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4	Rodolfo J. Galindo, MD, FACE ¹ , Guillermo E. Umpierrez, MD, CDE ¹ , Robert J. Rushakoff, MD ² ,
5	Ananda Basu, MD, FRCP ³ , Suzanne Lohnes, MA, RN, CDCES, CPT ⁴ , James H. Nichols, PhD,
6	DABCC, FAACC ⁵ , Elias K. Spanakis, MD ^{6,7} , Juan Espinoza, MD, FAAP ⁸ , Nadine E. Palermo, DO ⁹ ,
7	Dessa Garnett Awadjie, MSN, FNP ¹⁰ , Leigh Bak, MSN, APRN, ACNS-BC, CDCES ¹¹ , Bruce
8	Buckingham, MD ¹² , Curtis B. Cook, MD ¹³ , Guido Freckmann, MD ¹⁴ , Lutz Heinemann, PhD ¹⁵ ,
9	Roman Hovorka, PhD, FMedSci ¹⁶ , Nestoras Mathioudakis, MD, MHS ¹⁷ , Tonya Newman, JD ¹⁸ ,
10	David N. O'Neal, MD, FRACP ¹⁹ , Michaela Rickert, MS, PA-C, RDN, CDE ²⁰ , David B. Sacks, MB,
11	ChB, FACP, FRCPath ²¹ , Jane Jeffrie Seley, DNP, MPH, MSN, GNP, RN, BC-ADM, CDCES, CDTC,
12	FADCES, FAAN ²² , Amisha Wallia, MD, MS ²³ , Trisha Shang, BA ²⁴ , Jennifer Y. Zhang, BA ²⁴ , Julia
13	Han, BA ²⁴ , David C. Klonoff, MD, FACP, FRCP (Edin), Fellow AIMBE ²⁵
14	
15	Affiliations:
16	¹ Emory University School of Medicine, Atlanta, GA, USA
17	² University of California, San Francisco, San Francisco, CA, USA
18	³ University of Virginia School of Medicine, Charlottesville, VA, USA
19	⁴ University of California San Diego Medical Center, La Jolla, CA, USA
20	⁵ Vanderbilt University Medical Center, Nashville, TN, USA
21	⁶ University of Maryland School of Medicine, Baltimore, MD, USA
22	⁷ Division of Endocrinology, Baltimore Veterans Affairs Medical Center, Baltimore, MD, USA
23	⁸ Children's Hospital Los Angeles, Los Angeles, CA, USA
24	⁹ Brigham and Women's Hospital, Harvard Medical School, Boston, MA, USA
25	¹⁰ New York University Langone Health, New York, NY, USA
26	¹¹ Yale New Haven Hospital, New Haven, CT, USA
27	¹² Stanford University School of Medicine, Palo Alto, CA, USA

Continuous Glucose Monitors and Automated Insulin Dosing Systems in the Hospital

Consensus Guideline

- 26
- 27 ¹³Mayo Clinic Arizona, Scottsdale, AZ USA 28
- 29 ¹⁴Institute for Diabetes-Technology GmbH, Ulm, Germany
- ¹⁵Science Consulting in Diabetes GmbH, Neuss, Germany 30
- ¹⁶University of Cambridge, Cambridge, UK 31
- ¹⁷Johns Hopkins University School of Medicine, Baltimore, MD 32
- ¹⁸Neal, Gerber and Eisenberg LLP, Chicago, IL, USA 33
- ¹⁹University of Melbourne Department of Medicine, St. Vincent's Hospital, Fitzroy, Victoria, 34
- 35 Australia
- ²⁰Oregon Health and Sciences University, Portland, OR, USA 36
- ²¹National Institutes of Health, Bethesda, MD, USA 37
- ²²Weill Cornell Medicine, New York, NY, USA 38

- ²³Northwestern University Feinberg School of Medicine, Chicago, IL, USA
- 40 ²⁴Diabetes Technology Society, Burlingame, CA, USA
- 41 ²⁵Mills-Peninsula Medical Center, San Mateo, CA, USA
- 42
- 43 Corresponding Author:
- 44 David C. Klonoff, MD, FACP, FRCP (Edin), Fellow AIMBE
- 45 Mills-Peninsula Medical Center, San Mateo, CA, USA
- 46 Address: 100 South San Mateo Drive Room 5147, San Mateo, CA 94401
- 47 Phone Number: 1 (650) 218-1142
- 48 Email: dklonoff@diabetestechnology.org
- 49
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- 51 Hospital
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53 <u>Abstract</u>

This article is the work product of the Continuous Glucose Monitor and Automated Insulin 54 55 Dosing Systems in the Hospital Consensus Guideline Panel, that was organized by Diabetes 56 Technology Society and met virtually on April 23, 2020. The guideline panel consisted of 24 57 international experts in the use of CGMs (continuous glucose monitors) and AID (automated 58 insulin dosing) systems representing adult endocrinology, pediatric endocrinology, obstetrics 59 and gynecology, advanced practice nursing, diabetes care and education, clinical chemistry, 60 bioengineering, and product liability law. The panelists reviewed the medical literature pertaining to five topics: 1) continuation of home CGMs after hospitalization, 2) initiation of 61 CGMs in the hospital, 3) continuation of AID systems in the hospital, 4) logistics and hands-on 62 care of hospitalized patients using CGMs and AID systems, and 5) data management of CGMs 63 and AID systems in the hospital. The panelists then developed three types of recommendations 64 65 for each topic, including clinical practice (to use the technology optimally), research (to improve 66 the safety and effectiveness of the technology), and hospital policies (to build an environment

for facilitating use of these devices) for each of the five topics. The panelists voted on 78
proposed recommendations. Based on the panel vote, 77 recommendations were classified as
either strong or mild. One recommendation failed to reach consensus. Additional research is
needed on CGMs and AID systems in the hospital setting regarding device accuracy, practices
for deployment, data management, and achievable outcomes. This guideline is intended to
support these technologies for the management of hospitalized patients with diabetes.

73

74 Introduction

75 Continuous glucose monitors (CGMs) are becoming an important technology for improving 76 glycemic outcomes in diabetes. The opportunity for a patient (or by way of wireless communication, a caregiver or relative) to see real-time glucose concentrations tested 77 automatically and continuously is transforming the practice of diabetes care. Recent 78 79 generations of these devices offer improved accuracy, smaller form factors, extended sensor life, and new data presentation software for translating data into increasingly useful metrics on 80 81 various mobile platforms. Some new factory-calibrated CGMs have eliminated the need for 82 finger-stick blood glucose testing by users (except at certain times per individual product instructions, such as soon after insertion, when there appear to be errors or no readings at all, 83 when the CGM value does not match how the patient feels, or when an icon indicates the need 84 for testing blood glucose.) 85

CGMs for monitoring glucose concentrations and automated insulin dosing (AID) systems, that
 contain a CGM controlling a continuous subcutaneous insulin infusion (CSII) system (also known

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88	as an insulin pump), are cleared (class II) or approved (class III) by the United States Food and
89	Drug Administration (FDA) for home use (by prescription) by people who have diabetes.
90	However, many clinicians believe that CGMs have the potential to be utilized by hospitalized
91	patients in a variety of situations.
92	Escalating interest in utilizing CGMs and AID systems in a hospital setting has resulted in a need
93	for guidance on the continuation of these technologies in the hospital setting. This interest has
94	been stimulated by four trends in the application of CGM technology, including: 1)
95	improvements in the technology and human factors of CGMs, 2) an increasing number of
96	patients wearing these devices in ambulatory settings, 3) growing interest by clinicians to
97	understand and interpret their hospitalized patients' glucose concentrations, and 4) an
98	accumulation of published reports describing use of these products in investigational settings.
99	Diabetes Technology Society (DTS) previously organized guidance on the use of CGMs in the
100	hospital as "Consensus Statement on Inpatient Use of Continuous Glucose Monitoring" ¹ ,
101	published in 2017. Because of recent increasing interest in this topic, coupled with advances in
102	technology, DTS recognized a need for an updated consensus guideline on the use of CGMs and
103	AID systems in an acute care setting.
104	On April 23, 2020, DTS, led by Dr. David Klonoff, convened the Continuous Glucose Monitor and
105	Automated Insulin Dosing Systems in the Hospital: Consensus Guideline Panel. This
106	international panel consisted of experts in diabetes technology from the United States, Europe,
107	and Australia. The purpose of this meeting was to provide guidance for clinicians on how and
108	when to best use both subcutaneous CGMs and AID systems, as well as to promote clinical
109	research utilizing these devices.

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The panel was planned in late 2019 before the first case of Coronavirus Disease 2019 (COVID-110 111 19) was reported. Two weeks prior to the panel meeting, two CGM companies announced that during the pandemic, the FDA had told them that the Agency would not object if these 112 companies provided devices and technical support to hospitals who ordered CGMs for off label 113 use.^{2,3} Because some healthcare systems were interested in validating CGMs for use in their 114 hospitals to preserve PPE supplies and to minimize patient/provider contact, there was 115 additional urgency for the panel to develop new clinical guidance. Panelists discussed how the 116 117 pandemic has impacted inpatient glucose monitoring and how an urgent need has arisen for 118 alternative approaches to this monitoring.⁴ The traditional approach of testing capillary blood glucose (BG) every 1-2 hours in patients who are receiving intravenous insulin in an intensive 119 120 care unit (ICU) as well as frequent BG testing in non-ICU wards for patients receiving subcutaneous insulin is not workable during the pandemic. Other methods are needed to 121 122 decrease nurse contact with the patient for assisted monitoring of BG (AMBG)⁵ in order to: 1) 123 decrease risk of contagion from exposure to patients, 2) save time from donning and doffing personal protective equipment (PPE) wherever possible, and 3) preserve limited supplies of 124 PPE⁴. Despite limited guidance, established studies, or widespread support from the clinical 125 community to use CGMs in acute care⁶, some HCPs in the hospital diabetes community have 126 127 recently begun to prescribe CGMs in the hospital setting for investigational or off-label use for 128 COVID-19 patients.⁷

The Continuous Glucose Monitors and Automated Insulin Dosing Systems in the Hospital
 Consensus Guideline Panel included professionals from a variety of backgrounds. Members
 included experts in the use of CGMs from adult endocrinology, pediatric endocrinology,

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132	obstetrics and gynecology, advanced practice nursing, diabetes care and education, clinical
133	chemistry, bioengineering, and product liability law. The expert panel included representatives
134	from academia and government and observers from government (FDA), and industry (Abbott
135	Diabetes Care, Dexcom, Glytec, Medtronic, and Roche Diagnostics). One member represented
136	the College of American Pathologists, one represented the Endocrine Society, and one
137	represented the Association of Diabetes Care and Education Specialists.
138	The expert panel discussed the following five topics: 1) continuation of home CGMs after
139	hospitalization, 2) initiation of CGMs in the hospital, 3) continuation of AID systems in the
140	hospital, 4) logistics and hands-on care of hospitalized patients using CGMs and AID systems,
141	and 5) data management of CGMs and AID systems in the hospital. (Table 1) Panelists reviewed
142	available evidence on the inpatient use of diabetes technology, and discussed potential
143	opportunities, potential barriers, and recommendations associated with the use of these
144	devices in the hospital setting.
145	Recommendations were proposed by the panelists and then reviewed by the entire panel for
146	favorability. Recommendations receiving at least 80% favorable votes were classified as strong
147	recommendations, proposals receiving 60-79% favorable votes were classified as mild
148	recommendations, and proposals receiving less than 60% favorable votes were classified as
149	recommendations which failed to receive consensus support.
150	For each of the five topics of this guideline (Table 1), six categories of recommendations (two
151	for clinical practice, two for future research, and two for hospital policies) were developed for
152	the main stakeholders of CGM and AID system technology in the hospital. These types of

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153 recommendations included: 1) and 2) strong and mild recommendations that clinicians 154 (healthcare professionals, HCPs or nursing) should do to utilize the technology optimally, 3) 155 and 4) strong and mild recommendations that researchers and manufacturers need to do to improve the safety and effectiveness of the technology, and 5) and 6) strong and mild 156 157 recommendations that hospitals need to do to build an environment for facilitating use of these devices. We define "should" as a statement of good practice and "need" as a necessary step to 158 ensure patient safety or proper fulfillment of a procedure. These recommendations are 159 160 intended to promote the best use of CGMs and AID systems in the hospital.

