# **APPENDIX A. DATA COLLECTION FORMS**

## VA-EPC Self-Monitoring of Blood Glucose Article Screener

Article ID	<b>Reviewers:</b>	Assigned on:
Citation:		
First Author:		
*Complete Q7 & Q8 on ALL forms*		observational study, what is
Is the study a test of efficacy or effectiveness of SMBG alone or as part of a multi-component intervention?  (Check all that apply)  Alone	the duration of to (Circle one) < 12 weeks/not an RCT observational study 12 weeks or greater If >=12 wks, wr	/CCT or 0 ( <b>STOP</b> )
Multi-component		<u>Units</u>
2. Study design (Circle one)	<b>Duration</b>	Units 01. Days 04. Years 02. Weeks 05. NR 03. Months
RCT/CCT1		eets no other criterion,
Review article: systematic or M-A2 Observational Study (cohort,	(Circle one)	ed for background?
case control, etc)	Yes	0
other syst review	8. Are any of the si Veterans?	ubjects identified as
2a. Is this a crossover study?	(Circle one) Yes	1
(Circle one)	No	
Yes	Notes	
3. Is A1c reported as an outcome? (Circle one) Yes		
4. Is hypoglycemia reported as an outcome? (Circle one) Yes		
5. Is the frequency of SMBG testing reported? (Circle one) Yes		

# VA Self-Monitoring of Blood Glucose Project-Detailed Review Form

Article ID: Reviewer:		FINAL 12	-14-06
First Author:			
Study Number: of Description: (if me	ore than one <b>study</b> )		
Do you think that this article might include the state of the sta	ne same data as another	6. If reported, was the method of double blinding	
study?		appropriate?	(CIRCLE ONE)
<b>V</b>	(CIRCLE ONE)	Yes	1
Yes No.		No	2
If YES enter IDs:	2	Double blinding method not described  Not applicable	8
ID(s):		Not applicable	
2. Design:	(CIRCLE ONE)	7. If study was randomized, did the method of random	nization provide
RCT	( )	for concealment of allocation?	(CIRCLE ONE)
CCT		Yes	1
Other design		No	2
•	(5161)	Concealment not described	8
<i>2</i>	(CIRCLE ONE)	Not applicable (not randomized)	
Yes	1		
No	2	Q. And with drawals (IV) and draw outs (D) described?	
4. If the study was randomized, was method of	randomization	8. Are withdrawals (W) and dropouts (D) described? Yes, reason described for <b>all</b> W and D	
appropriate?	(CIRCLE ONE)	Yes, reason described for <b>some</b> W and D	
Yes		1	
No		Not described	
Method not described		Not applicable	9
Not applicable (not randomized)		, , , , ,	(CIRCLE ONE)
5. Is the study described (with respect to SMBC)	S)as: (circle one)	Yes No	
Double blind		1	
Single blind, patient		10 Commission (Transport	
Single blind, outcome assessment		10. Sample size: (Enter 999 for not reported)	
Single blind, not described		Γ., 11 - 1.	
Open		Enrolled: <sub>5</sub>	
Blinding not described		8 / 1 1	
Not applicable		Followed-up/analyzed:	

# VA Self-Monitoring of Blood Glucose Project-Detailed Review Form

11. What were the characteristics of the patient population?	If yes, please enter the following: Weight	Units	Units
A. Demographics: % women =	Mean weight		1. kilograms
(CHECK ALL THAT APPLY)	Median weight		2. pounds 3. NA
Caucasian	Weight Rangeto		4. ND
African Ancestry   Hispanic	weight Rangeto		999. NR
Other (Specify:) □	15. Was duration of diabetes reported?	40	
Demographics not reported□	Yes	(CIRCLE ONE)	)
25 of 12 of 15	No		
12. What was reported for the following questions regarding	If yes, please enter the following: Time	Units	
subjects' ages? (Enter number 999 for not reported)	Mean time		<u>Units</u>
Mean Age	Median time	2	. Hour 5. Year 8. ND
Median Age	Time Rangeto	3	. Week 9. NA . Month 999.NR
Age Range to	Time Rangeto		
	16. Which of the following co-morbidities were rep	orted on:	
13. Was BMI reported?	Myocardial infarction	(CHECK ALL THAT	APPLY)
Yes	CongestivelHeart Failure		
No	Peripheral \(\frac{2}{2}\) ascular disease		
	Cerobrovascular disease		
If yes, please enter the following: (Enter number 999 for not	Dementia		
reported)	Chronic pulmonary disease		
Mean BMI	Rheumatologic disease		
Median BMI	Peptic ulcer disease		
BMI Range to	Mild liver disease		
	Hemiplegia or paraplegia		
	Renal disease		
14. Was weight reported?	Malignancy, leukemia, lymphoma		
Yes	Moderate-severe liver disease		
No	AIDS1	<b>ப</b>	

