School-Based Prevention of Depressive Symptoms: A Randomized Controlled Study of the Effectiveness and Specificity of the Penn Resiliency Program

Jane E. Gillham
University of Pennsylvania and Swarthmore College

Karen J. Reivich, Derek R. Freres, Tara M. Chaplin, Andrew J. Shatté, Barbra Samuels, Andrea G. L. Elkon, Samantha Litzinger, and Marisa Lascher
University of Pennsylvania

Robert Gallop
West Chester University

Martin E. P. Seligman
University of Pennsylvania

The authors investigated the effectiveness and specificity of the Penn Resiliency Program (PRP; J. E. Gillham, L. H. Jaycox, K. J. Reivich, M. E. P. Seligman, & T. Silver, 1990), a cognitive–behavioral depression prevention program. Children (N = 697) from 3 middle schools were randomly assigned to PRP, Control (CON), or the Penn Enhancement Program (PEP; K. J. Reivich, 1996; A. J. Shatté, 1997), an alternate intervention that controls for nonspecific intervention ingredients. Children’s depressive symptoms were assessed through 3 years of follow-up. There was no intervention effect on average levels of depressive symptoms in the full sample. Findings varied by school. In 2 schools, PRP significantly reduced depressive symptoms across the follow-up relative to both CON and PEP. In the 3rd school, PRP did not prevent depressive symptoms. The authors discuss the findings in relation to previous research on PRP and the dissemination of prevention programs.

Keywords: depression, prevention, children, adolescence

Several cognitive–behavioral interventions show promise in preventing depressive symptoms in youths (for recent reviews,

Jane E. Gillham, Department of Psychology, University of Pennsylvania, and Psychology Department, Swarthmore College; Karen J. Reivich, Tara M. Chaplin, Andrew J. Shatté, Barbra Samuels, Andrea G. L. Elkon, Samantha Litzinger, Marisa Lascher, and Martin E. P. Seligman, Department of Psychology, University of Pennsylvania; Derek R. Freres, Annenberg School for Communication, University of Pennsylvania; Robert Gallop, Department of Mathematics Applied Statistics Program, West Chester University.

Barbra Samuels is now at the Treatment Research Institute (Philadelphia). Andrea G. L. Elkon is now at the Georgia Regional Hospital at Atlanta (Atlanta, Georgia). Samantha Litzinger is now at the Department of Psychology, University of Tennessee. Marisa Lascher is now at Lehman Brothers (New York).

The Penn Resiliency Program for Children and Adolescents is owned by the University of Pennsylvania. The University of Pennsylvania has licensed this program to Adaptive Learning Systems (King of Prussia, Pennsylvania). Andrew J. Shatté is employed part-time by Adaptive Learning Systems, and Karen J. Reivich, Andrew J. Shatté, and Martin E. P. Seligman own stock in Adaptive Learning Systems and could profit from the sale of this program. The authors do not have financial relationships with Adaptive Learning Systems.

This research was funded by National Institute of Mental Health Grant MH52270. We are grateful to the children and parents who participated in the project, and to the school teachers, counselors, administrators, and personnel who assisted us in carrying out the project.

Correspondence concerning this article should be addressed to Jane E. Gillham, Psychology Department, Swarthmore College, 500 College Avenue, Swarthmore, PA 19081. E-mail: jgillha1@swarthmore.edu or jgillham@psych.upenn.edu

see Horowitz & Garber, 2006; Merry, McDowell, Hetrick, Bir, & Muller, 2004). Among these are the Coping with Stress Course (Clarke & Lewinsohn, 1995), the Resourceful Adolescents Program (RAP; Shochet, Whitefield, & Holland, 1997), the LISA-T Program (Pössel, Horn, Groen, & Hautzinger, 2004), and the Penn Resiliency Program 1 (PRP; Gillham, Jaycox, Reivich, Seligman, & Silver, 1990). Despite this promise, however, depression prevention programs that demonstrate positive effects are rarely incorporated into school or clinical settings, and little is known about the effectiveness of most programs when delivered by the individuals who work in these settings.

