10 meters should be used, with at least 10 measurements taken by each meter (i.e., at least 100 measurements per concentration). Test strips should be taken from the same vial and/or package for each meter.

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We recommend you present the results as the mean value of all measurements per meter for each glucose concentration with the corresponding standard deviation (SD) and percent coefficient of variation (CV). In addition, for each glucose concentration range in Table 1, you should also provide the mean value, standard deviation (with 95% confidence intervals), and percent CV for data combined over all meters. You should describe the statistical procedures used in the analysis. You should provide the results based on all data, and if you wish to exclude any data points (outliers), a separate, additional data analysis with those points excluded and a full description of the method of outlier identification and the results of your investigations into those outliers, should be included

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Intermediate Precision Evaluation:

Intermediate precision measurement studies are bench studies designed to evaluate imprecision under simulated normal use conditions; for example, measurement by multiple operators over multiple days using multiple reagent system lots. These studies may be performed with prepared control solutions rather than whole blood samples.

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403 The total number of meters and individual operators in these studies is at the discretion of
404 the sponsor; however, a minimum of 10 meters should be used for each glucose
405 concentration. Intermediate precision should be evaluated over a minimum of 10 days,
406 taking at least 1 measurement per meter per day of a sample from each glucose
407 concentration interval listed in Table 1. This should produce a minimum of 10 measurements
408 per meter for each glucose concentration and 100 total measurements per glucose
409 concentration. You should use a minimum of 500 test strips from a minimum of 10 vials or
410 packages that cover a minimum of 3 manufacturing lots. These test strips should be taken
411 from the same vial and/or package for each meter.

412
413 For each glucose concentration in Table 1, you should present data for each test strip lot, as
414 well as for pooled lots, including the mean value of the measurements for each meter with
415 the corresponding standard deviation (SD) and percent coefficient of variation (CV). You
416 should also present the mean value, standard deviation (with 95% confidence intervals), and
417 percent CV for data combined over all meters. You should describe the statistical
418 procedures you use and provide results based on all data. If you wish to exclude any data
419 points, a separate, additional data analysis with those points excluded and a full description
420 of the method of outlier identification and the results of your investigations into those
421 outliers, should be included.

422 **B. *Linearity Evaluation Study***

423 You should evaluate the linearity of your BGMS across the entire claimed measuring
424 range. We recommend that studies include an evaluation of at least 11 evenly spaced
425 concentrations tested and analyzed according to the guideline “Evaluation of the Linearity
426 of Quantitative Measurement Procedures: A Statistical Approach,” CLSI document EP6-
427 A. Linearity studies should be performed using venous whole blood samples. Altered
428 venous whole blood samples such as those that are spiked, diluted, or glycolyzed are
429 acceptable in order to facilitate coverage of the entire glucose concentration range. You
430 should clearly identify the number of altered samples (spiked, diluted, or glycolyzed) within
431 your 510(k) submission.

432
433 You should submit a detailed description of the study design, target concentrations, a list of
434 all data collected in this study, summary of the results and conclusions drawn from the
435 study, and a description of the statistical analysis used.

436 **C. *Method Comparison/User Evaluation***

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438

1. *General Study Design*

439 When testing samples from the intended patient population, you should design your study to
440 accurately reflect system performance in the hands of the intended user. You should
441 perform a set of comprehensive clinical evaluations to assess system accuracy to support
442 the professional use of these devices in the intended use population.

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444 FDA recognizes that most study evaluations performed for 510(k) submissions occur in
445 idealized conditions, thereby potentially overestimating the total accuracy of the BGMS,
446 even when performed in the hands of the intended user. Nonetheless, it is important that you
447 design your study to most accurately evaluate how the device will perform in the intended
448 use population. Therefore, the study should be conducted in conditions that reflect the
449 expected use of the device, as well as = environmental conditions that are consistent with
450 the validated environmental conditions of the device (e.g., temperature, humidity, altitude,
451 etc.). You should fully describe the conditions of your study in your 510(k) submission.

452
453 You should evaluate device accuracy for each claimed sample type (e.g., arterial, venous,
454 capillary, heelstick whole blood, etc.) when the device is used by a POC operator.
455 Evaluation of each sample type should include a minimum of 350 patients (e.g., samples
456 from at least 350 patients for an arterial study, samples from at least 350 patients for a
457 capillary study, samples from at least 350 patients for a venous study, etc.). FDA
458 recommends sponsors perform their studies to support concurrent CLIA waiver at the time
459 of clearance by performing the studies as described in this guidance with consideration to
460 the aspects of study design described in FDA’s guidance entitled “Recommendations:
461 Clinical Laboratory Improvement Amendments of 1988 (CLIA) Waiver Applications for
462 Manufacturers of In Vitro Diagnostic Devices”
463 (<http://www.fda.gov/RegulatoryInformation/Guidances/ucm079632.htm>).¹⁰ Different sample
464 types (e.g., arterial and venous) can be acquired from the same patient and be used in the
465 different studies. Each sample should be fresh and measured on both the candidate device
466 (i.e., new device) and the comparator method. Samples do not have to be collected
467 specifically for your studies; however, to obtain CLIA waiver, the tests should be done per
468 the labeling instructions by untrained users typical of CLIA-waived users. Note that patient
469 information should be available for each sample to aid in the identification of potential
470 interfering factors. In order to robustly assess the accuracy of your device, it is important
471 that the glucose value on the comparator method be as reliable as possible. Therefore, more
472 than one comparator measurement may be taken and averaged for each sample in order to
473 allow a better estimate of the true glucose value of that sample. However, no
474 measurements should be excluded from the 510(k) submission and a justification should be
475 provided for any data that is excluded from the analysis. It is not necessary that POC
476 operators perform the comparator method measurements in the study.

477
478 For each claimed sample type, the samples tested should adequately span the claimed
479 glucose measuring range of the BGMS device. Though it may be difficult to obtain samples
480 at the extreme ends of the measuring range, the study for each sample matrix should
481 contain at least 10 unaltered samples < 80 mg/dL and at least 10 unaltered samples between
482 300 mg/dL and the upper limit of the claimed measuring range of the device. It may be
483 necessary to enroll more than 350 patients for each sample type in order to obtain the

¹⁰ For example, users should be untrained, and the studies should be performed in intended use settings in the midst of normal working conditions. Please note that we intend to accept 350 patient samples for each sample type for the purposes of CLIA waiver studies for these devices.

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484 necessary unaltered samples. Testing should be performed by the intended POC operators
485 (e.g., nurses, nurse assistants, etc.) to accurately reflect device performance in POC
486 settings; at least 9 operators should participate in each study (e.g., capillary, venous, and
487 arterial studies). Different operators may be used for each study. You should submit data
488 from all subjects; no data or subjects should be excluded from your analysis.

489
490 The subjects you enroll in the method comparison/user study should accurately reflect the
491 intended use population of your device. In your 510(k) submission, you should describe the
492 inclusion and exclusion criteria for enrolling study subjects, as well as the demographics of
493 the subjects that participated in the study. If your intended use population is broad but
494 includes patient sub-populations that might be particularly vulnerable to potential
495 interferences and/or health risks resulting from meter inaccuracy, you should identify and
496 include patients from these specific vulnerable sub-populations in your study. You should
497 define these sub-populations and provide a rationale for your definitions. For example,
498 vulnerable sub-populations could be defined as patients in specific hospital wards, units, or
499 departments— medical, neonatal, pediatric or surgical intensive care units (ICUs).
500 Vulnerable subpopulations could, for example, also be defined as categories of patients with
501 general types of medical conditions—cardiac, surgical, pulmonary, or oncology patients.
502 These sub-populations are provided as an example of common patient groups found in a
503 hospital setting, however, if you would like to discuss other sub-populations or other aspects
504 of your study with the Agency, we recommend that you submit a request for a Pre-
505 Submission meeting prior to conducting your testing. For information about the pre-
506 submission process, see FDA’s guidance entitled “Requests for Feedback on Medical
507 Device Submissions: The Pre-Submission Program and Meetings with Food and Drug
508 Administration Staff
509 ([http://www.fda.gov/downloads/MedicalDevices/DeviceRegulationandGuidance/GuidanceD](http://www.fda.gov/downloads/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocuments/UCM311176.pdf)
510 [ocuments/UCM311176.pdf](http://www.fda.gov/downloads/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocuments/UCM311176.pdf)).

