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4		Prefix	
5		Family name	Myers
6	Corresponding Author	Particle	
7		Given name	Valerie H.
8		Suffix	
9		Degrees	
10		Organization	Pennington Biomedical Research Center
11		Divison	Department of Behavioral Medicine
12		Address	6400 Perkins Road, Baton Rouge, Louisiana 70808-4124, USA
13		e-mail	MyersVH@pbrc.edu
<hr/>			
14		Prefix	
15		Family name	Boyer
16	Author	Particle	
17		Given name	Bret A.
18		Suffix	
19		Degrees	
20		Organization	Widener University
21		Divison	The Institute for Graduate Clinical Psychology
22		Address	Chester, PA, USA
<hr/>			
23		Prefix	
24		Family name	Herbert
25	Author	Particle	
26		Given name	James D.
27		Suffix	
28		Degrees	
29		Prefix	
30		Family name	Barakat
31	Author	Particle	
32		Given name	Lamia P.
33		Suffix	
34		Degrees	
35		Organization	Drexel University
36		Divison	Department of Psychology
37		Address	Philadelphia, PA, USA
<hr/>			
38		Prefix	
39		Family name	Scheiner
40	Author	Particle	
41		Given name	Gary

42 Suffix
43 Degrees

44 Organization Integrated Diabetes Services
45 Address Wynnewood, PA, USA

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49 **Abstract** This study investigated the prevalence of hypoglycemic fear (FH) and hypoglycemia-specific posttraumatic stress (PTS) among individuals with Type I diabetes. Over 25% of participants met diagnostic criteria for current PTSD. High percentages of participants endorsed PTS symptom clusters, suggesting that individuals may be experiencing distress without necessarily meeting diagnostic criteria. Hierarchical multiple regression analyses revealed that perceived threat of death from hypoglycemia and FH were significantly related to PTS. Number of recent hypoglycemic episodes did not predict PTS/PTSD. Depression and nonspecific anxiety did not contribute to the statistical prediction of PTSD, suggesting that symptomatology endorsed represents hypoglycemia-specific anxiety rather than global psychological distress. The hypothesis that greater PTS symptomatology would relate to poorer glycemic control was unsubstantiated. Perceived death-threat from hypoglycemia and nonspecific anxiety were the only variables that contributed to prediction of glycemic control suggesting that PTS did not represent a significant barrier for glycemic control in this sample.

50 **Keywords** Diabetes – Posttraumatic stress – Hypoglycemia – Glycemic control – Hypoglycemic fear
separated by ‘-’

Fear of Hypoglycemia and Self Reported Posttraumatic Stress in Adults with Type I Diabetes Treated by Intensive Regimens

Valerie H. Myers · Bret A. Boyer · James D. Herbert ·
Lamia P. Barakat · Gary Scheiner

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Abstract This study investigated the prevalence of hypoglycemic fear (FH) and hypoglycemia-specific posttraumatic stress (PTS) among individuals with Type I diabetes. Over 25% of participants met diagnostic criteria for current PTSD. High percentages of participants endorsed PTS symptom clusters, suggesting that individuals may be experiencing distress without necessarily meeting diagnostic criteria. Hierarchical multiple regression analyses revealed that perceived threat of death from hypoglycemia and FH were significantly related to PTS. Number of recent hypoglycemic episodes did not predict PTS/PTSD. Depression and nonspecific anxiety did not contribute to the statistical prediction of PTSD, suggesting that symptomatology endorsed represents hypoglycemia-specific anxiety rather than global psychological distress. The hypothesis that greater PTS symptomatology would relate to poorer glycemic control was unsubstantiated. Perceived death-threat from hypoglycemia and nonspecific anxiety were the only variables

that contributed to prediction of glycemic control suggesting that PTS did not represent a significant barrier for glycemic control in this sample.

Keywords Diabetes · Posttraumatic stress · Hypoglycemia · Glycemic control · Hypoglycemic fear

Diabetes Mellitus (DM) and its subsequent complications are the third leading cause of death in the United States (Strauss, 1996). Since most diabetes-related morbidity and mortality are associated with persistent hyperglycemia, or elevated blood glucose (BG) levels, the therapeutic goal of glycemic control is to maintain BG within the normative range (Diabetes Control and Complications Trial Research Group, 1993). For individuals with Type I diabetes, administration of exogenous insulin is necessary to achieve these normative levels (Rubin & Peyrot, 2001). Recent advances in treatment options that facilitate maintenance of “tight control” of BG levels include Multiple Daily Injection Regimens (MDI) using Glargine with per-meal Lispro or Aspart insulin in a basal/bolus format, and Continuous Subcutaneous Insulin Infusion (CSII) using an insulin pump (American Diabetes Association, 2001). Results from the DCCT (Diabetes Control and Complications Trial Research Group, 1993) showed that intensive therapy regimens, defined as either (a) three or more daily injections of insulin (MDI), or (b) treatment with an insulin pump (CSII), effectively delayed the onset and slowed the progression of diabetic complications. These results suggested that intensive therapy (MDI or CSII) as compared to the conventional therapy was significantly better at preventing complications associated with DM. These intensive therapies also showed improved glycemic control as measured by glycosylated hemoglobin. One adverse effect associated with intensive insulin therapies was an increased likelihood of having a severe hypoglycemic episode.

V. H. Myers (✉)
Department of Behavioral Medicine,
Pennington Biomedical Research Center,
6400 Perkins Road, Baton Rouge,
Louisiana 70808-4124, USA
e-mail: MyersVH@pbrc.edu

B. A. Boyer
The Institute for Graduate Clinical Psychology,
Widener University,
Chester, PA, USA

J. D. Herbert · L. P. Barakat
Department of Psychology, Drexel University,
Philadelphia, PA, USA

G. Scheiner
Integrated Diabetes Services,
Wynnewood, PA, USA

58 However, there were no significant differences between the
59 conventional and intensive therapies with regard to acute
60 medical complications directly related to a severe hypo-
61 glycemic state leading the DCCT to conclude that the bene-
62 fits associated with intensive therapy outweighed the risks.