161 Background

162 CGMs were developed for the outpatient setting, and their transition for use in hospitals has been the subject of ongoing scholarship, research, and consensus guidelines. The first CGM 163 became commercially available in 1999⁸. CGM technology has greatly improved since then and 164 165 several revolutionary developments in CGM technology have taken place over the past 5 years. These advances have all significantly reduced patients' burden of diabetes care. The result has 166 167 been improved patient satisfaction and self-care behaviors, increased clinician awareness, and a significant increase in CGM adoption, mostly by patients with Type 1 diabetes mellitus 168 169 (T1DM), but also in some patients with Type 2 diabetes mellitus (T2DM)⁹. Software for analyzing continuous glucose data streams has permitted the development of new CGM-based 170 glycemic metrics, which compared to hemoglobin A1c, illustrate multidimensional patterns of 171 glycemia more directly and with greater granularity¹⁰. Improvements in CGM technology have 172 also permitted integration with CSII systems to create AID systems. With the increasing 173

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174	popularity of AID systems that depend on CGMs, hospital HCPs will increasingly encounter
175	patients who will want to utilize their CGMs and AID systems for inpatient diabetes care.
176	AID systems are becoming more advanced and are more frequently utilized for outpatients to
177	successfully achieve glycemic outcomes in diabetes by facilitating increased time in range (TIR)
178	and decreased time in hypo- and hyperglycemia. Two AID systems are currently cleared or
179	approved by the FDA for home use in people with diabetes: 670G (Medtronic, Northridge,
180	California) and Tandem Control IQ (Tandem Diabetes Care, Inc., San Diego, California). Some
181	patients utilizing these AID systems and/or their physicians wish to continue the AID systems
182	even during a hospitalization, believing that the benefits of commercial AID systems outweigh
183	potential risks in this setting and noting that product use would not be off label if a patient is
184	self-managing using the device even if the patient is in the hospital while doing it.
185	CGM sensors can be invasive (intravascular blood sampling or sensing devices that remove
185 186	CGM sensors can be invasive (intravascular blood sampling or sensing devices that remove blood), minimally invasive (subcutaneous placement of a sensor), or non-invasive (transdermal
186	blood), minimally invasive (subcutaneous placement of a sensor), or non-invasive (transdermal
186 187	blood), minimally invasive (subcutaneous placement of a sensor), or non-invasive (transdermal CGMs that do not puncture the skin). They are measuring in different compartments, which can
186 187 188	blood), minimally invasive (subcutaneous placement of a sensor), or non-invasive (transdermal CGMs that do not puncture the skin). They are measuring in different compartments, which can lead to different values. ¹¹ The frequency of receiving a signal by a CGM ranges from every 1 to
186 187 188 189	blood), minimally invasive (subcutaneous placement of a sensor), or non-invasive (transdermal CGMs that do not puncture the skin). They are measuring in different compartments, which can lead to different values. ¹¹ The frequency of receiving a signal by a CGM ranges from every 1 to every 15 minutes, most commonly every 5 minutes. Invasive CGMs that are intended only for
186 187 188 189 190	blood), minimally invasive (subcutaneous placement of a sensor), or non-invasive (transdermal CGMs that do not puncture the skin). They are measuring in different compartments, which can lead to different values. ¹¹ The frequency of receiving a signal by a CGM ranges from every 1 to every 15 minutes, most commonly every 5 minutes. Invasive CGMs that are intended only for hospital use include two systems cleared by the FDA. They are 1) the GlucoScout (International
186 187 188 189 190 191	blood), minimally invasive (subcutaneous placement of a sensor), or non-invasive (transdermal CGMs that do not puncture the skin). They are measuring in different compartments, which can lead to different values. ¹¹ The frequency of receiving a signal by a CGM ranges from every 1 to every 15 minutes, most commonly every 5 minutes. Invasive CGMs that are intended only for hospital use include two systems cleared by the FDA. They are 1) the GlucoScout (International Biomedical, Austin, TX) ¹² and 2) the OptiScanner 5000 (OptiScan Biomedical Corporation,
186 187 188 189 190 191 192	blood), minimally invasive (subcutaneous placement of a sensor), or non-invasive (transdermal CGMs that do not puncture the skin). They are measuring in different compartments, which can lead to different values. ¹¹ The frequency of receiving a signal by a CGM ranges from every 1 to every 15 minutes, most commonly every 5 minutes. Invasive CGMs that are intended only for hospital use include two systems cleared by the FDA. They are 1) the GlucoScout (International Biomedical, Austin, TX) ¹² and 2) the OptiScanner 5000 (OptiScan Biomedical Corporation, Hayward, California) ¹³ . Both devices track glycemic patterns of blood that is withdrawn from

196	Rastatt, Germany) ¹⁶ , and 4) Optiscanner 5000 ¹³ . The Optiscanner 5000 has received FDA
197	clearance, but the Glucoclear, Glysure System, and Eirus products all have not received FDA
198	clearance. The Glucoclear and Eirus products have been discontinued, and Glysure Ltd. went
199	out of business in 2018. The Optiscanner 5000 is available in the US and Europe. One CGM with
200	a subcutaneous sensor was available in Europe for measuring glucose in hospitalized patients:
201	Sentrino Continuous Glucose Management System (Medtronic, Northridge, California) ¹⁷ .
202	However, at this time Sentrino is not a commercial product. There are no commercially
203	available non-invasive CGMs in the United States.
204	In the hospital special issues can arise that can impair proper function of CGMs. No CGM is
205	labeled to allow for exposure to X-Rays, CT scans, magnetic resonance imaging (MRIs),
206	diathermy, radiation therapy, or other types of radiation. Typically, the device is removed or
207	covered with a lead shield during these procedures. Some sites have covered their CGMs with a
208	lead shield and have not reported adverse events. Emerging data suggests there may be no
209	need for removal of the Dexcom G6 sensor (Dexcom, San Diego, California) during X-rays, CT
210	scans, radiation therapy, or when electrocautery is used during surgical procedures. ^{18–20} There
211	were no data errors observed when FreeStyle Libre Pro sensor was exposed to chest X-rays,
212	computed tomography (CT), radiotherapy (RT), and magnetic resonance imaging (MRI). ²¹ The
213	panel expected that each manufacturer will continue to determine and report the impact of
214	imaging studies and electrocautery on their particular devices.
215	An attractive feature of CGMs is that they can measure glucose concentrations automatically
216	and sound an alarm for readings that are outside of a prespecified safe target range. Table 2

217 contains a list of the five currently available subcutaneous home-use CGMs that have the

218	potential for hospital use: FreeStyle Libre 14 day system 22 , FreeStyle Libre 2 23 (both Abbott
219	Diabetes Care, Chicago, Illinois), Dexcom G6 ²⁴ , Medtronic Guardian Sensor 3 ²⁵ (Medtronic
220	Diabetes, Northridge, California), and Eversense (Senseonics, Inc., Germantown, Maryland) ²⁶ .
221	This table includes the devices' glucose sensing methods, technical features, and presence of
222	interference from chemical substances.
223	Continuation of home Continuous Glucose Monitors after hospitalization
224	Chair: Robert J. Rushakoff, M.D.
225	University of California, San Francisco, San Francisco, California, USA
226	POTENTIAL OPPORTUNITIES
227	Patient Considerations
227 228	Patient Considerations Standalone CGMs and AID systems are typically used in the outpatient setting. If a patient
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228 229 230 231	Standalone CGMs and AID systems are typically used in the outpatient setting. If a patient wearing either of these technologies is hospitalized, then policies are needed to continue these technologies. Some hospitals have policies for removing personal use devices like CGMs, CSII systems and AID systems from patients when they are admitted. It is within the FDA's
228 229 230 231 232	Standalone CGMs and AID systems are typically used in the outpatient setting. If a patient wearing either of these technologies is hospitalized, then policies are needed to continue these technologies. Some hospitals have policies for removing personal use devices like CGMs, CSII systems and AID systems from patients when they are admitted. It is within the FDA's authorized use for a patient to use their own device for self-management while in a hospital.

hospital and a subsequent section focuses on initiating a CGM in the hospital. Anyone with

diabetes who is using a CGM and who is not cognitively impaired is a candidate to continuewith this device in the hospital.

239 Benefits of CGMs

240	Several studies have demonstrated that CGMs in ambulatory settings improve patients'
241	satisfaction, ^{38,39} as well as control (e.g. better TIR and time in hypo- and hyperglycemia) ^{40,41} .
242	Continuation of an outpatient CGM during a hospitalization could improve patient satisfaction
243	and efficacy of glycemic monitoring by assisting the patient and the hospital staff to identify
244	glucose patterns, predict future glycemia with trend arrows and rate-of-change ⁴² , and
245	potentially prevent severe hypo- and hyperglycemic events. ⁴³ This would be particularly
246	relevant if staffing shortages exist or a patient is no longer aware of hypoglycemia. Accordingly,
247	asking patients to remove their CGMs in the hospital could potentially contribute to decreased
248	patient satisfaction and quality of care. CGM use in ICU and non-ICU settings has several
249	superior features over intermittent point of care (POC) testing for glucose monitoring during
250	continuous insulin infusion and subcutaneous insulin therapy, and possibly is a safer and less
251	costly approach that can reduce workload. Additionally, CGM technology could potentially
252	replace many uses of POC capillary BG testing in the hospital. ⁴³ However, if CGM readings turn
253	out to be inaccurate, then more confirmatory testing would be needed and that could increase
254	workload.

255 Pregnancy

The use of CGMs in pregnant patients with T1DM has been associated with improvement in
both maternal and fetal outcomes in five areas, including: 1) time in glycemic target range

258 without increase in hypoglycemia, 2) lower incidence of large for gestational age babies, 3) 259 fewer neonatal intensive care unit admissions, 4) reduced neonatal hypoglycemia, and 5) decreased LOS.^{44,45} The use of CGMs in pregnancy is considered off-label in the US, but not in 260 Europe. In recent years, patients and HCPs have identified real time continuous glucose 261 262 monitoring as a helpful adjunct. Although there is ongoing interest in the use of CGMs in 263 pregnancy, there is limited data about its use in the acute care setting. If an HCP intends to use such a device, then it would be important to avoid placing it near areas of potential obstetric 264 265 surgery.

266

267 **POTENTIAL BARRIERS**

Studies on substances that interfere with current subcutaneous CGMs are shown in Table 2. 268 The panel agreed that CGM results should be interpreted cautiously in patients using select 269 270 drugs known to cause interference with CGM sensing technologies. For these situations, 271 panelists recommended using more accurate glucose testing, such as laboratory analyzers or 272 AMBG⁵ using hospital BGMs (which, unlike home-use BGMs, require special cleaning and 273 disinfection procedures). Even though these devices are factory-calibrated and a limited set of studies have reported acceptable accuracy in critically ill patients⁴⁶, several potential scenarios 274 in the hospital (e.g., interfering substances, hypoxia, acidosis, and hypotension) would require 275 very careful use of this technology. The panel did not feel that current CGMs can now replace 276 277 capillary POC finger stick monitoring or other FDA cleared methods for monitoring BG in the 278 hospital.

279	Recommendations for Continuation of home Continuous Glucose Monitors after
280	hospitalization
281	Clinical Practice
282	Strong Recommendations
283	• HCPs should consult with an inpatient diabetes team if available, when continuing or
284	initiating a CGM or AID system.
285	HCPs should avoid relying on CGM data for glycemic management decisions in patients
286	with severe hypoglycemia or hyperglycemia (i.e. BG < 40 mg/dL or >500 mg/dL).
287	• HCPs should avoid using CGMs for management of 1) diabetic ketoacidosis until glucose
288	is in the CGM measurement range, and then CGMs should be used adjunctively or 2)
289	situations with rapidly changing glucose levels and fluid/electrolyte shifts.
290	• HCPs should avoid continuing or initiating CGMs to patients with skin infections near the
291	sensor site or placing sensors in areas with significant edema as well as patients treated
292	with vasoactive agents or poor tissue perfusion.
293	HCPs should use a CGM checklist for elective procedures during the pre-operative visits
294	to ensure proper documentation of devices and real time data reporting.
295	HCPs should advise pregnant women to continue the use of a CGM during a
296	hospitalization to identify glucose trends and prevent hypo- or hyperglycemia.
297	HCPs should instruct patients to bring supplies with them to the hospital for the
298	duration of any pre-planned admission or elective procedures.

299	•	HCPs should check capillary BG or serum BG concentrations after procedures for non-
300		critically ill patients and venous/arterial blood for critically ill patients to ensure the
301		patient's CGM is functioning properly.
302	•	HCPs should use trend arrows and rate of change to help prevent extreme glycemic
303		excursions and (when a CGM is used adjunctively) to help determine when a BG test is
304		required.
305	•	HCPs should set alarm thresholds for inpatient glycemic targets, such as predicting
306		hypoglycemia (typically BG < 80-85 mg/dL) or predicting hyperglycemia.
307	•	Nursing should document CGM and/or CSII system information in the electronic health
308		record (EHR) for all admissions or elective procedures.
309	Resear	ch
310	Strong	Recommendations
311	•	Researchers need to provide more data to support definitive recommendations on
312		improved outcomes for continuation of home/ambulatory CGM use after
313		hospitalization.
314	•	Researchers need to conduct studies on the roles of CGM and POC BG testing and
315		identify the optimal features of telemetry to inform nursing staff about actionable CGM
316		patterns.
317	•	Researchers need to perform further studies to assess the accuracy of CGMs during
318		pregnancy, labor & delivery, and the peripartum period.

319	• Researchers need to study the impact of lag time on glucose measurements (i.e.
320	situations with rapid changes in the glucose concentration) in the hospital.
321	Hospital Policies
322	Strong Recommendations
323	• Hospitals need to develop standard CGM data reports and workflows.
324	Hospitals need to implement policies for testing capillary BGs and calibrating CGMs if
325	the CGM requires calibration.
326	• Hospitals need to develop a system for automatic staff notification for CGM alarms that
327	predict impending or current hypoglycemia or hyperglycemia.
328	Hospitals need to develop specific guidelines for using CGMs and AID systems for their
329	affiliated nursing homes and skilled nursing facilities.
330	
331	Initiation of Continuous Glucose Monitors in the hospital
332	Chair: Guillermo E. Umpierrez, MD, CDE
333	Emory University School of Medicine, Atlanta, Georgia, USA
334	POTENTIAL OPPORTUNITIES
335	COVID-19
336	The current COVID-19 pandemic created the need for innovative approaches for glycemic
337	monitoring in the hospital ⁴ . Coincidentally, two weeks before this meeting, the FDA stated that
338	they would exercise enforcement discretion and they would not object to the use of CGMs in

339	the hospital during the crisis ^{2,3} . This policy was intended for the factory-calibrated CGMs
340	manufactured by Abbott Diabetes Care and Dexcom. Subsequently, these two manufacturers
341	provided CGM supplies to hospitals to help monitor glucose remotely. Immediately afterward,
342	several institutions started the process of implementing CGM use and realized that there was a
343	need for training, implementation, and resource utilization and not all hospitals have this
344	expertise. The announcement also resulted in new reports on the use of CGMs in the hospital.
345	During the panel discussion, there was a recognition that this "exceptional" situation did not
346	indicate "label approval" for CGM use in the hospital by regulatory bodies. Collaborative efforts
347	from Emory University and DTS have recently provided examples of practical implementation of
348	CGMs and use of diabetes technology in the hospital through creation of a website that
349	contains information about original articles, commentary, news, and protocols related to
350	COVID-19 and diabetes ⁴⁷ (covidindiabetes.org). Small pilot studies have provided unconfirmed
351	evidence of the feasibility of remote glucose monitoring during this global crisis ⁴⁰ .