511
512 Your study should include at least 50 patients from each vulnerable patient sub-population
513 you have defined. in order to collect sufficient performance data to support the use of your
514 device in these populations. Please note that in some instances, in order to fully characterize
515 your device in your intended use population, FDA suggests that you use more than 50
516 patients per subpopulation to ensure all conditions and medications have been evaluated.
517 Furthermore, if you intend for your study to also support a broad intended use population,
518 then you should ensure that your study population includes sufficient numbers of patients
519 outside of the vulnerable subpopulations you identified to support the use of your device in
520 the broader intended use population. This broader population might include in-patients
521 dispersed throughout various hospital departments. Depending on the number of specific
522 vulnerable sub-populations you identify, the collection of samples from more than 350
523 subjects for each sample type (venous, arterial, capillary) may be indicated to support the
524 use of your device in your intended use population. Your results should clearly indicate the
525 specific patient population associated with each sample and you should present the
526 combined results for your entire intended use population and, separately, for each vulnerable
527 patient subpopulation (if present).

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If you wish to claim suitability of your device with anti-coagulants, then within the 350 (minimum) samples you collect for each sample type you should include at least 50 to 75 patient samples per claimed anti-coagulant.

Your study should include a minimum of 10 test strip vials or packages that cover a minimum of 3 test strip lots. All test strips used in the study should have undergone typical shipping and handling conditions from the site of manufacture to a U.S. user prior to the study. You should describe these shipping and handling conditions in your 510(k) submission.

Method comparison and user performance studies for a BGMS should include multiple users and multiple blood glucose meters. Only auto-disabling, single use lancing devices should be used in these studies. You should incorporate your labeling instructions for cleaning and disinfection in your user study protocol to ensure that the meters are appropriately cleaned and disinfected during the course of this study, and include any additional measures necessary to mitigate the risk of potentially transmitting disease between healthcare providers and subjects during the study (for example, use of disposable gloves or other physical barriers). The study protocol should also include details on how often and when gloves worn by the trained health professionals should be changed between subjects. Meters should be cleaned and disinfected after each subject, using validated procedures for all studies performed. Refer to Section IV, above (Reducing the Risk of Bloodborne Pathogen Transmission) for additional information regarding the validation of cleaning and disinfecting of BGMSs.

BGMS test results are used by healthcare professionals to make critical decisions about treatment; therefore, it is important that the results are accurate so that medical decision errors are better avoided. In order to demonstrate that a BGMS is sufficiently accurate for use by health care professionals, you should demonstrate that 95% of all values are within +/- 12% of the comparator method for glucose concentrations ≥ 75 mg/dL and within +/- 12 mg/dL at glucose concentrations < 75 mg/dL. In addition, 98% of values should be within +/- 15% of the comparator method for glucose concentrations ≥ 75 mg/dL and within +/- 15 mg/dL at glucose concentrations < 75 mg/dL. The BGMS should be as accurate as possible to avoid critical patient management errors. Though we expect that BGMSs will be able to meet these criteria, there may be instances where meters may be determined to be substantially equivalent when performance does not meet these criteria because, for example, other features of the meter or its setting of use provide benefits that compensate for different performance. In instances where your BGMS is unable to meet these criteria, you should provide a clinical justification for all test results, including those that exceed the above mentioned criteria, and describe why the potential for that error would not affect patient safety when extrapolated to the intended use setting (e.g., when extrapolated to the volume of testing performed in the intended use setting). FDA will review your justification to determine whether the data suggest that patients may be put at risk or whether your justification and any proposed mitigations are adequate.

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572 Hematocrit and sodium values should be measured and recorded for each study subject to
573 help identify potential interference with the device and to inform investigations into outlier
574 results. Similarly, blood oxygen levels should be measured and recorded for each patient for
575 any arterial blood study. You should present these individual values in the 510(k) submission
576 along with the BGMS and comparator method results. It is not necessary that hematocrit,
577 sodium, and blood oxygen measurements be made by POC operators.

578
579 We expect that the measuring range of the meter will meet the clinical needs of the
580 intended use population. BGMSs intended for prescription-use in the hospital setting should
581 be able to measure blood glucose accurately down to 10 mg/dL and up to 500 mg/dL, or a
582 clinical justification should be provided for alternate measuring ranges. BGMSs intended for
583 use outside a hospital setting and which will not reasonably be used to test neonatal samples
584 should be able to measure blood glucose accurately down to 20 mg/dL. The BGMS device
585 should identify and provide an error code in situations where the measured glucose level
586 falls outside of the device's stated measuring range. For example, if BGMS XYZ has a
587 measuring range that can detect glucose concentrations down to 10 mg/dL, then blood
588 samples with glucose concentrations below 10 mg/dL should provide an appropriate error
589 code (e.g., "LOW - Less than 10").

590
591 You should describe the following in your 510(k) submission:

- 592
- 593 • Study setting, including the size, type, and location of each site and a justification of
594 how the selected study conditions simulate intended use conditions. Study sites
595 should be representative of where BGMSs are used in the U.S. and you should
596 include an explanation of why you believe each site is representative.
 - 597 • Criteria used to select study subjects.
 - 598 • Description of the patient demographics, including age, disease states, and all
599 medications for each patient.
 - 600 • Sample types collected (arterial, venous, capillary).
 - 601 • Number of test strip lots, number of test strip vials, and number of meters used in
602 the study.
 - 603 • Description of the shipping and handling conditions of the test strips prior to use in
604 the study.

605
606 *Accuracy at Extreme Glucose Values:*

607 Because the study described above using real patient samples may not provide a robust
608 evaluation of BGMS performance in the extreme upper and lower ends of the measuring
609 range, you should perform additional studies using blood samples altered to achieve glucose
610 concentrations of less than 80 mg/dL and greater than 300 mg/dL. This additional extreme
611 glucose value study should be performed separately from the method comparison/user
612 evaluation described above and may be performed in a laboratory setting, though untrained
613 intended users typical of users in a CLIA waived setting should perform the testing to
614 support CLIA waiver of the device.

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616 Your study of accuracy at extreme glucose values should include a minimum of 50 prepared
617 samples with glucose concentrations < 80 mg/dL and a minimum of 50 prepared samples
618 with glucose concentrations > 300 mg/dL. These samples should evenly cover the lower and
619 upper limits of the claimed measuring range. Samples may be altered by spiking or allowing
620 the samples to glycolyze in order to obtain appropriate glucose concentrations. Samples
621 should be measured on both the BGMS device and the comparator method. You should
622 analyze this data separately from the user evaluation data but using the same methods
623 described below for the user evaluation. FDA will apply the same review criteria to both
624 studies.

625

626

Neonatal Studies:

627

628 If your intended use population includes neonates, you should perform studies to support
629 performance in neonatal samples (defined as samples from subjects less than 28 days old).
630 Neonatal blood is known to differ from adult blood and these differences may have a direct
631 impact on the safety of blood glucose monitoring in that population. For example, neonatal
632 blood often has higher hematocrit levels (51 to 65%) and lower blood glucose concentrations
633 (20 to 80 mg/dL) compared to adult blood.

633

634

635 You should evaluate device performance with neonatal samples in direct comparison to the
636 comparator method by testing 100 -150 fresh neonatal blood specimens, including samples
637 from neonates less than 24 hours old. Samples should be collected and measured by at least
638 three POC users in a POC setting. Glucose concentrations should be measured with the
639 BGMS and the comparator method, and the hematocrit levels for each patient should also
640 be measured and reported in the study. You should present your results as described below
641 in the Data Analysis Subsection. Data from all subjects in the study should be submitted in
642 your 510(k), and no subjects should be excluded from the data analysis.

642

643

644 Since it may be difficult to obtain samples at the extreme low end of the measuring range
645 using real neonatal patient samples, you should perform additional studies using blood
646 samples (either adult blood or maternal cord blood) altered to achieve glucose
647 concentrations between 10 and 50 mg/dL. Blood specimens used in these additional studies
648 should be adjusted to at least two levels of hematocrit at or near 40% and 65%, in order to
649 simulate the high hematocrit levels of neonatal blood. This will allow you to provide a robust
650 evaluation of device performance in the extreme lower end of the measuring range for
651 simulated neonatal blood. These additional studies should be performed separately from the
652 neonatal studies described above and may be performed in a laboratory setting (e.g., at the
653 manufacturer's facility), however, untrained users typical of CLIA-waived users should
654 perform the testing to support CLIA waiver of the device.