63 One shortcoming of the DCCT study is that no distinc-
64 tions were made within the intensive therapy regimen group
65 regarding differences that may be related to regimen choice.
66 Specifically, the intensive regimen condition was a combined
67 sample of individuals who either utilized multiple daily self-
68 injections (MDI) or insulin pumps. No within group compar-
69 isons were made between these two intensive management
70 regimens to clarify whether there were any systematic differ-
71 ences related to choice of intensive regimen that may have
72 influenced the outcome of the study.

73 There are no set criteria establishing which individuals
74 should use MDI or insulin pumps. The National Institute for
75 Health and Clinical Excellence (2006) suggests that CSII
76 therapy should be used for those individuals who have failed
77 multiple-dose insulin therapy. However, the American Dia-
78 betes Association (2006) suggests that insulin pumps are a
79 good choice for almost any person who is willing to monitor
80 their diabetes management closely. Regardless of regimen
81 choice, both methods require frequent BG monitoring and
82 have similar feedback methods for BG. In the present study,
83 both MDI and CSII participants were instructed by their dia-
84 betes educator to monitor BG four times daily suggesting
85 that there were no differences in the amount of BG feedback
86 between the two intensive regimens.

87 CSII therapy has been associated with increased flexibil-
88 ity and lifestyle advantages (Wolf, Jacober, Wolf, Cornell,
89 & Floyd, 1989), more accurate insulin delivery (American
90 Diabetes Association, 2006), and improved/tighter glycemic
91 control (Champion, Sheperd, Rodger, & Dupre, 1980; Di-
92 abetes Control and Complications Trial Research Group,
93 1993). Clinical follow-up studies have also reported de-
94 creased rates of severe hypoglycemia for those using CSII
95 methods. However, randomized studies have not confirmed
96 this finding and less severe hypoglycemia has been found
97 to be more common with pump use (Hanas & Ludvigsson,
98 2006).

99 In general, tight glycemic control may increase the risk of
100 hypoglycemia (Irvine, Cox, & Gonder-Frederick, 1994), or
101 excessively low levels of BG. Hypoglycemic episodes can
102 be physically aversive, create negative mood states, and are
103 potentially life threatening (Gold, MacLeod, Frier, & Deary,
104 1995; Gonder-Frederick, Cox, Bobbitt, & Pennebaker, 1989;
105 Polonsky, Davis, Jacobson, & Anderson, 1992; Taylor &
106 Rachman, 1988). Many individuals with DM are knowl-
107 edgeable that the symptoms of hypoglycemia may signal
108 potential death (Cox, Irvine, Gonder-Frederick, Nowacek,
109 & Butterfield, 1987; Strauss, 1996).

110 Studies have shown fear of hypoglycemia (FH) to relate
111 to poorer glycemic control (Cox, Irvine, Gonder-Frederick,
112 Nowacek, & Butterfield, 1987), to higher trait anxiety, and
113 difficulty distinguishing between anxiety and hypoglycemia,
114 and past hypoglycemic experiences (Polonsky, Davis,
115 Jacobson, & Anderson, 1992), as well as higher per-
116 ceived stress, frequency of past hypoglycemic episodes,
117 and greater daily BG variability (Irvine, Cox, & Gonder-
118 Frederick, 1992). Additionally, some individuals compro-
119 mise their glycemic control by running their insulin lev-
120 els lower/BG levels higher (Surwit, Scovern, & Feinglos,
121 1982), or overtreat early signs of hypoglycemia (Cox, Irvine,
122 Gonder-Frederick, Nowacek, & Butterfield, 1987), in an at-
123 tempt to avoid these hypoglycemic sensations. For these in-
124 dividuals, FH may induce behaviors that increase risk for
125 the long-term medical complications associated with hyper-
126 glycemia, and reduce the efficacy of these regimens for op-
127 timal glycemic control. While an important literature has
128 begun to investigate FH, indicating that it may interfere with
129 self-management, more thorough investigation of this phe-
130 nomenon appears warranted, and may serve to guide clinical
131 intervention and optimize metabolic outcomes.

132 Taken together, these studies suggest that some individu-
133 als with Type 1 DM: 1) become hypervigilant and experience
134 intrusive ideation about the risk and threat of hypoglycemia
135 (Cox, Irvine, Gonder-Frederick, Nowacek, & Butterfield,
136 1987; Irvine, Cox, & Gonder-Frederick, 1992), 2) become
137 anxious when experiencing signals of hypoglycemia and/or
138 misconstrue anxiety symptoms as hypoglycemia (Polonsky,
139 Davis, Jacobson, & Anderson, 1992), and 3) show exces-
140 sive escape and avoidance behaviors when they perceive
141 the threat of hypoglycemia (Cox, Irvine, Gonder-Frederick,
142 Nowacek, & Butterfield, 1987; Surwit, Scovern, & Feing-
143 los, 1982). This pattern, which we have observed clinically,
144 raises questions as to whether this symptom pattern reflects
145 the posttraumatic stress symptom clusters (intrusive ideation,
146 anxious arousal, and avoidance) related to hypoglycemia.