In addition, when these programs do work, it is usually unclear whether the cognitive–behavioral therapy skills or other nonspecific factors (e.g., time spent in a structured after-school activity, attention from an adult, support from peers) are responsible. The majority of depression prevention studies compare cognitive–behavioral interventions with a no-intervention control. Only a few studies have compared prevention programs with attention control groups or alternate interventions. Pattison and Lynd-Stevenson (2001) found that PRP did not significantly reduce or prevent depressive symptoms relative to both no-intervention and attention control groups. The small sample size (N = 63) may have limited power to detect effects, however. Merry, McDowell, Wild, Bir, and Cunliffe (2004) compared RAP with a placebo control group consisting primarily of group arts and crafts activities and found

1 PRP materials are available for use in research. Requests for the curriculum should be made to info@pennproject.org.
some support for the efficacy of RAP relative to placebo, but the effect size was small, and effects were not found across different measures of depressive symptoms. The placebo condition used by Merry et al. controlled for several factors that are not specific to cognitive–behavioral interventions, including adult attention and opportunities to interact with peers. A stronger test of the cognitive–behavioral model would control for additional nonspecific factors, such as the discussion of stressors and topics relevant to depression.

Shatté (1997) and Reivich (1996) took this approach in their studies, which compared PRP with an alternate intervention (the Penn Enhancement Program [PEP]) and a no-intervention control. A total of 152 middle school students were randomly assigned to one of the three conditions. Group leaders were school teachers, school counselors, and psychology graduate students not affiliated with the research team. PRP and PEP both reduced depressive symptoms relative to the no-intervention control. PRP and PEP participants’ depression scores were not significantly different at most of the follow-up assessments. The lack of differences may reflect limited power, as very large cell sizes are often needed to detect differences between active interventions. In addition, it is possible that differences would have emerged had the researchers followed participants for more than 1 year postintervention.

In the present study, we evaluated PRP’s effectiveness as compared with PEP and a no-intervention control in a large sample (baseline N = 697) over a 3-year follow-up period. Our major goals were as follows: (a) to investigate the effectiveness of PRP when delivered in schools by school teachers, school counselors, and other group leaders not affiliated with a research team, and (b) to evaluate intervention specificity. We examined two conceptualizations of depression prevention: (a) reduced levels of symptoms across an extended period of time (Jaycox, Reivich, Gillham, & Seligman, 1994), and (b) reduced onset of symptoms above a certain threshold (Gillham, Shatté, & Freres, 2000).

Method

Participants

This study was approved by the Institutional Review Board and by school administrators and school boards in each of the participating school districts. Figure 1 shows the participant flow from recruitment through the 3-year follow-up. Starting in 1997, two consecutive cohorts of students participated in this study. Each year, the research team sent letters and consent forms describing the project to parents of students in three middle schools (henceforth referred to as Schools A, B, and C) in a suburban metropolitan area in the United States. Recruitment materials described the study as an investigation of two interventions designed to help students cope with day-to-day stressors that are common in adolescence. We informed parents that those children who reported elevated symptoms at baseline would first be offered spots in the project and that other children would be included as space allowed. In total, parents of approximately 4,000 children received recruitment materials. Because the response rate was lower than anticipated, we decided to evaluate PRP with all children for whom we received consent as long as they were not suffering from a depressive disorder at baseline.

A total of 718 children and their parents consented to participate in the project. Response rates ranged from 15% to 22% across the three schools. These children completed baseline measures, including the Children’s Depression Inventory (CDI; Kovacs, 2001). The 175 children with baseline CDI ≥ 13 were administered the depressive disorders section of the Diagnostic Interview for Children and Adolescents (Reich, 2000). Eight children scored positive for major depressive disorder on the Diagnostic Interview for Children and Adolescents and were referred to therapy. Thirteen families dropped from the study after baseline and prior to randomization.

The remaining 697 children (376 boys and 321 girls) entered the intervention phase. Within cohort and school, we stratified children by grade, gender, and baseline CDI score (on the basis of a median split) and then used a computer-generated random numbers sequence to randomly assign participants to one of three study conditions: PRP (n = 232), PEP (n = 231), or Control (CON; n = 234). Within each school and across all three schools combined, there were no significant between-conditions differences on baseline demographic characteristics or depressive symptoms.

At baseline, children’s mean age was 12.13 years (SD = 1.03). A total of 268 children (39%) were in sixth grade, 237 (34%) were in seventh grade, and 191 (27%) were in eighth grade (grade information was missing for 1 student). Age and grade did not significantly differ by school. Other demographic characteristics differed by school (see Table 1).