352 ICU Patients

There is strong evidence from large prospective and randomized studies indicating that optimal glucose management results in improved outcomes, reduced complications, and a decreased length of stay (LOS)^{48,49}. In the ICU setting, therapy with intravenous insulin infusion allows clinicians to maintain narrow glycemic targets. The panelists reviewed studies using CGMs in the ICU in adult populations (**Table 3**) and pediatric populations (**Table 4**).

In the ICU, bedside POC glucose using factory-calibrated BGMs (performed every 1-2 hours) has

been recommended as the preferred method to assess glycemic management and to guide

360	hyperglycemia treatment with intravenous insulin infusion. POC BG testing has drawbacks. This
361	testing method is labor-intensive. Also, POC testing does not provide: 1) a full 24-hour glycemic
362	profile, 2) predictions of hypoglycemic events, or 3) alarms for asymptomatic hypo- or
363	hyperglycemia. Although the use of POC glucose testing, compared to central laboratory
364	glucose testing, is approximately as convenient and generates faster results, another drawback
365	is that it costs more. Estimated mean total costs (including equipment, supplies and labor) can
366	be up to \$5.13 per POC test in a high-test volume nursing unit, and up to \$16.49 per POC test in
367	a low-test volume nursing unit, compared to \$3.78 for central laboratory glucose testing ⁸⁹ .
368	Moreover, the accuracy of POC glucose meters is not optimal, with only six of eighteen glucose
369	monitor systems (representing 90% of commercially available meters and intended for
370	outpatient use) meeting regulatory accuracy requirements ¹⁷ in a recent study. In 2018 the FDA
371	cleared the first POC glucose meter - the StatStrip Glucose (Nova Biomedical, Waltham,
372	Massachusetts)- for all hospitalized patients, including critically ill patients, to test capillary,
373	venous, and arterial blood specimens ⁹⁰ . However, not all hospitals use this system to measure
374	BG. While definitive validation of CGM accuracy in ICU patients is still forthcoming there
375	remains a potential role for CGMs to measure glucose concentrations in this population. ^{46,91,92}
376	Non-ICU Patients

Studies using older CGM technology that required regular recalibration have shown minimal
differences in mean daily glucose, premeal, fasting, or 2-hour postprandial glucose levels
between CGM and POC BG testing. In a pilot study, CGMs detected a higher number of
hypoglycemic events compared to POC BG testing, particularly nocturnal or asymptomatic

hypoglycemia⁹³. Few studies have been published on the use of newer factory calibrated CGMs
 in non-ICU settings.⁹⁴

383 A recent study of patients with T2DM admitted to general medicine and surgery wards and managed with basal-bolus insulin therapy, compared the FreeStyle Libre Pro (Abbott Diabetes 384 Care, Alameda, California)⁹⁵ to POC BG testing⁹⁶. This CGM system is a variant of the FreeStyle 385 Libre 14 day system, where glucose readings are available to the HCP but not to the patient. 386 387 The FreeStyle Libre Pro CGM, compared to POC BG testing, showed a tendency towards lower mean glucose with an estimated mean glucose difference of 12.8 mg/dL (Confidence Interval, CI 388 389 8.3-17.2). Accordingly, CGMs, compared to POC BG testing, were more sensitive at detecting 390 hypoglycemic events. The overall Mean Absolute Relative Difference (MARD) was 14.8%. The 391 percentage of glucose concentrations within the $\pm 15\%$ or 15 mg/dL, $\pm 20\%$ or 20 mg/dL, and $\pm 30\%$ or 30 mg/dL (where for CGM concentrations $\leq 100 \text{ mg/dL}$, the units of the range were 392 393 mg/dL and for CGM concentrations > 100 mg/dL, the units of the range were percent) was 394 62%, 76%, and 91%, respectively. A Clarke Error Grid analysis showed acceptable clinical accuracy with 98.0% of glucose concentrations falling into Zones A (75.1%, n=1,184) and B 395 (23.7%, n=374).⁹⁶ Panelists reviewed CGM studies in the non-ICU in adult populations (Table 5). 396 Evidence suggests that initiating the use of CGMs in the non-ICU settings provides better 397 398 glycemic monitoring, compared to standard 3-4 times daily POC BG testing, with improved 399 detection and potential prevention of hypo- and hyperglycemic events. Most of these events, particularly nocturnal and asymptomatic hypoglycemia, might otherwise be missed. Ongoing 400 hospital CGM studies listed on ClinicalTrials.gov⁹⁷ may provide some guidance (**Table 6**). 401

402 Glucose Telemetry

403	The hospital should possess the physical infrastructure to download the patient's CGM data for
404	the retrospective review of patterns in glycemia. CGM data can be automatically delivered to
405	the nursing station by way of automatic downloading into a monitor at the nursing station. A
406	recently published manuscript evaluated whether such a system for presenting CGM data,
407	called the "Glucose Telemetry System", can decrease hypoglycemia in the general wards/non-
408	ICU setting ⁴³ . This report is the first interventional randomized controlled trial (RCT) study of
409	CGM technology to improve outcomes in the non-ICU setting. The study included patients with
410	T2DM, who were at high risk for hypoglycemia. Participants were randomized to either the
411	"Glucose Telemetry System" (intervention group) or to POC BG testing (control group). For
412	patients in the "Glucose Telemetry System", nurses were instructed to proceed with
413	hypoglycemia prevention actions if the low glucose alerts were activated (for a setting of BG <
414	85 mg/dL). Participants in the control group were placed on "blinded" CGMs which were only
415	used to collect glucometric data. Overall, the subjects in the "Glucose Telemetry System"
416	experienced fewer events of hypoglycemia (BG < 70 mg/dL) and clinically significant
417	hypoglycemia (BG < 54 mg/dL) compared to the POC BG group. The outcomes of the
418	intervention versus control groups for these two levels of hypoglycemia were, respectively,
419	0.67 versus 1.69 events/ patient, p =0.024 (BG < 70 mg/dL) and 0.08 versus 0.75 events/patient,
420	p =0.003 (BG < 54 mg/dL). There was a reduction in percentage of time in hypoglycemic range
421	(BG < 70 mg/dL and < 54 mg/dL) in the glucose telemetry system group compared to POC group
422	(0.40% versus 1.88%, p =0.002 and 0.05% versus 0.82%, p =0.017).

423 **POTENTIAL BARRIERS**

424 Minimally Invasive CGMs

As discussed in previous consensus reports^{1,106} during the past 20 years, many studies have 425 426 been published on the initiation of subcutaneous CGMs in critically ill patients (Table 3 and 4). However, most of those studies were intended to focus only on accuracy data and not clinical 427 outcomes. In addition, it is difficult to reach conclusions from these reports because of different 428 429 study designs and small sample sizes. A recent systematic review by van Steen et al. analyzed 430 32 studies that assessed the accuracy of CGMs in the ICU. These authors reported moderate to good accuracy especially with intravascular devices¹⁰⁷. The authors included only five RCTs for 431 efficacy assessment and recognized methodological limitations ¹⁰⁷. Panelists noted that there is 432 currently insufficient data to provide definitive recommendations on improved outcomes based 433 on reports in the medical literature. 434

435 It is unclear whether CGMs will be able to fully replace POC BG testing and be approved as nonadjunctive use for treatment decisions in acute care. Panelists had concerns with the accuracy 436 437 of subcutaneous CGM values for the first hours after insertion to make treatment decisions or 438 even during the first 1-2 days of use. Panelists also had concerns with the unintentional added burden on nursing when: 1) a CGM has overreported low glucoses values and these false low 439 values have required POC confirmation, 2) new CGM technology must be learned during a 440 crisis, and 3) time is needed for troubleshooting. In addition, skin-related issues have been 441 mentioned in 19% of articles about recent CGMs.^{108–110} 442

443 Invasive CGMs

Although these systems were not the focus of the guideline, the panelists briefly considered the
 role of invasive CGMs. They noted that few intravascular invasive sensors are cleared for ICU

446	patients. Also, compared to subcutaneous CGM sensors, intravascular sensors tend to have			
447	three main disadvantages. First, these systems are invasive and some are associated with			
448	vascular complications, such as thrombosis, catheter occlusion, biofilm formation, or			
449	intravascular catheter-related infections ¹¹¹ . Second, they impose a higher implementation			
450	resource and care burden to patients and the ICU system. Third, they are not intended for non-			
451	ICU settings. Therefore, intravascular CGMs, compared to subcutaneous CGMs, are less			
452	attractive options.			
453	Recommendations for Initiation of Continuous Glucose Monitors in the hospital			
454	Clinical Practice			
455	Strong Recommendation			
456	HCPs should consider prescribing CGMs to reduce the need for frequent nurse contact			
457	for POC glucose testing and the use of PPE for patients on isolation with highly			
458	contagious infectious diseases (e.g. COVID-19).			
459	Mild Recommendation			
460	HCPs should avoid initiating CGMs in patients with severe hypoglycemia or			
461	hyperglycemia (i.e. BG < 40 mg/dL or >500 mg/dL) or during periods of rapid glucose			
462	fluctuations.			
463	Research			
464	Strong Recommendations			
465	• Researchers need to provide data to support initiation of CGMs for improving patient-			
466	centered outcomes.			

467	• Researchers need to provide data on hospital outcomes when initiating CGMs in the
468	hospital, including improved glycemic outcomes, detection and/or reduction of
469	hypoglycemia and hyperglycemia, reduction of ICU LOS, and cost-effectiveness.
470	• Researchers need to conduct studies on long term benefits for initiating CGMs in the
471	hospital after discharging patients with newly diagnosed diabetes or recurrence of
472	diabetic ketoacidosis (DKA) or other complications of diabetes.
473	• Manufacturers need to develop educational tools for patients, hospital staff, and HCPs.
474	Hospital Policies
475	Strong Recommendations
476	Hospitals need to develop plans, including process maps, protocols, staff educational
477	resources, and order sets for prescribing CGM use during hospitalizations prior to
478	implementing a CGM.
479	• Hospitals need to provide educational tools for patients, nurses, house staff, and
480	attending physicians when a patient in the hospital starts on a CGM.
481	
482	Continuation of Automated Insulin Dosing Systems in the hospital
483	Chair: Ananda Basu, MD, FRCP
484	University of Virginia School of Medicine, Charlottesville, Virginia, USA
485	POTENTIAL OPPORTUNITIES

486 Improved Glycemic Outcomes

487	Evidence about the potential glycemic benefits of continuing AID systems from the outpatient
488	into the inpatient setting is limited, and currently it is possible only to extrapolate data from
489	studies of AID systems initiated during a hospital stay. Several such studies of initiating AID
490	systems in the hospital have been performed in medical or surgical patients as well as in
491	patients on hemodialysis or women in the peripartum/postpartum period $^{112-119}$. In the largest
492	of these studies ¹¹² , Bally et al. reported that initiation of AID system technology in the hospital
493	for patients receiving noncritical care achieved a higher percentage of TIR when compared to
494	standard hospital management. The times in range were, respectively, 65.8 (±Standard
495	Deviation 16.8)% vs 41.5(±16.9)%, with a difference of 24.3 (±2.9)% [95%Cl 18.6 to 30.0;
496	P<0.001). Mean glucose levels were lower in the AID system arm compared to the group
497	treated with conventional subcutaneous insulin delivery (with the differences being 154 (\pm 29)
498	mg/dL vs 188 (\pm 43 mg/dL), p <0.001) and there was no significant difference in time spent in
499	hypoglycemia <54 mg/dL or < 70 mg/dL. AID systems have also been found to improve TIR in
500	women in the peripartum/postpartum period ¹¹³ and patients on hemodialysis ¹¹⁴ . AID system
501	management has reduced surgical site infections resulting in shorter postoperative
502	hospitalizations ¹¹⁵ . In a single center observational study that was performed in an ICU setting,
503	use of AID system management compared to standard sliding scale insulin therapy led to a
504	decreased frequency of blood sampling, reduced time required for achieving glycemic targets,
505	and a decreased nursing workload per admission of diabetes management from 68 (\pm 25)
506	minutes (AID system) to 33 (± 21) minutes (sliding scale) (p < 0.001). ¹¹⁶ In a randomized,
507	parallel-group trial, inpatients with T2DM in the United Kingdom received fully closed loop
508	insulin delivery without meal-time boluses which was found to be safe and effective ¹¹⁷ . In a

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509	two-center open-label, randomized controlled trial of fully automated insulin dosing in the
510	United Kingdom and Switzerland, this method was found to improve glycemic outcomes for
511	inpatients receiving nutritional support ¹¹⁸ .
512	Glycemic management in hospitalized patients aims to avoid both hypoglycemia and
513	hyperglycemia. Since inpatients with diabetes are often in a compromised state of health and at
514	risk for hypoglycemia because of interrupted nutrition, inadvertent insulin overdosages
515	associated with intensive insulin therapy, or unexpected improvements in insulin sensitivity,
516	hypoglycemia can be a serious problem for these patients. Special AID systems that can deliver
517	both insulin and glucose have been created exclusively for inpatient use. A clinical study in
518	Japan compared two such systems (differing in size and weight, but not algorithms)
519	manufactured by Nikkiso Co., Ltd., and used for perioperative glycemic management. The
520	newer (STG-55) and older (STG-22) AID system models ¹²⁰ both achieved similar glycemic control
521	without hypoglycemia, leading the investigators to conclude that the newer (as well as smaller
522	and lighter) system could potentially be used in routine practice for perioperative glycemic
523	management ¹¹⁹ . A study in Denmark assessed an intravenous AID infusion system delivering
524	both insulin and glucose based on a proprietary controller (Admetsys, Boston,
525	Massachusetts) ¹²¹ .