### VA Self-Monitoring of Blood Glucose Project-Detailed Review Form

Enter sample size and intervention/exposure data for each arm beginning with CONTROL/USUAL CARE for arm 1, then in order of first mention. For observational studies answer only columns denoted with asterisks (\*):

Arm/ Group	Sample size *	Components *  (check all that apply)	Total # of Visits	Frequency of SMBG	Number of Days per week	Duration of *	Units *	Co-therapies(s)
1	P PY CNTRL NENTERING  CASES NCOMPLETING	SMBG		Control				
2	P PY CNTRL NENTERING  CASES N COMPLETING	SMBG		GD BID TID QID QID PP Other Before/After meals NR				
3	P PY CNTRL NENTERING  CASES NCOMPLETING	SMBG		GD BID TID QID QID PP Other Before/After meals NR				
4	P PY CNTRL NENTERING  CASES	SMBG Exercise		GD BID TID QID QID PP Other Before/After meals NR				
	Enter a number for N entering and N completing or enter 9999 if not reported. If observational study, circle appropriate unit of measurement: P Persons PY People years CNTRL Control CASES Cases		Enter # of visits or contact s		Enter a number 997. Variable 998. ND 999. NA	Enter a number 997. Variable 998. ND 999. NA	Enter a number 1.Hour 2.Day 3.Week 4.Month 5.Year 8.ND, 9. NA	

## VA Male OP Project-Detailed Review Form- Diagnostic Studies

### **Outcomes**

17. Please check the type of outcomes measured. For case control e	enter	the
outcome that defines the study:		

	(CHECK ALL THAT APPLY)
HbA1c	
Fasting glucose	
Fructose	
BMI/Weight loss	
F ( 1.1	
Fast v. meal glucose	
Health related quality of life	🗖

### **Evaluation**

18. When, relative to the start of the intervention, were outcomes reported?

(Enter the number/code in the appropriate box)

	Control		Interve	ention
	Numbe	Units	Numb	Units
	r		er	
1 <sup>st</sup> follow-				
up				
up 2 <sup>nd</sup> follow-				
up				
up 3 <sup>rd</sup> follow-				
up				
up 4 <sup>th</sup> follow-				
up				
up 5 <sup>th</sup> follow-				
up				
up 6 <sup>th</sup> follow-				
up				
Additional				
follow-ups				

	<u>Units</u>
1. Hour	5. Year
2. Day	8. ND
3. Week	9. NA
4. Month	999. NR

#### **Adverse Events**

124 (0180 22 (01108	
19. Were any of the following advers	e events mentioned
, c	(Check all that apply)
Hypoglycemia	
Other adverse events	
No Adverse events	
Not described	
Not applicable	□

20. Is there a reference that needs to be checked?

	(Circle one)
Yes	1
No	2
If YES, which one(s):	

(Enter reference # and/or author or 9999 if don't know.)