At baseline, mean levels of depressive symptoms (M = 8.45, SD = 7.35, range = 0–45) were in the range typically found in community samples (Kovacs, 2001). Mean depressive symptoms differed by school, F(1, 694) = 9.52, p < .01. Students in School A (M = 7.03, SD = 7.41) reported lower levels of depressive symptoms than those in School B (M = 9.51, SD = 7.11) or School C (M = 8.71, SD = 7.34), both ps < .01, whereas children in Schools B and C did not differ on baseline depressive symptoms.

Intervention Conditions

PRP (Gillham et al., 1990) is a group intervention that teaches cognitive–behavioral and social problem-solving skills. PRP strives to teach students to think flexibly and accurately about the challenges and problems that they confront. Students learn about (a) the link between beliefs, feelings, and behaviors; (b) cognitive styles, including pessimistic explanatory styles; and (c) cognitive restructuring skills, including how to challenge negative thinking by evaluating the accuracy of beliefs and generating alternative interpretations. Students also learn a variety of techniques for coping and problem-solving, including assertiveness, negotiation, decision making, and relaxation. Students apply the cognitive and problem-solving techniques in their lives through group discussions and weekly homework assignments.

PEP (Reivich, 1996; Shatté, 1997) is a group intervention that focuses on many of the stressors associated with adolescent depression. Topics include peer pressure, ethical dilemmas, trust and betrayal, improving communication, friendships, family conflict, setting and achieving goals, self-esteem, and body image. Each session includes structured activities, role-playing, and guided discussion. PEP was designed to control for adult attention, group cohesion, social support, the discussion of depression-relevant topics, sharing thoughts and feelings with peers, and engagement in homework and in-session activities. A detailed description of PEP is available from the authors on request.

PRP and PEP groups met after school once each week for a total of twelve 90-min sessions. Group leaders were school teachers, school counselors, and graduate students in school psychology, education, and clinical psychology (not affiliated with the research team) who participated in a 30-hr training workshop and biweekly group supervision with the PRP and PEP developers. During the study, 19 PRP groups and 19 PEP groups were implemented, each containing between 6 and 14 children. Teachers and counselors led 25 groups, and graduate students led 11 groups. Because of scheduling difficulties and a shortage of group leaders, research team members led one PRP group and one PEP group. No difference was found in effects for school staff (teachers and counselors) versus graduate students and researchers. Parents of children in PRP and PEP received $5 per session that their child attended and a $15 bonus if their child attended at least eight sessions.
In meetings and correspondence with group leaders and potential participants, we described both PRP and PEP as interventions that aim to increase well-being and help to buffer students against the effects of common stressors. We informed leaders and parents that the programs use different strategies to achieve this goal: PRP teaches specific coping and problem-solving skills, whereas PEP is more discussion oriented.

Students in the CON condition did not participate in PRP or PEP but completed the same assessments as students in the active interventions. In all three conditions, students and their families were permitted to seek counseling or therapy.

Intervention Integrity and Attendance

Intervention sessions were audio-recorded. Following the intervention phase, we selected four lessons from each group for intervention adherence coding. We chose these lessons because they contained key content and included early, middle, and later sessions. Group leaders were not informed which lessons would be rated. Two research assistants rated each lesson on the presentation of 11–14 concepts, skills, or activities. For each content item, raters used a 7-point scale ranging from 1 (no coverage) to 4 (satisfactory coverage) to 7 (excellent coverage). Interrater reliability for the 1–7 ratings was good (PRP intraclass correlation [ICC] = .65; PEP ICC = .77). We computed two integrity scores for each group: (a) the degree covered score, calculated by averaging both raters’ integrity item scores for each session and then averaging the session averages, and (b) the percentage covered score, calculated by averaging the percentage of items covered satisfactorily (≥4) across the sessions. Degree covered and percentage covered scores were highly correlated (r = .83, p < .01). Integrity scores could not be computed for three PRP and two PEP groups because leaders forgot to tape sessions or experienced equipment malfunctions.
Average integrity scores for PRP groups were 4.9 (SD = 0.48) for degree covered and 80% (SD = 7.5%) for percentage covered. Average scores for PEP groups were 4.4 (SD = 0.36) for degree covered and 68% (SD = 5.7%) for percentage covered. Integrity scores were higher for PRP than PEP groups, likely reflecting the increased structure of the PRP curriculum. Mean integrity scores for PRP and PEP groups did not differ significantly by school.

