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The Live Well, Be Well Study: A Community-Based, Translational Lifestyle Program to Lower Diabetes Risk Factors in Ethnic Minority and Lower–Socioeconomic Status Adults

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The prevalence of type 2 diabetes continues to rise at an alarming rate in the United States. Approximately 25.6 million adults (11.3% of the US population aged 20 years or older) have diabetes, and another estimated 79 million have prediabetes.\(^1\) Greater risk of diabetes is observed for ethnic minority\(^1\) and lower–socioeconomic status (SES) groups\(^6\) compared with White adults of similar ages.

Several clinical trials have tested intensive lifestyle interventions or pharmacological agents in preventing or delaying type 2 diabetes in adults at risk.\(^7\)–\(^9\) These trials showed impressive diabetes risk reductions for lifestyle interventions associated with relatively modest amounts of weight loss and exercise.\(^7\)–\(^9\)

Translating this knowledge into lifestyle interventions delivered in real-world settings is thus a major priority.\(^10\)–\(^12\)

To reduce observed disparities in risk of diabetes, translational studies need to be community-based and designed for lower-SES and ethnic minority populations. Although many translational lifestyle interventions are available, most were designed for clinical settings.\(^13\)–\(^21\) Of community-based translations, only 3 were designed specifically for lower-SES or minority populations.\(^23\)–\(^25\) and only 1 of these—Project HEED, or Help Educate to Eliminate Diabetes—was evaluated with a randomized controlled trial design.\(^23\) HEED was successful in obtaining significant group differences in weight loss at 12 months, but no other significant clinical or behavioral changes were observed.

We conducted a randomized controlled trial of a low-intensity lifestyle intervention for lower-SES, ethnic minority, Spanish- and English-speaking adults. This was a collaborative project between the University of California, San Francisco, and the City of Berkeley Division of Public Health. Public health departments are a good venue for community-based translations to reduce disparities because they serve vulnerable populations most at risk for chronic disease and engage in chronic disease prevention.

OBJECTIVES. We evaluated a community-based, translational lifestyle program to reduce diabetes risk in lower–socioeconomic status (SES) and ethnic minority adults.

METHODS. Through an academic–public health department partnership, community-dwelling adults at risk for diabetes were randomly assigned to individualized lifestyle counseling delivered primarily via telephone by health department counselors or a wait-list control group. Primary outcomes (6 and 12 months) were fasting glucose level, triglycerides, high- and low-density lipoprotein cholesterol, weight, waist circumference, and systolic blood pressure. Secondary outcomes included diet, physical activity, and health-related quality of life.

RESULTS. Of the 230 participants, study retention was 92%. The 6-month group differences for weight and triglycerides were significant. The intervention group lost 2 pounds more than did the control group \((P=.03)\) and had decreased triglyceride levels \((\text{difference in change}, 23 \text{ mg/dL}; P=.02)\). At 6 months, the intervention group consumed 7.7 fewer grams per day of fat \((P=.05)\) and more fruits and vegetables \((P=.02)\) than did control participants.

CONCLUSIONS. Despite challenges designing effective translational interventions for lower-SES and minority communities, this program modestly improved some diabetes risk factors. Thus, individualized, telephone-based models may be a promising alternative to group-based interventions. (Am J Public Health. 2012;102:1551–1558. doi:10.2105/AJPH.2011.300456)
kilograms divided by height in meters squared). The diabetes risk appraisal, adapted from existing diabetes risk tools for use in community settings, used only self-reported variables and a simplified scoring system. Staff scored the diabetes risk appraisal and explained the risk score (0–3 points = low, 4–8 points = moderate, ≥ 9 points = high). Individuals with a moderate or high score (> 4) were educated about their diabetes risk and invited to complete an 8-hour fasting finger-stick test (Accu-chek; Roche Diagnostics, Indianapolis, IN) to determine fasting capillary glucose level.