147 Posttraumatic stress has been investigated following other
148 life-threatening or severe medical stresses, such as cancer
149 (Barakat, Kazak, Gallagher, Meeske, & Stuber, 2000; Boyer
150 et al., 2002; Erickson & Steiner, 2001; Mundy et al., 2000;
151 Neel, 2000; Pitman et al., 2001; Smith, Redd, Peyser, & Vogl,
152 1999; Widows, Jacobsen, & Fields, 2000), burns (Perez-
153 Jimenez, Graell-Berna, Perez-Sales, & Santodomingo, 1993;
154 Tarrier, 1995; Van Loey, Maas, Faber, & Taal, 2003), spinal
155 cord injury (Boyer, Knolls, Kafkalas, Tollen, & Swartz,
156 2000; Boyer, Tollen, & Kafkalas, 1998; Boyer, Ware, Knolls,
157 & Kafkalas, 2003; Kennedy & Evans, 2001; Lude, Kennedy,
158 Evans, & Beedie, 2005; Mona, Cameron, Lesondak, &
159 Norris, 2000; Nielsen, 2003a, 2003b; Radnitz et al., 1998;
160 Radnitz et al., 1995), and cardiac events (Bennett, 1999;
161 Doerfler, Pbert, & DeCosimo, 1994; Ginzburg et al., 2003).
162 Although two studies have examined PTS among parents

of children diagnosed with Type I DM (Landolt et al., 2002; Landolt, Vollrath, Laimbacher, Gnehm, & Sennhauser, 2005), no studies have investigated whether individuals with DM exhibit the full symptoms of PTS following potentially life-threatening aspects of the disease process and management.

Present study

Individuals utilizing intensive insulin regimens (i.e., MDI and CSII) have received less empirical investigation than the more traditional insulin delivery regimens. Attention to the experience of these individuals is important, because these treatment plans are becoming more widely utilized, and most closely mimic the natural endogenous insulin release of individuals without diabetes. Since the tight control attainable with these regimens is imperative, but can pose substantial risk for hypoglycemia, and since FH may serve as a barrier to successful self-management, understanding the scope and nature of FH is critical to optimal medical outcomes. The present study sought to assess the full symptom clusters of posttraumatic stress among individuals using self-selected MDI and CSII for Type I diabetes. In order to assess for relationship of any PTS symptoms to actual hypoglycemic experiences and/or appraisal of hypoglycemic experiences, patients' number of hypoglycemic episodes were queried, as well as complications and experiences incurred during or after hypoglycemic episodes, and perceived threat of death from hypoglycemia. Self-report of depression and nonspecific anxiety were also collected, and tested for relationship to hypoglycemia-related PTS/PTSD. Our primary hypotheses were (a) experiential history, appraisal factors (perceived death threat), psychological distress, and fear of hypoglycemia may relate significantly with PTS severity, as well as diagnostic levels of PTSD, and (b) experiential history, appraisal factors, psychological distress, fear of hypoglycemia, and PTS/PTSD may relate significantly to participants' glycemic control, as measured by glycosylated hemoglobin (HbA1c). Glycosylated hemoglobin is a blood assay test that measures average BG level over the past 6 weeks to 3 months, and often serves as a stable and reliable measure of glycemic control.

Method

Participants

A total of 90 participants (65 females [72.2 %] and 25 males [27.8 %]) participated in the study. Seventy-seven of the 90 participants (85.5 %) utilized insulin pumps. The average age for the sample was 43.2 years, and 82 participants (91.1 %) classified themselves as 'Caucasian'. The mean

number of months diagnosed with diabetes was 279 (23.25 years) (Range = 12-664 months, *s.d.* = 162.77) A total of 344 participants who met the inclusion criteria: (a) diagnosis of Type I DM; (b) had diabetes for at least 6 months duration; (c) were age 18 years or older; (d) were judged by their certified diabetes educator to be beyond any "honeymoon" period suggesting that the participant had stabilized with regard to their current regimen needs; and (e) used either an insulin pump or multiple daily self-injections in a basal/bolus format as their method of diabetes management were obtained from Integrated Diabetes Services located in a suburban location. The Integrated Diabetes Services (IDS) is a for-profit organization that provides individualized diabetes education and management services to children and adults, specializing in intensive blood glucose management and insulin pump services. Response rate was 26.1% of the eligible patients. A series of t-tests and chi-square tests were conducted to assess whether there were significant differences among the responder and non-responder groups. The t-test comparing the responder and non-responder groups on age was significant [*t*-test (1, 332) = 6.67, *p* < .001] with the responders being older than the non-responders. Chi square tests revealed a significant statistical difference for gender [$\chi^2 = 9.711$, *p* < .002]. The responder group contained a significantly higher number of females compared to the non-responder group.

Procedure

Each of the patients who met inclusion criteria was mailed a letter from the IDS, describing the study and requesting their participation. Verbal consent was attained by telephone before identities of patients were disclosed to non-IDS collaborators. A written Informed Consent Form and the questionnaire packet were then sent to those who had verbally consented. A follow-up telephone call was made 2 weeks after the original mailing date to all individuals who had not returned study materials, prompting them to participate if they so chose. The study protocol was approved by the university Institutional Review Board.

Measures

Demographics questionnaire

The demographics questionnaire is a self-report measure developed specifically for this study. Participants were instructed to provide information regarding their gender, age, ethnicity, date of DM diagnosis, general DM information including significant medical complications and/or hospitalizations, pump use/injection regimen, last glycosylated hemoglobin value, and history of hypoglycemic episodes.

