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Author(s)	Wong, CKH; Wong, WCW; Wan, EYF; Chan, AKC; Chan, FWK; Lam, CLK
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1 **Macrovascular and Microvascular Disease in Obese Patients with Type 2 Diabetes**
2 **Attending Structured Diabetes Education Program: A Population-based Propensity-**
3 **matched Cohort Analysis of Patient Empowerment Programme (PEP)**

4

5 Authors: Carlos K.H. Wong¹ PhD, William C.W. Wong¹ MD, Y.F. Wan¹ MSc, Anca K.C.
6 Chan¹ BSc, Frank W.K. Chan² MBBS, Cindy L.K. Lam¹ MD

7 ¹ Department of Family Medicine and Primary Care, The University of Hong Kong

8 ² Integrated Care Programs, Hospital Authority Head Office, Hong Kong Hospital Authority

9

10 **Running title:** PEP was effective in Diabetes

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12 **Correspondence Author and person to whom reprint requests should be addressed:**

13 Carlos King Ho Wong, PhD, MPhil, BSc

14 Department of Family Medicine and Primary Care, The University of Hong Kong

15 Address: 3/F, Ap Lei Chau Clinic, 161 Ap Lei Chau Main Street, Ap Lei Chau, Hong Kong

16 Contact: +852-25185688 (tel); +852-28147475 (fax) carlosho@hku.hk (email)

17

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23

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27

28

Abstract

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Patient Empowerment Programme (PEP) in primary care was effective in preventing diabetes-related complications in patients with diabetes. Nevertheless, the effect of PEP on glycaemic control, weight control, and complications was unclear in obese type 2 diabetic patients. We aimed to assess whether PEP reduced all-cause mortality, first macrovascular and microvascular disease events. A cohort of 6,372 obese type 2 diabetic patients without prior occurrence of macrovascular or microvascular disease events on or before baseline study recruitment date was linked to the administrative database from 2008 to 2013. Non-PEP participants were matched one-to-one with the PEP participants using propensity score method with respect to their baseline covariates. Cox proportional hazard regressions were performed to estimate the associations of the PEP intervention with the occurrence of first macrovascular or microvascular disease events and death from any cause, controlling for demographic and clinical characteristics. During a median 31.5 months of follow-up, 350 (PEP/non-PEP: 151/199) patients suffered from a first macrovascular or microvascular disease event while 93 patients (PEP/non-PEP: 34/61) died from any cause. After adjusting for confounding variables, PEP participants had lower incidence rates of all-cause mortality (hazard ratio (HR): 0.589, 95% confidence interval (CI) 0.380-0.915, P=0.018) and first macrovascular or microvascular disease events (HR: 0.782, 95% CI 0.632-0.968, P=0.024) than those with PEP. Enrolment to PEP was an effective approach in reducing all-cause mortality and first macrovascular or microvascular disease events in obese patients with type 2 diabetes.

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Introduction

Type 2 diabetes mellitus (T2DM) and obesity are evolving pandemics that had increased risk of developing comorbidities and complications, and thus imposed major health and economic burden to health care system worldwide [1]. Since 1970s, the term ‘diabesity’ has coined to describe the individuals with co-occurrence of diabetes and obesity, in which they had pathogenic inter-relationship [2]. Obesity confers one of the major risk factors of T2DM [3] and diabetes-related complications including macro- and microvascular diseases [4]. Nowadays, the vast majority of T2DM patients reported to be obese in the US where obesity was highly prevalent [5].

There were much evidence for the benefits of modest weight loss, equivalent to 5-10% loss of total body weight, in obese patients with T2DM[6,7]. Despite well-established benefits of weight loss, controversies are being focused on the optimal approaches for achieving treatment goals of weight management. Towards the means of effective management of obese T2DM patients, narrative reviews [2,6] have consolidated a broad range of therapeutic approaches including surgical approach via bariatric surgery, pharmacologic approach via anti-obesity and incretin-based anti-diabetic medications, and non-surgical-pharmacologic approach via intensive lifestyle modification. Still, conventional approach of community-based education and support in promoting healthy lifestyle and behavioural changes is one of the key strategies for improving the standard of diabetes care in primary care setting [8].

Currently, structured self-management education provides one of the most reliable pathways to sustained empowerment and healthy behavioural changes in diabetic patients managing their own condition[9]. Clinical benefit of structured diabetes education program delivered in a group or individual basis has been confirmed in systematic reviews [10-13] and meta-analyses [14-16], and resulted in significant improvements in weight control, glycemic control and cardiovascular risk factor control. Although the explicit changes in body weight after structured diabetes education have been well recognized in clinical trials, whether structured education would be associated with modest weight loss and a lower risk of macrovascular and microvascular complications remains questionable in ‘real-world’ setting.

90 Notably, recent studies [17-21] examined the effects on glycemic control, quality of life and
91 incidence of cardiovascular events and microvascular events of structured diabetes education
92 program, Patient Empowerment Programme (PEP), versus the usual clinical practice in
93 primary care setting. As yet, no randomised controlled trials, or population-based
94 observational cohort studies have been conducted to investigate the effect of structured
95 education on weight control, diabetes-related complications in diabetes patients.
96 Furthermore, diabetes patients who enrolled to PEP have access to additional weight
97 management program with exercise and nutrition empowerment sessions offered by trained
98 dietitians and physiotherapists. Nevertheless, no prior studies explored the effect of dual
99 program use on the diabetes patients, in which the effectiveness may be strengthened or
100 hampered.

