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A Cluster Randomized Trial of Adding Peer Specialists To Intensive Case Management Teams in the Veterans Health Administration

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Abstract

Use of Peer Specialists (PSs)—individuals with serious mental illness who use their experiences to help others with serious mental illness—is increasing. However, their impact on patient outcomes has not been demonstrated definitively. This cluster randomized, controlled trial within the Veterans Health Administration compared patients served by three intensive case management teams that each deployed two PSs for one year, to the patients of three similar teams without PSs (Usual Care). All patients (PS group=149, Usual Care=133) had substantial psychiatric inpatient histories and a primary Axis I psychiatric disorder. Before and after the year PSs worked, patients were surveyed on their recovery, quality of life, activation (health self-management efficacy), interpersonal relations, and symptoms. Patients in the PS group improved significantly more ($z=2.00$, $df=1$, $p=0.05$) than those receiving Usual Care on activation. There were no other significant differences. PSs helped patients become more active in treatment, which can promote recovery.

INTRODUCTION

As stated in the 2003 U.S. President's New Freedom Commission on Mental Health, Peer Specialists (PSs) are widely seen as important in recovery-oriented mental health care¹. PSs are individuals with serious mental illness (SMI) who draw upon lived experiences to provide services to others with SMI in clinical settings². Although research generally supports PSs' inclusion in clinical settings, implementation has been uneven and outcomes have not been demonstrated definitively. Despite this, the use of PSs is growing. Twenty-two states now have Medicaid reimbursement for PSs. The Veterans Health Administration (VHA) has hired about 250 Peer Specialists (PSs) across most of its medical centers and plans to hire 800 more over the next two years per an executive order by the Obama administration. However, impact within the VHA has not yet been evaluated. The current study is the first within VHA to evaluate PSs' impact on recovery outcomes for veterans with SMI.

The current emphasis on recovery in the VHA and the larger service system necessitate a broad range of outcomes when examining program impact beyond symptom stabilization and maintenance. An increasing number of studies are emphasizing the role of patient activation in SMI recovery³⁻⁶. Patient activation refers to the knowledge, skill, confidence, and attitudes for managing health and treatment⁷. Several studies in medical domains have found that individuals with higher activation are healthier, report a better quality of life, are more satisfied with treatment, and engage in more healthcare practices⁷⁻¹⁰. As recovery involves an active role for people with SMI in developing a life beyond the illness¹¹, activation can be a central construct in assessing the recovery impact of a new service like incorporating PSs.

PSs are in a unique position to help, consistent with social modeling theory, which states that similar others might have the most influence on behavior change¹². They offer systems knowledge and "street smarts," and teach successful coping strategies¹³ that can enhance the use of illness self-management strategies. Often PSs increase patients' involvement in treatment¹⁴ because they are able to empathize, access social services, appreciate clients'

strengths, be tolerant, flexible, patient, and persistent, and be aware of and responsive to clients' desires and goals¹⁵⁻¹⁷, even among persons who are homeless and have co-occurring psychiatric and substance disorders¹⁸. As a result, PSs have been successful in improving patient satisfaction with traditional mental health services¹⁹. PSs have re-established their own social networks, and can help patients do the same, often through mutual support activities. Finally, PSs actively model the possibility of recovery, addressing the loss of hope often evidenced by those with SMI.

In addition to direct work with patients, PSs can further impact the care process by moving services toward a recovery orientation in several ways²⁰. They often serve as an unofficial liaison to the non-consumer staff, interpreting, and in some cases mediating, between other staff and patients. They can challenge unacknowledged stigma²¹ and emphasize community integration over a singular focus on symptom stabilization^{22,23}. Thus, PSs aim to change how patients engage with the community (i.e., become less isolated) and with providers (i.e., becoming more active in his/her care).

Seventeen studies, all outside VHA, have tested the use of PSs in traditional clinical settings—five RCTs, eight quasi-experimental studies (comparison groups, not randomized), and four descriptive or correlational studies—typically comparing patients over time from different types of case management teams with or without PSs. Eleven (two RCTs, six quasi-experimental, and three correlational) showed some positive benefit to patients served by PSs compared to patients who were not, including less inpatient use²⁴⁻²⁸, better treatment engagement^{16,29,30}; greater satisfaction with life¹⁶, greater quality of life²⁵, greater hopefulness³¹, better social functioning²⁵, improved self-reported recovery^{23,32}, fewer days homeless³², and fewer problems and needs^{16,29,32}. The other six studies did not yield differential improvement in a variety of clinical outcomes over time³³⁻³⁸. One study, a quasi-experimental trial comparing patients from four Assertive Community Treatment teams with PSs to 16 teams without, did show that the presence of a PS was associated with an increase in psychiatric hospitalization days³².