On average, the 232 children assigned to PRP attended 6.71 (SD = 4.22) sessions, and the 231 children assigned to PEP attended 7.11 (SD = 4.43) sessions, t(461) = −1.01, ns. Thirty-seven children (16%) assigned to PRP and 35 (15%) assigned to PEP did not attend any sessions. Of these, 27 in each condition completed at least one follow-up assessment, and their data were included in analyses. PRP attendance differed by school, F(1, 231) = 5.58, p < .01. Attendance was higher in School A (M = 7.30, SD = 3.97) and School C (M = 7.38, SD = 4.04) than School B (M = 5.40, SD = 4.40), ps < .05. PRP attendance was not significantly different in Schools A and C. PEP attendance did not differ significantly by school.

Assessment Schedule and Depressive Symptom Measures

Children in all three conditions completed questionnaires at baseline and at follow-up assessments 2 weeks postintervention and every 6 months thereafter for 3 years. Families received $12 for each assessment.

This article focuses on results for the study’s primary outcome variable: depressive symptoms, as assessed with the CDI and the Children’s Depression Rating Scale—Revised (CDRS–R; Poznanski & Mokros, 1996). The CDI is a widely used measure of depressive symptoms in children and adolescents, which has adequate reliability and validity (Kovacs, 2001).
We used recommended cutpoints of CDI greater than or equal to 13 to categorize depressive symptoms as at least mild in intensity (referred to as elevated symptoms within), and greater than or equal to 19 to denote depressive symptoms as at least moderate in intensity (referred to as high symptoms within). These cutpoints are near the 70th and 90th percentiles, respectively, for this age group on the basis of the CDI standardization norms (Kovacs, 2001).

We also examined clinical symptoms with the CDRS–R. At each follow-up assessment, we invited children with a CDI score greater than or equal to 13 to participate in the CDRS–R, a semistructured interview assessing symptoms of depression in children and adolescents (Poznanski & Mokros, 1996). Trained research assistants conducted the interviews, and a second research assistant coded the interview tape for reliability purposes. Interviewers and coders were not informed of participants’ condition assignments. Interrater reliability was good (ICC = .79). A CDRS–R t score greater than or equal to 65 identified children likely to be suffering from a depressive disorder. Clinical psychologists working on the project contacted parents of these children to inform them of the assessment results and offer referrals. For analytic purposes, children with CDI scores less than 13 were assumed to have a CDRS–R t score less than 65.

During the 3-year follow-up, 235 (36%) of the 651 children who completed follow-up assessments scored greater than or equal to 13 on the CDI and were invited to interviews at one or more follow-up points. Of these 235 children, 62 (20 from PRP, 21 from PEP, and 21 controls) did not complete CDRS–R interviews and so were excluded from the CDRS–R analyses. Thus, CDRS–R analyses used data from 589 children: the 173 children with CDI scores greater than or equal to 13 who completed the interviews and the 416 children who consistently scored less than 13 on the CDI during the follow-up.

Data Analytic Strategy

**Power analyses.** Our goal was to detect small-to-medium intervention effects ($d \geq 0.30$) allowing for 20% attrition. We planned an initial cell size greater than or equal to 220 (final cell size of $n \geq 176$) to achieve a power of greater than or equal to .80 to detect a difference at $\alpha = .05$ (two-tailed). Attrition was higher than expected, however, and our final sample size at the 3-year follow-up assessment allowed us to detect effect sizes greater than or equal to .45 at a power of greater than or equal to .80 and $\alpha = .05$ (two-tailed).

**Inspection and transformation of data.** We evaluated assumptions of normality of continuous variables by examining histograms and calculating skewness and kurtosis scores at each assessment point. CDI scores were skewed at several assessment points so square root transformations were used for these data. Table 2 shows raw means to ease interpretation.

**Attrition and baseline differences.** We used analysis of variance (ANOVA) and chi-square analyses to compare baseline depressive symptoms and demographic characteristics of children who did and did not complete any follow-up assessments. The possibility of differential attrition was examined by testing for between-conditions differences in these measures. These children were evenly distributed across the study conditions and represented between 56% and 59% of the baseline samples in each condition. Children who did not complete any follow-up assessments were older than the children who completed follow-ups, $F(1, 663) = 6.99, p < .01$. We found no significant between-conditions differences in depressive symptoms or demographic characteristics among children who did not complete any follow-up assessments.