257 *Hypoglycemic fear survey-98*

258 The original Hypoglycemic Fear Survey (Cox, Irvine,
259 Gonder-Frederick, Nowacek, & Butterfield, 1987) is a 27-
260 item self-report questionnaire that contains two subscales.
261 The HFS-Worry subscale consists of 17 items which mea-
262 sure worries about hypoglycemia. The HFS-Behavior sub-
263 scale is 10 items and focuses on behaviors designed to
264 avoid hypoglycemia. Psychometric data on the measure-
265 ment indicate good internal reliability and temporal stabil-
266 ity (Cox, Irvine, Gonder-Frederick, Nowacek, & Butterfield,
267 1987). Responses to each item of the HFS items are on a
268 5 point Likert scale ranging from *never* (1) to *very often*
269 (5). Individual items are summed to produce each subscale
270 score. A revised version of the HFS was developed in 1998
271 (HFS-98), and was provided to the present study by the au-
272 thors of the instrument (Cox, 2001). The HFS-98 contains
273 similar items to the original form, however, six additional
274 items are included. Five additional items have been included
275 to the original Behavior subscale, and one additional item to
276 the Worry subscale. Similar to the original HFS, the HFS-98
277 is also on a 5 point Likert scale. However, items range from
278 *never* (0) to *always* (4). No published psychometric data are
279 yet available on the HFS-98. For the purposes of this study,
280 the Total Score for the HFS-98 was used.

281 *Posttraumatic diagnostic scale*

282 The Posttraumatic Diagnostic Scale (PDS) (Foa, 1995) is
283 designed specifically to correspond with DSM-IV criteria
284 for PTSD. Each of 18 items asks respondents to rate on a
285 4 point Likert scale how bothered they have been over the
286 past month by the DSM-IV PTSD criteria. In addition, nine
287 dichotomous items assess the degree to which the symp-
288 toms have interfered with functioning. Symptom Severity
289 Scores range from 0–51, with higher scores representing
290 higher severity of symptomatology. Participants were ori-
291 ented to complete the items regarding their experience with
292 hypoglycemia only. Participants were not instructed to com-
293 plete the PDS items for other trauma experiences such as
294 rape or victimization. Symptom cluster scoring, subjecting
295 the responses to the DSM-IV diagnostic criteria, was used to
296 determine symptoms consistent with current PTSD, and to
297 avoid false positives that occur more frequently when cut-off
298 scores are used with self-report measures of PTSD (Manne,
299 Du Hamel, Gallelli, Sorgen, & Redd, 1998). The PDS
300 shows good internal consistency (.78–.92) (Foa, Cashman,
301 Jaycox, & Perry, 1997), test-retest reliability of PTSD
302 diagnosis ($\kappa = .74$) and Total Symptom Severity
303 ($\kappa = .83$), and showed 82% agreement with the Struc-
304 tured Clinical Interview for the DSM-III-R (SCID) (Foa,
305 Cashman, Jaycox, & Perry, 1997). Overall, the psychomet-
306 ric properties of the PDS indicate that it is a valid and reliable

instrument for assessing both PTSD diagnoses and symptom 307
severity in a self-report format. While the PDS offers an 308
exceptional self-report format for assessing posttraumatic 309
stress symptoms, the adjunctive use of a clinical interview 310
is necessary for determining an actual diagnosis of PTSD. 311
Clinical interviews were not utilized in this study to avoid 312
extra burden for individuals who chose to participate, and 313
lifetime prevalence rates for PTSD were not assessed. 314

315 *Beck depression inventory-II*

The Beck Depression Inventory-II (BDI-II) is a 21-item self- 316
report measure of depression. Each item is rated on a 4 point 317
scale ranging from 0 to 3. The psychometric properties of the 318
BDI-II are sound (Beck, Steer, Ball, & Ranieri, 1996). The 319
psychometric evaluation of the BDI-II with primary care 320
medical patients has been demonstrated (Arnau, Meagher, 321
Norris, & Bramson, 2001), as well as with individuals with 322
diabetes (Lustman, Clouse, Griffith, Carney, & Freedland, 323
1997). 324

325 *Beck anxiety inventory*

The Beck Anxiety Inventory (BAI) is a 21-item self-report 326
instrument that assesses the severity of anxiety in adults and 327
adolescents. The BAI has demonstrated good psychometric 328
properties (Beck, Epstein, Brown, & Steer, 1988). 329

330 *Glycosylated hemoglobin scores*

Values from the last Glycosylated hemoglobin (HbA1c) 331
blood assay were collected by participants' self-report and 332
IDS chart reviews, representing the average blood glucose 333
value across a 6-week to three-month period. HbA1c rep- 334
resents a more stable and meaningful measure of glycemic 335
control than individual blood glucose tests, and is utilized 336
by most providers as the best measure of overall glycemic 337
control (American Diabetes Association, 2002; Sacks et al., 338
2002). 339

340 **Results**341 *Hypoglycemia-related experiences*

One of the goals of this study was to assess the impact of 342
hypoglycemia-related experiences that may relate to post- 343
traumatic stress (see Table 1). During the course of their 344
lifetime, over 97% of the sample reported having a low 345
BG episode, and 81.1% reported requiring assistance from 346
someone else during a low BG episode. More than 46% 347
of the sample required paramedic assistance during a low 348
BG episode, and over 45% reported a loss of consciousness. 349

Table 1 Participants' endorsement of hypoglycemia-related experiences

	Percentages		
	Total sample	MDI	CSII
Ever had a low BG episode	97.8%	100%	97%
Needed help from others	81.1%	84.6%	80.5%
Paramedic assistance	46.7%	46.2%	46.8%
Loss of consciousness	45.6%	38.5%	46.8%
Trip to the ER	38.9%	15.4%	42.9%
Fear of death from low BG	30%	46.2%	27.3%
Hypoglycemic seizure	25.6%	23.1%	26.0%
Hypoglycemic hospitalization	18.9%	15.4%	19.5%
Automobile accidents	11.1%	7.7%	11.7%

Note. MDI = Multiple Daily Injections; CSII = Continuous Subcutaneous Insulin Infusion.

was excluded as a predictor variable in the main regression analyses.