526 **COVID-19**

527 With the COVID-19 pandemic, increased mortality has been associated with hyperglycemia 528 both in patients diagnosed with diabetes prior to admission and those diagnosed with diabetes 529 during their admission¹²². There is a paucity of high-quality data about optimal monitoring and

therapy and associated outcomes in these patients. The need for improved glycemic
management for COVID-19 patients may accelerate the development of future novel glucose
monitoring technologies in the hospital setting, including possibly closed loop control for
intensively treated patients. During the pandemic, AID systems, if utilized, can also perhaps
reduce the risk of nursing exposure, the time needed for donning and doffing for any needed
BG monitoring, and the use of limited supplies of personal protective equipment.

536 Patient Satisfaction

Evidence about the potential benefits of using of AID systems in the inpatient setting is limited. 537 538 Even for the more traditional non-AID CSII system, the available data is based on retrospective studies, because no randomized clinical trials have been performed¹²³. One of these studies 539 reported that outpatients on CSII systems, who had reasonable control (mean hemoglobin A1c 540 7.5%)^{124,125}, were sufficiently confident to continue self-managing their diabetes and use their 541 542 own CSII systems during a hospitalization. Many of these CSII system users reported higher patient satisfaction (86%) when they were allowed to continue wearing their CSII system during 543 their inpatient stay¹²⁶. Similar outcomes are likely to be found with the use of AID systems. 544 Asking hospitalized patients with diabetes to remove their AID system could result in decreased 545 patient satisfaction, especially if their diabetes care is managed by healthcare professionals, 546 who have limited experience with inpatient and outpatient diabetes management. 547 548 Furthermore, a patient who must surrender their AID system upon hospitalization might express dissatisfaction with nocturnal POC BG testing. 549

550 **POTENTIAL BARRIERS**

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551 Patient-Related Factors

Although AID systems can be beneficial, five types of factors may preclude their use in the 552 inpatient setting.^{123,124,127} They can be divided into the following categories: 1) patient-related, 553 554 2) hospital-related, 3) device-related, 4) medication-related, and 5) surgical procedure-related. Examples of patient-related conditions in which AID systems should not be used are physical or 555 556 psychiatric conditions which can make patients incapable of self-managing an AID system in the hospital. Contraindications to CSII system and AID system therapy in the hospital are presented 557 558 in **Table 7**. Patients should be able to self-manage their AID systems and provide their pump 559 settings to the treating HCPs in case the AID system may need to be discontinued. Patients with severe metabolic decompensations, such as DKA¹²³, acute kidney injury, post-transplant T1DM 560 patients in acute rejection, or those with severe sepsis and hypovolemia, which may lead to 561 tissue hypo-perfusion, should also probably not use AID systems in the hospital. Skin infections 562 563 may represent another contraindication, especially if they are extensive, because they may 564 preclude CGM or pump placement. However, it is still unclear whether the above conditions can significantly affect the function of AID systems and more research is needed in this area. 565

566 Hospital-Related Factors

Examples of hospital related factors are situations where there are no policies in place that can safeguard the use of AID systems in the inpatient setting and delineate the roles of the patients, nurses, and HCPs^{124,127}. Because only limited information is currently available about the use of AID systems in the hospital, further research is needed in order to provide evidence-based recommendations¹²⁷. Another potential obstacle to the use of AID systems in the inpatient

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setting is the lack of nurses and HCPs who are adequately trained in the use and interpretation
of data from the AID systems. However, it is unclear whether AID systems do or do not lead to
increased workload for nursing and/or HCPs.

575 **Device-Related Factors**

Limitations related to device use include clinical scenarios where AID systems cannot be used 576 because of a device malfunction or insufficient medical supplies, either for the continuous 577 578 insulin infusion set or for the CGM components. A CGM can become compressed during a prolonged period of a prone position, such as with sleep or prone ventilation, and produce a 579 false low reading, which could also pose another limitation to their use ^{128,129}. For AID systems 580 581 that require the patient to select a meal-time bolus dose recommended by a bolus calculator, unexpected failure to reach postprandial glycemic targets could be due to manufacturer-582 specific pump settings resulting in a different dose recommendation by each pump brand.¹³⁰ 583

584 Medication-Related and Meal-Related Factors

585 Medications, such as glucocorticoids, which can cause severe insulin resistance and

586 uncontrolled hyperglycemia, may present a challenge for some AID systems, but others may

- ⁵⁸⁷ adapt well to changes in insulin resistance during periods of illness¹³¹. Other challenging
- 588 scenarios are nutritional interruptions, which are very common in a busy hospital

589 environment¹³¹. Nutrition in the inpatient setting is more complicated than in the ambulatory

- 590 environment. Patients may have nausea, vomiting, or other conditions that can affect nutrient
- absorption and therefore create irregular patterns in the glucose values. Insulin is not always
- administered at the right time before the meal is delivered. Meals can be interrupted or

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593	delayed and tube feedings and parenteral nutrition (either peripheral or total) can be suddenly
594	discontinued. Although the above scenarios are not absolute treatment-related
595	contraindications, they represent challenging situations for AID system use in the hospital. HCPs
596	should also be aware about the potential interactions of certain medications with subcutaneous
597	CGMs (Table 2). Additional studies are required to determine the effects, if any, of multiple
598	doses and combinations of potentially interfering medications on CGM accuracy.

599 Surgical Procedure-Related Factors

Surgical procedures can create additional barriers to the use of AID systems in the inpatient 600 setting^{123,125,132}. Surgical procedures can be broadly divided into two different categories, 601 602 elective or urgent. Elective surgeries can provide sufficient time for pre-admission preparation. 603 The endocrinology clinician or diabetes team would coordinate care between the different subspecialties that are involved such as the anesthesiology, surgical and inpatient diabetes 604 605 teams (if they are available and different from the primary endocrinologist) about the upcoming surgical procedure. The panel recognized that many hospitals do not have a diabetes 606 607 team or inpatient diabetes educator. Patients need to be instructed to insert the sensor and the insulin cannula away from the operative field and change the sites one day prior to the surgery. 608 609 Urgent surgeries do not allow for such planning. In the immediate preoperative period, for 610 either elective or urgent surgical procedures, the inpatient diabetes team should be notified, if 611 this has not been done earlier. Consent must be obtained from the patient about the use of an 612 AID system during surgery. Temporary higher glycemic targets may be needed to allow slightly 613 higher glucose values during surgery to decrease the risk of hypoglycemia in an unconscious patient. Ideally, the anesthesiology team would need to be familiar with the use of an AID 614

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615	system during the intraoperative period so the team can control or suspend the pump if
616	necessary because the unconscious patient will not be able to adjust the settings themselves.
617	However, it is unclear whether it would be realistic to expect an anesthesiologist to learn the
618	operation of an AID system and there is no data about anesthesiologists operating AID systems
619	during surgery. The basal insulin delivery rate is determined by an AID system controller. If the
620	team members are able to manage the AID system, then they should also have easy access and
621	proximity to the AID system intraoperatively. The use of an AID system during surgery is not
622	recommended if the insulin requirements are expected to fluctuate significantly
623	intraoperatively. In that case intravenous insulin delivery with insulin dosing software instead of
624	subcutaneous insulin delivery would be more appropriate with either an intravenous or
625	subcutaneous glucose sensor. AID systems can be continued during the operation if there are
626	no concerns regarding device malfunction. However, there is no good data available on the
627	safety or maximum safe duration of closed loop control during anesthesia. Even with control by
628	an AID system, BG concentrations should be monitored intraoperatively.
629	
630	Recommendations for Continuation of Automated Insulin Dosing Systems in the hospital
050	Recommendations for continuation of Automated Insulin Dosing Systems in the hospital
631	Clinical Practice
632	Strong Recommendations
633	HCPs should prescribe AID systems only for appropriate candidates, who will need to
634	have adequate knowledge and skills for using AID systems

635	 HCPs should reassess a decision periodically to transition use of outpatient AID systems
636	into the hospital in order to ensure that AID system continue to represent the best
637	treatment option for each patient
638	HCPs should prepare an alternative plan for diabetes management in case it becomes
639	inappropriate for a patient to continue using an AID system in the hospital
640	HCPs should discontinue AID systems in critically ill hospitalized patients (such as those
641	with hypovolemia or sepsis)
642	• HCPs should recognize glycemic patterns due to CGM compression, which can cause
643	false low readings
644	Mild Recommendation
645	HCPs should avoid initiating an AID system during a hospitalization
646	Research
647	Strong Recommendations
648	• Researchers need to conduct studies about whether continuing AID systems in the
649	hospital is beneficial to improve glycemic management or clinical outcomes
650	• Researchers need to provide data on hospital outcomes when using AID systems in the
651	hospital, including improved glycemic outcomes, detection and/or reduction of
652	hypoglycemia, reduction of ICU LOS, and cost-effectiveness
653	• Manufacturers need to research whether all types of CGMs and AID systems can be
654	used during radiological/imaging studies or diathermy

655 Hospital Policies

656	Strong Recommendations
657	Hospitals need to develop institution-specific protocols and order sets for the proper
658	use of AID systems during a hospitalization
659	Hospitals need to require that patients using AID systems bring with them sufficient
660	supplies for these devices during a hospitalization
661	Hospitals need to develop protocols for using AID systems during elective procedures
662	and surgeries
663	Recommendation Not Reaching Consensus
664	• HCPs should switch AID systems from "auto" mode to "manual" mode when a patient is
665	admitted to the hospital wearing an AID system
666	
667	Logistics and hands-on care of hospitalized patients using Continuous Glucose Monitors and
668	Automated Insulin Dosing Systems
669	Chair: Suzanne Lohnes, MA, RN, CDCES, CPT
670	University of California San Diego Medical Center, La Jolla, California, USA
671	POTENTIAL OPPORTUNITIES
672	Expectations for Patients and Hospital Staff and Practical Considerations for Use of CGMs and
673	AID systems in the Acute Care Setting
674	Continuation of CGM use can be a helpful adjunct to management in the acute care setting and
675	can increase patient satisfaction. However, because CGMs are not currently cleared by FDA for
676	the inpatient environment, a policy addressing practical considerations for use of CGMs and AID
677	systems in hospitalized patients is needed.

678

679 **POTENTIAL BARRIERS**

680 Necessary Hospital Responsibilities

- 681 It is important that key tasks, roles, and responsibilities, related to work system domains
- 682 (technology/data, tasks, personnel, structure/organization, and environment) are addressed for
- safe and effective implementation.¹³³ Below are listed potential responsibilities delineated by
- team members. It is helpful for diabetes team members to be interchangeable (e.g.
- subspecialty consultant with pharmacist or nurse with patient care technician). Furthermore, it
- is appropriate to predefine tasks, person assignments, policies, procedures, and a clear
- 687 organizational structure (e.g. determination of committee reporting) around monitoring and
- 688 interpretation of data, to facilitate use of CGMs and AID systems.

689 Necessary Patient Responsibilities

690 Patients who wish to continue use of CGMs or AID systems in the acute care setting should read

a detailed set of information and should review and sign a patient agreement about hospital

692 policy. The panel developed a sample patient agreement for the use of CGMs or AID systems in

- the hospital presented in **Figure 1**. This agreement is meant to be an example for a
- 694 subcutaneous non-implanted sensor. Each institution must develop their own agreement and
- 695 they should consider manufacturer labeling.
- 696 CGMs may be used for guidance about the direction and magnitude of changes in glucose
- 697 concentrations. The patient should notify hospital staff if they are observing glucose excursions
- out of range or if they experience symptoms of hypoglycemia. The patient should bring all
- supplies (infusion sets, sensors, receiver, etc.) needed for continuation of home use for the

duration of a hospitalization and be responsible for maintenance of their device and changing
sites as directed during a hospitalization. Device supplies may be stored per hospital policy and
will be returned to the patient upon discharge.

703 Necessary HCP Responsibilities

704 Inpatient caregivers must: 1) confirm that it is appropriate for a patient to continue using a

CGM or an AID system, 2) discuss hospital policy with the patient, and 3) review an agreement

with the patient. After the patient agreement is signed, the HCP should place an order for

inpatient use of a CGM or an AID system. A patient's ability to safely continue use of a CGM or

an AID system (which may change during the hospitalization) must be regularly assessed by

nursing staff and HCPs.¹³⁴ Daily documentation per institutional policy will be needed

throughout the hospitalization. If there is concern for patient's ability to use a CGM or an AID

711 system, then the caregiver will recommend an alternative treatment plan.

712 Necessary Nursing Responsibilities

713 In collaboration with other inpatient HCPs, it is important for nursing to assess the patient's suitability for using a CGM or an AID system and review hospital policies with the patient. It is 714 715 also important for nursing to assess the insertion site and document site changes in the EHR. Treatment decisions based on CGM data linked to insulin dosing software might lead to 716 717 unwanted outcomes unless the safety and efficacy of the system in the acute care setting can 718 be clearly established. For patients using AID systems in the hospital who are going to be transitioned to and/or discharged with subcutaneous multiple dose insulin therapy, if the 719 insulin dosing information (from "auto" mode) is not available in the EHR, then an estimate of 720 721 insulin requirements might be inaccurate and could lead to dysglycemia following discharge.