101

102 The main aim of this study was to test in a population-based propensity-matched cohort
103 study on whether this structured diabetes education program in primary care promoted
104 greater benefits on metabolic control and reduced macro- and microvascular diseases in
105 patients with diabetes. The exploratory aim was to evaluate whether weight management
106 program would improve macro- and microvascular diseases among diabetes patients who
107 have attended PEP. We hypothesized that diabetes patients with PEP attendance were more
108 effective than those without, and dual use of PEP and weight management program yielded
109 additional benefits when compared to standalone participation of PEP.

110

111 **Methods**

112

113 In 2010, the Hong Kong Hospital Authority has launched the Patient Empowerment
114 Programme (PEP) which provided tertiary wide primary care service to the patients. PEP is
115 a structured education programme which aims to enhance the quality of chronic disease
116 management, to equip participants with the knowledge, skills and self-awareness of their
117 own disease condition and to promote autonomous self-regulation to maximise their
118 potential for health and well-being. Through structural health education including skill
119 transfer, self-efficacy enhancement, mutual support groups, targeted treatment plan and
120 weight management, participants' lifestyle modification and risk factor management could
121 be enhanced effectively. Several medical experts in the non-government organisations
122 organised 6-7 PEP sessions (2 disease-specific sessions and 4-5 generic sessions) on
123 structural health education, disease-specific knowledge and lifestyle modification and post-

124 program follow-ups to enhance and maintain the participants' self-management. The total
125 contact time of disease-specific and generic sessions is 8-10 hours (2 hours per session) and
126 5 hours (2.5 hours per session), respectively. Disease-specific components were delivered by
127 experienced nurses through lecture-based learning sessions covering comprehensive
128 information about diabetes, responsibility of self-care management, medications in diabetes
129 control, and contingency management on hypo- and hyper-glycaemia. Each generic
130 component session covers the importance of self-management and behaviour modification,
131 healthy diet and regular exercise goal setting and problem-solving skills, sharing on self-
132 monitoring experience, stress coping management, psychosocial support and networking,
133 and communications with healthcare professionals. A detailed PEP setting and mode of
134 education delivery has been described in the previous study[17-21]. This study included
135 patients attended at least one session of PEP dated between 1 March, 2010 and 30 June,
136 2012.

137

138 **Subjects**

139

140 All patients with T2DM were sampled from a population-based cohort of patients attended
141 the general outpatient clinics in Hong Kong Hospital Authority, the largest public health
142 service provider in Hong Kong. The outcome evaluation included all obese patients (Body
143 mass index ≥ 27.5 kg/m² [22] at baseline) with T2DM who had attended at least one PEP
144 session. The T2DM subjects were identified with the International Classification of
145 Primary Care-2 (ICPC-2) code of 'T90', through the clinical management system database
146 of Hong Kong Hospital Authority. A total of 4,254 Diabetes subjects who had enrolled
147 into PEP and attended at least one PEP session between 1 March, 2010 and 31 March,
148 2012 were included in the evaluation of the incidence in macro- and microvascular events.
149 Out of 41,775 Diabetes subjects (PEP: 4,254, non-PEP: 37,221) within the database,
150 4,395 subjects (PEP: 326, non-PEP: 4,069) were excluded due to the prior diagnosis of
151 macrovascular or microvascular diseases before baseline. Each patient was observed from
152 baseline until the incidence of any macrovascular or microvascular disease events, death
153 from any cause, or date of last follow-up as censoring, or 31 December, 2013, whichever
154 came first. To evaluate the net effect of PEP on the post-intervention, 3,186 Diabetes
155 patients who have not ever participated in PEP on or before 31 December, 2013 were
156 matched to PEP subjects on propensity score matching (described below) as non-PEP
157 group.

158

159 Patients having history of co-morbidities and diagnosis of macro- and microvascular
160 disease events were defined according to the diagnosis coding system of *International*
161 *Classification of Diseases, Ninth Edition, Clinical Modification* (ICD-9-CM) and
162 *International Classification of Primary Care* (ICPC-2) in clinical management system
163 database of the Hong Kong Hospital Authority. The complementary use of ICPC-2 and
164 ICD-9-CM diagnosis coding systems were managed to identify the history of co-
165 morbidities and diagnosis of macro- and microvascular disease events in both the primary
166 and secondary care settings.

167

168 Ethics approval of this study was granted by institutional review board and clinical trial
169 registry (NCT01935349, ClinicalTrials.gov).

170

171 ***Macrovascular and Microvascular Diseases***

172

173 In the present study, four outcome events were our primary interests: 1) all-cause mortality,
174 2) first macrovascular event including coronary heart disease (CHD), stroke, or heart
175 failure, 3) first microvascular event including retinopathy, nephropathy or neuropathy, and
176 4) first composite macro- and microvascular event. The incidence of CHD was defined as
177 the earliest date of diagnosis with either ICPC-2 of K74-K76 or ICD-9-CM of 410.x-414.x
178 or 798.x. The incidence of stroke was defined as the earliest date of diagnosis with either
179 ICPC-2 of K89-K91 or ICD-9-CM of 430.x-438.x. The incidence of heart failure was
180 defined as the earliest date of diagnosis with either ICPC-2 of K77 or ICD-9-CM of 428.x.
181 The incidence of retinopathy was defined as the earliest date of diagnosis with either
182 ICPC-2 of F83 or ICD-9-CM of 249.5x, 362.03-362.06 or 366.41. The incidence of
183 nephropathy was defined as the earliest date of diagnosis with ICD-9-CM of 249.4x,
184 250.40-250.43, 581.x-585.x or 791.0. The incidence of neuropathy was defined as the
185 earliest date of diagnosis with either ICPC-2 of N94 or ICD-9-CM of 249.6x, 250.6x,
186 337.1, 355.x or 357.2.