Gender differences have been noted with regards to post-traumatic stress endorsement. Prevalence data suggest higher rates of PTSD in women than men (Tolin & Foa, 2002). Research suggests that men experience more trauma events, but women are more likely to develop PTSD (Gavranidou & Rosner, 2003), and that prevalence rates of PTSD in men compared to women has been higher for certain types of traumas (Resick & Calhoun, 2001). Chi-square analyses showed no significant differences between genders on PTSD criteria in the current sample $\chi^2(1, N = 90) = .756, p = .385$ or the three symptoms clusters: re-experiencing, avoidance, and arousal endorsement [$\chi^2(1, N = 90) = .037, p = .847, \chi^2(1, N = 90) = .013, p = .908, \chi^2(1, N = 90) = 1.274, p = .259$, respectively], and gender was, therefore, not entered into the subsequent regression equations.

Hypothesis a

A 2 × 2 chi square analysis was conducted to test whether MDI and CSII users differed regarding diagnostic levels of PTSD. In addition, a t-test was conducted with method of insulin delivery as an IV, and PDS total severity score as DV. The chi-square and t-tests assessing differences between participants using CSII and MDI were nonsignificant, suggesting that there were no statistically significant differences in PTS/PTSD related to methods of insulin administration. No significant differences were found for PTSD endorsement $\chi^2(1, N = 90) = 1.339, p = .247$, or on the three PTSD symptom clusters [$\chi^2(1, N = 90) = .091, p = .763, \chi^2(1, N = 90) = .678, p = .410, \chi^2(1, N = 90) = 1.339, p = .247$] and insulin regimen. For this reason the entire sample was aggregated for the analyses for hypotheses b and c.

Hypothesis b

It was hypothesized that experiential history (number of hypoglycemic episodes in last month), appraisal factors (perceived life-threat from hypoglycemia), psychological distress (nonspecific anxiety, depression), and fear of hypoglycemia may relate significantly to PTS severity, as well

350 Thirty percent of the total sample reported fear of death from
351 hypoglycemia, 25.6% experienced a hypoglycemic seizure,
352 38.9% required a trip to the emergency room, 18.9% reported
353 a hypoglycemia-related hospitalization, and 11.1% reported
354 having an automobile accident when BG was low.

355 Posttraumatic stress

356 A total of 23 participants (25.5%) met criteria for cur-
357 rent posttraumatic stress disorder based on symptom cluster
358 scoring of the PDS. Five individuals who utilized the self-
359 injection method (38.4%) and 18 individuals who utilized
360 insulin pumps (23.3%) met criteria for posttraumatic stress
361 disorder as measured by the PDS diagnostic score. Fifteen in-
362 dividuals (23.1%) were women and eight participants (32%)
363 were men (see Table 2). The percentage of participants meet-
364 ing the diagnostic criteria for one or more re-experiencing
365 symptom, three or more avoidance symptoms, or two or more
366 arousal symptoms are presented in Table 2. Within the total
367 sample, 65.5% met criteria on the re-experiencing cluster,
368 31.1% met criteria for avoidance symptoms, 54.4% met cri-
369 teria for the arousal cluster, and 95.5% reported interference
370 in functioning in at least one of the life domains of the PDS.
371 Simple correlations comparing time since diagnosis with the
372 PDS total severity and PDS diagnosis scores, as well as
373 fear of hypoglycemia were non-significant (Myers, Boyer,
374 Herbert, & Scheiner, 2004), therefore, time since diagnosis

Table 2 Participants' posttraumatic stress symptomatology

	Percentage of participants				
	Total sample	MDI	CSII	Men	Women
Current PTSD criteria	25.5%	38.4%	23.3%	32.0%	23.1%
PTSD symptom clusters					
Re-experiencing symptoms	65.5%	69.2%	64.9%	64.0%	66.1%
Avoidance symptoms	31.1%	38.4%	29.8%	32.0%	30.7%
Arousal symptoms	54.4%	69.2%	51.9%	64.0%	50.7%
Interference in functioning	95.5%	100%	94.8%	96.0%	95.4%

Note. MDI = Multiple Daily Injections; CSII = Continuous Subcutaneous Insulin Infusion.

414 as diagnostic levels of PTSD. Two hierarchical multiple re-
 415 gressions were conducted, with IVs for each entered in the
 416 following order: number of previous hypoglycemic episodes,
 417 perceived life-threat from hypoglycemia, anxiety, depres-
 418 sion, and fear of hypoglycemia. Number of episodes and fear
 419 of death from low BG were entered first into the regression
 420 due to the salience these factors may have on psychologi-
 421 cal distress and FH endorsement. FH was placed last in the
 422 analyses because it was conceptualized that it may uniquely
 423 contribute to PTS endorsement beyond the influence of the
 424 other predictors. Subscale scores of the HFS-98 were not
 425 included in the analysis because the subscales were not in-
 426 dependent of one another and the total HFS outcome, and
 427 were significantly correlated with one another with a strong
 428 magnitude ($R = .65-.85, p = .001$) (Cohen, 1988). One
 429 analysis used hierarchical regression with PDS total severity
 430 score as the dependent variable, and the other used hierarchi-
 431 cal logistic regression with the PDS score as the dependent
 432 variable. Hierarchical regressions were chosen based on cri-
 433 teria suggested by Tabachnick and Fidell (1996) that spec-
 434 ify hierarchical regressions should be chosen in order to 1)
 435 elucidate the proportion of variance attributable to each pre-
 436 dictor variable after controlling for the variance accounted
 437 for by other predictor variables already in the equation, and
 438 2) test specific hypotheses for a specific theoretical model.
 439 Since gender, method of insulin administration, and other
 440 non-PTSD related demographic variables (Myers, Boyer,
 441 Herbert, & Scheiner, 2004) were not significantly related
 442 to PTS\PTSD, these variables were not included in the re-
 443 gression analyses. The full hierarchical regression equation
 444 accounted for 64% of the variance in PTS total severity score
 445 ($F = 29.5, p = .001$) (see Table 3). However, perceived