Individuals who had a capillary blood glucose value between 106 and 160 milligrams per deciliter, who had a moderate to high diabetes risk appraisal score, and who were aged 25 years or older were told about the lifestyle program and study as well as the research process and screened for exclusion criteria. We excluded individuals with diabetes (physician diagnosis, use of insulin or other diabetes medications); diagnosis in past 6 months of myocardial infarction, congestive heart failure, or stroke; heart procedure or heart surgery in past 6 months; implanted defibrillator; hip or knee replacement in past 3 months; insufficient cognitive functioning; and pregnancy. We also excluded individuals not conversant in English or Spanish, with plans to move out of the area within 1 year, and not conversant in English or Spanish, with cognitive functioning; and insufficient cognitive functioning; and pregnancy. We also excluded individuals not conversant in English or Spanish, with plans to move out of the area within 1 year, and whose spouse or partner had already enrolled.

The diabetes risk appraisal, adapted from existing diabetes risk tools for use in community settings, used only self-reported variables and a simplified scoring system. Staff scored the diabetes risk appraisal and explained the risk score (0–3 points = low, 4–8 points = moderate, ≥ 9 points = high). Individuals with a moderate or high score (> 4) were educated about their diabetes risk and invited to complete an 8-hour fasting finger-stick test (Accu-chek; Roche Diagnostics, Indianapolis, IN) to determine fasting capillary glucose level.

Live Well, Be Well Intervention

The lifestyle program28 was designed for lower-SES, minority, and low-literacy adults and adapted from several interventions with established efficacy. It was delivered in Spanish and English and consisted of a 6-month active intervention phase and a 6-month maintenance phase. Trained health department counselors provided education and skills training to modify diet and physical activity through primarily telephone-based counseling (12 calls) with 2 in-person sessions and 5 optional group workshops. In-person and group sessions were held in neighborhood settings. Self-selected and attainable goal-setting and action plans were emphasized to enhance self-efficacy. Motivational interviewing techniques to develop and enhance participants’ motivation were used during the telephone calls. All program materials are available on the Live Well, Be Well Web site (http://iha.ucsf.edu/LiveWellBeWell).

Participation in each program component was tracked. The program consisted of 19 possible “contacts” for a total of 15 possible hours: 1 introductory session, which included a program binder; 1 in-person planning session; 12 telephone counseling calls (10 in active phase, 2 in maintenance phase); and 5 group workshops. Participation was calculated as the total number of contacts received. Minimum compliance was defined as completion of the introductory session, the planning session, and at least 8 telephone calls over the active phase.

We assessed all outcomes at baseline with 6- and 12-month follow-ups. Trained, bilingual research assistants administered questionnaires and performed clinical and anthropometric measurements with standardized protocols. At baseline, we obtained demographic information and a brief medical history. All questionnaires were translated into Spanish. We selected 7 primary clinical and anthropometric outcomes related to diabetes risk. Laboratory measures included fasting serum glucose, triglyceride, and low- and high-density lipoprotein (LDL and HDL) cholesterol levels measured by enzymatic calorimetric methods (Quest Diagnostics, San Jose, CA). Weight in pounds was measured on a digital scale (Detecto Balance Beam Scale; Cardinal Scale Manufacturing Co, Webb City, MO). Measured waist circumference with a Gullick II (FitnessMart; Country Technology, Inc, Gays Mills, WI) tape spring-tension measure at the between the lower ribs and the anterior superior iliac spine (mean of 2 measurements). Systolic blood pressure was measured with an Omron (Omron Healthcare, Inc, Lake Forest, IL) automated blood pressure monitor after sitting for 5 minutes (mean of 2 measurements). Participants provided additional consent for serum banking at baseline and 12 months for metabolic biomarker assays. We measured serum fasting insulin level by radioimmunoassay (Millipore, St. Charles, MO) and calculated homeostasis model assessment-insulin resistance (HOMA-IR).29

Secondary behavioral risk factor outcomes targeted diet and physical activity. We assessed dietary intake with the Modified Block Food Frequency Questionnaire available in English and Spanish.30–32 Questionnaires were scored by NutritionQuest (Berkeley, CA). We used 4 measures targeted specifically in the Live Well, Be Well program: (1) total kilocalories per day; (2) total fat per day; (3) total fiber per day; and (4) daily frequency of consumption of fruits, fruit juices, and vegetables. Missing values were assigned to daily energy intake values that did not fall within the range of 500 to 5000 kilocalories per day (9 instances). Physical activity was measured with the CHAMPS Physical Activity Questionnaire.33 Because Live Well, Be Well emphasized increased participation in all physical activities, especially walking, we used 3 measures: (1) hours per week in any physical activity, (2) metabolic equivalent hours per week in any physical activity, and (3) hours per week walking.