# SMBG Project- Randomized Controlled Trials Quality Measurement

		PILOT	03/14/07
First Author:			
1. Treatment Allocation		5. Was the care provider blinded? Yes	П
a. Was a method of randomization performed		No.	
Yes		Don't know	······
No		Don't know	
Don't know			
b. Was the treatment allocation concealed?			
Yes		( Was the metions blinded?	
No		6. Was the patient blinded?	П
Don't know	□	Yes	
		No	
		Don't know	
2. Were the groups similar at baseline regarding	the most important	t	
	•		
prognostic indicators? Yes	_		
prognostic indicators?		7. Were point estimates and measures	of variability presented for the
prognostic indicators? Yes		7. Were point estimates and measures oprimary outcome measures?	v -2
yes		7. Were point estimates and measures of primary outcome measures?  Yes	
yes		7. Were point estimates and measures of primary outcome measures?  Yes	
yes		7. Were point estimates and measures of primary outcome measures?  Yes	
yes		7. Were point estimates and measures of primary outcome measures?  Yes	
yes		7. Were point estimates and measures of primary outcome measures?  Yes	
yes		7. Were point estimates and measures of primary outcome measures?  Yes  No  Don't know	
yes		7. Were point estimates and measures of primary outcome measures?  Yes	-to-treat analysis?
yes		7. Were point estimates and measures of primary outcome measures?  Yes  No  Don't know	-to-treat analysis?
yes		7. Were point estimates and measures of primary outcome measures?  Yes  No  Don't know	i-to-treat analysis?
yes		7. Were point estimates and measures of primary outcome measures?  Yes	i-to-treat analysis?
yes		7. Were point estimates and measures of primary outcome measures?  Yes  No  Don't know	i-to-treat analysis?

# **APPENDIX B. PEER REVIEW COMMENTS TABLE**

## **Peer Review Comments Table 1.**

Reviewer	Section	Comment	Change
Pogach	Background	The investigators frame the background in terms of targets and measures. I would suggest that the background by Guerci in the ASIA study frames the question better: "Theoretically, SMBG can improve compliance with recommendations on diet and exercise and medication regimens. The American Diabetes Association has recommended that the optimal frequency of SMBG for patients with type 2 diabetes should be adequate to facilitate reaching glucose goals. This hypothesis is based on the fact that lifestyle changes are facilitated by SMBG. Under these conditions, we should expect an improvement of glycemic control SMBG increases patient management costs, and because of the high prevalence of type 2 diabetes, efforts to establish the efficacy of SMBG in type 2 diabetes mellitus are of greater relevance."	This suggested change was made, however, reference to targets was kept in this revision as the key questions from VA concern targets and not general improvements in glycemic control.
Pogach	Background	If the investigators want to include a discussion of targets, their reliance on ADA Clinical Practice Recommendations is incomplete, and needs to take into account other guidelines and be more complete in describing the ADA recommendations. The authors frame the ADA recommendations to bias the reviewer towards tight control for most. "The Association (ADA) recommends an A1c goal of <7% for "patients in general" but adds that, "for the individual patient," intensive therapy to achieve an A1c as close to normal (<6%) without hypoglycemia is the goal, although the latter recommendation is based on weaker or incomplete evidence.4". To be evidence explicit and transparent, the investigators need to note (to be evidence explicit) that multiple guidelines, including the ADA, American Geriatric Society, and VHA-DOD discuss the need for less stringent targets based upon life expectancy (AGS and VA) or age (ADA >65 years of age), comorbid conditions, and side effects (including hypoglycemia). The ADA "in general" thus refers to individuals who are younger without contraindications. Moreover, the NHLBI study permits an A1c between 7.0-7.9%(expected mean 7.5%) in the control group.	We deemphasized the focus about targets and the ADA, but retained the text about VA performance measures as targets, since the key questions given to us by VA concern efficacy at achieving target glycemic control levels.
Aron	Introduction	This evidence review is being performed by VA. Therefore, it is quite surprising that the recommendations of the American Diabetes Association are so prominently stated. The recent article in the New York Times related to conflicts of interest in determining performance measures should give us pause. I realize that this is in the introduction and meant to provide context, but I would rather have seen studies cited, e.g., DCCT and UKPDS rather than the ADA (or any other advocacy organization).	Text about ADA has been deemphasized.
Pogach	Background	I don't understand why performance measurement is pertinent to the introduction. Only NCQA recommends public reporting for A1c <7% (see Pogach, Engelgau, Aron JAMA 2007). Thus, I would recommend removing references to performance measures as being not relevant.	The text regarding performance measures is retained because VA's questions to us were framed in terms of target levels.
Aron	Study Identification/ Study Selection	Some of the criteria for study inclusion were not explicit. I am referring here specifically to the statement that studies not included in other meta-analyses/reviews were included in this one. The reasons why are not included.	The reasons were indicated in Table 1, and no change was made in the text.
Pogach	Study Identification/ Study Selection	I am not satisfied with the investigators' explanation that "we included studies rejected by Balk and/or by Welschen for a variety of reasons (italics mine)".	
Pogach	Study Identification/ Study Selection	If the investigators believe that their inclusion is still justified, in contrast to the AHRQ Evidence Synthesis (Balk report) the investigators should provide an explicit explanation of the reasons why they disagreed.	
Pogach	Study Identification/ Study Selection	The investigators frame the meta-analysis by noting that it is to address SMBG in individuals on oral hypo-glycemic medications. It is unclear to me whether the Kwan study included individuals on insulin; the Cho study did include 7 out of 40 control groups on insulin (4 insulin only) and 11 of 40 intervention group (6 insulin only). If these studies are included, this needs to be noted as a limitation of generalization of the study findings. In addition, the willingness and ability to use the internet to download meter results may prevent generalization to other populations with lower Socio-economic position.	We agree and the articles by Cho and Kwon were removed from the analysis.
Aron	Study Identification/ Study Selection	P17. "Initial screening of the articles resulted in 13 RCTs that measured the effect of SMBG compared to a group not receiving SMBG and monitored A1c levels with at least three months of follow-up. Two were excluded; one because the trial presented duplicate data, the other because the trial compared a control group of SMBG to an intervention group of SMBG plus other components. (Figure 1)" Unfortunately, this is not the case. The Cho study states: "We performed a diabetes education program again to standardize every patient's education for diabetes management and the method and frequency of self-monitoring of blood glucose (SMBG) according to glucose control." The control group used SMBG. The only difference was that the experimental group had the internet intervention. Why is this study included?	