As a group, these studies do not show a clear advantage for incorporating PSs. One reason could be that most studies have either focused on symptoms and hospitalizations, rather than on the full range of recovery domains. In addition, prior studies may have not addressed common implementation barriers of employing PSs such as ill-defined PS roles and non-PS resistance, which have been well documented within VHA^{2,39}. To overcome these difficulties, we conducted a randomized trial -- PEER (PEers Enhancing Recovery) -- using organizational implementation strategies to incorporate PSs into traditional VHA case management teams and test the impact of PSs on a broad range of recovery outcomes. Six full-time PSs were randomly assigned to work for a year on three of six VHA Assertive Community Treatment teams, an evidence-based, intensive case management model⁴⁰, while the other three teams delivered Usual Care. In VHA, these teams are called MHICM (Mental Health Intensive Case Management) and are expected to adhere to Assertive Community Treatment practice (e.g., low caseloads, majority of contact in community). In PEER, it was hypothesized that the involvement of PSs would lead to greater gains—at the individual patient level—in recovery, quality of life, patient activation, and to a lesser extent, symptoms.

METHODS

Study Sites and Assignment to Study Condition

PEER was a cluster randomized controlled trial⁴¹ on MHICM teams in six medium to large cities in Southwestern US. Using a random sequence generator by the study PI, three MHICM teams were assigned to receive two PSs each. The PSs were then hired after site randomization, which was done (versus patient randomization) to minimize potential contamination. To receive MHICM services in VHA, veterans must have at least 30 psychiatric inpatient days or three psychiatric admissions in the past year and have a primary Axis I psychiatric disorder. MHICM teams provide intensive, flexible community support to reduce psychiatric symptoms, substance abuse, and inpatient hospitalization; improve community adjustment and quality of life; enhance satisfaction with services; and reduce treatment costs⁴². The randomization yielded two groups from which patients were recruited: teams A (n=135), B (n=53), and C (n=64) were randomly assigned receive PSs and teams D (n=89), E (n=75), F (n=52) were assigned to Usual Care. It was not possible to blind either participating patients or data collectors to study condition because VHA required different informed consent forms in PS and Usual Care teams, and patients knew when their team had a PS or not.

PS Intervention on the MHICM Teams

PEER used an organizational change framework—the Simpson Transfer Model⁴³—to guide the implementation of PSs through four stages: Exposure (e.g., training in PSs), Adoption (deciding to hire PSs), Implementation (deploying PSs), and Practice (monitoring and refining PS services). Research staff collaborated with MHICM staff at each stage and assisted the MHICM teams to hire, train, and supervise the PSs with ongoing consultation (See ³⁹ for more details of PEER’s implementation process using the Simpson Transfer Model).

Consistent with previous PS descriptions^{44,45}, PSs developed strong relationships with veterans and conducted all types of case management duties including delivering medication, accompanying veterans to appointments, developing recovery plans, meeting with veterans individually, leading and co-leading groups, engaging veterans into services, and helping other MHICM staff, all while drawing upon their own lived experiences. They participated fully in all MHICM activities including team meetings, meetings between veterans and psychiatrists, team-organized events and outings, and charting in the medical record. The PSs “floated”—i.e., they did not have their own caseload—because the teams have case managers assigned administrative responsibility for each patient and that they wanted the PSs to work with multiple veterans more readily.

Training and supervision—Prior to the providing services, PSs attended a 30-hour training conducted by the Peer-to-Peer Resource Center of the Depression and Bipolar Support Alliance on recovery, basic counseling skills, and psychosocial rehabilitation. The training and certification examination is based on the Georgia model of Medicaid-reimbursed peer specialists⁴⁶. PSs also received a two-day training in Illness Management and Recovery (IMR⁴⁷) by staff from the ACT Center of Indiana. IMR is an evidence-based,

manualized approach to helping consumers with SMI learn information and skills to better manage illness and achieve goals. Illness Management and Recovery has been used successfully by PSs on ACT teams previously^{23,28} and it was believed this could provide some structure for their work. Finally, MHICM staff provided training on MHICM rules such as emergency procedures and clinical charting. PSs were supervised weekly by both an “internal” supervisor (MHICM staff) and an “external” supervisor (PEER principal investigator, a licensed clinical psychologist). Research staff also met monthly with each MHICM team leader and PS supervisor to troubleshoot any implementation issues.

