Explanatory models of illness in schizophrenia: comparison of four ethnic groups

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Background Explanatory models of illness may differ between ethnic groups and influence treatment satisfaction and compliance.

Aims To compare explanatory models among people with schizophrenia from four cultural backgrounds and explore their relationship with clinical and psychological characteristics.

Method Explanatory models, insight, treatment compliance, health locus of control, quality of life, treatment satisfaction, therapeutic relationships and symptomatology were assessed in UK Whites and Bangladeshis, African–Caribbeans and West Africans.

Results When biological and supernatural causes of illness were compared, Whites cited biological causes more frequently than the three non-White groups, who cited supernatural causes more frequently. When biological and social causes were compared, Whites cited biological causes more frequently than African–Caribbeans and Bangladeshis, who cited social causes more frequently. A biological explanatory model was related to enhanced treatment satisfaction and therapeutic relationships but not treatment compliance.

Conclusions Explanatory models of illness contribute to patient satisfaction with treatment and relationships with clinicians.

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Explanatory models of illness encompass a person’s ideas about the nature of their problem, its cause, severity, prognosis and treatment preferences (Kleinman, 1980). Dissonance between patients’ and professionals’ explanatory models may affect help-seeking behaviour (MacCarthy, 1988), treatment compliance (Foulks et al, 1986), satisfaction (Callan & Littlewood, 1998) and culturally sensitive clinical practice (Bhui & Bhugra, 2002). In the context of cultural background, there is interest in explaining the higher rates of psychotic illness, involuntary admissions and dissatisfaction with services among first- and second-generation immigrants reported in the UK and The Netherlands (e.g. Harrison et al, 1997; Parkman et al, 1997; Bhugra & Bhui, 1998; Selten et al, 2001; Bhui et al, 2003). However, the generalisability of research has been limited by isolated samples in different countries (e.g. Weiss et al, 1986), lack of standardised instruments (e.g. Callan & Littlewood, 1998) and heterogeneous samples (e.g. Lloyd et al, 1998).

METHOD

Aims

(a) To use a standardised measure to assess qualitatively the explanatory models among patients with schizophrenia in a local White British group and three second-generation ethnic groups living in similar socio-economic conditions in East London.

(b) To compare quantitatively the explanatory models across the four groups.

(c) To investigate the association between explanatory models and clinical and psychological characteristics in all of the patients.

The four groups were of African–Caribbean, Bangladeshi, West African and UK White origin. Second-generation patients were selected because: there are consistent differences in service use and outcome among this group; the findings would be less confounded by individual histories of migration and schooling because all interviewees have grown up in the same educational system and background culture; all interviews could be conducted in English because this is most relevant for clinical practice; and the material could be elicited in English and analysed without considering specific connotations in different languages. The inclusion criteria were selected to reduce the heterogeneity of the groups and increase the possibility of detecting specific differences despite relatively small sample sizes for each ethnic group.

Sample

The sample consisted of four groups of outpatients from four distinct cultural backgrounds aged between 18 and 48 years who met DSM–IV (American Psychiatric Association, 1994) criteria for a diagnosis of schizophrenia, were in the care of a community mental health team, had no known organic impairment and had no significant formal thought disorder as assessed on the Brief Psychiatric Rating Scale (BPRS; Overall & Gorham, 1962). The four groups were recruited in East London and the three groups of non-UK origin were all second-generation immigrants. Patients were identified as second generation if they were either born in the UK with both parents of the specified origin or moved to the UK before the age of 12 years. Participants were included when clinical and self-report of ethnicity were in full agreement. In accordance with the central limit theorem, 30 participants per group were recruited so that the mean could be approximated closely by the normal distribution (Mukhopadhyay, 2000). Samples A, B and C comprised individuals of second-generation African–Caribbean, Bangladeshi and West African (from Nigeria or Ghana) ethnic origin. Sample D consisted of White English participants, all born in the UK.

Participants were recruited from secondary mental health services (i.e. four community mental health teams, a day hospital and a psychology service). Service managers, psychiatrists and support workers were consulted to identify those clients meeting the inclusion criteria. Diagnosis was clinical rather than derived from case notes. On clinicians’ recommendations, patients were contacted by letter, telephone or in person at clinics according to the
individual patient’s preferences and circumstances. All participants provided written informed consent to take part in the study.

**Measures**

A modified version of the Short Explanatory Model Interview (SEMI; Lloyd et al., 1998) enabled a qualitative assessment of patients’ conceptualisations of their illness within a structured framework that allowed for eventual quantification of data. Questions originally developed for use in primary care that were not relevant for people with chronic illness were omitted from the scale. The modified version was divided into three sections on the nature of the problem and its consequences, aetiology and treatment. Each patient interview was audio-recorded and transcribed.