722	Standard approaches to documentation are also needed. The panel recognized a spectrum of
723	practice for nursing documentation and institutional requirements. Nursing should document
724	all AID system device settings, including any insulin boluses in "manual" mode, in the inpatient
725	progress notes and/or in the patient's bedside log which is scanned into the EHR. Additionally,
726	the frequency that this information is documented (i.e. every shift vs. daily) may vary based on
727	individual hospital resources and policies.
728	Specialty Consultation
729	When using CGMs or AID systems in the acute care setting, specialty consultation, if available, is
730	required and the request for consultation should be documented. While some institutions have
731	inpatient diabetes support available for in-person consultation and ongoing management, the
732	panel recognizes there are circumstances in which inpatient diabetes expertise may not be
733	readily available. The panel suggested consideration for telemedicine consultation with a
734	diabetes specialist if necessary. It is useful to document the patient's ability to use the
735	technology to assist with glucose management.
736	Recommendations for Logistics and hands-on care of hospitalized patients using Continuous
737	Glucose Monitors and Automated Insulin Dosing Systems
738	Clinical Practice

- 739 Strong Recommendations
- HCPs **should** inquire about and document the medication and supplement history of
- 741 patients who use CGMs to determine whether there are any agents that can interfere
- 742 with glucose measurements

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743	•	HCPs should ensure that off-label use of CGMs and AID systems is consistent with
744		medical practice and appropriate precautions have been taken to protect patients
745	•	Nursing should document hands-on training of CGM use and AID system therapy
746		through a technology certification program
747	•	Nursing should confirm that the patient is appropriate to continue using a CGM or an
748		AID system and also review the agreement and hospital policy with the patient
749	•	Nursing should inspect the insertion site every shift with attention to skin integrity and
750		signs of erythema or infection, and should document site changes
751	•	Nursing should know device basics, institutional policies, HCPs roles, and whom to
752		contact if questions arise
753	•	Nursing should administer a patient competency assessment or survey to assess patient
754		ability to safely assist with managing a CGM or an AID system
755	•	Nursing should set expectations and clarify that there will be a need to continue
756		checking POC capillary glucose even when using a CGM.
757	•	Nursing should measure POC BG concentrations to confirm or supplement CGM
758		readings (usually a minimum of 4 times daily: before each of three meals and at bedtime
759		if patients are eating, or every 6 hours if patients are fasting) as well as at patient
760		request; however, the CGM glucose, trend arrows, and rate of change may be used to
761		help determine if and when a BG test is required.
762	Resea	rch

763 Strong Recommendations

764	• Researchers need to conduct further studies on the best logistics and hands on care for
765	patients using CGMs and AID systems to achieve the best outcomes
766	• Manufacturers need to research interoperable components for AID systems that are
767	compatible with hospital EHRs
768	Hospital Policies
769	Strong Recommendations
770	• Hospitals need to provide interpreter services to translate CGM and AID system
771	agreements
772	Hospitals need to state in their policy and patient agreement documents that treatment
773	decisions will be based on hospital-calibrated BGM readings (or laboratory readings) and
774	not on CGM readings, barring a need to isolate a patient with a severe and highly
775	contagious infection
776	• Hospitals need to maintain their CGM and AID system policy and patient agreement
777	documents in easily accessible electronic files stored in the EHR order set for CGMs and
778	AID systems
779	Hospitals need to develop policies for when to discontinue or temporarily suspend the
780	use of CGMs and AID systems
781	Hospitals need to survey their HCPs, nursing, and patients to improve outcomes and
782	satisfaction

783

784

785 the hospital Chair: James H. Nichols, PhD, DABCC, FAACC 786 Vanderbilt University Medical Center, Nashville, TN, USA 787 788 **POTENTIAL OPPORTUNITIES Policies and Procedures** 789 790 As previously noted, there is a distinction between CGM glucose values and laboratory glucose 791 values, and CGM data is currently not part of the laboratory information system. Rather, CGM data is analogous to ICU vital sign monitoring data rather than lab values like serum potassium 792 793 and sodium. Because of this distinction, it is important to consider where in the medical records this data should reside and how they should be displayed, such as in reports, tables, or graphs. 794 Given this known difference between CGM glucose values and lab glucose values,¹³⁵ criteria 795 796 should also be developed on when to check or cross-reference CGM values with a POC or 797 laboratory glucose test. A related question is whether or not clinical decisions should be made 798 on the basis of CGM data, or whether clinicians should always obtain a laboratory or POC 799 glucose test for treatment decision making. Finally, criteria should be established as to whether a minimum number of laboratory or POC BG tests must be performed while patients are using 800 CGMs or AID systems in the hospital. Manufacturers of some CGMs have recommended a 801 802 calibration frequency, but those recommendations are intended for outpatient use, and might

Data management of Continuous Glucose Monitors and Automated Insulin Dosing Systems in

803 not be adequate for inpatient use.

As part of the standardization of summary metrics, we should also develop clear criteria for 804 805 values or trends that require a clinical intervention. The panel discussed creating a framework for clinical action based on CGM data. This includes understanding what data and trends are 806 807 actionable, as well as what the appropriate clinical interventions might be. Critical values are 808 considered to be imminently life-threatening test results that require immediate contact by the 809 ordering HCPs. CGMs can trend the rise and fall of glucose concentrations, and can predict critical hypo- or hyperglycemia. Data management systems can be set to alarm when CGM glucose 810 811 trends reach or cross certain critical values. These alarms should lead to clinician and patient 812 notification so that appropriate actions may be taken in a timely fashion. The panel noted that data and security are major concerns in Germany and the rest of Europe. In 813

Europe, every manufacturer uses a different data scheme and interface to download their data, which can be confusing.

816 Information Technology (IT) Infrastructure

817 The Health Insurance Portability and Accountability Act of 1996 (HIPAA) protects health 818 information, promotes transparency, trust, and patient welfare in medical practice. Since CGMs 819 and AID systems collect protected health information (PHI), when they are used by institutions 820 and clinicians to make medical decisions, institutions have a responsibility to treat it like all other PHI, meaning they must ensure the integrity, security, and appropriate availability of that 821 data. Documenting CGM results and data in the EHR designates it as part of the medical record, 822 823 and it becomes subject to HIPAA. The IT department is needed to assist with licenses to 824 download the data, and install the software into each hospital system.

825	Healthcare facilities should adopt the Unique Device Identifier (UDI) system to track devices in
826	the EHR. In 2013, the FDA issued guidelines for the implementation of a global UDI system to
827	adequately identify and track medical devices across their lifecycle , from distribution to patient
828	use ¹³⁶ . The UDI final rule established a timeline for all qualifying medical devices in the US to be
829	compliant with UDI labeling by 2022 ¹³⁷ . Diabetes technologies like BGMs, CGMs, CSII systems,
830	and AID systems are all required to bear a UDI. Institutions should rapidly move toward UDI
831	adoption and integration into the EHR, and ensure that CGM and AID system data is associated
832	with the correct UDI for safety and quality assurance.

833 **Data**

Panelists recognized that there is limited evidence on how CGM data is integrated into EHRs at this time. With the near-universal adoption of EHRs among inpatient facilities in the United States ¹³⁸, integrating device data into the EHR is important for quality and consistency. Several groups have explored the integration of these data into the EHR ^{139–141}, but many questions still remain regarding best practices for the acquisition, storage, display, and use of that data.

Distinctions should be made when recording CGM data in the EHR, since CGM data differs from laboratory glucose results. CGMs measure glucose within interstitial fluid, while laboratory instruments measure glucose in plasma, serum, or whole blood. This means that CGM data may not agree with laboratory glucose measurements collected at the same time.¹³⁵ While individual CGM data points may be less precise than lab instrumentation generated values, a major advantage offered by CGMs is the presentation of multiple data points over time. These create an opportunity to evaluate glucose patterns as well as trends in the rate of change, percent of

time spent hypo- or hyperglycemic or within target range, and estimate stability/instability of the
glucose concentration over time. These summary patterns may be more valuable than individual
data points and provide a synthesis of the patient's overall glycemic status.

849 Data Patterns

As EHR integrations of CGM data become more common, HCPs with a wider variety of backgrounds in training and experience with CGM data interpretation will have access to this data. Some might be less familiar with its use and interpretation. It is important that standardized, clear, and interpretable summary metrics be established in order to facilitate the clinical use of CGM data in the hospital setting.

855 When considering how to integrate device data, the first decision is how to source data. There are two main options: 1) obtaining the data directly on a platform provided by the manufacturer 856 857 (e.g. Abbott, Dexcom, or Medtronic) and 2) obtaining the data from a third-party aggregator, e.g. 858 Tidepool (Tidepool, Palo Alto, California) or Glooko (Glooko, Inc., Mountain View, California). Each of these approaches has advantages and disadvantages, as well as associated costs and 859 860 technical requirements. It may be reasonable to use a hybrid approach, connecting directly with 861 a few manufacturers that have significant market share, and then using an aggregator to capture other devices. 862

The next decision is what data to extract. There are several options for extracting, storing and displaying CGM data, and at varying levels of complexity (**Table 8**). Static reports (view only documents, typically PDFs) are the simplest, and some CGM manufacturers have already developed mechanisms to bring the CGM reports found on their provider platforms into the EHR.

Structured summary data are predefined and standardized, and can be added to existing data tables in the EHR for charting, trending, etc. Structured continuous data refers to the hundreds of daily individual blood glucose measurements, and is the most complex to manage, but potentially offers the most flexibility and control.

Data storage and display will be dictated by the type of data extracted from the device. Reports and structured summary data can be stored in native EHR data tables, but continuous glucose readings would likely overwhelm those tables, and would best be stored in a separate environment. In terms of displaying the data, this can be accomplished in a variety of ways described in **Table 8**.

A consensus list of core data elements should be developed and standardized across all models 876 and manufacturers. Data standards and ontologies are critical for ensuring interoperability across 877 information systems¹⁴². A core set of data elements and definitions developed and applied by the 878 entire CGM industry would facilitate storage and use of CGM data. Finally, core data elements 879 would ideally be submitted to the appropriate governing bodies for inclusion in existing 880 healthcare ontologies and common data models, such as Systematized Nomenclature of 881 Medicine—Clinical Term (SNOMED-CT), Logical Observation Identifiers Names and Codes 882 (LOINC), and Observational Medical Outcomes Partnership (OMOP). 883

Patient-reported outcomes (PROs) are any reports of the status of a patient's health condition that come directly from the patient, without interpretation of the patient's response by a clinician or anyone else.¹⁴³ PROs can be leveraged for research, clinical care, and quality improvement. While several groups are actively working on the development of PROs in diabetes, there is still

890	Atypical Scenarios
889	technology-specific PROs will enable a more holistic approach to patient care and research.
888	significant work to be done ¹⁴⁴ . The development, dissemination, and implementation of diabetes

891 Guidelines should address the use of CGMs and AID systems for diagnoses other than diabetes, 892 where glucose monitoring is valuable. In pediatrics, several clinical situations require close 893 monitoring of BG concentrations and tight glycemic control, such as the titration of glucose 894 infusion rates in premature infants on total parenteral nutrition. Early detection of hypoglycemia in infants with inborn errors of metabolism (e.g., fatty acid oxidation disorders, ketotic 895 896 hypoglycemic disorders, and disorders of gluconeogenesis) could be another critical use for CGMs 897 in the hospital setting. In these diseases, infants are often allowed to become hypoglycemic as a challenge in order to draw critical diagnostic labs. CGM measurements could make that process 898 less stressful for parents and HCPs and safer for patients. 899

900 Economic Analysis

901 Panelists had concerns with the costs of some CGMs and AID systems being a limiting factor (i.e., batteries, sensors, transmitters, and/or a monitor or smartphone), but found that some CGMs 902 903 are affordable. Panelists considered questions about the reimbursement for these devices. Who 904 is responsible for covering their costs and consumable components? What if the patient has a device from one manufacturer, but the hospital only stocks supplies from a different 905 906 manufacturer? Panelists also discussed the economic implications of CGM and AID system use in 907 hospitalized patients. Inpatient hypo- and hyperglycemia, which might prove to be reduced with 908 structured CGM or AID system programs, have been associated with increased LOS,

readmissions, and costs^{48,145}. In patients undergoing cardiac surgery, studies suggested potential
 cost saving with intensive glycemic management (targeting 100 -140 mg/dL)¹⁴⁶. Finally, panelists
 acknowledged the need for well-powered studies comparing the use of CGMs vs POC BGMs on
 hospitalization costs. ¹⁴⁷

913 **POTENTIAL BARRIERS**

914 **Regulatory Considerations**

915 The Clinical and Laboratory Improvement Amendment of 1988 (CLIA) sets a minimum quality 916 standard for any laboratory test performed in the US for patient care or clinical decision making. Externally attached patient-dedicated monitoring devices like pulse oximetry capnography are 917 not subject to CLIA¹⁴⁸. CGMs and AID systems are also automatic monitoring devices that are 918 wearable and continuously or intermittently detect glucose concentrations in interstitial fluid or 919 tissue fluid. There is no sample collection and analysis in a separate instrument that can be 920 921 calibrated or validated with a Quality Control sample. As such, a CGM is more of a monitoring 922 device than a laboratory instrument, and should not be subject to CLIA.