We measured 5 secondary health-related quality-of-life outcomes hypothesized to improve with lifestyle changes. Participants rated their overall health from “poor” to “excellent” on a scale from 1 to 5. Three Medical Outcomes Study measures were (internal consistency reliability in parentheses) the Psychological Distress II and Psychological Well-Being II indexes (0.90 and 0.78, respectively)34 and the Sleep Problems Index (0.77).35 We used the Perceived Stress Scale (0.86) to assess stress.36 Sample size estimates assumed equal allocation to experimental groups, correlation between repeated outcomes equaling 0.75, unit standardized outcomes, 90% retention at 12 months, 80% power, and 2-tailed α equal to .05. Modeled outcomes included difference scores computed by subtracting outcome values assessed at baseline from each corresponding follow-up value. Given the study sample size of 230, the minimum detectable effect corresponded to a group difference in the effect of time equal to 0.28 SDs. This is generally considered a small to small-to-medium effect, suggesting good power.

Eligible participants were stratified by self-reported race/ethnicity (African American, Latino, other) and age categories (25–39, 40–64, ≥ 65 years). We generated stratum-specific sequential identification numbers to randomly allocate individuals to experimental groups in
blocks of 4. Study staff was blinded to the linkage between identification numbers and group assignment. At the end of the baseline visit, the participant opened a sealed, opaque envelope preprinted with the sequential identification number to determine the experimental group assignment. Because of the behavioral nature of the intervention, counselors administering the intervention and participants were not blind to group assignment.

**Statistical Methods**

A randomization check compared experimental groups at baseline via \( \chi^2 \) and \( t \) tests. Intention-to-treat linear regressions modeled repeated change scores: outcomes assessed at each follow-up (6 or 12 months) minus the corresponding baseline value. All models included effects of experimental groups, assessment time, and the groups-by-time interaction. Initially, an unstructured residual covariance structure was specified for the model of each outcome, which was subsequently relaxed in 2 steps to consider separate covariance structures for (1) each experimental group and (2) each combination of experimental group and participant sex. The final residual covariance structure was chosen by reference to deviance statistics. The likelihood-based approach to model estimation allowed models to be fit to all available data and invoked the assumption that missing values occurred randomly, conditional on observed values.\(^{37}\)

**RESULTS**

We randomly assigned 238 individuals to an intervention or a control group between July 2006 and July 2008 (Figure 1) with follow-up assessments from December 2006 through August 2009. Of the 119 individuals assigned to the intervention group, 6 were excluded because of use of diabetes medications at enrollment or for interim pregnancy, leaving 113 for analysis. From the control group, 2 were excluded because of use of diabetes medications, leaving 117 for analysis. Twelve-month study retention was 105 (93%) in the intervention group and 107 (91%) in the control group.

Baseline characteristics indicated a primarily ethnic minority and lower-SES sample, and the average age was older than 55 years, and 73% were women (Table 1). Almost one third were Spanish-speaking, and 77% were from an ethnic minority group. About 20% had no health insurance, about 40% were employed, and approximately 30% reported financial hardship within the prior year. Educational attainment was diverse; about 38% had a high-school diploma or less education. More than half (55%) of the sample were obese, and an additional 32% were overweight.

Approximately 72% of the intervention group participants were minimally compliant. Of 19 possible program components, a mean (SD) of 12.5 (4.9) were completed. Of 12 total possible telephone calls, a mean of 8.9 (3.8) were completed. The average number of workshops attended of 5 was 1.7 (1.5). The mean (SD) hours of contact received was 9.0 (3.3) of a possible 15.