446 death threat from hypoglycemia ($\beta = .25, partial R = .35,$
 447 $p = .001$) and FH ($\beta = .47, partial R = .48, p = .001$)
 448 were the only variables to significantly contribute to the pre-
 449 diction of PTS total severity scores. By Cohen's standards,
 450 R^2 values between 0.2 and 0.49 are small effects, R^2 values
 451 between 0.5 and 0.79 are medium effects, and large effects
 452 are represented by R^2 s 0.8 and higher (Cohen, 1988; Rosnow
 453 & Rosenthal, 1996). The effect sizes for the first hierarchi-
 454 cal analysis ranged from $R^2 = .002$ to .640. The effect size
 455 for previous hypoglycemic episodes was small ($R^2 = .002$),
 456 however, the other variables had large effects. The full hierar-
 457 chical regression equation accounted for 53.6% of variance
 458 in PTSD diagnosis scores ($F = 19.2, p = .001$), with per-
 459 ceived death threat from hypoglycemia ($\beta = .22, partial$
 460 $R = .28, p = .009$) and HFS total score ($\beta = .45, partial$
 461 $R = .42, p = .001$) contributing significantly to the predic-
 462 tion of PTSD diagnosis score. Effect sizes for this analysis
 463 ranged from $R^2 = 0.000$ to 0.536, with the majority of the
 464 predictor variables demonstrating large effects (see Table 4).

465 A t -test was conducted to compare individuals' current
 466 PTSD diagnosis on measures of depression and anxiety
 467 [t -test (1, 88) = -4.69, $p = .001$]. Individuals who met
 468 current PTSD according to the PDS symptom cluster and
 469 severity scores criteria reported significantly higher BDI-II
 470 and BAI scores than participants who did not meet current
 471 PTSD criteria. Additional correlations between the PDS to-
 472 tal severity score and both the BDI-II and BAI were statisti-
 473 cally significant [$r_{(BDI-II)} = .499, p = .001; r_{(BAI)} = .633,$
 474 $p = .001$]. This suggests that there is a positive correlation

Table 3 Hierarchical regression analysis summary for experiential history, appraisal factors, psychological distress, and fear of hypoglycemia predicting PTS severity ($N = 89$)

Measure	<i>B</i>	<i>SEB</i>	β	R^2	ΔR^2	ΔF
Step 1 ^a						
Previous hypoglycemic episodes	.012	.073	.011	.002	.002	.217
Step 2 ^b						
Perceived life threat	5.039	1.481	.251*	.278	.275	32.73*
Step 3 ^c						
Anxiety	.171	.102	.167	.520	.242	42.87*
Step 4 ^d						
Depression	.107	.079	.112	.528	.008	1.422
Step 5 ^e						
Fear of hypoglycemia	.223	.044	.473*	.640*	.112	25.83*

^aNumber of low blood glucose episodes in last month, ^bEndorsed whether believe would die from a hypoglycemic event, ^cTotal score on the Beck Anxiety Inventory, ^dTotal score on the Beck Depression Inventory, ^eTotal score of the Hypoglycemia Fear Survey.

* $p < .01$.

Table 4 Hierarchical regression analysis summary for experiential history, appraisal factors, psychological distress, and fear of hypoglycemia predicting PTS diagnosis ($N = 89$)

Measure	<i>B</i>	<i>SEB</i>	β	R^2	ΔR^2	ΔF
Step 1 ^a						
Previous hypoglycemic episodes	-.0276	.043	-.049	.000	.000	.016
Step 2 ^b						
Perceived life threat	2.344	.871	.225*	.219	.219	24.09*
Step 3 ^c						
Anxiety	.101	.060	.197	.434	.215	32.36*
Step 4 ^d						
Depression	.0112	.047	.023	.435	.000	.050
Step 5 ^e						
Fear of hypoglycemia	.110	.026	.451*	.536*	.102	18.21*

^aNumber of low blood glucose episodes in last month, ^bEndorsed whether believe would die from a hypoglycemic event, ^cTotal score on the Beck Anxiety Inventory, ^dTotal score on the Beck Depression Inventory, ^eTotal score of the Hypoglycemia Fear Survey.

* $p < .01$.

475 between higher scores on the PDS and higher scores on the
476 BDI-II and BAI.

477 *Hypothesis c*

478 It was expected that the same factors may relate significantly
479 to participants' glycemic control, as measured by HbA1c. A
480 hierarchical regression was conducted with IVs entered in the
481 following order: number of previous hypoglycemic episodes,
482 perceived life-threat from hypoglycemia, anxiety, depres-
483 sion, and fear of hypoglycemia, and total PTS severity score,
484 and HbA1c as the DV. Although the full model accounted for
485 18.8% of the variance in HbA1c, and remained a significant
486 prediction of HbA1c scores ($F = 3.12, p = .008$), only per-
487 ceived death-threat from hypoglycemia (F change = 4.46,
488 $p = .038$) and BAI score (F change = 10.2, $p = .002$)
489 accounted for significant F change in prediction of HbA1c
490 ($R^2 = .154, F = 5.12, p = .003$), and only BAI score
491 contributed significantly to the prediction ($\beta = .34$, partial
492 $R = .33, p = .002$). Effect sizes for this analysis ranged
493 from $R^2 = .002$ to 0.188, with the majority of variables
494 demonstrating medium and large effects (see Table 5).

