

## Abbott Diabetes Care Investigator Initiated Study Proposal Form

Study Title	[study title]				
Investigator(s)	[Investigator(s) responsible for the research]				
Institution(s)	[Institution(s) involved with the research. Ensure to indicate planned number of sites and countries. and indicate how many site(s) and countries where data will be collected in this study.]				
Contact information	[contact details: physical and e-mail address]	1			
Plain language abstract	[Describe the overall research aims and proje				
Study Timeline	[Please provide approximate anticipated time Milestones Ethical approval Recruitment start Data collection complete Publication submission	eline for this study] Date MMM/YYYY MMM/YYYY MMM/YYYY MMM/YYYY			
Objectives	[Specific objectives or hypotheses.]				
Trial design	[Description of trial design including type of trial (eg, parallel group, crossover, factorial, single group), allocation ratio, and framework (eg, superiority, equivalence, noninferiority, exploratory).]				
Eligibility criteria	[Inclusion and exclusion criteria for participants. If applicable, eligibility criteria for study centres and individuals who will perform the interventions. For those with diabetes, please include details around diagnosis and medication.]				
Interventions	[Interventions for each group with sufficient detail to allow replication, including how and when they will be administered. Please include clear description regarding how CGM will be used in this study]				
Outcomes	<ul> <li>[Primary outcome, secondary outcomes, and other exploratory outcomes, including the specific measurement variable, analysis metric (e.g., change from baseline, final value, time to event), method of aggregation (e.g., median, proportion), and time point for each outcome.]</li> <li>For example: <ul> <li>Primary outcome: Change of TIR (baseline vs. 6 months)</li> <li>Secondary outcomes: Change of HbA1c, TAR, TBR, body weight</li> <li>Exploratory outcomes: questionnaires (qualify of life, diabetes distress scale, etc.)</li> </ul> </li> </ul>				
Participant timeline	[Time schedule of participant activities. A sch For example: • Visit 1 (-21 days) -> enrolment, conse				

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		drawing, physical exam, blinded CGM inst	tallation for baseline a	ssessment,	
		and questionnaires)			
	• Visi	it 2 (-7 days) -> Download blinded CGM			
	• Vis	it 3 (0 day) -> randomization, start unblind	led CGM, education pe	r study	
		group			
	• Visi	it 4 (90 days) -> blood draw, physical exam	, and questionnaires		
	• Visi	it 5 (180 days) -> blood draw, physical exar	m, and questionnaires,	), blinded	
		CGM installation			
	• Visi	it 6 (194 days) -> Download blinded CGM			
		1 × 2			
	Visit 1	Visit 2	Visit 5	Visit 6	
	(-21 days)	(-7 Visit 3 Visit 4 days) (o days) (90 days)	(180 days)	(194 days)	
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Sample size	[Estimated number of participants needed to achieve study objectives and how it was				
	determined, including clinical and statistical assumptions supporting any sample size				
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