Although all veterans in the PS group had access to the PSs, medical records showed that 43% of enrolled veterans had zero PS contacts, 11% had one contact, 10% had 2–4 contacts, 18% had 5–12 contacts, and 18% had greater than 13 contacts. However, the number of contacts was underestimated because PSs sometimes recorded contacts in the medical record incorrectly or not at all. Therefore, the analyses focused on the intent to treat approach—comparing all veterans from both groups—which is nonetheless consistent with how PSs are typically used (i.e., float and provide a range of services to some and not others) and the hypothesis that PSs can impact the recovery-orientation of a whole team.

Further, the impact of PSs was evaluated over and above the outpatient services patients were receiving in MHICM and the VHA more broadly. According to VHA administrative data on Behavioral Health outpatient service contacts tracked through a central electronic medical record (includes psychiatric, social work, substance use, and rehabilitation services), patients in the PS and Usual Care groups were receiving similar numbers of contacts (per month) in the year prior to study enrollment ($M_{PS}=1.61$, $SD=2.44$ in the PS group vs. $M_{UC}=1.73$, $SD=2.45$ in the Usual Care group, $p=0.52$). As could be expected with a case management service like MHICM in which patients join the team in a more acute state and then generally improve over time, the number of contacts a year after study enrollment decreased similarly in both groups to $M_{PS}=1.54$, $SD=2.67$ vs. $M_{UC}=1.26$, $SD=2.00$ (within group test, $p=0.03$, interaction, $p=0.12$).

Data Collection

Procedures—Start dates for PSs varied during the project because of different rates of hiring PSs, and patient data collection extended between October 2006 and May of 2011. The inclusion criterion was simply being a current patient on one of the six teams. All 468 veterans ($N_{PS}=252$ in the PS group, $N_{UC}=216$ in the Usual Care group) who were active patients at the time from the six MHICM teams were eligible. All participants gave written informed consent and scored 100% on a 10-item true/false quiz on the details of their consent⁴⁸. Participants in the PS group also signed separate consent forms to potentially receive services from a PS. Trained interviewers administered the patient outcome measures at baseline by reading the questions aloud and recording the patient’s responses. The one-year follow-up assessment repeated the same measures used at baseline. The study was approved by each of the participating VHA medical centers’ local IRBs.

Measures

Demographic variables—These variables were gathered as part of the interview process and included age, gender, race, ethnicity, living situation, education level, and age at which the individual began taking psychiatric medication or was first hospitalized for psychiatric reasons.

Patient outcomes—Perceptions of the recovery orientation of the program were assessed with the Recovery Self-Assessment (RSA), a 36 item survey that assesses domains of recovery-orientated practice (e.g., focus on life goals, involvement of patients in their own care)⁴⁹. The RSA has high internal consistency and is thought to represent a more recovery-oriented or recovery-supportive environment⁵⁰.

Two measures of self-reported recovery were included. The Mental Health Recovery Measure (MHRM) is a 30-item, 5-point behaviorally-anchored self-report measure based upon recovery experiences of persons with psychiatric disabilities^{51,52}. The MHRM total score has good validity, correlating strongly with the Empowerment Scale⁵³ and Community Living Skills Scales⁵⁴, yet assessing unique aspects of recovery. The Illness Management and Recovery Scale⁵⁵ (IMR) has 15 items (rated on 5-point behaviorally anchored scales) that assess progress toward goals, knowledge about mental illness, involvement with significant others and self-help, time in structured roles, impairment in functioning, symptom distress and coping, relapse prevention and hospitalizations, use of medications, and alcohol and drug use. A total IMR score is made of the mean of the items and has demonstrated good internal consistency, stability (test-retest after two weeks), and convergent validity, correlating with the Recovery Assessment Scale⁵⁶ and the Colorado Symptom Index⁵⁷.

Subjective ratings of overall quality of life and the quality of social relationships, daily life, and family interactions was assessed using a combination of selected scales from the Quality of Life Instrument-Brief Version (QOLI^{58–60}), which been used extensively with a wide range of populations including those who are homeless, have a dual diagnosis, and are ethnic minorities. Because of low internal consistencies of subscales in our sample, a factor analysis was conducted which indicated that a larger scale that included the items from the overall quality of life, social relationships, daily life, and family interactions scales would be more reliable.