A total of 396 participants did not complete the final 3-year follow-up assessment. These children were evenly distributed across the study conditions and represented between 56% and 59% of the baseline samples in each condition. Children who did not complete the 3-year follow-up assessment were older, came from households with lower incomes, and were less likely to have parents who were married than children who completed this assessment, $F(1, 662) = 5.01, p < .05$; $F(1, 438) = 5.83, p < .05$; and $\chi^2(1; N = 672) = 10.28, p < .01$. We found no significant between-conditions differences in depressive symptoms or demo-
ographic characteristics among children who did not complete the final 3-year follow-up.

We found no significant differences by condition on baseline depressive symptoms or demographic variables in the baseline sample. In the follow-up sample, parents’ marital status varied by condition, consistent with the outcome analyses reported within.
Intervention effects. PRP did not significantly reduce depressive symptoms over the follow-up period relative to CON, \(t(622) = -1.29, ns, d = -0.09\ (95\% CI = -0.23, 0.05)\); or PEP, \(t(623) = -0.31, ns, d = -0.02\ (95\% CI = -0.16, 0.11)\). PEP also did not reduce depressive symptoms significantly relative to CON, \(t(628) = -0.97, ns, d = -0.07\ (95\% CI = -0.21, 0.07)\). ANCOVAs comparing pairs of conditions at each assessment revealed no significant intervention effects and only one trend. PRP tended to prevent depressive symptoms relative to CON at the 12-month follow-up, \(F(1, 324) = 3.11, p < .10, d = -0.18\ (95\% CI = -0.37, 0.02)\).

PRP prevented elevated symptoms (onset of CDI \(\geq 13\)) relative to CON. \(\chi^2(1, N = 325) = 5.29, p < .05, \text{odds ratio} = .61\ (95\% CI = 0.40, 0.95)\), but not PEP, \(\chi^2(1, N = 330) = 0.03, \text{ns, odds ratio} = .96\ (95\% CI = 0.60, 1.54)\). PRP prevented elevated symptoms relative to CON, \(\chi^2(1, N = 333) = 4.64, p < .05, \text{odds ratio} = .64\ (95\% CI = 0.42, 0.98)\). Among children with baseline CDI scores less than 13, 33 (20\%) of 161 in PRP, 36 (21\%) of 169 in PEP, and 52 (32\%) of 164 in CON scored greater than or equal to 13 at least once during the follow-up. There were no significant intervention effects on high (onset of CDI \(\geq 19\)) or clinical symptoms (CDRS–R \(t \geq 65\)).

School effects. Intervention effects differed by school. A MM ANOVA predicting follow-up CDI scores from condition, school, and the School \(\times\) Condition interaction (with baseline symptoms covaried) revealed no main effect of condition, a significant main effect for school, \(F(1, 617) = 3.16, p < .05\), and a significant School \(\times\) Condition interaction, \(F(4, 619) = 2.61, p < .05\). Pairwise comparisons indicated that follow-up CDI scores were significantly worse in School C than School A (\(p < .05\)), and tended to be worse in School B than School A (\(p < .10\)). Follow-up CDI scores in School B and School C did not significantly differ.

To locate the source of the School \(\times\) Condition interaction, we conducted MM ANOVAs predicting CDI score from condition, school, and the School \(\times\) Condition interaction for each pair of schools. These analyses revealed no School \(\times\) Condition interaction for the School A and School B comparison but marginal and significant School \(\times\) Condition interactions for the comparisons of School C to Schools A and B, \(F(2, 419) = 2.89, p = .057\), and \(F(2, 445) = 4.16, p < .05\), respectively. An analysis predicting CDI from condition, School A + B versus C, and their interaction revealed no significant main effect of condition but a significant School (A + B vs. C) \(\times\) Condition interaction, \(F(2, 609) = 5.11, p < .01\). To reduce the number of analyses and retain statistical power, we examined outcomes for Schools A and B (combined) and School C.

Schools A and B

Attrition and baseline differences. We found no evidence for differential attrition in Schools A and B. One variable—parents’ marital status—differed by condition in the baseline sample and in the follow-up sample. \(\chi^2(6, N = 411) = 15.02, p < .05\), and \(\chi^2(6, N = 392) = 17.10, p < .01\), respectively. Parents of PEP participants were more likely to be divorced and less likely to list marital status as ‘other’ than parents of PRP participants or controls. Parents of PRP participants and controls did not differ on marital status. Analyses covarying marital status were consistent with those reported within.