### **Peer Review Comments Table 1. Continued**

Reviewer	Section	Comment	Change
Pogach	Study Identification/ Study Selection	The investigators note that "Initial screening of the articles resulted in 13 RCTs that measured the effect of SMBG compared to a group not receiving SMBG and monitored A1c levels with at least three months of follow-up. Two were excluded; one because the trial presented duplicate data, the other because the trial compared a control group of SMBG to an intervention group of SMBG plus other components. (Figure 1)." By these criteria, the Kwon (2004) and Cho (2006) articles should be excluded, since the control group and intervention group each received the same number of monitoring strips and received the same instructions on monitoring. The intervention being tested was therefore the "Internet Based Blood Glucose Monitoring System", which essentially increased the frequency of access to the diabetes team; electronic case management in a sense. It's my perspective that the investigators are obligated to remove these studies from the main analysis.	We agree and the articles by Cho and Kwon were removed from the analysis.
Pogach	Study Identification/ Study Selection	The investigators note that "Eligible study designs included controlled clinical trials, RCTs, and systematic reviews/meta- analyses. Observational studies, case reports, non-systematic reviews, letters to the editor and other similar contributions were excluded." This review separately comments on observational studies done in veterans, but not observational studies of non- veterans. The investigators need to be consistent; either remove them or separately discuss all observational studies. I suggest excluding them as not being relevant to the meta-analysis as defined. In addition, the investigators, in their criteria for inclusion, do not include observational studies. None the less, they include older retrospective VA studies. If they choose to include VA studies, they should modify their inclusion/exclusion criteria to include others. Otherwise (and given that meta- analyses of RCTs have significant limitations as well), I would exclude them.	We have revised the methods and results to indicate that the observational studies in veterans were searched for and reported on as evidence regarding the effectiveness of SMBG in the VA patient population and delivery
Aron	Study Identification/ Study Selection	P13 "Eligible study designs included controlled clinical trials, RCTs, and systematic reviews/meta-analyses. Observational studies, case reports, non-systematic reviews, letters to the editor and other similar contributions were excluded." However, in discussing studies in veterans, observational studies were included. It is not clear why they were included here and not elsewhere. The reasons should be made explicit. That also raises the question about using observational studies in non-veterans.	system, as opposed to the efficacy evidence from RCTs.
Aron	Study Identification/ Study Selection	Inconsistencies aside, it is an interesting philosophical issue what the appropriate control group should be in studies like this. Individuals with diabetes have free access to SMBG, i.e., can do it without a prescription. What is usual care in this regard?	We agree this is an interesting question. We agree that the Cho and Kwon studies aren't comparing SMBG to no SMBG, so as indicated above, we deleted these. We interpreted VA's main interest as SMBG vs. no SMBG at all.
Pogach	Data Synthesis	A significant positive aspect of this study is to adjust for baseline A1c. This is welcome, and should be commented upon in more detail (see also data synthesis).	We have added text about this.
Pogach	Data Synthesis	The reviewer's perspective is that adjusting for baseline HbA1c is an appropriate consideration and can be defended (see Bloomgarden Z et al Lower Baseline Glycemia Reduces Apparent Oral Agent Glucose-Lowering Efficacy: A meta-regression analysis Diabetes Care 2006 29: 2137-2139. This should be commented upon in greater detail.	
Aron	Data Synthesis	It is an interesting issue whether or not to adjust for baseline A1c. I would have liked to see both adjusted and unadjusted analyses.	Only unadjusted pooled results are presented in Figure 2. Figure 3 presents the pooled result of studies adjusting for baseline levels of A1c at the individual study level. The metaregression analysis assesses the relationship between baseline A1c and efficacy of SMBG. So all three kinds of analyses are already included in the report - unadjusted, adjusted at the individual study level, and adjusted at the pooled analyses level.