The mental health version of the Patient Activation Measure⁴ (PAM) is a single 13-item scale designed to assess patient's knowledge, skill, and confidence in health self-management. Respondents endorse items (e.g., "I know what each of my prescribed medications do") on a scale from 1 ("disagree strongly") to 4 ("agree strongly"). This version has similar psychometric properties as the original 13-item PAM and correlates with related constructs in other samples of people with SMI^{4,5}. The BASIS-R is a brief yet comprehensive instrument assessing a range of psychiatric symptoms and problems. It is valid and reliable in both inpatient and outpatient settings in populations with SMI⁶¹. In addition to the total score, the BASIS-R scales that were used were Interpersonal Relationships (5 items), Depression (6 items), and Psychosis (6 items). All items have five response options ranging from 0 to 4, with higher scores indicating more problems. The total

BASIS-R assesses an overall level of impairment, while the individual subscales assess more targeted domains.

Data Analyses

First, demographic variables (e.g., age, gender, race, Hispanic) were compared between the PS and Usual Care groups with chi-squares for categorical data and analyses of variance for age. Second, baseline scores of all patient outcome measures were compared between the PS and Usual Care groups with regression models that corrected for nesting of subjects within sites nested within treatment. Third, comparisons between the PS and Usual Care groups were completed with a series of regressions testing the interaction of group (PS vs Usual Care) and time (baseline vs follow-up) for each of the nine patient outcome measures. Like at baseline, measures were analyzed with mixed effect hierarchical regressions which accounted for the nesting of site under treatment, subjects within sites, and subjects over time. Models included site, age, race, ethnicity and the BASIS-R Total score, which was used as a marker for severity of mental illness at both assessment points. Models of the BASIS-R Total score and its subscales did not include the BASIS-R Total score as a covariate. Finally, an additional set of analyses compared the time effect separately for the PS and Usual Care groups for each patient outcome measure. These analyses were also completed with mixed effect regressions that accounted for nesting of subjects within sites, demographics, and the BASIS-R total score (for the non-BASIS outcomes). Since one PS site started earlier, those patients were not included in the analyses of the RSA, PAM, and QOLI. Due to the 1:1 nature of the data collection, there was no missing data. Stata (v12, 2012 College Station, TX) was used for all analyses.

RESULTS

Enrollment and response rates

Of the 468 current patients, 62 were deemed ineligible because MHICM staff believed they were inappropriate (e.g., too ill at the time to consent, $N_{PS}=26$, $N_{UC}=16$) or died before enrollment ($N_{PS}=8$, $N_{UC}=12$). Of 406 eligible veterans ($N_{PS}=218$, $N_{UC}=188$), 72 refused ($N_{PS}=40$, $N_{UC}=32$) and 49 either could not be located, did not speak English, or were in another study ($N_{PS}=29$, $N_{UC}=23$), leaving 282 enrolled veterans or 69% of the total patient census of the teams: 149 in the PS group (68%) and 133 in the Usual Care group (71%).

Follow-up assessment rates across the three PS teams ranged from 71–95% (overall 81%), resulting from deaths ($n=10$), moving ($n=7$), and loss to follow-up ($n=10$). The follow-up assessment rates across the three Usual Care teams ranged from 80 – 88% (overall 82%), resulting from deaths ($n=3$), moving ($n=4$), and loss to follow-up ($n=10$). The follow-up rates of the two groups were similar (chi-sq=1.52, $df=1$, $p=0.22$), leaving 122 in the PS group and 116 in the Usual Care group. Comparison of the demographic variables and available patient outcomes at baseline between participants that completed the follow-up interviews ($n=238$) and those that did not ($n=44$) did not differ, with one exception. Females were more likely to complete the interviews at both times than were males (100% vs. 83%, $p=0.02$).