Intervention effects. In Schools A and B, PRP significantly reduced depressive symptoms over the follow-up period relative to CON, \(t(636) = -2.59, p < .05, d = -0.24\ (95\% CI = -0.42, -0.06)\), and PEP, \(t(629) = -2.03, p < .05, d = -0.18\ (95\% CI = -0.36, -0.01)\). PRP did not significantly reduce depressive symptoms relative to CON, \(t(640) = -0.59, ns, d = -0.05\ (95\% CI = -0.24, 0.13)\). See Table 2 for means and ANCOVAs by assessment point.

PRP prevented elevated symptoms (onset of CDI \(\geq 13\)) relative to CON, \(\chi^2(1, N = 205) = 8.03, p < .01, \text{odds ratio} = .38\ (95\% CI = 0.19, 0.75)\), but not relative to PEP, \(\chi^2(1, N = 212) = 1.38, ns, \text{odds ratio} = .66\ (95\% CI = 0.32, 1.33)\). PEP tended to prevent elevated symptoms relative to CON, \(\chi^2(1, N = 207) = 2.92, p < .10, \text{odds ratio} = .58\ (95\% CI = 0.31, 1.09)\). Among children with baseline CDI less than 13, 16 (15\%) of 105 in PRP, 23 (22\%) of 107 in PEP, and 32 (32\%) of 100 controls scored greater than or equal to 13 on the CDI at least once during the follow-up (see Figure 2). There were no significant intervention effects on high symptoms. PRP tended to prevent clinical symptoms relative to CON, \(\chi^2(1, N = 237) = 2.74, p < .10, \text{odds ratio} = .51\ (95\% CI = 0.23, 1.14)\), but not PEP, \(\chi^2(1, N = 243) = 0.01, ns, \text{odds ratio} = .96\ (95\% CI = 0.40, 2.30)\). No difference was found between PEP and CON, \(\chi^2(1, N = 232) = 2.37, ns, \text{odds ratio} = .53\ (95\% CI = 0.24, 1.20)\). Across the follow-up period, 11 (9\%) of 124 children in PRP, 11 (9\%) of 119 in PEP, and 18 (16\%) of 113 controls reported clinical symptoms on the CDRS–R.

School C

Attrition and baseline differences. In School C, there were no between-conditions differences at baseline, and there was no evidence for differential attrition.

Intervention effects. PRP did not reduce depressive symptoms over the follow-up period relative to CON, \(t(585) = 1.29, ns, d = 0.14\ (95\% CI = -0.08, 0.36)\). Contrary to our expectations, PEP significantly reduced depressive symptoms relative to PRP, \(t(594) = 2.08, p < .05, d = 0.24\ (95\% CI = 0.01, 0.47)\). No difference was found between PEP and CON, \(t(592) = -0.86, ns, d = -0.10\ (95\% CI = -0.32, 0.12)\). See Table 2 for means and ANCOVAs by assessment. There were no significant intervention effects on elevated, high, or clinical symptoms (see Figure 2).

Examining the School \(\times\) Condition Interaction

We looked at the pattern of results on other measures (demographics, intervention integrity, attendance) to see whether any of these variables could explain the different effects across schools. Group integrity scores did not differ by school. For most of the variables that differed by school, mean levels for School C were between the means of Schools A and B. One exception was race/ethnicity. Compared with Schools A and B, School C included a higher proportion of Caucasian participants and a lower proportion of participants who indicated their race was “other.” MM ANOVAs predicting follow-up CDI scores revealed no significant Race \(\times\) Condition interaction, however. Thus, none of the variables we examined explained the differential effects across schools.
Discussion

The full sample findings provide little evidence for PRP’s effectiveness or specificity. PRP prevented the onset of elevated symptoms relative to a no-intervention control but not relative to PEP (an intervention that controlled for noncognitive-behavioral therapy specific factors). PRP did not reduce mean levels of depressive symptoms over the follow-up and did not prevent high or clinical levels of symptoms relative to either comparison group. PRP’s effects varied by school, however. In two schools, PRP reduced depressive symptoms over the
follow-up relative to CON and PEP. In contrast, PRP was not effective in the third school.

In Schools A and B, PRP’s effects on depressive symptoms were long lasting. PRP participants reported fewer depressive symptoms than controls through the 30-month follow-up. Effect sizes were moderate at several assessment points. In addition, PRP prevented the onset of elevated symptoms and tended to prevent clinical levels of symptoms relative to no intervention control.

These findings indicate that school teachers and counselors can be trained to implement the intervention effectively and that schools are a viable setting in which to teach depression prevention skills. Given that depression is both very common and undertreated in youths (Hirschfeld et al., 1997), the importance of these findings should be underscored. Skills-based depression prevention programs, like PRP, may be an important addition to the educational goals of our school systems. By offering programs like PRP, schools may play a vital role in preventing a growing national health concern.