### **Peer Review Comments Table 1. Continued**

Reviewer	Section	Comment	Change
Aron	Conclusions	To reiterate, it is not clear why observational studies are included and I don't see how one can draw the conclusion that veteran patients may not be receiving the full possible benefits of SMBG. I happen to agree with the conclusion, but that comes more from my experience in clinic than from these studies.	The reason for including observational VA studies has now been made clear.
Pogach	Conclusions	In multiple sections of the report the investigators state that "The results of the studies with Veterans do not negate the evidence from RCTs that the addition of SMBG and education can result in a decrease in A1c levels of about 0.3% absolute at six months and up to one year. As previously noted, I do not know why observational studies are included at all, and recommend that that the observational studies be removed.	Observational studies were included as the only available evidence of effectiveness in VA patients.
Pogach	Conclusions	The investigators, on multiple occasions state "that these studies do raise the question of whether veteran patients are receiving the full possible benefits of SMBG." It should be removed. Further, these statements indicate to me a pre-conceived bias, especially since the issue of SMBG efficacy, in individuals who are diet controlled or stable is controversial, and cannot be fully resolved by a meta-analysis. Furthermore, and this is more pertinent to the issue, the investigators indicated that "we draw no conclusion about the effect of frequency of SMBG monitoring on A1c values, and judge the strength of the evidence to be very low."	We disagree with the suggestion to remove the statement about effectiveness of SMBG in Veterans, as there is evidence to support no effectiveness.
Pogach	Future Research	One important limitation of the meta-analysis is that earlier studies from the early mid-90s used SMBG methodology that was much more inconvenient than current methodology. Glucose meters from that era required substantially more blood, transfer to the monitoring strip was more cumbersome, and data feedback from the meters less user friendly if present at all. All of these factors may have contributed to inconclusive results from early studies, and emphasizes the need for research in this area.	We have added this to future research
Pogach	Future Research	The investigators note: "The evidence is insufficient to draw conclusions about which components of SMBG (additional-education, algorithms or other techniques to adjust medication) and frequency of testing are most associated with better results. More research is needed." Agree, this limitation is important and should be better highlighted.	We added additional text on this.
Pogach	Future Research	"However, observational studies in the VA do not report differences in A1c levels between Veterans using or not using SMBG supplies. This raises the question about implementation: more research is needed to understand if implementation of SMBG in a typical VA clinic setting is sufficient for Veterans to receive the full benefit reported in clinical trials." The more pertinent issue is efficacy not effectiveness (see item 2). Please delete this statement.	We disagree, and note that VA's key question to us concerned effectiveness as well as efficacy.
Pogach	Future Research	"Additionally, data are needed about the cost-effectiveness of SMBG in a VA setting.". Unless I am mistaken doesn't cost effectiveness analysis depend upon efficacy data? This seems premature to me. Even if such data were available, it would also involve a number of assumptions that would have to be based upon Markov modeling.	We agree this would involve modeling, but disagree that such an effort is premature. Our analysis of efficacy data support that SMBG is efficacious, therefore a CEA analysis may help better determine which variables are most important in determining cost effectiveness and the identification of these important variables could then target new studies.
Pogach	Future Research	Impact of SMBG on medication adherence should be evaluated. Non-compliance with oral-antiglycemic medications is a recognized issue among veterans and among non-veterans. It is also possible the system interventions to improve adherence may not need to incorporate increased frequency of SMBG.	We have added this to future research.
Pogach	Future Research	I have noted my comments about the Cho/Kwon study design in the previous section. Nonetheless, although I have some reservations about the study design for the purpose of this meta-analysis given the author's inclusion/exclusion criteria, I think that the study design is actually more relevant to what is now considered usual care; e.g., most persons with type 2 diabetes with training in SMBG and some supplies. (Key question 4). This might be mentioned under future research; i.e., that usual care (infrequent) for SMBG be the control group for persons with diabetes on oral agents.	We added this to future research.