187

188 ***Baseline Covariates***

189

190 Demographic, biometric data and disease characteristics, and treatment modalities and
191 enrolment of co-intervention [23] for diabetes at baseline were treated as the covariates of

192 patients. Demographic characteristics of patients included sex, age, smoking status, alcohol
193 status, and educational level. Biometric data included body mass index (BMI), hemoglobin
194 A1c (HbA1c) level, blood pressure (BP), lipid profile, triglyceride and estimated glomerular
195 filtration rate (eGFR) on the date within three-month period of baseline. Disease
196 characteristics included the duration of T2DM, history of hypertension, family history of
197 T2DM, insulin, oral anti-diabetic drugs, anti-hypertensive drugs and lipid-lowering agents
198 used, Charlson Comorbidity Index[24] and the enrolment of co-intervention.

199

200 ***Propensity Score Matching***

201

202 A propensity score is the conditional probability of being intervention given the observed
203 covariates [25]. The technique aims to form comparable PEP intervention and non-PEP
204 groups by logistic regression with relevant baseline characteristics of each patient
205 summarized into a single-index variable (the propensity score) and match patients in the non-
206 PEP comparison pool to patients in the PEP intervention group based on the value of the
207 propensity score [26-28]. Correspondingly, the propensity score was generated for each
208 patient, modelling PEP intervention as a dependent variable and baseline covariates of
209 patients (including sex, age, smoking status, alcohol status, educational level, HbA1c level,
210 BMI, BP, triglyceride, total cholesterol-to-high density lipoprotein cholesterol ratio, low
211 density lipoprotein cholesterol, eGFR, the level of duration of T2DM, history of hypertension,
212 family history of diabetes mellitus, the use of insulin, oral anti-diabetic drugs, hypertensive
213 drugs and lipid-lowering agent, Charlson Comorbidity Index and enrolment of co-
214 intervention for diabetes) as independent variables. The propensity score mapping was made
215 by using the “psmatch2” command [29] with the nearest neighbour without replacement
216 approach in the STATA.

217

218 ***Data Analysis***

219

220 Descriptive statistics were used to calculate the baseline characteristics of demographic
221 and clinical data in PEP and non-PEP groups after propensity score matching. Differences
222 in baseline characteristics between PEP and non-PEP groups were tested for matched-pairs
223 [30] using independent t-test for continuous variables or chi-square test for categorical

224 variables. Independent t-test was used to assess the differences in HbA1c, systolic BP,
225 diastolic BP, LDL-C and BMI between PEP and non-PEP groups at different time points.
226 The cumulative incidence rate and incidence rate of all-cause mortality, macrovascular and
227 microvascular disease events with the corresponding 95% confidence interval (CI) were
228 reported in both groups based on the assumption that the observed incident cases followed
229 a Poisson distribution.

230

231 Multivariable Cox proportional hazards regression was performed to estimate the effect of
232 PEP on the dependent variable of macrovascular event, microvascular event, first
233 composite event and all-cause mortality, accounting for all baseline characteristics of
234 patients. For each model, survival curves were estimated by Kaplan-Meier method and
235 their differences between PEP and non-PEP groups were compared using the log-rank test.
236 Hazard ratio (HR) and the corresponding 95% CI were reported for each variable in the
237 regression models. Predictive accuracy of Cox models was assessed and compared using
238 Harrell's discrimination C-index, ranging from zero to one. A value of 0.5 indicates no
239 predictive discrimination, and values of 0 or 1.0 indicate perfect separation of patients [31].
240 Goodness-of-fit of Cox regression model were assessed using Akaike information criterion
241 and Bayesian information criterion. Similar analyses were pursued on the subgroup
242 analysis of the effect of weight management on dependent variables among PEP
243 participants.

244

245 All statistical analyses were performed using STATA Version 13.0. All significance tests
246 were two-tailed and those with a p-value less than 0.05 were considered statistically
247 significant.

248

249 **Results**

250

251 Table 1 shows cohort baseline characteristics after 1:1 propensity score matching. Out of
252 4,254 diabetes subjects, 3,186 (74.9%) were successfully matched with non-PEP
253 participants using the demographic and clinical characteristics. As expected, the two
254 groups had similar baseline demographic and clinical characteristics, as indicated in the
255 insignificance of all the p-values (≥ 0.05).

256

257 Comparisons of PEP and non-PEP participants in five of the clinical parameters (HbA1c,
258 systolic BP, diastolic BP, LDL-C and BMI) at different time points are displayed in Figure
259 1. Both groups did not show any significant difference in all of the parameters at baseline
260 but PEP participants had smaller means in all clinical measurement after baseline by
261 observation, when compared with non-PEP participants.

262

263 Table 2 and Figure 2 present Kaplan-Meier survival curves and the number of all-cause
264 mortality, macro- and microvascular disease, and composite events at a median follow-up
265 of 29.5 to 31.5 months (range, 0.5 to 46.5 months). PEP participants generally suffered
266 from fewer death, macro- and microvascular disease events than the non-PEP participants.
267 Specifically, 95 deaths (34 PEP participants and 61 non-PEP participants) were resulted
268 during a total of 8,200 person-years for PEP groups and 8,164 person-years for non-PEP
269 groups. In addition, 350 first macrovascular or microvascular disease events (151 PEP
270 participants and 199 non-PEP participants) occurred during a total of 7,972 person-years
271 for PEP participants and 7,926 person-years for non-PEP participants. This also coincides
272 with the results obtained if macrovascular or microvascular disease events were
273 considered individually.