An important direction for future research is to determine whether PRP can prevent clinical levels of depressive symptoms when delivered by school teachers and counselors. The present findings provide more evidence for the prevention of mild-to-moderate depressive symptoms than for the prevention of clinical depression. This could reflect several factors, including (a) a limit of the PRP intervention, (b) the fact that most of our group leaders did not have clinical training and thus may have been unprepared to help the children at greatest risk, or (c) the low attendance of many PRP participants. The lack of findings for clinical depression also could be a function of statistical power as only a small proportion of children reported clinical levels of symptoms during the follow-up period.

**PRP’s Inconsistent Effects**

Despite the positive findings in Schools A and B, PRP did not prevent depression in School C. We were unable to identify the source of the differential effects observed in the present study. Group leaders in the three schools received similar training and supervision, and adherence to the PRP and PEP protocols did not vary significantly by school. Children’s baseline depression scores, demographic characteristics, and attendance also did not account for the differential effects observed. We also are not aware of any differences in school services, curriculum, or in events that transpired within schools that could account for the discrepancy. Subtle differences among the schools in program endorsements, students’ needs, or general school climate may have enhanced PRP’s effectiveness in two schools and/or hindered its effectiveness in the third.

The inconsistency in PRP’s effects within the present study mirrors its inconsistent effects across studies. PRP has reduced and prevented depressive symptoms relative to control in several studies (e.g., Gillham, Reivich, Jaycox, & Seligman, 1995; Yu & Seligman, 2002). For example, in the original evaluation of PRP, PRP halved the rate of moderate to severe symptoms. Effect sizes were moderate to large ($d > 0.60$) 12–24 months postintervention (Gillham et al., 1995). However, other studies failed to replicate these findings (e.g., Pattison & Lynd-Stevenson, 2001; Roberts, Kane, Bishop, & Matthews, 2004). A review of PRP research suggests that PRP’s effects are strongest when group leaders are the intervention developers, members of their research team, and graduate students who receive extensive training and supervision from the intervention developers. PRP’s effects on depression are often smaller or nonsignificant when group leaders are school teachers, clinicians, or other researchers not affiliated with the PRP laboratory (Gillham, Hamilton, Freres, Patton, & Gallop, 2006). In other words, as research on PRP moves along the path from efficacy to real world implementation, intervention effects wane or become inconsistent. This pattern corresponds with recent findings for the RAP intervention (Harnett & Dadds, 2004) and with the observation that psychological interventions that yield substantial effects in research settings often produce small or no effects when delivered by community providers (Weisz, Donenberg, Han, & Weiss, 1995). A critical priority for prevention research is the development of effective dissemination strategies.

**Specificity**

The School A and School B findings provide some support for specificity as mean depression scores were lower in PRP than PEP participants over the follow-up. Contrary to our expectations, however, PEP participants reported fewer depressive symptoms over the follow-up than PRP participants in School C. Because PRP was not beneficial relative to CON in School C, it is unclear whether PEP’s superiority reflects the limited value of cognitive-behavioral skills or a more general failure of the PRP intervention in that school.

**Limitations**

The study’s limits include the low recruitment rate, substantial attrition, incomplete assessment of clinical depression, and the inclusion of graduate students as group leaders. In each school, fewer than 25% of students enrolled in the study. Enrollment rates were so much lower than we expected that we revised our original plan to select children with elevated symptoms and offered the program to all children who were not clinically depressed at the start of the study. The children who enrolled reported a wide range of symptoms at baseline, but we did not evaluate the interventions as delivered to most school students in a school setting as in the recent study by Spence, Sheffield, and Donovan (2005). Our participants’ demographic characteristics reflected the communities from which they came, but it is likely that these children (and their parents) differed in other important ways from those who did not sign up for the study. This limits the extent to which we can generalize from our findings.

Fewer than 50% of the original baseline sample completed the 3-year follow-up. Attrition was due to vicsissitudes across schools but did not seem to vary by condition. For example, in School A, because of a change in administration, school staff stopped reminding children to attend assessment sessions. School B has had a high student attrition rate in general, with 10%–20% of students leaving the school each year. We tried to follow participants who moved through the mail, but compliance with these assessments was low, and we were often unable to find forwarding addresses. Attrition and noncompletion of assessments are common problems in longitudinal depression prevention studies. Attrition is often 40% or more in studies with 3 and 4 years of follow-up (e.g., Roberts et al., 2004; Spence et al., 2005). In our current work, we
are implementing several strategies to minimize attrition, including increased incentives for complete assessments and strategies for tracking participants, such as requesting contact information for relatives.