Sample characteristics

As shown in Table 1, both groups were mostly male (90% in the PS group, 86% in the Usual Care group) and older (average age of PS group = 54.6 SD=9.19, Usual Care group = 51.9, SD=11.13). The majority of the participants were White, although the PS group had more African-Americans (27%) than the Usual Care group (11%). Both groups were similar in that about a third had a high school education or less (PS group= 34%, Usual Care group=31%) and most lived alone (PS group=84%, Usual Care group=76%). Most participants in both groups began psychiatric treatment in their late 20s, although the PS group received medications at a younger age on average (M=26.9, SD=8.0) than participants in the Usual Care group (M=29.5, SD=11.1). At baseline the PS group had significantly higher (better) MHRM scores (85 vs. 78, $p=0.005$) and lower (better) BASIS-R Depression (26 vs 29, $p=0.02$) and total BASIS Scores (1.28 vs. 1.55, $p=0.09$) (see Table 2).

At baseline, both PS and Usual Care groups had worse Psychosis and Depression subscale ratings than a national sample of 2,656 psychiatric inpatients and 3,222 outpatients⁶². Both groups had Interpersonal Relationship ratings that were worse than the outpatient sample, but better than the inpatient sample. Baseline scores on the PAM indicate that both groups were in the lowest of four possible levels of activation, suggesting that they tend to be more passive recipients of treatment⁶³.

Leading up to the start of the study, all teams were similarly adherent to Assertive Community Treatment fidelity as documented by the FY 2005 MHICM Monitoring data collected by the Northeast Program Evaluation Center⁴². For example, all teams had provider caseload sizes below the limit of 15 (ranged from 10 to 14) and met the minimum criteria of at least 60% of contact made in the community (ranged from 84–97%). Also, all teams had moderate to high fidelity (ranged 3.8 to 4.1, out of a high of 5) as measured by the DACTS, the established scale to measure fidelity of this type of case management team^{64,65}. The teams' DACTS data showed that they were similarly adherent during the study period.

Patient outcomes

As shown in Table 2, analyses of the scores indicated that patients in the PS group improved more on the PAM scale than those in the Usual Care group, although the absolute value still reflects a low level of activation. There were no significant differences between treatment groups on the other outcome measures. The additional analyses by individual group showed several significant differences over time. The Usual Care group improved over time on the IMR scale, BASIS-R Total score, the BASIS-R Interpersonal scale, the BASIS-R Depression scale, and the Quality of Life scale. The PS group improved over time on the BASIS-R Interpersonal scale.

At the individual scale level for baseline and follow-up measures, the intrapersonal dependency varied from 0–0.83; with the median value equaling 0.59 for the PS group participants and 0.56 for the Usual Care participants. Across the two groups the lowest rhos occurred on the Interpersonal Relations subscale of the BASIS-R and the overall measure of Quality of Life. At the site level, the inter-person dependency varied from 0–0.04 with a median value of 0.017 with the highest level for the MHRM scale.

DISCUSSION

As the first study of PSs in VHA, the PEER project sought to evaluate the use of PSs using a comprehensive organizational implementation strategy and a wide range of patient outcomes. One of the study's hypotheses about impacts on patient activation was supported, while the other hypotheses were not. While receiving the same level of outpatient treatment, individuals receiving treatment on teams with PSs for a year improved more on patient activation than those in the Usual Care group. Other measures of recovery, quality of life, and symptoms did not differentially change between the PS and Usual Care groups. Using a sample that was typical of a VHA population, this study adds to the literature examining PSs on Assertive Community Treatment teams, and the activation findings are consistent with previous studies that also showed PSs improved treatment participation^{29,30}. These findings are also consistent with a national survey of PS duties, in which promoting active participation was among the most commonly provided type of support⁴⁵.

The impact PSs can have on patient activation could improve how patients engage in their mental health and medical care. Even a one point improvement on the PAM (about what was found in this study) has been found to predict improved health care utilization⁶⁶. Yet, activation still may be lower in those with SMI as those with SMI receive less appropriate medical care and have worse health outcomes⁶⁷⁻⁶⁹. Thus, in this study, although there was a small improvement in activation, the absolute value still places the PS-assigned patients at a low level, which could be why the other outcome measures also did not show improvement. Other studies of patient activation suggest that while those with SMI may be ready and capable to engage with their providers, providers and patients may need assistance⁵, which PSs could provide. However, this study suggests that more PS services may be needed than was provided to elevate low active patients to a higher level of activation.