274

275 *Multivariable Cox Regression Analysis*

276

277 Multivariable Cox regression analyses of all-cause mortality, macro- and microvascular
278 disease events as dependent variables are shown in table 3. After adjusting for
279 confounding variables, PEP participants had a lower incidence rate of all-cause mortality
280 than the non-PEP participants (HR: 0.589, 95% CI 0.380-0.915; P=0.018). Log-rank test
281 further suggested that there was a significant difference in the survival times between the
282 two groups (chi-square statistic=8.47; P=0.004). Additionally, a lower risk of first
283 macrovascular or microvascular disease event was observed among the PEP groups than
284 the non-PEP groups (HR: 0.782, 95% CI 0.632-0.968; P=0.024) and the difference in
285 survival time was significant (chi-square statistic=5.82; P=0.016). However, if the
286 macrovascular or microvascular disease events were studied alone, those two groups were
287 not significantly different in incidence rates (macrovascular diseases: HR: 0.828, 95% CI
288 0.619-1.108; P=0.205; microvascular diseases: HR: 0.761, 95% CI 0.567-1.021; P=0.069).

289

290 *Subgroup Analysis*

291

292 Among those 3,186 PEP participants, 94.0% (n=2994) had not participated in the weight
293 management program. A higher risk of death, but not statistically significant, was
294 observed among PEP participants who participated the weight management program than
295 those who did not (HR: 1.824, 95% CI 0.516-6.442; P=0.351). This result was further
296 confirmed by the corresponding log-rank test (chi-square statistic=0.13; P=0.716).
297 Moreover, participation of weight management program was not associated with a lower
298 incidence risk of macrovascular or microvascular disease events (HR: 0.861, 95% CI
299 0.420-1.765; P=0.682). Similar findings were obtained for the incidence of macrovascular
300 and microvascular disease events individually.

301

302 **Discussions**

303

304 The major findings in this propensity matched cohort study revealed that lower composite
305 macro- and microvascular complication and all-cause mortality were associated with PEP
306 participation in a median of 31.5 months. Compared with non-participants, PEP
307 participants had a reduction in composite macro- and microvascular complication by one-
308 quarter (PEP/non-PEP: 151/199, HR=0.782) and all-cause mortality by half (PEP/non-PEP:
309 34/61, HR=0.589), after adjusting for demographic and clinical characteristics. Results of
310 structured education program were promising, having reduced occurrence of death from
311 any cause and diabetes-related complication events, mainly attributable to the sustainable
312 improvement in glycemic control at various follow-up assessments. Moreover, the
313 additional component of weight management program was not associated with a
314 significant reduction in the mortality, macro- and microvascular events in diabetes
315 patients who attended PEP. Once diabetes patients had participated weight management
316 program in addition to PEP, effectiveness may be reduced due to potentially excessive
317 intervention.

318

319 Macro- and microvascular complications have seldom been reported in the structured
320 diabetes education literature. Besides evidence of prior observational studies from PEP
321 [18,19], the role of structured diabetes education in the incidence of macro- and
322 microvascular complication has only been investigated in the cost-effectiveness analysis of
323 diabetes education and self-management for ongoing and newly diagnosed (DESMOND)
324 [32], using the Sheffield Type 2 Diabetes Model for the long-term incidence of macro- and

325 microvascular complications. It was worthwhile noting that the Sheffield Type 2 Diabetes
326 Model replicated the predicted risk of macro- and microvascular complications among
327 T2DM patients, indicating that the effects of structured diabetes education on observed
328 events of microvascular complication have not been shown in the literature. The results of
329 current study investigated not only the effects of PEP on observed composite complication
330 events, but also the effects of PEP on observed composite macro- and microvascular
331 events. Interestingly, the decreased risk for composite events for PEP participants
332 compared with non-PEP participants was mainly driven more by the occurrence of
333 microvascular events and less by the occurrences of macrovascular events. **Although there
334 was no evidence of a significant reduction in macrovascular events or microvascular
335 events separately among PEP group compared with non-PEP group, the incidence of
336 microvascular event might play an slightly more important role on incidence of composite
337 events in PEP patients.**

338

339 ***Comparison with previous studies***

340

341 It was noteworthy to compare findings of current study with previous studies which
342 investigated the effects of lifestyle intervention for diabetes in the prevention and control
343 of macro- and microvascular complications. The randomized controlled trial focusing on
344 intensive lifestyle modification such as Look AHEAD (Action for Health in Diabetes) trial
345 [33] demonstrated that the lifestyle intervention group had modest weight loss compared
346 to usual care referring to diabetes education program but occurrence of cardiovascular
347 events were not significantly less (HR=0.95, P=0.51) in lifestyle intervention group after a
348 decade of follow-up. By contrast with lifestyle therapeutic approach, results from surgical
349 approach significantly reduced the incidence of macro- and microvascular events.
350 Evidence from long-term follow-up (at least 10 years) observational studies [34,35]
351 consistently showed that bariatric surgery has considered as highly effective approach in
352 reducing risk of macrovascular (HR=0.39-0.68) or microvascular diseases (HR=0.22-0.44)
353 event, and composite event (HR=0.36) when compared to diabetes patients receiving
354 usual care. Despite such effective therapeutic approach, adverse events following bariatric
355 surgery were estimated to be 0.3%-1.0% [36] in a meta-analysis of 32 studies reporting
356 results of bariatric surgery.