Because of budget and staffing constraints, CDRS–R interviews were only conducted with children with elevated CDI scores (CDI ≥ 13). CDI manual norms suggest that 8% of children who score below this cutpoint may have depressive disorders (Kovacs, 2001). In addition, only about 75% of children who were invited to interviews actually participated. Thus, we probably failed to detect clinical levels of symptoms in several children. During the study, we became concerned about another possible limitation of the triggered interview procedure—that it influenced questionnaire responses. Children may have become aware that elevated questionnaire scores triggered interviews, and children who wanted to avoid interviews may have endorsed few symptoms of depression on the CDI. It is striking that during a developmental phase when depression rates generally increase (Angold, Costello, & Worthman, 1998), depressive symptoms declined in all three conditions in our sample. In the end, the triggered interviews may have limited our ability to find intervention effects on both depression measures.

The inclusion of graduate students as group leaders is also somewhat problematic. The graduate students who led groups were not affiliated with our research laboratory and most planned to pursue careers as school psychologists or school counselors. Still, these graduate students may differ from the school personnel who would deliver an intervention like PRP once disseminated. An important finding is that analyses revealed no significant differences in intervention effects for graduate students versus school teachers and counselors.

Future research may also benefit from more sophisticated analyses of intervention implementation. Our intervention integrity scoring system weighted specific content items equally, but it is likely that some skills are more beneficial or require more explanation and discussion than others. An examination of the relationship between specific content covered and intervention outcome could yield valuable information about the mechanisms of PRP’s effects and help set priorities for group leader training and supervision.

Our integrity coding system was designed to measure leaders’ adherence to the intervention protocol and probably did not fully capture the variability in leaders’ competence, or skill, in intervention delivery (McGlinchey & Dobson, 2003). Although competence and adherence are related, competent leaders may sacrifice adherence at times to allow for more in-depth discussion of group members’ specific problems, review of concepts that seem confusing to group members, or more attention to group process. An assessment of competence could help to explain our differential findings; it may be that PRP group leaders in School C adhered to the manual but did not show true competence.

Clinical Significance

We found little evidence for PRP’s effectiveness or specificity in preventing depression in the full sample. However, in two of the three schools, PRP reduced depressive symptoms over a 2.5-year period and prevented elevated depressive symptoms. This suggests that PRP can have long-lasting effects when delivered by school teachers, counselors, and other providers in a school setting. Although PRP’s effects are inconsistent, we believe that research that evaluates interventions in real world settings will lead to the development of dissemination procedures that enable schools to include powerful depression prevention programs as part of their educational mission.

References


Received December 5, 2005
Revision received May 31, 2006
Accepted July 10, 2006

---

**Members of Underrepresented Groups: Reviewers for Journal Manuscripts Wanted**

If you are interested in reviewing manuscripts for APA journals, the APA Publications and Communications Board would like to invite your participation. Manuscript reviewers are vital to the publications process. As a reviewer, you would gain valuable experience in publishing. The P&C Board is particularly interested in encouraging members of underrepresented groups to participate more in this process.

If you are interested in reviewing manuscripts, please write to the address below. Please note the following important points:

- To be selected as a reviewer, you must have published articles in peer-reviewed journals. The experience of publishing provides a reviewer with the basis for preparing a thorough, objective review.

- To be selected, it is critical to be a regular reader of the five to six empirical journals that are most central to the area or journal for which you would like to review. Current knowledge of recently published research provides a reviewer with the knowledge base to evaluate a new submission within the context of existing research.

- To select the appropriate reviewers for each manuscript, the editor needs detailed information. Please include with your letter your vita. In the letter, please identify which APA journal(s) you are interested in, and describe your area of expertise. Be as specific as possible. For example, “social psychology” is not sufficient—you would need to specify “social cognition” or “attitude change” as well.

- Reviewing a manuscript takes time (1–4 hours per manuscript reviewed). If you are selected to review a manuscript, be prepared to invest the necessary time to evaluate the manuscript thoroughly.

Write to Journals Office, American Psychological Association, 750 First Street, NE, Washington, DC 20002-4242.