654

655

2. Data Analysis

656

Data exclusion and outliers:

657

658 You should present all data in the 510(k) submission, including cases in which the meter
displays an error code, a 'High' or 'Low' message, or no result. All outliers (e.g., data

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points that do not conform to minimum accuracy criteria) should also be included in your 510(k) submission. You should investigate all outlier results and describe the results of these investigations, providing explanations for the occurrence of outliers when possible. To help inform your investigations into outlier results, you should collect information regarding patient medications, hematocrit measurements, oxygen levels, and sodium levels during your study. You should include the following in your description of the results:

Analysis of Results:

You should present the difference between individual study subject results and results of the comparator method (or mean of the comparator measurement, if multiple replicates are measured using the comparator method) by plotting the data on an X-Y graph. The plot should include the regression line and line of identity. Your summary of results should include the slope and y-intercept, along with 95% confidence intervals, calculated using a suitable analysis procedure (e.g., Linear Regression, Deming regression), and the estimate of the deviation (standard error). Difference plot of Y-X vs X analysis may also be presented. You should describe all statistical methods used and clearly identify and describe any outliers in the analysis.

Tabular data presentation:

You should present results in the following tabular format for each sample matrix. In Table 2 and Table 3 below, X= the number of samples within the specified difference from the comparator method, and Y= total number of samples.

Summary of data within specified mg/dL of the comparator method.

Table 2. For glucose concentrations <75 mg/dL:

Within +/- 5 mg/dL	Within +/- 10 mg/dL	Within +/- 12 mg/dL	Within +/- 15 mg/dL	Exceeds +/- 15 mg/dL
X/Y (%)	X/Y (%)	X/Y (%)	X/Y (%)	X/Y (%)

Table 3. For glucose concentrations ≥75 mg/dL:

Within +/- 5%	Within +/- 10%	Within +/- 12%	Within +/- 15%	Within +/- 20%	Exceeds +/- 20%
X/Y (%)	X/Y (%)	X/Y (%)	X/Y (%)	X/Y (%)	X/Y (%)

D. Interference Evaluation

You should evaluate the effect of potentially interfering endogenous and exogenous substances and conditions, such as icterus, lipemia, and varying hematocrit levels, as well as the effect of common medications on your device's performance. Conditions that are known to interfere with glucose monitoring test systems, such as ketoacidosis, should be

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696 included in the labeling as limitations unless you have provided data demonstrating that these
697 conditions do not interfere with your device.

698
699 **1. Endogenous/Exogenous Substances**

700 *Study design:*

701 You should perform interference testing using samples containing glucose concentrations
702 across the range of the device. Specifically, testing should be performed in samples with
703 target glucose values of approximately between 50 - 70 mg/dL, 110-130 mg/dL, and 225-
704 270 mg/dL to evaluate clinically relevant decision points.

705
706 You should evaluate each potentially interfering substance at clinically relevant
707 concentrations. When performing your studies, you should test all substances at the highest
708 concentration that could potentially be observed in a whole blood sample; if interference is
709 observed, you should perform dilutions of the interferent to determine the concentration at
710 which interference begins to occur. For example, if interference is observed with 20 mg/dL
711 acetaminophen, additional testing should be performed with samples containing lower
712 concentrations of acetaminophen, such as 15 mg/dL, 10 mg/dL and 5 mg/dL, to determine
713 the lowest concentration of acetaminophen where interference is first observed. If the
714 results from the additional testing determine that interference is not observed in the sample
715 containing 5 mg/dL acetaminophen and interference is observed in the sample containing 10
716 mg/dL acetaminophen, then 5 mg/dL is the highest concentration of acetaminophen where
717 no interference is observed.

718
719 The substances listed below in Table 4 represent known or potential interferents for current
720 blood glucose measurement technologies and comprise the minimal list of substances that
721 should be tested for interference.

722
723 **Table 4. List of Known or Potential Interferents for BGMSs:**

Interferent	Recommended Test Concentration
Acetaminophen	20 mg/dL
Ascorbic acid	6 mg/dL
Conjugated Bilirubin	50 mg/dL
Unconjugated Bilirubin	40 mg/dL
Cholesterol	500 mg/dL
Creatinine	15 mg/dL
Dopamine	0.09 mg/dL
EDTA*	0.1 mg/dL
Galactose	60 mg/dL
Gentisic acid	1.8 mg/dL
Reduced Glutathione	4.6 mg/dL
Hemoglobin	1000 g/dL
Heparin*	300 IU/dL

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Interferent	Recommended Test Concentration
Ibuprofen	50 mg/dL
L-Dopa	0.75 mg/dL
Maltose	480 mg/dL
Mannitol	1800 mg/dL
Methyldopa	2 mg/dL
Salicylic acid	60 mg/dL
Sodium	180 mmol/L
Tolbutamide	72 mg/dL
Tolazamide	9 mg/dL
Triglycerides	1500 mg/dL
Uric acid	23.5 mg/dL
Xylose	600 mg/dL
Sugar Alcohols**	0.09 mg/dL

724 *The inclusion of EDTA and Heparin in this table refers to their use as therapeutic substances and
725 not as anticoagulants for sample preparation. Separate studies should be performed to validate the
726 use of these substances as anticoagulants used for sample preparation (as described in Section C,
727 above).

728 **All common sugar alcohols, including but not necessarily limited to, sorbitol, xylitol, lactitol, isomalt,
729 maltitol should be independently tested.

730
731

732 In addition to the list of potential interferents provided in Table 4, you should conduct an
733 interference risk analysis and carry out bench studies to evaluate interference from
734 additional drugs commonly used in your intended use population. These bench studies of
735 additional drugs should be conducted in the same manner described in this Section.

736

737 You should provide a reliable estimate of the interference predicted for each potential
738 interferent. To do this, we recommend the following method of measuring and calculating
739 interference. First, blood samples should be generated at each target glucose concentration
740 described above. Each glucose sample should be tested in replicates with the comparator
741 method (we suggest at least 4 replicates in order to reduce standard error) to establish the
742 glucose concentration in the sample. The glucose samples should then be split into a test
743 sample to which a specific amount of potential interferent is added and a control sample
744 containing solvent/vehicle in lieu of the potential interfering substance. Both control samples
745 and test samples should be measured in replicates on the BGMS. At least three test strip
746 lots should be used for this evaluation. Each of the control and test samples should be tested
747 on your BGMS in replicates of 30 across the three lots (10 replicates per lot of test strips for
748 a total of 30 replicates per sample). The mean of replicates should be calculated for each
749 control and test sample. The relative bias (mg/dL) and percent bias should be calculated
750 using the results of the control sample relative to test sample for each concentration of
751 potential interferent. These results should be submitted with 95% confidence intervals as
752 part of your 510(k) submission.

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754 For BGMSs, the degree of acceptable interference may vary by substance tested and the
755 intended patient population of your device. Therefore, you should report in your 510(k)
756 submission the interference testing data as well as the expected imprecision of the system at
757 that glucose concentration. If interferences are observed, you should propose appropriate
758 labeling to address any observed interferences; the labeling language appropriate for the
759 observed interference will be discussed during the review of the 510(k) submission.

760

761 As new drugs are developed that could potentially interfere with your device, or new
762 interfering substances are identified for other BGMSs, you should evaluate these new drugs
763 or substances for potential interference with your device. For example, if a new drug
764 intended to treat cardiac complications in diabetic patients is approved, you should conduct a
765 careful evaluation to determine whether the new drug interferes with your device. You
766 should report to FDA if significant new interferences are observed with your device or with
767 any cleared glucose monitoring devices that are on the market. New drugs/potential
768 interferences should also be evaluated when new or significantly modified technology is
769 introduced.

770

771 *Data Analysis:*

772 You should provide raw data sets as well as a summary table for all interference results.
773 Please note that the summary tables should be presented separately for each test strip lot
774 and for all lots pooled for each glucose level tested. Table 5 below provides a sample format
775 of a summary table.