Table 2 presents baseline mean scores and 6- and 12-month change scores for the intervention and control groups and the significance of between-group comparisons for the coprimary clinical outcomes. No baseline group differences were seen in these measures.

Group differences in 6-month change for weight and triglycerides were significant. The intervention group lost 1.9 pounds more than
TABLE 1—Demographic and Socioeconomic Characteristics of Study Participants: Live Well, Be Well Program, Berkeley, Oakland, and Richmond, California, July 2006–August 2009

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Intervention Group (n = 113), % or Mean (SD)</th>
<th>Control Group (n = 117), % or Mean (SD)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Female</td>
<td>73</td>
<td>74</td>
<td>.87</td>
</tr>
<tr>
<td>Race/ethnicity</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>African American</td>
<td>23</td>
<td>23</td>
<td>.74</td>
</tr>
<tr>
<td>Non-Hispanic White</td>
<td>22</td>
<td>23</td>
<td></td>
</tr>
<tr>
<td>Latino</td>
<td>35</td>
<td>39</td>
<td></td>
</tr>
<tr>
<td>Asian</td>
<td>18</td>
<td>13</td>
<td></td>
</tr>
<tr>
<td>Native American/Pacific Islander</td>
<td>1</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Multiethnic/mixed</td>
<td>2</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Age, y</td>
<td>58 (16)</td>
<td>55 (17)</td>
<td>.29</td>
</tr>
<tr>
<td>Language of interview</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>English</td>
<td>72</td>
<td>66</td>
<td>.34</td>
</tr>
<tr>
<td>Spanish</td>
<td>28</td>
<td>34</td>
<td></td>
</tr>
<tr>
<td>Immigrant to the United States</td>
<td>47</td>
<td>48</td>
<td>.93</td>
</tr>
<tr>
<td>Of immigrants, speak English</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Not at all/poorly/fairly well</td>
<td>55</td>
<td>68</td>
<td>.16</td>
</tr>
<tr>
<td>Well/very well</td>
<td>45</td>
<td>32</td>
<td></td>
</tr>
<tr>
<td>Educational achievement</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt; High school</td>
<td>21</td>
<td>25</td>
<td>.14</td>
</tr>
<tr>
<td>High school/GED</td>
<td>20</td>
<td>11</td>
<td></td>
</tr>
<tr>
<td>Some college/tech</td>
<td>27</td>
<td>22</td>
<td></td>
</tr>
<tr>
<td>≥ Bachelor’s degree</td>
<td>32</td>
<td>42</td>
<td></td>
</tr>
<tr>
<td>Health insurance type</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Any private</td>
<td>63</td>
<td>61</td>
<td>.97</td>
</tr>
<tr>
<td>Public</td>
<td>15</td>
<td>17</td>
<td></td>
</tr>
<tr>
<td>None</td>
<td>22</td>
<td>22</td>
<td></td>
</tr>
<tr>
<td>Employed full- or part-time</td>
<td>35</td>
<td>43</td>
<td>.22</td>
</tr>
<tr>
<td>Financial hardship in past y&lt;sup&gt;a&lt;/sup&gt;</td>
<td>30</td>
<td>32</td>
<td>.8</td>
</tr>
<tr>
<td>Family history of diabetes</td>
<td>53</td>
<td>45</td>
<td>.24</td>
</tr>
<tr>
<td>BMI, kg/m²</td>
<td>30.1 (5.3)</td>
<td>29.9 (6.1)</td>
<td>.78</td>
</tr>
<tr>
<td>BMI categories&lt;sup&gt;b&lt;/sup&gt;</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Normal</td>
<td>10</td>
<td>17</td>
<td>.2</td>
</tr>
<tr>
<td>Overweight</td>
<td>31</td>
<td>32</td>
<td></td>
</tr>
<tr>
<td>Obese</td>
<td>59</td>
<td>50</td>
<td></td>
</tr>
<tr>
<td>Hypertension&lt;sup&gt;c&lt;/sup&gt;</td>
<td>50</td>
<td>44</td>
<td>.36</td>
</tr>
<tr>
<td>Arthritis&lt;sup&gt;d&lt;/sup&gt;</td>
<td>35</td>
<td>34</td>
<td>.85</td>
</tr>
</tbody>
</table>

Note. BMI = body mass index; GED = general equivalency diploma.