357

358 ***Strengths and Limitations of this study***

359

360 There were several strengths in this study. First, as a result of the large patient load and
361 clinical information fully available in the administrative database of Hong Kong Hospital
362 Authority, the study was able to carry out propensity score matching using important
363 baseline covariates. Secondly, owing to similar culture and natural course of T2DM
364 patients with obese in Chinese population, the results would be presumably generalizable
365 to other Chinese populations in primary care setting.

366

367 The study also had some limitations. Firstly, current study was performed as non-
368 randomized study design but instead sourced from the clinical data of routine clinical
369 practice in ‘real-world’ setting. For instance, those who joined PEP may have more health
370 consciousness and motivation compared to those who did not join. We cannot rule out the
371 possibility that PEP participants tended to have better skills and self-awareness, resulting
372 in lower incidence of macro- and microvascular complications. These baseline
373 characteristics were not measurable to isolate the effect of confounding variables on the
374 outcomes. To adjust for confounding variables, the administrative database was lacking in
375 the lifestyle and psycho-social factors such as quality of life and self-efficacy measures,
376 which might result in less robust control for the unbalanced baseline covariates when
377 selecting controls through propensity score matching.

378

379 **Conclusion**

380

381 Results of this propensity score matched cohort study provided evidence that structured
382 diabetes education program was an effective approach in reducing not only HbA1C levels
383 but also all-cause mortality and first microvascular or microvascular disease events in
384 diabetes patients. However, dual use of structured education program and weight
385 management program was not associated with reduction in event occurrences, partly due
386 to potentially excessive program intervened on diabetes patients.

387

388 **Competing interest**

389

390 None declared

391

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393

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398 preparation of the manuscript.

399

400 **Author Contributions**

401

402 C.K.H.W. wrote the manuscript and researched data. F.W.K.C. and A.C. contributed to
403 acquisition of data and reviewed/edited the manuscript. W.C.W.W. and C.L.K.L.
404 contributed to study design. Y.F.W. and A.K.C.C reviewed/edited the manuscript,
405 contributed to statistical analysis and interpretation of results. W.C.W.W. and C.L.K.L.
406 reviewed/edited the manuscript.

407

408

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420

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422

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553

554 **Figure Legend:**

555

556 Figure 1 Comparisons of PEP and non-PEP participants in HbA1c, SBP, DBP, LDL-C and
557 BMI at baseline, 12-month, 24-month and 36-month follow-up

558

559 Figure 2 Kaplan-Meier Survival Curves for All-cause Mortality, Macrovascular and
560 Microvascular Disease Events

Figure 1 Comparisons of PEP and non-PEP participants in HbA1c, SBP, DBP, LDL-C and BMI at baseline, 12-month, 24-month and 36-month follow-up