776

777 **Table 5. Recommended Summary Table Format:**

778

Test Strip Lot #(s)

Interferent	Mean Glucose Value (Comparator)	Interferent Concentration (mg/dL)	Control Sample Mean	Test Sample Mean	Bias (mg/dL)	% Bias	Confidence Interval around % Bias
Acetaminophen	60 mg/dL	20 mg/dL					
	120 mg/dL	20 mg/dL					
	250 mg/dL	20 mg/dL					

779

780 In your 510(k) submission, you should include a detailed description of the study design, a list
781 of all data collected in this study, the summary tables indicated above, and a description of
782 the conclusions drawn from the study.

783

784 **2. Hematocrit**

785 *Study design:*

786 Because a reasonably sized method comparison study may not include the full range of
787 hematocrit values expected in the intended use population, you should perform a separate
788 study to determine how much analytical error is contributed by varying hematocrit levels.

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789 This should constitute a bench study designed to evaluate the effect of hematocrit on the
790 performance of your BGMS to assess whether your device can safely be used across the
791 claimed hematocrit range in the intended use population. The observed hematocrit levels
792 may be very broad in the intended use population for this type of device; the intended patient
793 population may reasonably be expected to have hematocrit levels between 10 and 65%.
794 Therefore, we recommend a minimum hematocrit range of 10-65% as the claimed range for
795 BGMSs.

796
797 You should evaluate hematocrit interference by measuring samples containing various
798 glucose concentrations. The samples should be prepared to contain designated levels of
799 hematocrit that span the claimed hematocrit range for the device. Blood samples may be
800 altered by spiking or allowing them to glycolyze to obtain desired glucose concentrations.
801 Specific percentages of hematocrit may be achieved for each sample by manipulating the
802 plasma to packed cell ratio following centrifugation. Hematocrit levels tested should span
803 the claimed range in 5% intervals, as such 5% intervals allow for a more accurate
804 assessment of bias from hematocrit interference than using broader intervals. Additionally, a
805 sample having a nominal hematocrit of 42% should be tested. For example, if your claimed
806 hematocrit range is from 10-65%, you should test samples at 10, 15, 20, 25, 30, 35, 42, 50,
807 55, 60 and 65% hematocrit. The samples should also span the claimed measuring range for
808 blood glucose. Samples should include 5 different blood glucose concentrations evenly
809 spread and targeted to the following ranges: 30 – 50, 51 – 110, 111 – 150, 151 – 250, and
810 251 – 400 mg/dL.

811
812 Each sample should be tested on the comparator method in multiple replicates (we
813 recommend a minimum of 4 replicates). A mean of the comparator measurements
814 ($\text{Mean}_{\text{Comp}}$) should give greater confidence in the true glucose concentration of the sample.
815 You should test a minimum of 3 test strip lots to evaluate interference from hematocrit.
816 Each sample should be tested on your new BGMS in replicates of 30 (10 replicates per lot
817 of test strips for a total of 30 replicates per sample).

818
819 *Data Analysis:*

820 An analysis should be performed for each of the 5 blood glucose concentrations tested and
821 each test strip lot. The bias should first be determined with respect to the comparator
822 method and then with respect to the nominal hematocrit samples, so that the hematocrit
823 effect can be isolated.

824
825 *(1) Estimation of Bias to Comparator Method*

826 For each sample, you should calculate the average of 30 replicates of your new BGMS
827 ($\text{Mean}_{\text{BGMS}}$). Using the $\text{Mean}_{\text{BGMS}}$ and the estimate of the true glucose concentration in
828 the sample, $\text{Mean}_{\text{Comp}}$, you should estimate a bias and percent bias as $(\text{Mean}_{\text{BGMS}} -$
829 $\text{Mean}_{\text{Comp}})$ and $(\text{Mean}_{\text{BGMS}} - \text{Mean}_{\text{Comp}}) / \text{Mean}_{\text{Comp}}$, correspondingly, for each sample.
830 The results should be presented as in the table below and in graphical format appropriate
831 for each specific glucose concentration range.
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For glucose concentrations less than 75 mg/dL, the analysis should be presented as a graph where the X-axis represents hematocrit values and the Y-axis represents the absolute bias values. For glucose concentrations greater than or equal to 75 mg/dL, the analysis should be presented as a graph where the X-axis represents hematocrit values and the Y-axis represents percent bias values.

Table 5. Example table of bias calculated versus the comparator method for the hematocrit evaluation on a BGMS with 120 mg/dL glucose:

Hematocrit (%)	Average of Comparator measurements (Mean _{Comp})	Number of measurements for BGMS	Average of BGMS measurements (Mean _{BGMS})	% Bias (Mean _{BGMS} -Mean _{Comp})/Mean _{Comp}
10	118.0	30	127.6	8.1%
15	118.4	30	127.6	7.8%
20	122.4	30	130.4	6.5%
25	120.7	30	127.1	5.3%
30	123.7	30	129.5	4.7%
35	121.5	30	127.1	4.6%
42	119.7	30	124.6	4.1%
50	121.3	30	125.4	3.4%
55	120.8	30	122.7	1.6%
60	120.1	30	119.5	-0.5%
65	118.1	30	116.0	-1.8%
70	117.5	30	115.6	-1.6%

(2) Estimation of Bias due to Hematocrit

In order to isolate the effect of hematocrit on device performance, the bias relative to a sample having a nominal hematocrit (42%) should be determined. This nominal hematocrit is representative of the average hematocrit value of the intended use population, and BGMSs are designed to perform optimally with such samples; therefore, bias due to hematocrit is considered 0% (or 0 mg/dL) for the sample with hematocrit value equal to the average (42%). The estimate bias due to hematocrit for each sample should be calculated by subtracting the bias at the average hematocrit (42%) from the bias of each sample.

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Table 6. Example table of bias due to hematocrit calculated for the nominal hematocrit value of 42% on a BGMS with 120 mg/dL glucose:

Hematocrit (%)	Average of Comparator measurements (Mean _{Comp})	Number of measurements for BGMS	Average of BGMS measurements (Mean _{BGMS})	% Bias (Mean _{BGMS} -Mean _{Comp})/Mean _{Comp}	% Bias due to hematocrit
10	118.0	30	127.6	8.1%	4.0%
15	118.4	30	127.6	7.8%	3.7%
20	122.4	30	130.4	6.5%	2.4%
25	120.7	30	127.1	5.3%	1.2%
30	123.7	30	129.5	4.7%	0.6%
35	121.5	30	127.1	4.6%	0.5%
42	119.7	30	124.6	4.1%	0.0%
50	121.3	30	125.4	3.4%	-0.7%
55	120.8	30	122.7	1.6%	-2.5%
60	120.1	30	119.5	-0.5%	-4.6%
65	118.1	30	116.0	-1.8%	-5.9%
70	117.5	30	115.6	-1.6%	-5.7%

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You should include in your 510(k) submission a detailed description of the study design, a list of all data collected in this study, the summary tables indicated above, and a summary of the conclusions drawn from the study.

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3. Oxygen

Study design:

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A typical professional use setting can include patients with a broad range of blood oxygen levels. If you intend for your BGMS to be used in patients with a broad range of blood oxygen levels, you should conduct a study using a validated method appropriate to the sample type to demonstrate the range of blood oxygen levels with which your device can be used. You should supplement the results of this study by collecting data on the blood oxygen levels of patients in your Method Comparison/User Evaluation Study (Subsection C, above), as appropriate, and conducting an analysis for any oxygen effects on BGMS performance. If you believe that blood oxygen levels do not affect the performance of your device you should provide a comprehensive justification for this, which should be supported by any analysis of interference of blood oxygen levels on device performance, as evaluated in your Method Comparison/User Evaluation Study (Subsection C, above).

875

E. *Flex Studies*

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Generally, the risk of an erroneous result may be greater for POC tests than laboratory-based tests. This is because there are fewer controls in place in POC settings to mitigate risks and the users may be untrained and may not know how to identify or address an

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879 incorrect result. You should demonstrate that your BGMS design is robust (i.e., insensitive
880 to environmental and usage variation) and that all known sources of error have been
881 assessed through a detailed risk assessment and are effectively controlled. In general, flex
882 studies should be used to demonstrate robust design while risk management should be used
883 to demonstrate the identification and effective control of error sources, although the two are
884 not mutually exclusive.