<sup>a</sup>Responded yes to “In the past 12 months, was there ever a time when you did not have enough money to meet your daily needs?”

<sup>b</sup>BMI categories defined as follows: for non-Asian participants, normal (< 25.0 kg/m²); overweight (25.0 to ≤ 29.9 kg/m²); and obese (≥ 30.0 kg/m²). For Asian participants, normal (< 23.0 kg/m²); overweight (23.0 ≤ BMI ≤ 24.9 kg/m²); and obese (≥ 25.0 kg/m²).

<sup>c</sup>Hypertension classified if either systolic > 140 mm Hg or diastolic > 90 mm Hg, or if participant reported using any blood pressure medication.

<sup>d</sup>Responded yes to “Has a health professional ever told you that you had arthritis or other joint problems?”

Approximately half of the study participants consented to optional blood banking for the insulin assays. Among those with insulin data, the baseline HOMA-IR among the intervention group was 1.18 ± 0.55 and among the control participants was 1.29 ± 0.70. The 12-month HOMA-IR in the intervention group was 1.21 ± 0.54 and in the control group was 1.34 ± 0.63 with no difference between groups.

Table 3 presents baseline mean scores and 6- and 12-month change scores for the intervention and control groups, as well as between-group comparisons, for the secondary behavioral and health-related quality-of-life outcomes. No baseline group differences were found.

The intervention group consumed 7.7 fewer grams per day of fat at 6 months (P = .05). Intervention group members reported more frequent consumption of fruits and vegetables than did the control group at 6 months (P = .02) and 12 months (P = .04). No significant group differences were observed in total calories, dietary fiber, or physical activity. However, some within-group changes were significant: within both groups at 6 months, total consumed calories decreased by 264 kilocalories per day for the intervention group (P < .001) and 217 kilocalories per day for the control group (P < .01). Total fiber intake decreased in both groups at 12 months (P < .05 for the intervention group and P < .01 for the control group). Within the control group, metabolic equivalent hours per week in physical activity increased at 12 months (P < .05).

The intervention group had better psychological well-being than did the control group at 6 months (P = .05) and 12 months (P = .04). The intervention group also had greater improvement in self-rated health at 6 months (P < .05) and fewer reports of sleep problems at 12 months (P = .05). There were no group differences in perceived stress or psychological distress, but there was a within-group change: psychological distress was reduced at 12 months in the intervention group (P = .05).
The intervention group had significant reductions in triglycerides and a significant reduction in total fat intake (7.7 g/day less for the intervention group at 6 months) was notable because reduction in fat consumption was a strong predictor of lower diabetes risk in the Diabetes Prevention Program. We found changes in consumption of fruits and vegetables of 0.6 servings per day at 6 months. The observed effect sizes of these 2 dietary changes were comparable to those found in a systematic review of other physical activity interventions. The improvement in self-rated health was notable because self-rated health consistently predicts future health.

No significant group differences were found in fasting glucose or LDL- or HDL-cholesterol levels, waist circumference, and systolic blood pressure. We have 3 possible explanations: (1) some clinical risk factors were in the normal range at baseline; (2) for some factors, although risk reductions were observed in the intervention group, reductions also were seen in the control group; and (3) the intervention may not have been intensive enough to achieve change.