Although CGMs and AID systems should not be subject to CLIA, quality control is still an important consideration for inpatient CGM and AID system use. Previous consensus panels have stressed the need for clear safety and quality protocols to be in place¹. There is known variation between sensors, both between brands and within brands. Also, calibration errors can lead to significant deviations in glucose values. Currently some hospitals using CGMs require a patient agreement, which outlines that the patient can still use their CGM, but hospital BGM testing is still mandatory. See **Figure 1** for a sample agreement. In Germany, laboratory quality control guidelines require twice daily internal testing and quarterly external testing for hospital lab
meters¹⁴⁹. This is a prerequisite for the use of data for diagnostic or therapeutic decisions. With
CGMs, there is no sample and no control materials, so these procedures cannot be applied to
CGMs, which is why some BG monitoring is still mandatory in the hospital. One possible path
forward is for manufacturers to develop a mechanism to perform quality control procedures for
CGMs. Otherwise, CGMs in the hospital may be limited to adjunctive use only.

936 Off-label use of prescription drugs and devices is common in modern medical practice, and has 937 been recognized as "an accepted and necessary corollary of the FDA's mission to regulate in 938 this area without directly interfering with the practice of medicine" by the United States Supreme Court ¹⁵⁰. A manufacturer may not market unapproved uses of a medical device, but a 939 940 physician may in their independent judgement decide to use a cleared device in an off-label manner. While off-label use is seen as accepted practice, it does not shield physicians from 941 942 liability, and there is potential tort exposure. Whether a hospital would also be liable under 943 those circumstances would probably depend on what sort of control it exerted over the 944 physician. If it is for an employed physician, then the hospital might be liable for the physician's actions under a theory of respondeat superior, which is a doctrine that states that an employer 945 is responsible for the acts of an employee. If the physician is an independent 946 947 contractor, then hospital liability for the physician's actions would be more difficult to 948 establish. One way to evaluate the liability or legal risk of off-label use is to consider whether or not that action may expose the practitioner to a claim of negligence or malpractice. 949 950 Negligence can be thought of as a breach in duty (for example, to a patient), or as the failure to 951 act reasonably in light of foreseeable consequences.

952

953 Data privacy and security

954 Another potential risk is around the data itself, and whether it is being stored and protected 955 with the proper precautions for PHI. Overall, this should not be seen as an obstacle provided it 956 is consistent with standard practice. Tracking UDIs may also be an appropriate risk mitigation step that can address some safety and quality concerns. Software whose sole purpose is to 957 958 store and summarize data may not be considered a medical device, but there are still privacy and cyber-security concerns with these products ^{151,152}. Document retention policies are 959 960 important in order to protect HCPs and hospitals from possible legal actions. In situations 961 where the hospital is developing custom institutional ("home-brewed") software, it is important to follow cybersecurity risk management standards and realize that not all insurance 962 policies cover cyber security breaches related to custom developed software. Risk management 963 teams should be in close communication with their insurance brokers to ensure appropriate 964 coverage for that type of activity. 965

Finally, it may be important to develop maturity models for diabetes technology. Maturity models are tools developed in the information technology field to provide guidance to organizations for assessing their current level of development in a particular topic, as well as a roadmap for systemic and structured improvement ¹⁵³. Healthcare IT maturity models have been developed to cover a variety of topics, ranging from continuity of care and healthcare analytics, to telemedicine and mobile technology ¹⁵⁴. Diabetes technology integration would greatly benefit

972	from a maturity model to help guide implementation at healthcare institutions in a systematic
973	way.
974	Recommendations for Data management of Continuous Glucose Monitors and Automated
975	Insulin Dosing Systems in the hospital
976	Clinical Practice
977	Strong Recommendation
978	HCPs should develop a set of core data elements and definitions for CGM data for
979	inclusion in common data models and the her
980	Mild Recommendation
981	Nursing should contact an HCP immediately when CGM results cross critical value
982	thresholds set by the institution
983	Research
984	Strong Recommendations
985	• Researchers need to conduct further studies on the best data management practices of
986	CGMs and AID systems
987	• Researchers need to develop and validate robust glucose telemetry systems for both
988	ICU and non-ICU populations
989	• Researchers need to develop a diabetes technology maturity model that helps
990	institutions understand the requirements to successfully integrate diabetes-related data
991	and technology
992	• Researchers need to develop, disseminate, and validate CGM- and AID system-specific

993 PROs Measures to improve patient care

994	•	Manufacturers need to research methods for quality control for CGMs and AID systems,
995		which is critical as part of inpatient use of CGMs and AID systems
996	•	Manufacturers need to research optimally expanded device labeling in order to
997		overcome clinical inertia and align practice with regulatory policy
998	•	Manufacturers need to research systems for integration of CGM data following initial
999		upload into the cloud (e.g. the Eversense CGM) subsequently into the EHR
1000	•	Manufacturers need to research secure communications systems for protecting data
1001		from wireless wearables, telemedicine systems, and Bring-Your-Own-Device portable
1002		computers used by HCPs (also known as "data in motion")
1003	Mild F	Recommendation
1004	•	Researchers need to develop computerized insulin decision support system that will
1005		integrate with CGMs
1006	Hospi	tal Policies
1007	Stron <u>c</u>	g Recommendations
1008	•	Hospitals need to develop appropriate security protocols, dedicated data storage,
1009		visualization tools, and adequate cyber insurance coverage (also known as"data at rest")
1010	•	Hospitals need to integrate AID system data into the EHR system for nursing and HCPs
1011		to have easy access to this information
1012	•	Hospitals need to determine the number of laboratory or POC BG tests that must be
1013		performed while patients are using CGMs or AID systems in the hospital.
1014	•	Hospitals need to adopt the UDI (Unique Device Identifier) system for healthcare
1015		facilities to track devices in the EHR

1016	•	Hospitals need to identify CGM data reports in the patient's EHR to distinguish them
1017		from laboratory glucose results
1018	•	Hospitals need to present clear criteria to clinicians to identify data that will require
1019		intervention
1020	•	Hospitals need to implement CGM- and AID system-specific PROs to improve patient
1021		care
1022	•	Hospitals need to develop a universal platform for their EHRs that can be used by all
1023		CGMs to present core data elements, summary glucometrics, consistent formats, and
1024		uniform interfaces across all CGM products
1025	•	Hospitals need to arrange for CGM results to be automatically uploaded into the EHR
1026	•	Hospitals need to manage CGM data with the same safety and security measures as all
1027		other PHI
1028	•	Hospitals need to develop policies for CGM and AID system use with atypical scenarios
1029		outside of diabetes, when glucose monitoring is valuable
1030		
1031	Concl	usion
1032	This c	onsensus guideline for subcutaneous CGMs and AID systems was created to provide
1033	recon	nmendations to clinicians, researchers, and hospitals for promoting the safe and effective
1034	use of	f CGMs and AID systems in the hospital environment. Through a consensus process, an
1035	intern	national expert panel voted on 78 recommendations. 77 of the recommendations were
1036	classif	fied as either strong or mild, and 1 failed to reach consensus (Table 9). The panel's
1037	recom	nmendations are intended to support clinical practice, future research, and improved

- 1038 hospital policies, to facilitate the use of these tools. The success of this guideline will be the
- 1039 impact to clinicians, researchers, manufacturers, and hospitals in the management of
- 1040 hospitalized patients with diabetes.

1041 Abbreviations

1042

1043 AMBG, assisted monitoring of blood glucose; BG, blood glucose; BGM, blood glucose monitoring system; CGM, continuous glucose monitor; CI, Confidence Interval; AID, automated 1044 1045 insulin dosing; CLIA, Clinical and Laboratory Improvement Amendment of 1988; COVID-19, 1046 coronavirus disease 2019; CSII, continuous subcutaneous insulin infusion; DKA, diabetic 1047 ketoacidosis; DTS, Diabetes Technology Society; EHR, electronic health record; FDA, United 1048 States Food and Drug Administration; GO, Glucose Oxidase; HCP, healthcare professional; 1049 HIPAA, Health Insurance Portability and Accountability Act of 1996; ICU, intensive care unit; IT, 1050 information technology; LOS, length of stay; MARD, Mean Absolute Relative Difference; MDI, 1051 multiple daily injections; MRI, Magnetic Resonance Imaging; PHI, protected health information; 1052 POC, point of care; PPE, personal protective equipment; PROs, Patient-reported outcomes; RCT, 1053 randomized controlled trial; SAP, sensor-augmented pump; TIR, Time in Range; T1DM, Type 1 1054 Diabetes Mellitus; T2DM, Type 2 Diabetes Mellitus; UDI, Unique Device Identifier; YSI, YSI 2300 1055 STAT Plus Glucose and Lactate Analyzer

1056

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Tables and Figure

- 1555 **Table 1.** The five topics discussed at the Continuous Glucose Monitors and Automated Insulin
- 1556 Dosing Systems in the Hospital Panel

Topic 1: Continuation of home Continuous Glucose Monitors after hospitalization

Topic 2: Initiation of Continuous Glucose Monitors in the hospital

Topic 3: Continuation of Automated Insulin Dosing Systems in the hospital

Topic 4: Logistics and hands-on care of hospitalized patients using Continuous Glucose Monitors and Automated Insulin Dosing Systems

Topic 5: Data management of Continuous Glucose Monitors and Automated Insulin Dosing Systems in the hospital

1558	Table 2. List of Currently Available Subcutaneous CGM Devices and their Interferences

CGM System	Glucose Sensing Methods	Technical Features ⁴	Known Interferences from Chemical Substances
Abbott Diabetes Care FreeStyle Libre 14 day system ²⁸	Glucose Oxidase (GO) + Redox Sensing Membrane	No required calibration; Warm- up 1 hours; 14 days of sensor wear; Range 40- 500 mg/dL; No predictive alerts; Requires scanning at least every 8 hrs	Ascorbic Acid Salicylic Acid
Abbott Diabetes Care FreeStyle Libre 2 ^{29,30}	GO + Redox Sensing Membrane	No required calibration; Warm- up 1 hours; 14	Ascorbic Acid

		days of sensor wear; Range 40- 400 mg/dL; No predictive alerts; Optional alarms for hypoglycemia, hyperglycemia, and signal loss; Requires scanning at least every 8 hrs	
Dexcom G6 ^{31,32}	GO + Perm- selective membrane coating	No required calibrations; Warm-up 2 hours; 10 days of sensor wear; Range 40- 400 mg/dL; Hypoglycemia predictive alerts	Hydroxyurea
Medtronic MiniMed Guardian Sensor 3 ^{34,35}	GO	Requires 2-4 calibrations/d; Warm-up 2 hours; 7 days of sensor wear; Range 40- 400 mg/dL; Predictive alerts	Acetaminophen
Senseonics Eversense ^{36,37}	Non-enzymatic electrochemical fluorescent-based polymer	Required 2 calibrations/d; Implantable; Warm-up 24 hrs; 90-180 days of sensor wear; Predictive alerts for Hypoglycemia and Hyperglycemia; Conditional MRI compatibility	Mannitol Tetracycline

Authors	Population	CGM Type	CGM Manufacturer	Performan ce Measure ment	Comparator
Goldberg, 2004 ⁵⁰	ICU (n: 22)	CGMS	Medtronic MiniMed	Accuracy	Capillary by POC
Vriesend orp, 2005 ⁵¹	OR, SICU, n: 8	CGMS and GlucoDay	Medtronic MiniMed and A. Menarini Diagnostics (A. Menarini Diagnostics Ltd., Florence, Italy)	Accuracy and Feasibility	Arterial by Blood Gas Analyzer
Corstjens, 2006 ⁵²	MICU, n: 45	System Gold	Medtronic MiniMed	Accuracy	Arterial by Blood Gas Analyzer, YSI (YSI 2300 STAT Plus Glucose and Lactate Analyzer, YSI Life Science, Yellow Springs, OH) and POC
De Block, 2006 ⁵³	MICU (n: 50)	Glucoday	A. Menarini Diagnostics	Reliability	Arterial
Price, 2008, ⁵⁴	Mixed ICU, n: 17	Guardian	Medtronic MiniMed	Accuracy	Arterial by Blood Gas Analyzer and POC
Holzinger , 2009 ⁵⁵	MICU (n: 50)	System Gold	Medtronic MiniMed	Accuracy and Reliability	Arterial by Blood Gas Analyzer
Rabiee, 2009 ⁵⁶	SICU/Burn (n: 19)	Dexcom STS	Dexcom	Accuracy and Reliability	Capillary by POC and Serum by Lab
Yamashit a, 2009 ⁵⁷	ICU (n: 50)	STG 22	Nikkiso Co., Ltd., (Nikkiso Co., Ltd., Tokyo, Japan)	Accuracy	Arterial by Blood Gas Analyzer
Logtenbe rg, 2009 ⁵⁸	Cardiac Surgery ICU; (n=30)	Paradigm	Medtronic MiniMed	Accuracy and	Capillary, Arterial, and Venous by POC

				glycemic control	
Holzinger , 2010 ⁵⁹	ICU, mechanical ventilation (n=24)	Guardian	Medtronic MiniMed	Glycemic control (% time at glucose < 110mg/dL), LOS, mortality	Arterial by Blood Gas Analyzer and blinded Medtronic MiniMed System Gold CGM
Jacobs, 2010 ⁶⁰	ICU (n: 29)	Guardian RT	Medtronic MiniMed	Accuracy and Feasibility	Capillary by POC
Brunner, 2011, ⁶¹	MICU, n; 174	Guardian & System Gold	Medtronic MiniMed	Accuracy and Reliability	Arterial by Blood Gas Analyzer
Lorencio, 2012 ⁶²	ICU (n: 41)	Guardian	Medtronic MiniMed	Accuracy	Arterial by Blood Gas Analyzer
Kalmovic h, 2012 ⁶³	Peri- Operative Cardiac Surgery, n: 32	System Gold Blinded	Medtronic MiniMed	Accuracy and Feasibility	Venous by Blood Gas Analyzer
Kopecký, 2013 ⁶⁴	Cardiac ICU; (n=24)	Guardian RT	Medtronic MiniMed	Accuracy and Glycemic control	Arterial by Blood Gas Analyzer and Computer (enhanced Model Predictive Control) algorithm alone
Leelarath na, 2013 ⁶⁵	Neurosurgica I ICU (n: 24)	FreeStyle Navigator	Abbott Diabetes Care	Glycemic control	Arterial by Blood Gas Analyzer
Rodríguez - Quintanill a, 2013 ⁶⁶	CCU (n: 16)	Guardian RT	Medtronic MiniMed	Time to normoglyc emia	Venous and Capillary by POC
Schuster, 2014 ⁶⁷	SICU (n: 24)	Guardian	Medtronic MiniMed	Accuracy	Capillary by POC