885

886 Most risk control measures should be fail-safe mechanisms or failure alert mechanisms.
887 Examples of fail-safe mechanisms are lock-out functions to ensure that a BGMS does not
888 provide a result when test conditions are inappropriate, such as when there is a component
889 malfunction or operator error. Other examples are measures within the BGMS to prevent
890 operator error, such as guides or channels that prevent improper strip placement. We
891 recommend that BGMS design incorporate fail-safe mechanisms whenever technically
892 practicable. If fail-safe mechanisms are not technically practicable for some risks, failure
893 alert mechanisms should be used. Failure alert mechanisms notify the operator of any
894 BGMS malfunction or problem. They may include measures such as internal procedural
895 controls or electronic controls. Devices with such mechanisms allow the operator to correct
896 the error, or put the operator on notice that the results will be unreliable due to the error. For
897 example, in cases where the result exceeds the reportable range (i.e., extremely high or low
898 glucose result) and the result is a critical value, the device should give a message such as
899 "high" or "low."

900

901 Flex studies, or studies that stress the operational boundaries of a BGMS, should be used to
902 validate the insensitivity of the test system to performance variation under stress conditions.
903 Where appropriate, flex studies should also be used to verify and/or validate the
904 effectiveness of control measures at operational limits.

905

906 In order to identify all relevant flex studies for your BGMS device, we recommend that you
907 conduct a systematic and comprehensive risk analysis that identifies all potential sources of
908 error, including test system failures and operator errors, and identify which of these errors
909 can lead to a risk of a hazardous situation. You should then identify control measures,
910 including fail-safe mechanisms and failure alert mechanisms that will reduce risks for these
911 sources of error. When the control measures have been implemented, you should (1) verify
912 that each control measure has been properly implemented, and (2) verify and/or validate the
913 effectiveness of each control measure. When appropriate, flex studies should be used to
914 verify and/or validate the effectiveness of these control measures.

915

916 Below, we have identified flex studies that we believe are important for you to perform and
917 recommend including in the 510(k) submission of your BGMS. At the same time, we
918 encourage you to continue to perform risk analyses to determine whether your device
919 includes any unique or new features that should be validated through additional flex studies.

920

921 If your BGMS does not perform adequately in flex studies, we recommend that you either
922 provide a justification, determined by means of thorough risk analysis, as to why adequate

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923 performance in that flex study is not necessary, or alternatively, you should indicate an
924 additional implemented validated control mechanism. FDA will review such justifications to
925 determine whether the proposed mitigation strategies are adequate to protect patients.

926

927 In the case of the following flex studies, verification should include performance testing;
928 however, it is sufficient if you provide information indicating that flex studies have been
929 conducted in accordance with an FDA-recognized industry standard. We recommend you
930 include information regarding the type of testing performed, the reference standard
931 followed, the acceptance criteria, and whether the BGMS passed testing requirements.

932 The flex studies we recommend performing in this manner are:

933

- 934 • Mechanical Vibration Testing
- 935 • Shock Testing
- 936 • Electromagnetic compatibility (EMC) Testing
- 937 • Electrostatic Discharge/Electromagnetic Interference Testing

938

939 We have also identified additional flex studies (described below) that manufacturers should
940 perform in order to demonstrate adequate system performance in intended use settings.

941 Unless otherwise indicated, we recommend that you clearly identify all flex studies
942 performed on your device in your 510(k) submission. A detailed description of the following
943 attributes should be included in your 510(k) submission for each study:

944

- 945 • Study goal
- 946 • Study protocol
- 947 • Methods used to apply samples to test strips
- 948 • Sample type and any anticoagulants used
- 949 • Study results
- 950 • Conclusions made from the study

951

952 The recommended flex studies as well as recommended study designs are outlined below in
953 Subsections 1-8. These flex studies should be performed using fresh venous or capillary
954 whole blood samples, not control solutions.

955

956 **1. Test Strip Stability Testing**

957 You should perform studies that assess test strip performance throughout the test strip
958 stability claims, including both closed and open vial claims. Two studies should be performed
959 to support test strip stability: 1) closed vial stability (shelf life) should be performed to assess
960 the recommended shelf life and conditions when the vial is stored closed throughout the
961 claimed expiration dating, at different combinations of temperature and humidity spanning
962 the recommended storage conditions; and 2) open vial stability should be performed to

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963 mimic conditions under which an individual would actually use the strips where the vial is
964 opened and closed throughout its claimed open vial life and stored at different combinations
965 of temperature and humidity throughout the recommended storage conditions. We suggest
966 that you submit only the study protocols for these test strip stability assessments, the
967 acceptance criteria, and the conclusions of any studies which have been completed.
968

969 These studies (shelf life and open vial stability) should be designed to span both the claimed
970 temperature range and humidity range at various time points throughout the duration of the
971 respective claim. The time points that are assessed (e.g., 1 month, 3 months, 2 years) should
972 be specified in the protocol. Combinations of real-time and accelerated stability studies are
973 acceptable. However, if accelerated studies are provided, real-time studies should be
974 ongoing and the protocols and acceptance criteria should be provided for both study types.
975

976 Separate testing of test strip and meter shipping and storage conditions is not necessary if
977 the temperature and humidity studies outlined here use only packaged blood glucose meters
978 and blood glucose test strips that have undergone appropriate storage conditions and the
979 longest possible shipping duration (both as specified by the manufacturer).
980

981 You should perform adequate precision and accuracy evaluations at each identified time
982 point. The following are provided only as examples of such studies. Through these
983 evaluations, you should demonstrate that the CV and accuracy calculated in these studies
984 are within the labeled performance of the BGMS.
985

986 *Precision Evaluation:*

987 Precision with Control Materials

988 This study should be completed over 5 days and use glucose controls. At least two
989 meters should be included in this study and at least 10 measurements should be taken
990 per glucose control level, per meter.
991

992 Precision with Whole Blood Samples

993 This study should use whole blood samples spanning the claimed measuring range of the
994 BGMS. Samples may be altered by spiking with glucose or allowing the samples to
995 glycolyze in order to evaluate the extreme end of the system's measuring range. At
996 least two meters should be included in this study and at least 10 measurements should be
997 taken per glucose level, per meter.
998

999 *Accuracy Evaluation:*

1000 This study should be performed using whole blood samples that span the claimed measuring
1001 range of the BGMS. It is acceptable for samples to be spiked with a known concentration
1002 of glucose or allowed to glycolyze to achieve the desired concentration in order to evaluate
1003 the extreme ends of the system's measuring range. Glucose concentrations (e.g., 30-50,
1004 100-150, 200-300, 350-500 mg/dL) should be measured with the BGMS and compared to
1005 values obtained with the comparator method.
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1007 **2. System Operating Conditions Testing**

1008 You should perform a study to assess the performance of your BGMS when used under
1009 various operating temperature and humidity conditions. These studies should be designed to
1010 represent actual use conditions experienced by BGMS users. Tested temperature and
1011 humidity ranges should not only cover the operating ranges that adequately reflect the
1012 intended use environment, and that are specified in the device labeling, but should also stress
1013 the BGMS by including ranges outside of the claimed operating range. Testing should
1014 incorporate the four extreme temperature and humidity combinations (high temperature/low
1015 humidity; high temperature/high humidity; low temperature/low humidity; low
1016 temperature/high humidity), or other testing combinations, if a suitable rationale can be
1017 provided. Measurements made on whole blood samples with your candidate device should be
1018 compared to values obtained using the candidate device at a nominal condition (such as 23°C,
1019 40% relative humidity).

1020
1021 We also encourage manufacturers to consider ways in which temperature and/or humidity
1022 detectors might be incorporated into test strip containers to alert users when strips have not
1023 been handled correctly or stored according to recommended and validated conditions.