First, our baseline venous fasting glucose screening can be done. Nevertheless, our sample was at moderate to high risk on other risk factors: about 85% were overweight (more than half were obese), 78% of women and 50% of men had elevated waist circumference, 35% fulfilled metabolic syndrome criteria, and almost 50% had a family history of diabetes. Second, although the intervention group had significant improvements in LDL and HDL cholesterol and a significant reduction in total caloric intake, similar control group improvements precluded observing between-group differences. This may have occurred because diabetes prevention educational materials were part of outreach and recruitment, and control group participants may have changed behavior to some extent without the program. Indeed, at follow-up, 17% of the control group reported having participated in another lifestyle program at some point during the year. In addition, completing the food frequency questionnaire may have raised awareness about diet and portion sizes. Control group improvements have been observed in other diabetes risk reduction interventions. The DEPLOY (Diabetes Education & Prevention with a Lifestyle Intervention Offered at the YMCA) study found significant weight loss in the control group YMCA setting and similarly had provided diabetes education during recruitment. Project HEED found significant control group weight loss; qualitative analyses indicated that control group participants believed they benefited from learning that they were at risk and being
given information about diabetes. Generally, however, such minimal interventions are not effective. Third, the Live Well, Be Well program may not have been sufficiently intensive to achieve broad changes. This underscores the substantial challenge to design practical, sustainable programs for underserved populations that obtain risk reductions comparable to those of the Diabetes Prevention Program. One recent community-based translation, Healthy-Living Partnerships to Prevent Diabetes (HELP PD), found significant reductions in fasting glucose level and several other clinical risk factors, but it was a high-intensity program offered to relatively well-educated participants with high levels of risk. Designing such programs to be more intensive to achieve greater risk reduction might jeopardize the likelihood of adoption and sustainability by community-based organizations.

Live Well, Be Well conformed to contemporary recommendations for translating diabetes prevention lifestyle programs—namely, to use individually tailored goals, self-monitoring, counselors, and other participants to provide support and a problem-solving approach to overcome barriers. Live Well, Be Well also conformed to criteria for an "individually-adapted health behavior change program," strongly recommended by the Task Force on Community Preventive Services to increase physical activity, and behavior change strategies were similar to those in the Diabetes Prevention Program. Instead, the Live Well, Be Well program was a 10-week peer-led group-based program offered in community settings to lower-SES Latino and African American overweight adults with prediabetes. Participants were thus at higher risk than were those in Live Well, Be Well. Ninety participants were randomly assigned with 73% study retention. Significant differences in baseline characteristics were noted; however, such minimal interventions are not effective.

### TABLE 3—Effect of Intervention and Control Group on Changes From Baseline in Behavioral and Health-Related Quality-of-Life Outcomes: Live Well, Be Well Program, Berkeley, Oakland, and Richmond, California, July 2006 to August 2009

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Intervention Group (n = 113)</th>
<th>Control Group (n = 117)</th>
<th>Between-Group Comparison of Change</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Baseline, Mean (SE)</td>
<td>6-Mo Within-Group Change, Mean (SE)</td>
<td>12-Mo Within-Group Change, Mean (SE)</td>
</tr>
<tr>
<td></td>
<td>Baseline, Mean (SE)</td>
<td>6-Mo Within-Group Change, Mean (SE)</td>
<td>12-Mo Within-Group Change, Mean (SE)</td>
</tr>
<tr>
<td>Baseline, Mean (SE)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total calories, kcal/day</td>
<td>1870.51 (78.12)</td>
<td>-264.29*** (50.57)</td>
<td>-301.57*** (64.69)</td>
</tr>
<tr>
<td>Total fat, g/day</td>
<td>71.49 (3.58)</td>
<td>-12.95*** (2.41)</td>
<td>-14.35*** (2.89)</td>
</tr>
<tr>
<td>Total dietary fiber, g/day</td>
<td>17.84 (0.90)</td>
<td>-11.33 (0.70)</td>
<td>-1.97* (0.77)</td>
</tr>
<tr>
<td>Daily frequency fruits and vegetables</td>
<td>3.01 (0.15)</td>
<td>0.25 (0.16)</td>
<td>0.12 (0.14)</td>
</tr>
<tr>
<td>Physical activity, h/week</td>
<td>7.99 (0.63)</td>
<td>0.74 (0.62)</td>
<td>0.68 (0.67)</td>
</tr>
<tr>
<td>Physical activity, metabolic equivalent, h/week</td>
<td>25.59 (2.07)</td>
<td>3.00 (2.20)</td>
<td>2.24 (2.11)</td>
</tr>
<tr>
<td>Walking, h/week</td>
<td>4.40 (0.38)</td>
<td>0.37 (0.41)</td>
<td>0.57 (0.46)</td>
</tr>
<tr>
<td>Self-rated health</td>
<td>3.09 (0.08)</td>
<td>0.31*** (0.07)</td>
<td>0.13 (0.07)</td>
</tr>
<tr>
<td>Perceived stress</td>
<td>2.26 (0.07)</td>
<td>-0.01 (0.06)</td>
<td>0.01 (0.06)</td>
</tr>
<tr>
<td>Sleep problems</td>
<td>2.15 (0.07)</td>
<td>0.06 (0.06)</td>
<td>0.01 (0.06)</td>
</tr>
<tr>
<td>Psychological well-being</td>
<td>3.74 (0.07)</td>
<td>0.01 (0.05)</td>
<td>0.07 (0.06)</td>
</tr>
<tr>
<td>Psychological distress</td>
<td>2.09 (0.07)</td>
<td>-0.06 (0.05)</td>
<td>-0.11* (0.05)</td>
</tr>
</tbody>
</table>