Boom, 2014 ⁶⁸	MICU/SICU (n: 156)	FreeStyle Navigator	Abbott Diabetes Care	Accuracy and Glycemic Control	Arterial by Blood Gas Analyzer, and POC
Kosiboro d, 2014 ¹⁷	Cardiac ICU (n: 21)	Sentrino	Medtronic MiniMed	Accuracy and Reliability	Central Venous by POC or Lab
Umbrello, 2014 ⁶⁹	MICU (n=6)	OptiScanne r 5000	OptiScan Biomedical	Glycemic control	Central Venous by Blood Gas Analyzer or Lab (reported elsewhere)
Van Hooijdon k, 2015 ⁷⁰	ICU (n: 50)	Sentrino	Medtronic MiniMed	Accuracy and Reliability	Arterial by Blood Gas Analyzer
Sechterb erger, 2015 ⁷¹	Cardiac ICU, n: 8	FreeStyle Navigator	Abbott Diabetes Care	Accuracy	Arterial by Blood Gas Analyzer
Punke, 2015 ⁷²	SICU, n: 14	Sentrino	Medtronic MiniMed	Accuracy	Arterial by Blood Gas Analyzer
De Block, 2015 ⁷³	MICU (n=35)	GlucoDay S	A. Menarini Diagnostics	Accuracy and glycemic control	Arterial by Blood Gas Analyzer and Blinded Microdialysis- Based CGM
Ballestero s, 2015 ⁷⁴	MICU, n: 18	Soft Sensor	Medtronic MiniMed	Accuracy	Capillary by POC
Nohra, 2016 ⁷⁵	SICU, n: 23	Optiscanne r 5000	Optiscan Biomedical	Accuracy	Central Venous by YSI
Wollersh eim, 2016 ⁷⁶	MICU, n: 20	Sentrino	Medtronic MiniMed	Accuracy and Feasibility	Arterial, Central Venous, or Venous by Blood Gas Analyzer
Gottschal k, 2016 ⁷⁷	Extracorpora l Cardiac Life Support, n: 25	Sentrino	Medtronic MiniMed	Accuracy	Arterial by Blood Gas Analyzer

Righy Shinotsuk a, 2016 ⁷⁸	ICU (n: 88)	OptiScanne r 5000	Optiscan Biomedical	Accuracy	Arterial by YSI
Schierenb eck, 2017 ⁷⁹	Cardiac ICU, n: 26	Freestyle Libre Subcutane ous-CGM vs Eirus Intravascul ar	Abbott Diabetes Care and Maquet Getinge Group	Accuracy	Arterial by Blood Gas Analyzer and Capillary by POC
Song, 2017 ⁸⁰	OR, ICU, n: 22	Guardian	Medtronic MiniMed	Accuracy and Reliability	Arterial by Blood Gas Analyzer
Rijkenber g, 2017 ⁸¹	Mixed ICU, n: 155	FreeStyle Navigator	Abbott Diabetes Care	Accuracy and Reliability	Arterial by Blood Gas Analyzer
Ancona, 2017 ⁴⁶	ICU, n: 8	FreeStyle Libre CGM	Abbott Diabetes Care	Accuracy and Feasibility	Arterial by Blood Gas Analyzer or Capillary by POC
Bochicchi o, 2017 ⁸²	ICU (n: 243)	OptiScanne r 5000	OptiScan Biomedical	Accuracy	Arterial, Central Venous, or Venous by YSI
Nukui, 2019 ⁸³	Acute Stroke, n: 39	FreeStyle Pro CGM	Abbott Diabetes Care	Accuracy and Efficacy	Capillary by POC

1562 **Table 4.** CGM Studies in the ICU in Pediatric Populations

Author, Year	Population	Type of CGM	CGM Manufactu rer	Performance Measurement	Comparator
Bridges, 2010 ⁸⁴	ICU, n: 47	Guardian	Medtronic MiniMed	Accuracy	Arterial, Venous, and Capillary by iSTAT POC and Lab

Steil, 2011 ⁸⁵	Cardiac ICU, n: 311	Guardian	Medtronic MiniMed	Accuracy and hypoglycemia prevention	Arterial by POC and Lab
Prabhudesai, 2015 ⁸⁶	ICU, n: 19	Guardian	Medtronic MiniMed	Accuracy	Arterial by Lab
Kotzapanagiotou, 2019 ⁸⁷	ICU, n: 16	FreeStyle Libre	Abbott Diabetes Care	Accuracy	Arterial by Blood Gas Analyzer Capillary by POC, Biochemical Serum by Lab
Sopfe, 2020 ⁸⁸	Stem Cell Transplant ation n: 29	FreeStyle Libre Pro	Abbott Diabetes Care	Accuracy	Central Venous by Lab

1564 **Table 5.** CGM Studies in the Non-ICU in Adult Populations

	Patient Population	CGM Type	CGM Manufactu rer	Performance Measurement	Comparator
Dungan, 2012 ⁹⁸	T1DM and T2DM (n: 58), on Intravenous (IV) or subcutaneous insulin	iPro syste m	Medtronic MiniMed	Accuracy	Capillary by POC
Burt, 2013 ⁹⁹	T1DM and T2DM, on basal bolus insulin (n:26)	Syste m Gold	Medtronic MiniMed	Accuracy and glycemic control	Capillary by POC
Schaupp, 2015 ¹⁰⁰	T2DM, on basal bolus insulin (n:84)	iPro2 syste m	Medtronic MiniMed	Accuracy	Capillary by POC

Gómez, 2015 ⁹³	T2DM, on basal bolus insulin (n=38)	iPro2 syste m	Medtronic MiniMed	Glycemic control and Hypoglycemia detection	Capillary by POC
Spanakis, 2018 ¹⁰¹	T2DM, on insulin therapy (n:5)	Dexco m G4 CGM with Share2 applic ation	Dexcom	Glucose telemetry system feasibility	None
Singh, 2019 ¹⁰²	T2DM, on basal-bolus insulin (n: 13)	Dexco m G4 Platinu m CGM	Dexcom	Feasibility and Prevention of hypoglycemia	Blinded CGM
Nair, 2020 ¹⁰³	Surgical Ward (n: 10)	Dexco m G6 Blinde d	Dexcom	Accuracy	Capillary by POC
Shehav- Zaltman, 2020 ¹⁰⁴	T1DM on CSII (n: 1) and T2DM on basal bolus (n: 3), COVID-19 Wards (n: 5)	Guardi an	Medtronic MiniMed	Feasibility	None
Galindo, 2020 ⁹⁶	T2DM, on basal-bolus insulin (n: 97)	FreeSt yle Libre Pro CGM	Abbott Diabetes Care	Accuracy and Hypoglycemia detection	Capillary by POC
Singh, 2020 ⁴³	T2DM, on basal-bolus insulin (n: 72)	Dexco m G6	Dexcom	Prevention of hypoglycemia	Capillary by POC

Ushigome , 2020 ¹⁰⁵	Diabetes (unknown type) with COVID-19 (n: 1)	Dexco m G4 Platinu m	Dexcom	Safety and Effectiveness	Lab
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1566 **Table 6.** Ongoing hospital CGM studies listed on ClinicalTrials.gov

Dexcom intervention trial (NCT03877068)
CGM in Hospitalized Veterans/ Glucose Telemetry System (NCT03508934)
Scripps Digital Diabetes (NCT04269655)
Green Line From Hospital to Territory (GreenLightHT) (NCT03764709)
Use of Wearables for Early Detection of Complications After Major Acute Abdominal Surgery
(NCT04257344)
DRIVE—Perioperative Period (DRIVE-Periop) (NCT04033705)
Flash Glucose Measurement in Patients on Total Parenteral Nutrition (NCT03871660)
Early Glargine (Lantus) in Diabetic Ketoacidosis Management in Children With Type 1
Diabetes (NCT03107208)
Reducing Emergency Department Visits and Improving Glucose Control in Uncontrolled Type
2 Diabetes Using CGM Sensors at Hospital Discharge (NCT04277780)
CGM in hospitalized patients with diabetes (NCT04230694)
Remote Continues Glucose Monitoring During the COVID-19 Pandemic in Quarantined
Hospitalized Patients (CGM-ISO) (NCT 04430608)

The Use of a Continuous Glucose Monitoring System (Dexcom G6) in Hospitalized Patients

for Acute Care (NCT04385862)

Wireless Assessment of Respiratory and Circulatory Distress - Continuous Glucose

Monitoring (WARD-CGM) (NCT04473001)

Reliability of the Freestyle Libre CGM in the Inpatient Setting During the COVID-19 Surge

(NCT04417270)

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7 Table is up-to-date as of August 8, 2020

1568 **Table 7**. Contraindications to CSII system and AID system therapy in the hospital

Impaired level of consciousness (except during short-term anesthesia)
Patient's inability to correctly demonstrate appropriate CSII system settings
Critical illness requiring intensive care
Psychiatric illness that interferes with a patient's ability to self-manage diabetes
Diabetic ketoacidosis and hyperosmolar hyperglycemic state
Refusal or unwillingness to participate in self-care
Lack of CSII system supplies
Lack of trained health care providers, diabetes educators, or diabetes specialists
Patient at risk for suicide
Health care decision

- 1570 Table has been reproduced with permission from Umpierrez and Klonoff, Diabetes Care,
- 1571 2018¹²³. "Insulin pump therapy" in the title of the table has been changed to "CSII system and
- 1572 AID system therapy". "Pump" in the second and seventh bullets has been changed to "CSII
- 1573 system".

4 4	Table 0. COM data tata anti- a sa da ta sa sa da ta
1574	Table 8. CGM data integration complexity across three key domains

Data Extraction (from least to most complex)	Data Storage (from least to most complex)	Data Display (from least to most complex)
 Static, standard reports Custom reports Structured summary data Structured continuous data Device metadata 	 Web storage, linked to EHR Native EHR data tables External storage and computing environment 	 Text and graphic reports Structured data fields with native analytics Embedded analytics displayed from a web service Native integration of manufacturer analytics platform

Table 9. 78 proposed recommendations for the guideline voted on by the panel

1577

Continuation of home Continuous Glucose Monitors after hospitalization

Clinical Practice: Strong Recommendations

- HCPs **should** consult with an inpatient diabetes team if available, when continuing or initiating a CGM or AID system.
- HCPs should avoid relying on CGM data for glycemic management decisions in patients with severe hypoglycemia or hyperglycemia (i.e. BG < 40 mg/dL or >500 mg/dL).
- HCPs **should** avoid using CGMs for management of 1) diabetic ketoacidosis until glucose is in the CGM measurement range, and then CGMs should be used adjunctively or 2) situations with rapidly changing glucose levels and fluid/electrolyte shifts.
- HCPs **should** avoid continuing or initiating CGMs to patients with skin infections near the sensor site or placing sensors in areas with significant edema as well as patients treated with vasoactive agents or poor tissue perfusion.
- HCPs **should** use a CGM checklist for elective procedures during the pre-operative visits to ensure proper documentation of devices and real time data reporting.
- HCPs **should** advise pregnant women to continue the use of a CGM during a hospitalization to identify glucose trends and prevent hypo- or hyperglycemia.
- HCPs **should** instruct patients to bring supplies with them to the hospital for the duration of any pre-planned admission or elective procedures.
- HCPs **should** check capillary BG or serum BG concentrations after procedures for noncritically ill patients and venous/arterial blood for critically ill patients to ensure the patient's CGM is functioning properly.

•	HCPs should use trend arrows and rate of change to help prevent extreme glycemic					
		unctively) to help determine when a BG test is				
	•	required.				
•	HCPs should set alarm thresholds for inpatient glycemic targets, such as predicting					
	hypoglycemia (typically BG < 80-85 mg/c					
•	Nursing should document CGM and/or (
-	health record (EHR) for all admissions or					
Resear	rch: Strong Recommendations					
•		to support definitive recommendations on				
	improved outcomes for continuation of					
	hospitalization.					
•	•	the roles of CGM and POC BG testing and				
		ry to inform nursing staff about actionable				
	CGM patterns.	,				
•	Researchers need to perform further stu	idies to assess the accuracy of CGMs during				
	pregnancy, labor & delivery, and the per					
•	Researchers need to study the impact of					
	situations with rapid changes in the gluc					
Hospit	al Policies: Strong Recommendations	· · ·				
•	Hospitals need to develop standard CGN	1 data reports and workflows.				
•	Hospitals need to implement policies for	r testing capillary BGs and calibrating CGMs if				
	the CGM requires calibration.					
•	Hospitals need to develop a system for a	automatic staff notification for CGM alarms				
	that predict impending or current hypog	lycemia or hyperglycemia.				
•	Hospitals need to develop specific guide	lines for using CGMs and AID systems for				
	their affiliated nursing homes and skilled	nursing facilities.				
Initia	tion of Continuous Glucose Monito	rs after hospitalization				
		-				
	I Practice: Strong Recommendation	Clinical Practice: Mild Recommendation				
•	HCPs should consider prescribing	HCPs should avoid initiating CGMs in				
	CGMs to reduce the need for frequent	patients with severe hypoglycemia or				
	nurse contact for POC glucose testing hyperglycemia (i.e. BG < 40 mg/dL or					
	and the use of PPE for patients on >500 mg/dL) or during periods of					
	isolation with highly contagious rapid glucose fluctuations.					
_	infectious diseases (e.g. COVID-19).					
Resear	rch: Strong Recommendations					
•		oport initiation of CGMs for improving patient-				
	centered outcomes.					
٠	-	spital outcomes when initiating CGMs in the				
	hospital, including improved glycemic ou					
	hypoglycemia and hyperglycemia, reduction of ICU LOS, and cost-effectiveness.					