1024
1025 **3. Altitude Effects**

1026 Relative to sea level, high altitude comprises a complex set of environmental differences and
1027 can induce multiple physiological changes, any or all of which might interfere with BGMS
1028 performance. For example, high altitude often involves extremes of temperature and
1029 humidity and can result in changes to hematocrit and blood pressure. The intended use
1030 environment of BGMSs in the United States includes high altitude conditions and, therefore,
1031 manufacturers should conduct studies to assess the effects of altitude on their BGMS, or
1032 should provide a justification for why altitude does not have an effect on the performance of
1033 their BGMS.

1034
1035 An altitude effects study should compare results from whole blood samples with your
1036 candidate device relative to values obtained using the candidate device at a nominal
1037 condition (such as sea level). These studies should also include a pressure change. Studies
1038 based on oxygen tension instead of pressure change are not adequate, because oxygen
1039 tension is only one component that changes with altitude. Altitude pressure changes can be
1040 accomplished by physically increasing altitude (e.g., in an airplane, on a mountain), or by
1041 simulating increasing altitudes and atmospheric conditions in a pressurized chamber. Results
1042 should support the altitude labeling claim for your device. You should provide your definition
1043 for terms such as “sea level.” The definition of sea level should not extend above 500 feet.
1044 You should test your BGMS at a minimum of 10,000 feet above sea level.

1045
1046 **4. Error Codes for Samples Outside the Measuring Range**

1047 You should perform adequate analyses to demonstrate that your meter provides the
1048 appropriate error codes when measured glucose concentrations are outside of the BGMS’s
1049 claimed measuring range, and include these results in your 510(k) submission.

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5. Short Sample Detection

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Blood glucose measurement from short samples (samples of reduced sample volume) can lead to inaccurate results. To avoid the risk of inaccurate results, BGMSs should be able to detect that a short blood sample that has been applied to the test strip and should not provide a result to the user. Short sample detection systems should not rely on visual verification by the user.

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The volume required to classify a test sample as a short sample is dependent upon your BGMS. In your short sample detection studies, you should include blood samples with known glucose concentrations in the following three ranges: 50-65, 100-120, and 200-250 mg/dL. You should test blood samples with your candidate device at each of the glucose concentrations listed above. Results obtained from the candidate device should be compared to results using the candidate device at a nominal condition (such as the claimed minimum sample volume). Blood samples with serially reduced volumes should be measured on the device until an error is either generated by the BGMS or the test result falls outside of the device's stated performance range. In your 510(k) submission, you should describe the results from the candidate device under both test and nominal conditions, as well as include the sample volumes tested for each glucose concentration range.

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6. Sample Perturbation Study

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Sample perturbation occurs when a user has applied an appropriate volume of blood to the test strip for glucose measurement but an event, such as wicking of blood away from the test strip, flicking of the test strip, or flicking of the meter, occurs during the start of measurement and alters the volume of the initial sample application. You should adequately demonstrate how your BGMS handles sample perturbation through a sample perturbation study.

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In a sample perturbation study, a sample should be applied to the test strip and after the BGMS has begun to read the sample, but before the measurement is complete, the test strip should be perturbed. The sample perturbation study should incorporate blood samples with known glucose concentrations in the following three ranges: 50-65, 100-120, and 200-250 mg/dL. In your 510(k) submission, you should describe your protocol, including your specific method of perturbing the test sample, as well as the candidate device results compared to results using the candidate device under a nominal condition (such as strips with no perturbation).

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7. Intermittent Sampling

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Intermittent sampling occurs when a short sample is applied to a test strip, a glucose measurement begins, and the user adds more sample to the test strip before the glucose measurement is complete. You should adequately demonstrate how your BGMS handles intermittent sampling by conducting an intermittent sampling study.

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1093 The intermittent sampling study should incorporate blood samples with known glucose
1094 concentrations in the following three ranges: 50-65, 100-120, and 200-250 mg/dL. You
1095 should perform intermittent sampling studies that are representative of actual events. For
1096 instance, approximately one half of the sample should be applied to the test strip prior to the
1097 start of sample measurement, then the other half of the sample should be applied to the strip
1098 after a set period of time, such as once the sample starts reading. For systems that allow a
1099 second sample of blood without producing an error message, different time delays
1100 throughout the claimed period of second application should be tested once the sample starts
1101 reading, but before the measurement is complete. You should describe how the device
1102 responds to this scenario, including whether a result is reported by the device, whether the
1103 result is accurate (relative to the nominal condition, such as with the minimum claimed
1104 sample volume), and when an error code is reported.

1105
1106 **8. Testing with Used Test Strips**

1107 You should perform a study to demonstrate how your BGMS performs when a used test
1108 strip is inserted. We recommend that BGMSs be designed to automatically recognize the
1109 insertion of used test strips. Insertion of used test strips into a blood glucose meter should
1110 not provide glucose measurement results to the user. If an automatic used test strip
1111 recognition function has been incorporated into your BGMS, you should perform a study to
1112 demonstrate the functionality of this recognition system. In your 510(k) submission, you
1113 should provide the study protocol, acceptance criteria, and results of your used test strip
1114 study.

1115 **F. Meter Calibration and Quality Control Material**

1116 We recommend you review FDA’s guidance entitled “Guidance for Industry and FDA Staff
1117 - Assayed and Unassayed Quality Control Material,”
1118 (<http://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocuments/ucm079179.htm>) and submit the recommended information to support clearance of any
1119 assayed glucose quality control material you choose to provide with your BGMS. For a
1120 description of more points to consider regarding calibration and quality control materials,
1121 please refer to FDA’s guidance entitled “Points to Consider for Review of Calibration and
1122 Quality Control Labeling for In Vitro Diagnostic Devices,”
1123 (<http://www.fda.gov/downloads/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocuments/UCM094139.pdf>). At least two levels of quality control material should be
1124 available for use with your system.
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1128 Your 510(k) submission should describe how your BGMS recognizes and distinguishes
1129 control materials from patient specimens, either automatically or manually by the user, as
1130 well as explain how the system compensates for differences between test strip lots (e.g.,
1131 how the meter is calibrated or coded for each test strip lot).
1132

1133 **VII. Test Strip Lot Release Criteria**

1134
1135 Your test strip lot release criteria should be sufficient to ensure consistent quality of the BGMS
1136 test strips. You should provide a description of the lot release criteria and a summary of the
1137 sampling scheme in your 510(k) submission. In addition, you should explain how the system
1138 compensates for differences between strip lots or strip types.

1139
1140 We recommend that you select a sampling scheme appropriate for the operation of your BGMS
1141 to test each outgoing test strip lot or batch. Your test strip lot release criteria should be designed
1142 to ensure that all released lots conform to the labeled BGMS device performance *in the hands*
1143 *of the intended user*. Therefore, these criteria typically should be tighter than the criteria used
1144 to evaluate total error in the performance studies, in order to achieve targeted performance in
1145 the intended user population.

1146

1147 **VIII. Third Party Test Strips**

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1149 Third party test strips refer to test strips manufactured and distributed by a company other than
1150 the company that manufactures and distributes the BGMS. Third party test strip manufacturers
1151 should ensure that they are aware of any design changes to the meter because such changes
1152 could affect compatibility of the strip with the meter. Because test strips and meters work as
1153 integral systems, third party test strip manufacturers should sufficiently address in their 510(k)
1154 submissions how they will mitigate the risk of incorrect results due to meter design changes.
1155 One way to effectively ensure that the third party test strip manufacturer is made aware of any
1156 design changes to the meter is by having in place an agreement between the third party test
1157 strip manufacturer and the manufacturer.

1158

1159 **IX. Software**

1160
1161 For software descriptions of BGMSs, their components, and accessories, we recommend that
1162 you follow FDA’s guidance entitled “Guidance for the Content of Premarket Submissions for
1163 Software Contained in Medical Devices,”
1164 ([http://www.fda.gov/downloads/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocu](http://www.fda.gov/downloads/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocuments/ucm089593.pdf)
1165 [ments/ucm089593.pdf](http://www.fda.gov/downloads/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocuments/ucm089593.pdf)). Generally, we consider blood glucose meters to be moderate level of
1166 concern devices because glucose results will be the basis for treatment, including determination
1167 of insulin dosage by the patient or health care provider. Incorrect glucose results or failure of
1168 the software to detect an error could result in improper therapeutic management. (Also, see
1169 Section V, above, regarding software descriptions in your 510(k) submission).