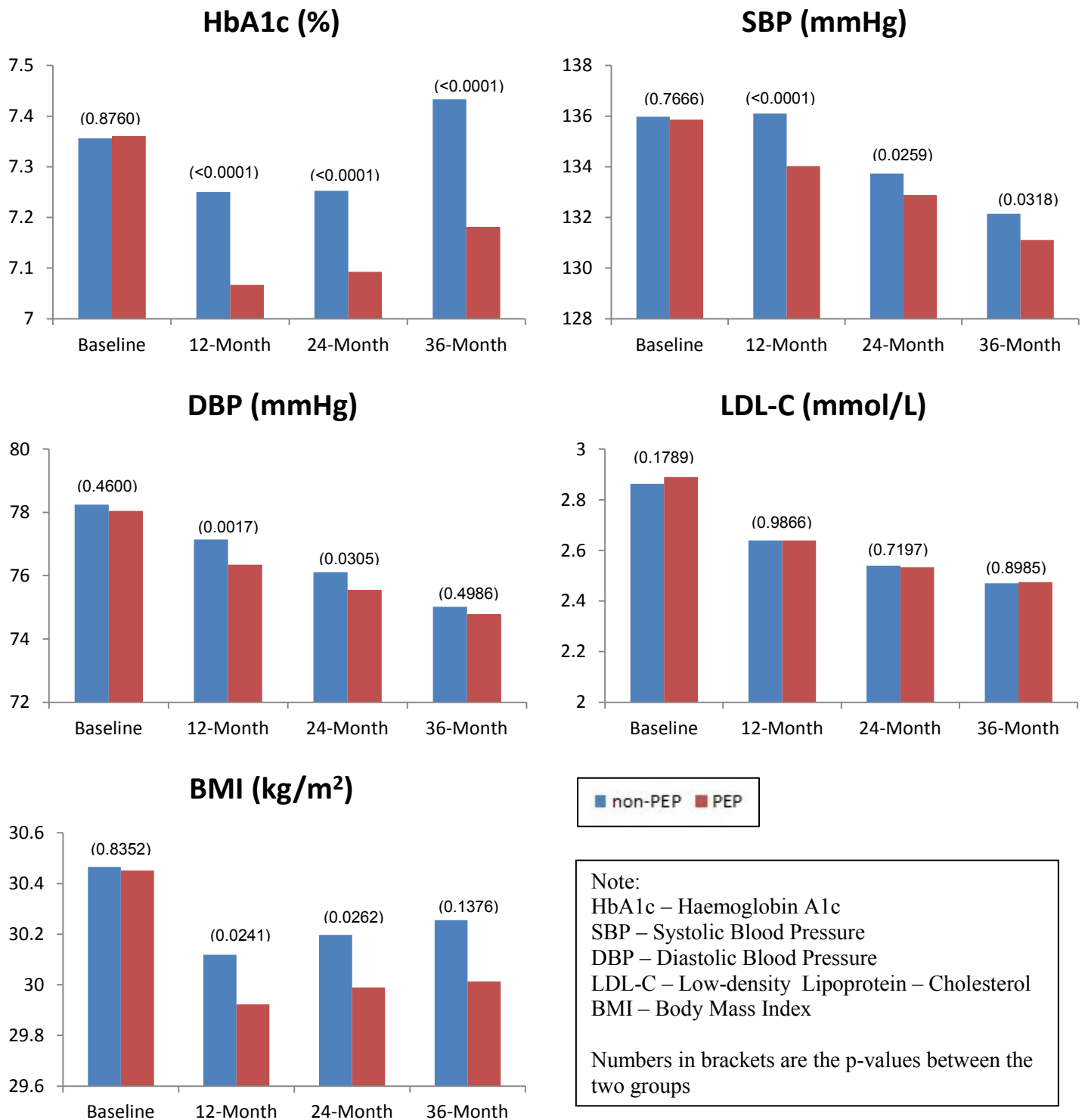


Figure 2 Kaplan-Meier Survival Curves for All-cause Mortality, Macrovascular and Microvascular Disease Events