495 **Discussion**

496 This study represents the first attempt to evaluate the full
497 scope of PTSD symptom clusters among individuals with

498 Type I diabetes. Based upon findings regarding FH, and
499 upon clinical observations regarding the role of anxious
500 arousal in the fearful avoidance of low BG, the explicit focus
501 was on reactions to hypoglycemia. Since hypoglycemia
502 represents a potentially life-threatening experience associ-
503 ated with salient and distressing symptoms, a hypoglycemic
504 episode may easily be perceived as life-threatening, even if
505 not conceptualized as meeting criteria A of the PTSD diag-
506 nosis (American Psychiatric Association, 2000). These data
507 indicate that one out of four participants reported symptoms
508 consistent with current PTSD in response to hypoglycemic
509 experience, and that a high proportion of individuals met
510 re-experiencing, avoidance, and arousal criteria for PTSD.
511 These results suggest that for a subset of individuals with
512 Type I diabetes, the medical sequelae of diabetes, particularly
513 hypoglycemia, may be sufficient to induce PTSD symptoms.
514 In addition, nearly two-thirds (65.5%) met criteria on the re-
515 experiencin symptom cluster, nearly one-third (31.1%) met
516 criteria for avoidance symptoms, and over half (54.4%) met
517 criteria for the arousal symptom cluster. Additionally, over
518 95% of the sample reported interference in functioning in at
519 least one of the life domains of the PDS. This suggests that
520 a large percentage of the total sample may be experiencing
521 emotional distress in each of these domains, but may not
522 meet full current diagnostic criteria.

523 Furthermore, individuals reporting a perceived threat
524 of death from hypoglycemia and reporting specific FH
525 were more likely to meet diagnostic criteria for PTSD,
526 and reported more severe PTS symptomatology. In con-
527 trast, the number of hypoglycemic episodes reported
528 within the past month did not predict either PTS/PTSD.
529 This finding parallels results regarding PTS and cancer,
530 in which subjective appraisal showed stronger rela-
531 tionship to PTS than did medically-determined severity
532 of the treatment conditions (Barakat, Kazak, Gallagher,
533 Meeske, & Stuber, 2000; Tedstone & Tarrier, 2003). The
534 greater relationship between PTS/PTSD and perceived death
535 threat than between PTS/PTSD and number of hypo-
536 glycemic experiences appears dramatic, since 38.9% of pa-
537 tients had required emergency room services, 46.7% had
538 required paramedic services, 45.6% had lost consciousness,
539 and 25.6% had experienced a hypoglycemia-related seizure
540 within the past month. Even though the actual acuity of hypo-
541 glycemic experiences was high, the appraisal of impending
542 death from hypoglycemia showed more potent association
543 with PTS/PTSD symptomatology.

544 It is also noteworthy that, whereas perceived death threat
545 and FH related significantly to PTS/PTSD, depression and
546 nonspecific anxiety were not significantly associated with
547 PTS/PTSD. It appears that despite significant simple cor-
548 relations between anxiety and PTS, and between depres-
549 sion and PTS, these phenomena are not accounting for the
550 hypoglycemia-related PTS when subjected to more rigorous

Table 5 Hierarchical regression analysis summary for experiential history, appraisal factors, psychological distress, and fear of hypoglycemia predicting glycosylated hemoglobin ($N = 88$)

Measure	<i>B</i>	<i>SEB</i>	β	R^2	ΔR^2	ΔF
Step 1 ^a						
Previous hypoglycemic episodes	-.0069	.016	-.043	.002	.002	.163
Step 2 ^b						
Perceived life threat	.450	.355	.151	.052	.050	4.465*
Step 3 ^c						
Anxiety	.0547	.023	.366*	.154	.103	10.20*
Step 4 ^d						
Depression	.0244	.018	.023	.174	.015	1.536
Step 5 ^e						
Fear of hypoglycemia	-.0037	.011	-.055	.178	.009	.865
Step 6 ^f						
PTS severity	-.0235	.024	-.162	.188*	.009	.934

^aNumber of low blood glucose episodes in last month, ^bEndorsed whether believe would die from a hypoglycemic event, ^cTotal score on the Beck Anxiety Inventory, ^dTotal score on the Beck Depression Inventory, ^eTotal score of the Hypoglycemia Fear Survey, ^fTotal Severity score on the Posttraumatic Diagnostic Scale.

* $p < .05$.

multiple regression analyses. The appraisal of lethality of hypoglycemia and FH showed the strongest association with PTS, and were the only variables to contribute significantly to the statistical prediction of PTS. Therefore, the symptomatology presented here represents hypoglycemia-specific anxiety, rather than representing global psychological distress. This finding adds more data to the controversy regarding PTS/PTSD as a legitimate symptom expression subsequent to medical stressors (Andrykowski, Manne, Cordova, & Coyne, 2003). Investigators have questioned whether symptoms reported in inquiries of PTSD are better accounted for by general distress, such as depression and nonspecific anxiety (Coyne, Palmer, & Cook, 2003).

The relationship between HFS and PDS total score raises methodological questions regarding the nature of this relationship. FH may represent a developmental precursor, acting as a risk factor for the development of full PTSD symptomatology. In contrast, HFS may be measuring a subset of the same symptoms assessed by the PDS. If the latter were the case, the relationship between HFS and PDS scores would represent a measurement confound rather than an association between two distinct constructs. A content analysis of the HFS and PDS items suggest that nearly all the HFS items represent a more detailed inquiry of hypoglycemia-specific avoidance and intrusive ideation, paralleling only two symptom clusters of the PDS. For a more rigorous investigation of this issue, future research should subject the HFS and PDS items to factor analytic assessments to test for item clustering. If, however, HFS scores represent measurement of a subset of the PTSD symptoms, similar to the Revised Impact of Events Scale's (Weiss & Marmar, 1997) measurement of intrusive ideation and avoidance but not arousal, then the regression results presented here require different interpretation. That is, if FH and PTS represent aspects of the same phenomenon, then perceived death-threat emerges as the primary predictor of PTS/PTSD.