### **Peer Review Comments Table 1. Continued**

Reviewer	Section	Comment	Change
Aron	Future Research	This section seems pretty generic for the most part. More problematic is that SMBG is viewed completely in isolation. Most diabetes interventions are complex and involve more than activity. Moreover, other outcomes are relevant, e.g., behavior change. Finally, what does pramlintide have to do with this? That seemed to come out of the blue.	We have revised the future research section and also deleted the reference to pramlintide.
Pogach	Future Research	I substantially disagree with the language of the research implications.  "Our review of existing data support the beneficial effect of SMBG on A1c levels in the context of a clinical trial. Although improvement in A1c is modest, it is equivalent to that achieved with some of the newer medical therapies for diabetes, such as pramlintide.44,45" As noted previously, I believe that there is a bias by including the Cho and Kwan studies. However, based upon the main analysis of this study, it is probably most pertinent to note that the benefit of SMBG [including bundled interventions] for persons on oral hypoglycemic agents is similar to that found for diabetes education interventions, many of which included SMBG (Norris et al, Diabetes Care, 2002). Better designed prospective clinical trials, especially for individuals with stable glycemic control (e.g., at their target A1c) are necessary.  Mentioning a specific medication is inappropriate. Please delete.	We have dropped the use of pramlintide as a reference for efficacy and have inserted the diabetes education.
Pogach	Future Research	I would recommend, as noted previously, that future research include alternative study designs to reflect the fact that SMBG is considered usual care for patients on medication (though not on diet alone).	We made this change.
Pogach	Future Research	Use of SMBG in context of VHA Health Buddy would be an appropriate area of investigation.	We added this to future research.
Pogach	Overall Evaluation	The investigators were thorough in their identification of possible trials for inclusion in their report, but the reviewer has concerns that the included randomized trials articles from Cho and Kwan did not meet the stated inclusion criteria. This introduces biases which are not fully addressed in their discussion/and conclusions. This is a significant flaw of the study as written, and it needs to be more fully addressed. If the investigators wish to justify their inclusion, then the reviewer suggests that the meta-analysis should be presented with and without these studies to permit comparison with the AHRQ evidence synthesis.	We agree that leaving in Cho and Kwon introduced biased and have therefore removed them from the analyses in this revision.