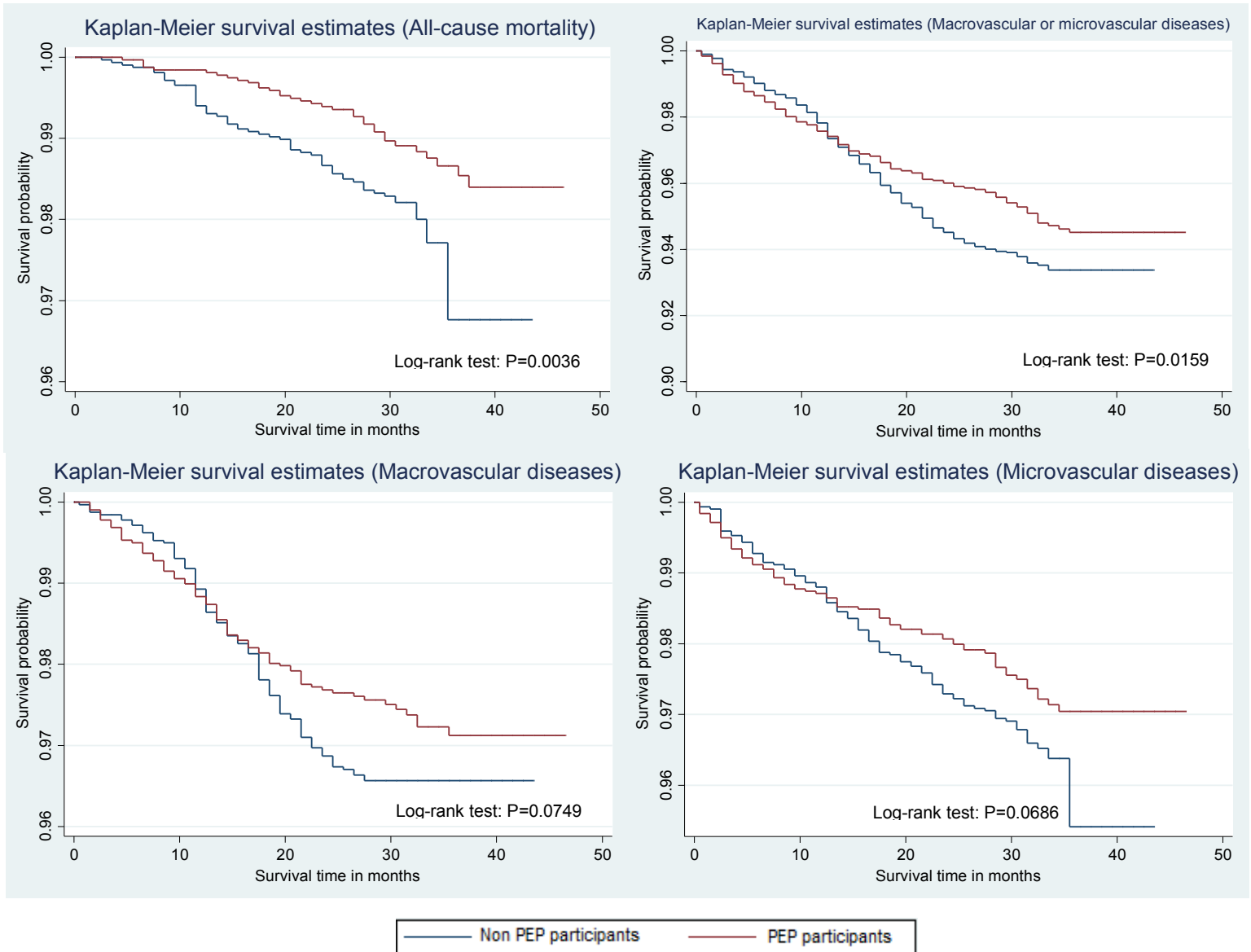


Table 1. Socio-demographic and Clinical Characteristics

Factor	PEP Participants vs non-PEP Participants				PEP Participants			
	Total (N=6,372) % (N)	PEP (N=3,186) % (N)	Non-PEP (N=3,186) % (N)	P-value	Total (N=3,186) % (N)	WM attended (N=2,994) % (N)	WM not attended (N=192) % (N)	P-value
Socio-demographic								
Sex				0.410				0.403
Female	61.7 (3,932)	61.2 (1,950)	62.2 (1,982)		62.2 (1,982)	62.4 (1,868)	59.4 (114)	
Male	38.3 (2,440)	38.8 (1,236)	37.8 (1,204)		37.8 (1,204)	37.6 (1,126)	40.6 (78)	
Age (mean±SD), year	61.60±10.64 (6,372)	61.64±9.75 (3,186)	61.57±11.47 (3,186)	0.785	61.64±9.75 (3,186)	60.49±8.93 (192)	61.71±9.79 (2,994)	0.092
Smoking status				0.755				0.645
Non-smoker	95.8 (6,103)	95.9 (3,054)	95.7 (3,049)		95.7 (3,049)	95.7 (2,864)	96.4 (185)	
Smoker	4.2 (269)	4.1 (132)	4.3 (137)		4.3 (137)	4.3 (130)	3.6 (7)	
Alcohol status				0.803				0.585
Non-drinker	79.8 (5,084)	79.9 (2,546)	79.7 (2,538)		79.7 (2,538)	79.8 (2,388)	78.1 (150)	
Drinker	20.2 (1,288)	20.1 (640)	20.3 (648)		20.3 (648)	20.2 (606)	21.9 (42)	
Educational level				0.379				0.024*
No formal education/ Primary	53.7 (3,421)	53.1 (1,693)	54.2 (1,728)		54.2 (1,728)	54.7 (1,639)	46.4 (89)	
Secondary/ Tertiary	46.3 (2,951)	46.9 (1,493)	45.8 (1,458)		16.3 (519)	16.4 (490)	15.1 (29)	
Laboratory results at baseline (mean±SD)								
BMI, kg/m ²	30.46±2.90 (6,372)	30.45±2.91 (3,186)	30.47±2.88 (3,186)	0.835	30.45±2.91 (3,186)	31.06±3.07 (192)	30.41±2.89 (2,994)	0.003*
HbA1c, %	7.36±1.13 (6,372)	7.36±1.09 (3,186)	7.36±1.17 (3,186)	0.876	7.36±1.09 (3,186)	7.19±0.93 (192)	7.37±1.10 (2,994)	0.023*
Systolic blood pressure, mmHg	135.92±16.04 (6,372)	135.86±16.62 (3,186)	135.98±15.43 (3,186)	0.767	135.86±16.62 (3,186)	134.35±17.07 (192)	135.96±16.59 (2,994)	0.194
Diastolic blood	78.14±10.51 (6,372)	78.05±10.87 (3,186)	78.24±10.15 (3,186)	0.460	78.05±10.87 (3,186)	78.34±11.22 (192)	78.03±10.85 (2,994)	0.701

Factor	PEP Participants vs non-PEP Participants				PEP Participants			
	Total (N=6,372)	PEP (N=3,186)	Non-PEP (N=3,186)	P-value	Total (N=3,186)	WM attended (N=2,994)	WM not attended (N=192)	P-value
	% (N)	% (N)	% (N)		% (N)	% (N)	% (N)	
pressure, mmHg								
Triglyceride, mmol/L	1.76±1.06 (6,372)	1.75±0.98 (3,186)	1.77±1.12 (3,186)	0.425	1.75±0.98 (3,186)	1.83±0.94 (192)	1.74±0.99 (2,994)	0.246
TC/HDL-C ratio	4.15±1.10 (6,372)	4.16±1.11 (3,186)	4.14±1.10 (3,186)	0.626	4.16±1.11 (3,186)	4.26±1.14 (192)	4.15±1.11 (2,994)	0.187
LDL-C, mmol/L	2.88±0.80 (6,372)	2.89±0.81 (3,186)	2.86±0.78 (3,186)	0.179	2.89±0.81 (3,186)	2.92±0.75 (192)	2.89±0.81 (2,994)	0.583
eGFR, ml/min/1.73m ²	85.02±20.98 (6,372)	84.94±19.98 (3,186)	85.09±21.93 (3,186)	0.782	84.94±19.98 (3,186)	84.15±17.89 (192)	85.00±20.11 (2,994)	0.568
clinical								
Duration of T2DM, year	5.82±5.40 (6,372)	5.72±5.57 (3,186)	5.91±5.21 (3,186)	0.155	5.72±5.57 (3,186)	5.30±4.61 (192)	5.75±5.63 (2,994)	0.277
Duration of T2DM, year				0.583				0.707
≤5 years	60.3 (3,840)	60.4 (1,923)	60.2 (1,917)		60.2 (1,917)	60.0 (1,796)	63.0 (121)	
5-10 years	23.1 (1,471)	22.6 (721)	23.5 (750)		23.5 (750)	23.6 (708)	21.9 (42)	
>10 years	16.7 (1,061)	17.0 (542)	16.3 (519)		16.3 (519)	16.4 (490)	15.1 (29)	
History of hypertension	82.9 (5,282)	82.6 (2,633)	83.1 (2,649)	0.595	83.1 (2,649)	83.5 (2,499)	78.1 (150)	0.055
Family history of diabetes mellitus				0.927				0.076
Yes	42.3 (2,697)	42.6 (1,356)	42.1 (1,341)		42.1 (1,341)	42.4 (1,270)	37.0 (71)	
No	8.7 (554)	8.6 (275)	8.8 (279)		8.8 (279)	8.9 (267)	6.3 (12)	
Unknown	49.0 (3,121)	48.8 (1,555)	49.2 (1,566)		49.2 (1,566)	48.7 (1,457)	56.8 (109)	
Insulin used	1.5 (97)	1.4 (46)	1.6 (51)	0.609	1.6 (51)	1.6 (49)	1.0 (2)	0.524
Oral anti-diabetic drugs used	85.2 (5,429)	85.1 (2,712)	85.3 (2,717)	0.860	85.3 (2,717)	85.4 (2,556)	83.9 (161)	0.565
Anti-hypertensive drugs	87.7 (5,589)	87.6 (2,792)	87.8 (2,797)	0.849	87.8 (2,797)	87.9 (2,633)	85.4 (164)	0.300