Limitations of the study should be noted. First, the study was a cluster randomized trial as whole teams were assigned to the PS or Usual Care groups. The impact of the individual teams may have exerted influence on the findings, although the mixed effect hierarchical regressions helped address this issue. Second, the analyses presented were intent-to-treat analyses, but about half the veterans in the PS group did not receive any PS services. Thus, these analyses may be a conservative underestimate of the true impact of the PSs. Although attempts were made to track all PS contacts, a combination of factors (PS unfamiliarity with charting, the electronic medical record not having codes for PS services) made the tracking unreliable, underestimating the total number of PS contacts, precluding analyses by PS contacts. Third, the study involved a small number of PSs (n=6). Although the findings are suggestive of impacts in the VHA, it still represents a small sample of the more than 250 PSs VHA-wide. Fourth, data collectors were not blind to condition, however, the survey instruments were all structured, mitigating somewhat this limitation. Fifth, having the PSs serve on MHICM teams may have also limited the impact that could have been detected over and above VHA's most intensive outpatient service. In addition, PSs' need to "fit in" may have led them to abandon their peer role, which would have reduced the differences between the two groups. However, some studies have shown that PSs on ACT outside of the VA have had an impact and may be an important component of helping ensure a recovery-oriented approach^{23,70,71}. Sixth, one of the teams had a much larger group of patients than the others,

which may have impacted the results. Finally, it is possible that patients improved in domains unmeasured here, for example, Hope, which is a construct that has been associated with peer support in other studies³¹. Future studies that closely track the amount and nature of PS contacts, vary the settings in which they work, assess other constructs such as Hope, and involve larger samples of PSs could help address the limitations of the current study.

Implications for Behavioral Health

The hiring of PSs is rapidly increasing, especially in VHA. Clinical managers and policy makers often struggle with how to best utilize PSs. This study of PSs, the first in VHA, found significantly greater improvements on patient activation among patients who were served by intensive case management teams with PSs, although their overall level of activation remained low. This finding is consistent with other research on PSs. Although PSs can play a variety of roles, clinical managers who hire PSs may want to ensure that improving patient activation is included in their range of duties.

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Table 1Description of the study subjects at Baseline¹

| Demographic variables | PS (n=122) | Usual Care (n=116) | PS vs Usual Care |
|--|------------------|--------------------|-----------------------------|
| | N (%) | N (%) | X ² (df), p |
| Race | | | |
| African-American non-Hispanic | 32 (26%) | 12 (10%) | 9.71(4), 0.046 ² |
| White non-Hispanic | 61 (50%) | 68 (59%) | |
| Biracial non-Hispanic | 8 (7%) | 10 (9%) | |
| African-American Hispanic | 1 (1%) | 1 (1%) | |
| White Hispanic | 15 (12%) | 19 (16%) | |
| No Race/ethnicity information | 5 (4%) | 6 (5%) | |
| Hispanic | 16 (13%) | 20 (17%) | 0.79 (1), .374 |
| Male | 110 (90%) | 100 (87%) | 0.60 (1), .437 |
| Education | | | |
| <=HS | 42 (36%) | 36 (34%) | 0.91 (2), .634 |
| Some post high school training | 61 (57%) | 60 (52%) | |
| College degree or more | 14 (9%) | 9 (9%) | |
| Current Marital Status | | | |
| Never Married | 49 (42%) | 39 (37%) | 0.97(2), .616 |
| Currently Married (includes those with previous divorces & current partners) | 24 (21%) | 27 (26%) | |
| Divorced or separated | 44 (38%) | 39 (37%) | |
| Lives Alone | 102 (87%) | 88 (84%) | 0.51 (1), .475 |
| | Mean (sd) | Mean (sd) | t-test (df), p |
| Age | 54.59 (9.19) | 51.89 (11.13) | 2.05 (236), p=.042 |
| Age first meds | 26.89 (8.04) | 29.49 (10.28) | 2.09 (215), p=.004 |
| Age first hospitalization | 27.00 (8.65) | 30.20 (10.76) | 2.40(211), p=.017 |

¹Since only 4 people reported being employed (2 in each condition), it was too few to include in the table.

²This significant difference is due to the over representation of African-American subjects in the PS group. With African-American subjects excluded, X² (df), p = 0.44(3), p=0.98.