- Researchers need to conduct studies on long term benefits for initiating CGMs in the hospital after discharging patients with newly diagnosed diabetes or recurrence of diabetic ketoacidosis (DKA) or other complications of diabetes. Manufacturers **need** to develop educational tools for patients, hospital staff, and HCPs. Hospital Policies: Strong Recommendations Hospitals **need** to develop plans, including process maps, protocols, staff educational resources, and order sets for prescribing CGM use during hospitalizations prior to implementing a CGM. Hospitals need to provide educational tools for patients, nurses, house staff, and attending physicians when a patient in the hospital starts on a CGM. Continuation of Automated Insulin Dosing Systems in the hospital Clinical Practice: Strong Recommendations Clinical Practice: Mild Recommendation HCPs **should** prescribe AID systems HCPs **should** avoid initiating an AID • ٠ only for appropriate candidates, who system during a hospitalization. will need to have adequate knowledge and skills for using AID systems. HCPs should reassess a decision periodically to transition use of outpatient AID systems into the hospital in order to ensure that AID system continue to represent the best treatment option for each patient. HCPs **should** prepare an alternative • plan for diabetes management in case it becomes inappropriate for a patient to continue using an AID system in the hospital. HCPs **should** discontinue AID systems in critically ill hospitalized patients (such as those with hypovolemia or sepsis). HCPs should recognize glycemic patterns due to CGM compression, which can cause false low readings. **Research: Strong Recommendations** Researchers **need** to conduct studies about whether continuing AID systems in the • hospital is beneficial to improve glycemic management or clinical outcomes.
 - Researchers need to provide data on hospital outcomes when using AID systems in the hospital, including improved glycemic outcomes, detection and/or reduction of hypoglycemia, reduction of ICU LOS, and cost-effectiveness.

• Manufacturers **need** to research whether all types of CGMs and AID systems can be used during radiological/imaging studies or diathermy.

Hospital Policies: Strong Recommendations

- Hospitals need to develop institution-specific protocols and order sets for the proper use of AID systems during a hospitalization.
- Hospitals **need** to require that patients using AID systems bring with them sufficient supplies for these devices during a hospitalization.
- Hospitals need to develop protocols for using AID systems during elective procedures and surgeries.

Recommendation Not Reaching Consensus

• HCPs **should** switch AID systems from "auto" mode to "manual" mode when a patient is admitted to the hospital wearing an AID system.

Logistics and hands-on care of hospitalized patients using Continuous Glucose Monitors and Automated Insulin Dosing Systems

Clinical Practice: Strong Recommendations

- HCPs **should** inquire about and document the medication and supplement history of patients who use CGMs to determine whether there are any agents that can interfere with glucose measurements.
- HCPs **should** ensure that off-label use of CGMs and AID systems is consistent with medical practice and appropriate precautions have been taken to protect patients.
- Nursing **should** document hands-on training of CGM use and AID system therapy through a technology certification program.
- Nursing **should** confirm that the patient is appropriate to continue using a CGM or an AID system and also review the agreement and hospital policy with the patient.
- Nursing **should** inspect the insertion site every shift with attention to skin integrity and signs of erythema or infection, and should document site changes.
- Nursing **should** know device basics, institutional policies, HCPs roles, and whom to contact if questions arise.
- Nursing **should** administer a patient competency assessment or survey to assess patient ability to safely assist with managing a CGM or an AID system.
- Nursing **should** set expectations and clarify that there will be a need to continue checking POC capillary glucose even when using a CGM.
- Nursing should measure POC BG concentrations to confirm or supplement CGM readings (usually a minimum of 4 times daily: before each of three meals and at bedtime if patients are eating, or every 6 hours if patients are fasting) as well as at patient request; however, the CGM glucose, trend arrows, and rate of change may be used to help determine if and when a BG test is required.

Research: Strong Recommendations

- Researchers **need** to conduct further studies on the best logistics and hands on care for patients using CGMs and AID systems to achieve the best outcomes.
- Manufacturers **need** to research interoperable components for AID systems that are compatible with hospital EHRs.

Hospit	al Policies: Strong Recommendations					
•	Hospitals need to provide interpreter se	rvices to translate CGM and AID system				
	agreements.					
•	Hospitals need to state in their policy and patient agreement documents that					
	treatment decisions will be based on hospital-calibrated BGM readings (or laboratory					
		ring a need to isolate a patient with a severe				
	and highly contagious infection.	This a need to isolate a patient with a severe				
•		nd AID system policy and patient agreement				
	-	c files stored in the EHR order set for CGMs				
	and AID systems.					
•	-	hen to discontinue or temporarily suspend				
•	the use of CGMs and AID systems.	nen to discontinue or temporarily suspend				
	-	ring and nationts to improve outcomes and				
•		sing, and patients to improve outcomes and				
_	satisfaction.					
		se Monitors and Automated Insulin				
	ng Systems in the hospital					
Clinica	l Practice: Strong Recommendation	Clinical Practice: Mild Recommendation				
•	HCPs should develop a set of core	 Nursing should contact an HCP 				
	data elements and definitions for CGM	immediately when CGM results cross				
	data for inclusion in common data	critical value thresholds set by the				
	models and the EHR.	institution.				
Resea	rch: Strong Recommendations	Research: Mild Recommendation				
•	Researchers need to conduct further	 Researchers need to develop 				
	studies on the best data management	computerized insulin decision				
	practices of CGMs and AID systems.	support system that will integrate				
•	Researchers need to develop and	with CGMs.				
	validate robust glucose telemetry					
	systems for both ICU and non-ICU					
	populations.					
•	Researchers need to develop a					
	diabetes technology maturity model					
	that helps institutions understand the					
	requirements to successfully integrate					
	diabetes-related data and technology.					
•	Researchers need to develop,					
	disseminate, and validate CGM- and					
	AID system-specific PROs Measures to					
	improve patient care.					
•	Manufacturers need to research					
	methods for quality control for CGMs					
	and AID systems, which is critical as					
	part of inpatient use of CGMs and AID					
	systems.					
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•	Manufacturers need to research	
	optimally expanded device labeling in	
	order to overcome clinical inertia and	
	align practice with regulatory policy.	
•	Manufacturers need to research	
	systems for integration of CGM data	
	following initial upload into the cloud	
	(e.g. the Eversense CGM)	
	subsequently into the EHR.	
•	Manufacturers need to research	
	secure communications systems for	
	protecting data from wireless	
	wearables, telemedicine systems, and	
	Bring-Your-Own-Device portable	
	computers used by HCPs (also known	
	as "data in motion").	
Hospit	tal Policies: Strong Recommendations	
•	Hospitals need to develop appropriate se	curity protocols, dedicated data storage,
	visualization tools, and adequate cyber in	nsurance coverage (also known as"data at
	rest").	
•	Hospitals need to integrate AID system da	ata into the EHR system for nursing and HCPs
	to have easy access to this information.	
•	Hospitals need to determine the number	of laboratory or POC BG tests that must be
	performed while patients are using CGMs	s or AID systems in the hospital.
•	Hospitals need to adopt the UDI (Unique	Device Identifier) system for healthcare
	facilities to track devices in the EHR.	
•	Hospitals need to identify CGM data repo	orts in the patient's EHR to distinguish them
	from laboratory glucose results.	
•	Hospitals need to present clear criteria to	o clinicians to identify data that will require
	intervention.	
•	Hospitals need to implement CGM- and A	AID system-specific PROs to improve patient
	care.	
•	Hospitals need to develop a universal pla	tform for their EHRs that can be used by all
	CGMs to present core data elements, sun	nmary glucometrics, consistent formats, and
	uniform interfaces across all CGM produc	cts.
•	Hospitals need to arrange for CGM result	s to be automatically uploaded into the EHR.
•	Hospitals need to manage CGM data with	n the same safety and security measures as
	all other PHI.	
٠	Hospitals need to develop policies for CG	M and AID system use with atypical
	scenarios outside of diabetes, when gluce	aso monitoring is valuablo

 Hospitals need to develop policies for CGW and AID system use with scenarios outside of diabetes, when glucose monitoring is valuable.

1580	Figure	 Continuous Glucose Monitors or Automated Insulin Dosing System Patient Agreement 				
1581	I	currently have a continuous glucose monitor and/or				
1582	insulir	insulin pump in place and wish to maintain this therapy during my admission to the				
1583	Hospit	tal. I understand and agree as follows:				
1584						
1585	Patient	's Continuous Glucose Monitor				
1586	1.	I may continue to wear my continuous glucose monitor (CGM) during my hospital				
1587		stay but my blood glucose will also be monitored using a hospital-approved blood				
1588		glucose meter and treatment decisions will be based on these results.				
1589	2.	I will keep a back-up supply of all CGM supplies including, without limitation,				
1590		sensors and dressings.				
1591	3.	I will change the CGM sensor every 7-14 days depending on the device instructions.				
1592	4.	I will notify my nurse immediately if my CGM indicates my glucose reading is trending				
1593		out of target (i.e., trending low or high) so that my blood glucose can be tested to				
1594		confirm the trending and appropriate treatment initiated according to the				
1595		prescriber's order.				
1596	5.	I will allow my nurse to assess the sensor site every shift.				
1597	6.	If I need any surgery or procedure, then the hospital might need to remove my				
1598		sensor. If I elect to leave my CGM sensor on during any surgery or procedure it may				
1599		present a risk of damage to my CGM sensor during the surgery or procedure.				
1600	7.	If I need an MRI scan, then I will remove the sensor prior to the procedure so that				
1601		the transmitter and receiver can be either secured by staff or sent home with a				
1602		designated family member/significant other.				
1603		If I need an X-ray or CT scan, then my CGM will be covered by a lead apron.				
1604	9.	Any of my CGM supplies stored by hospital staff will be returned to me prior to my				
1605		discharge.				
1606						
1607	Patient	's Automated Insulin Dosing System				
1608	1.	I can manage my own automated insulin dosing system (insulin pump and				
1609		continuous glucose monitors) and the medical condition for which the				
1610		automated insulin dosing system is prescribed.				
1611		a. If my physical or mental condition changes, my caregivers at the hospital may				
1612		re- assess my capability to manage my own pump. If it is determined that I can				
1613		no longer safely manage my pump, the hospital will remove the pump and				
1614		administer insulin by injection or IV as determined by my provider.				
1615		b. Hospital personnel will not operate my pump, except in the above-				
1616		described situation.				
1617	2.	Only family members/significant others who usually assist me with the operation of				
1618		my pump will do so during my hospital stay. I will keep a back-up supply of all insulin				
1619		pump supplies including, without limitation, insertion sets, infusion tubing and				
1620		dressings.				
1621	3.	My insulin will be kept in my personal medication bin and my nurse will get it for				

1622		me when needed for an insulin infusion set change.	
1623	4.	I will change my insulin infusion set every 48-72 hours (2-3 days) or earlier as needed.	
1624	5.	If I change my insulin pump settings, I will immediately communicate that with	
1625		my health care team.	
1626	6.	I will only make changes to the basal rate, unless in auto mode, after discussion with	
1627		my provider. I will notify my nurse immediately if I have any problem with my insulin	
1628		pump.	
1629	7.	If I need any surgery, procedure, radiation therapy, or diagnostic imaging (e.g. MRI or	
1630		x- rays), the hospital may need to disconnect my insulin pump and an alternative	
1631		insulin regimen will need to be prescribed.	
1632	8.	If I need diagnostic imaging (e.g. MRI or x-rays), I may need to remove the pump	
1633		prior to the procedure, and it will be secured by staff outside of the imaging area.	
1634	9.	Regarding the CGM part of my automated insulin dosing system, if I need an MRI	
1635		scan, then I will remove the sensor prior to the procedure so that the transmitter	
1636		and receiver can be either secured by staff or sent home with a designated family	
1637		member/significant other. If I need an X-ray or CT scan, then my CGM will be	
1638		covered by a lead apron.	
1639	10	. The hospital staff will monitor my blood glucose with a hospital-approved blood	
1640		glucose meter.	
1641	11	. I will report all bolus doses of insulin to my nurse for documentation purposes.	
1642	12	. I will allow my nurse to assess the insertion site every shift.	
1643	13	. If my blood glucose values are erratic and cannot be controlled, my insulin pump may	
1644		be discontinued, and an alternative insulin regimen will be provided for me.	
1645	14	. Prior to being discharged from the hospital, I will confirm with my nurse that the	
1646		pump is working correctly and that there are no problems with medication delivery or	
1647		the delivery site on my body. In the event that there are problems, they will be	
1648		corrected prior to my discharge from the hospital.	
1649	15	. Any of my unused insulin and pump supplies that I have brought with me to the	
1650		hospital will be returned to me prior to my discharge.	
1651	16	. My physicians and other health care providers may terminate my use of the insulin	
1652		pump if they observe any contraindication to its use or for any reason that they	
1653		believe medically necessary.	
1654			
1655	By signing below, I acknowledge that I have read, understood, and agreed to the above and		
1656	that al	l of my questions have been answered.	
1657			
1658	Patient	Signature:	
1659	Nurse/Provider Signature:		
1660	Nurse/Provider Print Name:		
1661	Unit/Se	rvice:	