Factor	PEP Participants vs non-PEP Participants				PEP Participants			
	Total (N=6,372)	PEP (N=3,186)	Non-PEP (N=3,186)	P-value	Total (N=3,186)	WM attended (N=2,994)	WM not attended (N=192)	P-value
	% (N)	% (N)	% (N)		% (N)	% (N)	% (N)	
used								
Lipid lowering agents used	43.1 (2,745)	43.9 (1,400)	42.2 (1,345)	0.164	42.2 (1,345)	42.0 (1,257)	45.8 (88)	0.295
Charlson Comorbidity Index	3.79±1.25 (6,372)	3.79±1.18 (3,186)	3.79±1.32 (3,186)	0.952	3.79±1.18 (3,186)	3.73±1.19 (192)	3.80±1.18 (2,994)	0.429
Enrolment of co-intervention on/before baseline	17.5 (1,113)	17.6 (560)	17.4 (553)	0.817	17.4 (553)	17.5 (525)	14.6 (28)	0.295

Note:

PEP = Patient Empowerment Programme; WM = Weight Management; BMI = Body mass index; HDL = High-density lipoprotein; TC = Total cholesterol; LDL = Low-density lipoprotein; eGFR = estimated glomerular filtration rate; T2DM = Type 2 Diabetes Mellitus

* Significant differences (P < 0.05) by independent t-test or by chi-square test, as appropriate

Table 2. Number and incidence rate of all-cause mortality, macrovascular and microvascular disease events at a median follow-up of 31.5 months

Event	Cumulative incidence		Incidence rate (Cases/ 100 person-years)			Median follow-up periods (Months)
	Cases with event	Rate	Estimate	95% CI*	Person-years	
Total (N= 6,372)						
All-cause mortality	95	0.0149	0.581	(0.470,0.710)	16,364	31.5
Composite Macrovascular or Microvascular Diseases	350	0.0549	2.202	(1.977,2.445)	15,898	31.5
Macrovascular Diseases	189	0.0297	1.172	(1.011,1.352)	16,123	31.5
Microvascular Diseases	185	0.0290	1.147	(0.988,1.325)	16,125	31.5
PEP Participants (N=3,186)						
All-cause mortality	34	0.0107	0.415	(0.287,0.579)	8,200	30.5
Composite Macrovascular or Microvascular Diseases	151	0.0474	1.894	(1.604,2.221)	7,972	29.5
Macrovascular Diseases	82	0.0257	1.015	(0.807,1.260)	8,080	30.5
Microvascular Diseases	79	0.0248	0.977	(0.773,1.218)	8,087	30.5
Non-PEP Participants (N=3,186)						
All-cause mortality	61	0.0191	0.747	(0.572,0.960)	8,164	31.5
Composite Macrovascular or Microvascular Diseases	199	0.0625	2.511	(2.174,2.885)	7,926	31.5
Macrovascular Diseases	107	0.0336	1.330	(1.090,1.607)	8,044	31.5
Microvascular Diseases	106	0.0333	1.319	(1.080,1.595)	8,038	31.5

Note:

PEP = Patient Empowerment Programme; CI = Confidence Interval

* The 95%CI was constructed based on Poisson Distribution

Table 3. Multivariable Cox proportional hazard regression on the dependent variable of all-cause mortality, macrovascular and microvascular disease events, adjusted for the socio-demographic and clinical characteristics

	PEP factor				AIC	BIC	Harrell's C-statistic
	HR†	s.e.	95%CI	P-value			
PEP Participants vs non-PEP Participants (N= 6,372)							
All-cause mortality	0.589	0.132	(0.380,0.915)	0.018*	1,420	1,589	0.896 (0.866,0.927)
Composite Macrovascular or Microvascular Diseases	0.782	0.085	(0.632,0.968)	0.024*	5,889	6,058	0.700 (0.670,0.729)
Macrovascular Diseases	0.828	0.123	(0.619,1.108)	0.205	3,139	3,308	0.751 (0.714,0.789)
Microvascular Diseases	0.761	0.114	(0.567,1.021)	0.069	3,101	3,270	0.706 (0.665,0.747)
	WM factor				AIC	BIC	Harrell's C-statistic
	HR†	s.e.	95%CI	P-value			
PEP with WM session attended vs PEP without WM session attended (N= 3,186)							
All-cause mortality	1.824	1.174	(0.516,6.442)	0.351	453	604	0.916 (0.870,0.962)
Composite Macrovascular or Microvascular Diseases	0.861	0.315	(0.420,1.765)	0.682	2,348	2,499	0.697 (0.654,0.741)
Macrovascular Diseases	1.198	0.515	(0.516,2.783)	0.675	1,266	1,417	0.759 (0.703,0.815)
Microvascular Diseases	0.402	0.290	(0.098,1.650)	0.206	1,230	1,381	0.716 (0.659,0.773)

Note:

WM = Weight Management; PEP = Patient Empowerment Programme; HR = Hazard Ratio;

CI = Confidence Interval; AIC = Akaike information criterion; BIC = Bayesian information criterion

† HR > 1 indicates greater risk for event

* Significant difference (P < 0.05)