# **APPENDIX C. EVIDENCE TABLE**

**Evidence Table 1. Randomized Controlled Trials Evaluating the Self-Monitoring of Blood Glucose** 

	l lab.	<u> </u>	11100		Controlled Trials Evaluating the Self-Monitoring of Blood Glucose  Delphi List Quality Criteria  Arm/ Group											
Author, Year	Sample Size Enroll/ Follow- up	Dur. of Diabetes inYears	Mean Age	Mean Weight (kg) / BMI	% Women / Race	Method of Randomization Allocation Concealment Similarity at Baseline between groups	Eligibility criteria specified Outcome assessor blind Care provider blind Patients blinded	Point estimates & measures of variability for primary outcome variable  Did analysis included intention to treat analysis	Sample Size entering	Components	# Visit	Freq of SMBG Times/ Week	Dur. of Tx	Outcome	Adverse Events	
			54			Yes	Yes	Yes	25	Exercise Counseling/Edu	20	Control	62 wks	Alc		
Wing RR et al., 1986 19	50 / 45	NR		54	54	98 / NR	78% / NR	No No	No No	No	25	SMBG Exercise Pt Control led	20	5.4	62 wks	Fasting Glucose BMI/Weight loss
						N	No	V.	(0)	Counseling/Edu	4	C 4 1	<i>C</i> 4			
Fontbonne A	208 /			73 / 27	42% /	No No	Yes No	Yes	68	Counseling/Edu	4	Control	6 mths	A1c BMI/Weight Loss		
et al., 1989 <sup>20</sup>	164	13	55		NR	Yes	No No	No	68	SMBG Counseling/Edu	4	7.5	6 mths		ND	
						No	Yes	Yes	83	NR	NA	Control	1 year			
Rutten G et al., 1990 <sup>23</sup>	149 / 127	8.1	63	75 / NR	65% / NR	No No	No No	No	66	SMBG Dietician Counseling/Edu	Vari- able	NR	1 year	A1c BMI/Weight Loss	ND	
						Yes	No Yes	Yes	14	Dietician Counseling/Edu	8	Control	44 wks	Alc		
Muchmore DB et al., 1994 <sup>24</sup>	29 / 23	5	59	99 / 34	61% / NR	No	No No	Yes	15	SMBG	8	3	44 wks	BMI/Weight Loss HRQOL*	ND	
						Yes	No									
					70% /	No	Yes	Yes	22	NR	2	Control	4 mths	A1c	Hypogly- cemia	
Jaber LA et al., 1996 25	45 / 39	6	62	90 / 33	Ancestr	No	No No	No	23		NR	8	4 mths	Fasting Glucose		
					У	Yes	No			Counseling/Edu				HRQOL*		
	64 / 64	NR	50		45% / NR	No	Yes	Yes No	32	Counseling/Edu 19	19	Control	18 mths	A1c Fasting Glucose	Hypogly- cemia	
Kibriya MG, et al., 1999 <sup>27</sup>				60 / 24		No	No		22	SMBG Pt Control led Counseling/Edu	7	1				
Ct al., 1999						No	No No		32				18 mths			
						Yes	Yes	Yes	110+	Counseling/Edu	6	Control	24 wks	A1c		
Schwedes U, et al., 2002 <sup>29</sup>	250 / 223	5.3	60	89 / 31	48% / NR	No Yes	No No	No	113+	SMBG Dietician	6	12	24 wks	BMI/Weight Loss HRQOL*	ND	
							No	***	244+	Counseling/Edu	-	0 . 1	6 1	TINQUE		
Guerci B, et	988 /				45% /	No No	Yes No	Yes	344+	Counseling/Edu	5	Control	6 mths	A1c	Hypogly-	
al., 2003 30	689	8.1	62	83 / 30	NR	Yes	No No	Yes	345+	SMBG Counseling/Edu	5	6	6 mths	Fasting Glucose	Other	
Davidson				82.3 /	74% / African	No	Yes	Yes	45	Dietician Other	13	Control	6 mths	Alc		
MB, et al., 2005 <sup>32</sup>	89 / 88	5.6	50	32.5	Ancestry, Hispanic, Other	No Yes	Yes Yes	Yes	43	SMBG Dietician	13	36	6 mths	BMI/Weight Loss	ND	
					Other		No V		152	Other	NID	C-ntn-1	12			
Farmer A et	Farmer A et d., 2007 <sup>33</sup> 453		66	NR / 31.3	Yes         Yes         Yes           43% / Yes         No         Yes           NR         Yes         No         Yes			Yes	152 150	Usual Care SMBG	NR NR	Control 6	12 mths 12 mths	Alc	Hypogly-	
al., 2007 <sup>33</sup>		3				Yes	SMRG	NR	12 mths	BMI/Weight Loss	ceima					
110 11 10				***		1 2 1 2 1 2 2 2 2 2 2	No entering cample size repor									

ND=Not Described, NR=Not Reported, NA=Not applicable, \*HRQOL=Health Related Quality of Life, \*No entering sample size reported, this is the sample size completing the trial