Table 2

Measures at Baseline and Follow-up Interviews

| Measures | PS Mean (sd) | | Usual Care Mean (sd) | Differences at Baseline ¹ | | Statistical results from mixed model analyses ² | | |
|---|--------------------|--------------------|----------------------|--------------------------------------|-------------|--|-------------|---------------------|
| | Total (n=122) | Total (n=116) | | t-test, p value | Interaction | Treatment | Pre-post | |
| Measures available at all sites | | | | | | | | |
| Mental Health Recovery Measure | | | | | | | | |
| Baseline | 85.00 (17.88) | 78.30 (18.79) | | 2.59, 0.01 | | -0.58, 0.56 | 0.36, 0.72 | 0.55, 0.58 |
| Post | 85.66 (15.90) | 80.52 (16.54) | | | | | | |
| Change | 0.52 | 2.40 | | | | | | |
| Illness Management and Recovery Scale | | | | | | | | |
| Baseline | 3.51 (.58) | 3.39 (.56) | | 1.63, 0.10 | | -0.16, 0.87 | 0.38, 0.70 | 0.71, 0.48 |
| Post | 3.56 (.56) | 3.48 (.54) | | | | | | |
| Change | 0.06 | 0.08 ³ | | | | | | |
| BASIS-R Total score | | | | | | | | |
| Baseline | 26.15 (14.28) | 29.34 (14.24) | | -2.38, 0.02 | | 0.67, 0.50 | -0.99, 0.32 | -2.40, 0.016 |
| Post | 24.57 (14.05) | 26.76 (15.11) | | | | | | |
| Change | -1.54 | -2.72 ³ | | | | | | |
| BASIS-R, Interpersonal Relationships scale | | | | | | | | |
| Baseline | 1.67 (.92) | 1.63 (.87) | | 0.36, 0.72 | | -0.89, 0.38 | 0.21, 0.84 | -1.00, 0.32 |
| Post | 1.48 (.85) | 1.54 (.96) | | | | | | |
| Change | -1.17 ³ | -1.10 ³ | | | | | | |
| BASIS-R, Depression scale | | | | | | | | |
| Baseline | 1.28 (.89) | 1.55 (.88) | | 2.37, 0.02 | | 1.34, 0.18 | -1.24, 0.22 | -2.05, 0.04 |
| Post | 1.27 (.91) | 1.41 (.92) | | | | | | |
| Change | -.01 | -.14 ³ | | | | | | |
| BASIS-R, Psychosis scale | | | | | | | | |
| Baseline | 1.03 (.95) | 1.15 (.91) | | -0.96, 0.34 | | 0.28, 0.52 | -0.76, 0.45 | -0.76, 0.45 |
| Post | 1.01 (.99) | 1.09 (.97) | | | | | | |
| Change | -.03 | -.08 | | | | | | |

| Measures | PS Mean (sd) Total (n=122) | Usual Care Mean (sd) Total (n=116) | Differences at Baseline ¹ t-test, p value | Statistical results from mixed model analyses ² | | |
|--|-------------------------------|---------------------------------------|---|--|--------------|-------------|
| | | | | Interaction | Treatment | Pre-post |
| Patient Activation Measure⁴ | | | | | | |
| Baseline | 39.31 (5.41) | 39.09 (5.37) | 0.49, 0.63 | 2.00, 0.05 | -1.98, 0.05 | -1.32, 0.19 |
| Post | 40.34 (5.50) | 38.76 (5.01) | | | | |
| Change | 0.85 | -0.24 | | | | |
| Recovery Self-Assessment⁴ | | | | | | |
| Baseline | 3.86 (.53) | 3.81 (.61) | 0.66, 0.51 | 0.56, 0.58 | -0.89, .38 | -0.14, 0.89 |
| Post | 3.93 (.61) | 3.83 (.52) | | | | |
| Change | 0.04 | 0.01 | | | | |
| Quality of Life-Brief, Subjective overall quality of life⁴ | | | | | | |
| Baseline | 4.64 (1.04) | 4.54 (1.03) | 1.27, 0.21 | 0.03, 0.98 | -2.76, 0.006 | 1.01, 0.31 |
| Post | 4.72 (1.04) | 4.71 (.90) | | | | |
| Change | -0.08 | 0.20 ³ | | | | |

¹Independent group t-tests comparing participants in the PS and Usual Care groups at baseline.

²Z-tests of whether the regression co-efficient for effects are significantly different from 0, which included demographic variables, site of care and BASIS-R total score as covariates.

³This change is significantly different from 0, indicating that there was a change in means over time within this treatment condition. This test does not compare the change between groups, only the change in the group identified.

⁴PS group n=66

* Analyses of the BASIS-R total and subscale did not include the BASIS-R total score as a covariate.