aResidual covariance unstructured within each combination of experimental groups and respondent sex.

bResidual covariance unstructured within each intervention group.

*P < .05; **P < .01; ***P < .001.
weight loss were reported (at 12 months, the intervention group lost 7.2 pounds compared with 2.4 pounds in the control group; P < 0.01); however, they found no other clinical or behavioral changes. Their results are notable because the program was less intensive than Live Well, Be Well and was delivered by trained peer educators.

Live Well, Be Well is the only community-based translation that used an individually tailored, primarily telephone-based model rather than a group-based model. The use of telephone counseling and neighborhood settings for in-person sessions made it convenient to participate, possibly allowing people to enroll who otherwise could not have participated. Indeed, completion of telephone calls was substantially higher than workshop attendance; thus, a group-based approach may not have been feasible for this population. Also, in participant interviews after program completion, telephone calls were rated as the most useful program feature. Use of public health department infrastructure and staff for intervention delivery has not been tested previously in diabetes risk reduction studies.

Our approach addressed 4 translation priority areas by (1) focusing on vulnerable, understudied groups; (2) having few exclusion criteria, thus being more generalizable; (3) being a partnership between researchers and a public health department, thus reflecting their shared perspectives; and (4) being designed to be sustainable by embedding the program in the public health department's chronic disease prevention program.

Limitations
The study had several limitations. Implementation in 1 city-level public health department setting limited generalizability. However, agencies with larger service areas (e.g., county-level health departments) might find the telephone-based counseling model more attractive than in-person health education workshops. Additionally, because we had difficulty recruiting men, our sample included only 26% men. Use of fasting screening tests limited generalizability to adults available in the morning, similar to other studies. Finally, the significant between-group differences in weight and triglyceride levels were small and may have limited clinical benefit.

Conclusions
Our community-based translational study indicated that the Live Well, Be Well intervention was associated with small changes in a few important diabetes risk factors in lower-SES ethnic minority adults, thus providing a promising approach for future translational efforts to reduce disparities. Because so few community-based models for delivering lifestyle interventions are available, our results suggest that individually tailored programs with telephone counseling should be considered along with the more traditional group-based approaches. Testing lifestyle programs that are integrated into a health department's chronic disease prevention infrastructure and delivered by public health department counselors in local community venues provides a novel and sustainable goal for translational research.

Future research could adapt Live Well, Be Well and explore the relative effectiveness of variations in program delivery organization (health department, peer educators), delivery mode (group- vs telephone-based), intensity (number of contacts, duration), and risk-level eligibility (overweight, other diabetes risk factors) in terms of risk reduction and program compliance.

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Contributors
A.M. Kanaya and A.L. Stewart had full access to all the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis. J. Santoyo-Olsson and S. Gregorich performed data analysis, and S. Gregorich led interpretation of the study results. M. Grossman and T. Moore helped with participant recruitment, study visits, and critical revision of the article. All authors helped to conceptualize ideas, interpret findings, and write and review drafts of the article.

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Human Participant Protection
The research protocol was approved by the University of California, San Francisco, Committee on Human Research, and written consent was obtained from all study participants.

References