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X. Labeling

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The 510(k) submission must include labeling in sufficient detail to satisfy the requirements of 21 CFR 807.87(e). Final labeling must also satisfy the requirements of 21 CFR 809.10. Distinct labeling (e.g. user manual, quick start guide (optional), package inserts for both test strips and controls, and box and container labels for the meter, test strips, and control materials) appropriate for the intended user of the BGMS should be provided for each device component.

The following items are intended to further assist you in complying with the requirements of 21 CFR 809.10 for all labeling. You should refer to that regulation for the complete list of labeling requirements for *in vitro* diagnostic devices.

1. All device labels and labeling must contain the proprietary and common names of the device (21 CFR 809.10(a)(1) and 21 CFR 809.10(b)(1)). The various test system components should be named in such a way that they are recognizable as belonging to the same system, or family of products, and to distinguish them from those components intended for single-patient use only (for example, ABC blood glucose test system, ABC blood glucose meter, ABC blood glucose test strips, etc.) to aid in identification of system components.
2. You must include the intended use of the product in your label and labeling (21 CFR 809.10(a)(2) and 21 CFR 809.10(b)(2)).
3. You must include the symbol statement “Rx only” or “℞ only” or the statement “Caution: Federal law restricts this device to sale by or on the order of a ___”, the blank to be filled with the word “physician”, “dentist”, “veterinarian”, or with the descriptive designation of any other practitioner licensed by the law of the State in which the practitioner practices to use or order the use of the device, in your label and labeling (21 CFR 809.10(a)(4) and 21 CFR 809.10(b)(5)(ii)).
4. Labeling must include the chemical, physical, physiological, or biological principles of the procedure, as per 21 CFR 809.10 (b)(4). The discussion of these principles should include identification and biological source of the enzyme and a description of the reaction. Labeling should clarify whether results are determined in terms of whole blood or plasma equivalents. BGMSs intended for use in the U.S. should report results in terms of plasma equivalents.
5. The labeling must provide instructions for specimen collection and preparation, including special precautions regarding specimen collection. as per 21 CFR 809.10(b)(7). Instructions should include a statement to users on the importance of thoroughly washing and drying the skin before taking a sample because contaminants on the skin may affect results.
6. You must include a statement of limitations of the procedure in your labeling (21 CFR 809.10(b)(10)). Labeling must state known extrinsic factors or interfering substances affecting results, as per 21 CFR 809.10(b)(10). This should include, but is not limited to, the following:

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- 1211 a. Testing conditions that may cause clinically significant errors (due to bias or
1212 imprecision) with your device (e.g., specific drugs, oxygen therapy, testing with
1213 venous, arterial, or neonatal blood, high altitude, or EMC interference). Sponsors
1214 should indicate the most extreme conditions (e.g., the highest altitude, highest and
1215 lowest temperatures, etc.) at which the device should be used based on the results of
1216 performance testing.
- 1217 b. Clinical situations, patient populations, or conditions in which the BGMS performance
1218 may not be acceptable. For example, FDA recommends statements such as the
1219 following: inaccurate results may occur in severely hypotensive individuals or in
1220 dehydrated patients or patients in shock; inaccurate results may occur for individuals
1221 experiencing a hyperglycemic-hyperosmolar state, with or without ketosis.
- 1222 c. Limitations against alternative site testing and use for tight glycemic control (unless
1223 appropriate studies are performed and included in the 510(k) submission). Labeling
1224 should also state that results from alternative sampling sites (if used) should not be
1225 used to calibrate continuous glucose monitoring systems (CGMS) or entered into
1226 insulin dose calculators for dosage recommendations.
- 1227 7. Labeling must provide appropriate storage instructions adequate to protect stability of the
1228 product (21 CFR 809.10 (b)(5)(iv)). This type of information should be provided for all
1229 components of the system, including control solutions, test strips, etc.
- 1230 8. Labeling must describe details of calibration and quality control procedures (21 CFR
1231 809.10(b)(8)(v) and 21 CFR 809.10(b)(8)(vi)). This is to help ensure optimal performance of
1232 the system.
- 1233 9. Labeling must include expected values (21 CFR 809.10(b)(11)). FDA recommends that the
1234 expected values in the package insert should be those for non-diabetics. FDA does not
1235 recommend including additional ranges adjusted for diabetics because such ranges are
1236 individually determined by a clinician. The expected values should be cited from in-house
1237 studies or up-to-date reference sources.
- 1238 10. Labeling must include specific performance characteristics (21 CFR 809.10(b)(12)).
1239 Sponsors should briefly describe all studies and summarize results in the package inserts.
1240 FDA recommends that this include performance data summaries from in-house and user
1241 studies. For presentation of accuracy, in particular, see the suggested representations below
1242 for an example. Performance should be presented separately for each anatomical site,
1243 matrix (arterial, capillary, etc.), and any additional specific claims (e.g. neonatal).
- 1244 We recommend the following types of presentations to show the results of your accuracy
1245 studies in user manuals and package inserts.
- 1246

1247 **Suggested Representation of Accuracy for Prescription-use Only Devices – Example:**

1248

1249 The [XYZ] meter and [XYZ] reagent strips for the [XYZ] monitoring system were tested on
1250 capillary blood samples from 350 patients, and the results were compared to the comparator
1251 method (e.g., YSI). The tables show differences in glucose values between the XYZ device

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1252 and the comparator method. Table 8 below represents samples for glucose results lower than 70
1253 mg/dL (by the XYZ device). Table 9 below table represents samples for glucose results greater
1254 than or equal to 70 mg/dL.

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Table 8. Glucose results lower than 75 mg/dL

Difference range between ABC laboratory comparator method and the XYZ device	Within +/- 5 mg/dL	Within +/- 10 mg/dL	Within +/- 12 mg/dL	Within +/- 15 mg/dL
The percent (and number) of samples for which the difference between the XYZ device and ABC laboratory comparator method were within the difference range shown in the top row.	90% (126/140)	95% (133/140)	96% (135/140)	98% (137/140)

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Table 9. Glucose results greater than or equal to 75 mg/dL

Difference range between ABC laboratory comparator method and the XYZ device.	Within +/- 5%	Within +/- 10%	Within +/- 12%	Within +/- 15%	Within +/- 20%
The percent (and number) of samples for which the difference between the XYZ device and ABC laboratory method were within the difference range shown in the top row.	80% (168/210)	95% (199/210)	96% (202/210)	98% (206/210)	100% (210/210)

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The tables above show that 347 (137+210) of the 350 samples met the defined acceptance criteria.

Note: When glucose meter results are compared to the laboratory results, differences below 70 mg/dL are expressed in mg/dL, while those greater than or equal to 70 mg/dL are expressed in percent.

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11. You must describe the principles of operation for the instrument as well as service and maintenance information (21 CFR 809.10(b)(6)). Labeling should include a list or summary of error messages, descriptions of what those error messages mean, and appropriate troubleshooting procedures for those error messages.
12. Label and labeling must include statements of warning or precautions as appropriate to the hazard presented by the product on the outer container and the insert (21 CFR 809.10(a)(4) and 21 CFR 809.10(b)(5)(ii)).

You should clearly and prominently state the important warnings for your devices, for example, in a section titled **Important Safety Instructions**. You should stress the risk of disease transmission when using BGMSs and reference any relevant public health

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1278 notifications, standard practice guidelines, or other resources available to users. At a
1279 minimum, the following warnings should be included:

- 1281 • Users need to adhere to Standard Precautions when handling or using this device.
1282 All parts of the glucose monitoring system should be considered potentially infectious
1283 and are capable of transmitting blood-borne pathogens between patients and
1284 healthcare professionals. For more information, refer to “2007 Guideline for
1285 *Isolation Precautions: Preventing Transmission of Infectious Agents in*
1286 *Healthcare Settings*,”
1287 <http://www.cdc.gov/hicpac/2007ip/2007isolationprecautions.html>.
- 1288 • The meter should be cleaned and disinfected following the manufacturer’s
1289 instructions after use on each patient. This Blood Glucose Monitoring System may
1290 only be used for testing multiple patients when Standard Precautions and the
1291 manufacturer’s cleaning and disinfection procedures are followed.
- 1292 • Only auto-disabling, single use lancing devices may be used with this device.