Nonetheless, investigators and clinicians must be cautious regarding the interpretation of these preliminary data, and further investigation of PTS/PTSD among individuals with diabetes appears warranted. Reviews of issues important for consideration in the diagnosis of PTSD are available (Herbert & Sageman, 2004; McNally, 2003), and have noted that general distress may inflate the endorsement of PTSD items (McNally, 2003). The findings presented here, indicating stronger relationships between perceived death-threat from hypoglycemia than between general distress (i.e., depression and nonspecific anxiety), suggest a hypoglycemia-specific phenomenon. In addition, the finding of a stronger relationship between perceived death-threat and PTS than between actual hypoglycemic history and PTS emphasizes the cognitive processing elements in the emotional processing of the hypoglycemic experience (Foa & Riggs, 1995; Foa & Rothbaum, 1998).

Another issue relevant to PTSD diagnostic conceptualization involves the phenomenon of an ever present, potentially life-threatening event, such as hypoglycemia for those with diabetes. Individuals with Type I diabetes face the threat of hypoglycemia for the duration of their insulin treatment across the course of their life. As such, these data raise a more phenomenological question. Does this PTS represent "posttraumatic stress" or rather ever-present "peri-traumatic stress?" Just as the diagnostic criteria for adjustment disorders have been revised to respect the fact that "stressors" to which one must adjust may be chronic and ongoing, PTSD scholars are currently grappling with the issue of how best to conceptualize individuals' response to ongoing life threat and trauma.

It is also noteworthy that the percentage of participants reporting diagnostic levels for the avoidance symptom cluster was the lowest of the three PTSD symptom clusters. This finding raises several questions. First, does the medical necessity of DM management and the guidance of the medical treatment team serve to prevent avoidance of hypoglycemia-related triggers? Second, as discussed below, this sample represents individuals utilizing the most advanced of all available regimens, and may represent the least avoidant patients. Greater levels of avoidance symptoms might be found among those who are less pursuant of intensive basal/bolus regimens (i.e., those employing traditional regimens like R and NPH or Lente insulins). Third, would the patients who are most avoidant in response to stimuli that trigger hypoglycemia-related distress be less likely to complete the questionnaires and participate in the study? If the answer to this third question is "yes," then the PTS prevalence reported here may be an underestimation of the actual levels of symptomatology in this population.

Either way, it is clear that PTS/PTSD symptoms need to be investigated more thoroughly among varied patient populations, and contrasted among those using different regimens or displaying different initiative in pursuit of advanced or intensive regimens.

Although it was hypothesized that greater PTS symptomatology would relate to higher HbA1c values (i.e., poorer glycemic control), this was not found. Of all the variables considered, only perceived death-threat from hypoglycemia and nonspecific anxiety (i.e., BAI score) contributed to prediction of HbA1c. It is unclear why FH did not impact glycemic control. Fear of hypoglycemia has been, in other studies, a major barrier for appropriate diabetes management with intensive therapies. However, these results suggest a lack of impact. Furthermore, only nonspecific anxiety accounted for a significant change in variance accounted for by the regression equation. As such, the strength of the PTS → HbA1c relationship did not represent an important barrier for glycemic control among these participants.

Therefore, the potential interference of PTS/PTSD on glycemic control requires further investigation.

Limitations

Limitations of this study include the discrepancy in the number of responders versus non-responders and the reliance on a convenience sample. Convenience samples can yield small, non-representative groups, and mailed survey studies have a poor response rate. Only 26% of the original solicitation sample chose to participate in this study. This suggests that the representativeness of this sample to the overall Type I diabetes population is limited, and draws into question the generalizability of these findings. This study also did not measure previous trauma history or adherence with self-management activities, preventing assessment of these factors' possible role in the findings. Despite these limitations, there were notable strengths. This study assessed the experiences of persons with diabetes using the most current regimens for tight control of blood glucose. Although these findings may not be generalizable to the entire Type I population, it addressed a subpopulation of persons with Type I diabetes receiving less attention in the literature.

Conclusion and clinical and research implications

These findings highlight the extent of the intrusive experiences caused by hypoglycemic episodes among individuals with diabetes using intensive insulin regimens. These data indicate that 25.5% of individuals using CSII or MDI basal/bolus regimens report symptoms consistent with current PTSD specific to hypoglycemia. Sixty-five percent reported diagnostic levels of intrusive re-experiencing symptoms, 54.4% reported diagnostic levels of arousal symptoms, 31.1% reported diagnostic levels of avoidance symptoms, and over 95% reported that their hypoglycemia-specific anxiety interfered in at least one area of life functioning (e.g., occupational, interpersonal). In multiple regression analyses, perceived death-threat from hypoglycemia and fear of hypoglycemia statistically predicted PTS/PTSD, while depression and nonspecific anxiety did not contribute significantly to the prediction of PTS/PTSD.

Research should further investigate hypoglycemia-specific PTS/PTSD among a broader population of individuals with diabetes, the potential relationship between PTS and glycemic control, and the relationships among general anxiety, depression, PTS, and perceived threat from hypoglycemia. These data raise the question of whether fear and anxiety specific to hypoglycemia should be screened carefully among individuals using intensive basal/bolus insulin regimen. To date, interventions to reduce FH, or related PTS, have been largely uninvestigated, leaving clinicians with limited empirical data to guide intervention. It may be, however,

that perceived threat from hypoglycemia, hypoglycemia fear and PTS should be a high priority for assessment and treatment among those with diabetes, as is the case for depression and general anxiety. Until the relationships among perceived threat, general anxiety, hypoglycemia-specific anxiety, and glycemic control are more well-investigated, however, it appears warranted to assess these phenomena on an individual basis among patients, in order to prevent any potential negative impact upon both psychological adjustment and medical outcomes.

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