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1294 In the section describing **how to obtain a blood sample** (see also item 4, above, regarding
1295 sample collection), you should re-iterate the risk of bloodborne pathogen transmission and
1296 state that only an auto-disabling, single use lancing device should be used. We recommend
1297 that you incorporate Standard Precautions and practices in your instructions. Include any
1298 graphics demonstrating correct blood draw procedures and ensure that the pictures show
1299 users wearing gloves.

1300
1301 In addition, we recommend that you refer users to the following practice guidelines:

1302
1303 “*Biosafety in Microbiological and Biomedical Laboratories (BMBL)*,”
1304 <http://www.cdc.gov/biosafety/publications/bmb15/>

1305
1306 CLSI (Clinical Laboratory Standards Institute) Document M29-A3: *Protection of*
1307 *Laboratory Workers From Occupationally Acquired Infections*.

1308
1309 You should stress that the operator should remove their gloves, clean their hands, and wear
1310 a new pair of clean gloves before testing each patient.

- 1311
1312 13. You must include a step-by-step outline of procedures (21 CFR 809.10(b)(8)). Labeling
1313 must list any points that may be useful in improving precision and accuracy, as per 21 CFR
1314 809.10(b)(8).

1315
1316 FDA recommends that the user manual should contain detailed instructions for how users
1317 are to perform **cleaning and disinfection procedures** for the meter **between patients**.
1318 This information should be based on the validation studies performed as described above in
1319 Section IV. You should also include the following:

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- 1321 • An explanation of why the cleaning and disinfection should be performed.
- 1322 • The recommended frequency of cleaning and disinfection, i.e., between each
- 1323 patient.
- 1324 • The materials needed for cleaning and disinfection and how they can be purchased
- 1325 or prepared.
- 1326 • A detailed procedure describing what parts of the device should be cleaned and
- 1327 disinfected, what should not be cleaned and disinfected (avoided), the amount of
- 1328 time the disinfectant needs to remain on the meter (contact time), etc. You should
- 1329 include graphics/photographs to assist the user. Again, be sure that all graphics show
- 1330 the user wearing gloves.
- 1331 • A statement that, after cleaning and disinfection, users' gloves should be removed,
- 1332 hands cleaned, and a new pair of clean gloves worn before proceeding to the next
- 1333 patient.
- 1334 • A contact telephone number for technical assistance or questions should be
- 1335 prominently listed in the cleaning and disinfection section.

1336
1337 We recommend you also include the references below:

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1339 *"FDA Public Health Notification: Use of Fingertick Devices on More than One*
1340 *Person Poses Risk for Transmitting Bloodborne Pathogens: Initial Communication,"*
1341 <http://www.fda.gov/MedicalDevices/Safety/AlertsandNotices/ucm224025.htm>

1342
1343 *"Infection Prevention during Blood Glucose Monitoring and Insulin Administration,"*
1344 <http://www.cdc.gov/injectionsafety/blood-glucose-monitoring.html>

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Appendix 1. Sources of Error to Consider for BGMSs

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Table 10 below lists sources of error associated with the design, production, and use of BGMSs. We do not intend for this to be a complete list. You should consider all sources of error based on your knowledge of your specific device. Documents such as CLSI EP-18A [7] and ISO 14971 [1] also provide lists of preanalytical, analytical, and post-analytical errors to consider.

Table 10 – Examples of Sources of Error

Category	Source of error or failure
Operator	<p>Failure to follow procedure correctly, for example:</p> <ul style="list-style-type: none"> • Sample contamination • Incorrect specimen collection (e.g., poor lancing technique and incorrect volume) • Application of an insufficient amount of blood to the strip or incorrect application of blood to strip • Use of a sample from an alternate site not validated by the manufacturer • Application of the specimen to the strip more than once (for example, if the user believes not enough specimen was added the first time) • Incorrect insertion of strip into meter • Inaccurate timing • Use of contaminated, outdated, or damaged strips or reagents, including calibrators or quality control materials • Failure to understand or respond to meter output • Errors in meter maintenance or cleaning • Errors in calibration or failure to calibrate or otherwise adjust the meter or check performance with quality control materials, as directed by labeling • Incorrect saving or use of stored data • Improper storage or handling of the meter, calibrators, quality control materials, or test strips, or improper maintenance of the meter • Inadvertent changes of parameters (such as units of measurement) • Incorrect incorporation of results into overall treatment plan (prescription-use) • Use of strips not validated for use on the meter

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Reagent	<ul style="list-style-type: none"> • Expired strips or reagents • Damaged or contaminated strips • Failure of strips, calibrators, or quality control materials to perform adequately • Incorrect manufacturing; product fails to conform with specifications • Incorrect dimensions of reagent strip • Interference with chemical reaction on strip (e.g., reducing substances) • Inadequate design of container for strips or other reagents; failure to prevent deterioration; failure of desiccant used to keep strips dry
Environmental	<ul style="list-style-type: none"> • DEVICE EFFECTS <ul style="list-style-type: none"> • Temperature • Humidity • Altitude; hyperbaric oxygen therapy conditions • Electromagnetic radiation • Visible light; sunlight • HUMAN FACTORS <ul style="list-style-type: none"> • Lighting, glare off meter surfaces • Distractions, visual and auditory • Stressful conditions • Limited manual dexterity
Software	<ul style="list-style-type: none"> • Confusing or obscure user prompts and feedback • Incorrect mathematical algorithm • Undetected or unrecognized signal errors • Timing failure • Incorrect storage of test results in memory, including matching result with correct patient or time of test • Other software failures
Hardware	<ul style="list-style-type: none"> • Electronic failure • Physical trauma or vibration • Damage to the device from incorrect strip dimensional tolerances (third party manufacturer) • Electrostatic discharge • Electromagnetic/radiofrequency interference • Battery reliability, lifetime, and replacement • Component(s) failure

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	<ul style="list-style-type: none">• Incorrect manufacture
System	<ul style="list-style-type: none">• Physical trauma or vibration• Incorrect calibration/adjustment (between lots of strips)• Calibration failure, interference, instability, or use beyond the recommended period of stability• Labeling not geared to intended user• Meter or operation complexity not geared to intended user• Inadequate training
Clinical	<ul style="list-style-type: none">• Interference from endogenous substances• Severe conditions (e.g., dehydration, hypoxia, hyperglycemic-hyperosmolar state, hypotension or shock, ketoacidosis)• Interference from other exogenous substances (e.g., maltose intravenous solutions)

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Appendix 2. Special 510(k)s and BGMSs

What is a special 510(k) and how does it apply to your blood glucose meter submission?

A special 510(k) submission is an alternative to the traditional method of demonstrating substantial equivalence for certain modifications to a manufacturer's own previously cleared device, the Agency believes that the rigorous design control procedure requirements outlined in the Quality System Regulation (QS reg) [See 21 CFR 820] produce highly reliable results that can form, in addition to the other 510(k) content requirements, a basis for the substantial equivalence determination.

As such, under the special 510(k) option, a manufacturer who is intending to modify his/her own legally marketed device will perform and present the risk analysis and the necessary verification and validation activities, to demonstrate that the design outputs of the modified device meet the design input requirements. Once the manufacturer has ensured the satisfactory completion of this process, a "Special 510(k): Device Modification" may be submitted.

Eligibility for a Special 510(k):

To determine whether a modified BGMS device is eligible to be submitted as a special 510(k), you should consult the FDA guidance entitled "The New 510(k) Paradigm - Alternate Approaches to Demonstrating Substantial Equivalence in Premarket Notifications - Final Guidance,"

(www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocuments/ucm080187.htm). Sponsors should also consult the document on FDA's website "How to Prepare a Special 510(k),"

(<http://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/HowtoMarketYourDevice/PremarketSubmissions/PremarketNotification510k/ucm134573.htm>).

As noted above, to be eligible for a special 510(k), the manufacturer should be modifying their own legally marketed device. This usually means that the candidate device and predicate device are part of the same device design file. *Similarities* between the candidate and predicate devices alone do not necessarily mean that the candidate device is a modification of the predicate device.

We recommend that you contact the Office of In Vitro Diagnostics and Radiological Health (OIR) to discuss any specific questions you have regarding your BGMS device's eligibility to be submitted as a special 510(k).