Psychopathology was observer rated using the 24-item version of the BPRS (Overall & Gorham, 1962). Patient insight into their illness was observer rated using the Schedule for Assessment of Insight into Psychosis (SAI; David et al., 1992), which has three sub-scales: treatment compliance, awareness of illness and relabelling of symptoms. Ratings were based on information from the patient and additional information from the keyworker. Quality of life was self-rated with the Manchester Short Assessment of Quality of Life (MANSA; Priebe et al., 1999). A modified version of the Helping Alliance Scale (HAS; Priebe & Gruyters, 1993) was self-rated by the patient who assessed their primary therapeutic relationship, typically with their keyworker. Treatment satisfaction was self-rated using the Patient Care Satisfaction Questionnaire (PCQS; Barker et al., 1998) and was used in data analyses. For the four causal categories, ratings were collapsed to form four categories: biological (physical illness/substance misuse); social (interpersonal problems/stress/negative childhood events/personality); supernatural (supernatural); and non-specific (do not know/mental illness/other). Although the heterogeneity of responses in the non-specific category renders it clinically meaningless, it is retained for the sake of completeness of the data. A power calculation in detecting differences between the groups of explanatory models and treatment preferences in categorical analyses indicated that: a large effect size (i.e. a difference in proportions when translated into a fourfold product moment r ranges between 0.37 and 0.39) would be detected with 87% power on a two-tailed significance test where $P = 0.05$; a medium effect size (when translated $r=0.25$) would be detected with 49% power; and a small effect size (when translated $r=0.10$) with only 12% power (Cohen, 1988). Interrater agreement in classifying the raw data into categories was calculated using the $k$ statistic. The four causal categories were compared on socio-demographic, clinical and psychological variables. The therapeutic relationship scores were recorded into a score below or above the midpoint on the scale, indicating a generally positive or a generally negative therapeutic relationship. Differences between the groups according to explanatory model and ethnicity were analysed using analysis of variance and $\chi^2$ or Fisher’s exact test. Where multiple tests were conducted, the Bonferroni-adjusted significance levels are reported.

**RESULTS**

A total of 180 patients were identified by clinicians as meeting the inclusion criteria, 131 were sent letters informing them about the study and requesting their participation. Of these, 72 were interviewed, 10 did not consent and 49 did not respond at all. Fourteen patients were contacted by telephone: thirteen were interviewed and one did not consent. Thirty-five patients were approached at clinics, of whom thirty-four were interviewed and one did not consent. Of those patients who were approached by telephone or in person, the non-consent rate was 8.4%. The total number of people interviewed, all in face-to-face interviews, represented 66% of the potential sample.

Thirty African–Caribbeans, Bangladeshis and UK Whites and 29 West Africans were interviewed. Their socio-demographic and clinical history characteristics are presented in Table 1. Bangladeshis were younger and had a shorter duration of illness than African–Caribbeans and UK Whites. They were also less likely to live alone than West Africans.

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Socio-demographic characteristics of the four groups</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>A</td>
</tr>
<tr>
<td>Age: mean (s.d.)</td>
<td>35.7 (6.5)</td>
</tr>
<tr>
<td>Male: %</td>
<td>57</td>
</tr>
<tr>
<td>Length of illness, years: mean (s.d.)</td>
<td>12.9 (8.5)</td>
</tr>
<tr>
<td>Admissions, n: mean (s.d.)</td>
<td>3.7 (2.4)</td>
</tr>
<tr>
<td>Involuntary admissions, n: mean (s.d.)</td>
<td>1.8 (1.8)</td>
</tr>
<tr>
<td>Length of hospitalisation in months: mean (s.d.)</td>
<td>13.9 (14.7)</td>
</tr>
<tr>
<td>Unemployed: %</td>
<td>90</td>
</tr>
<tr>
<td>Living alone: %</td>
<td>41</td>
</tr>
</tbody>
</table>

Bonferroni-adjusted post hoc multiple comparisons ($P < 0.05$): *sample B v. A, D; †B v. C.
EXPLANATORY MODELS OF ILLNESS IN SCHIZOPHRENIA

Table 2 Content analysis of responses to questions on concept, cause, treatment, severity and course of illness

<table>
<thead>
<tr>
<th>Concept</th>
<th>Cause</th>
<th>Treatment</th>
<th>Severity</th>
<th>Course</th>
</tr>
</thead>
<tbody>
<tr>
<td>Do not know</td>
<td>Do not know</td>
<td>Do not know</td>
<td>12%</td>
<td>Do not know</td>
</tr>
<tr>
<td>Schizophrenia/psychosis</td>
<td>Drugs/alcohol</td>
<td>Medication/brain scan</td>
<td>19.7%</td>
<td>Not severe</td>
</tr>
<tr>
<td>Depression/manic depression</td>
<td>Interpersonal</td>
<td>Practical help</td>
<td>5.4%</td>
<td>Moderately severe</td>
</tr>
<tr>
<td>Mental illness</td>
<td>Supernatural</td>
<td>Counselling/understanding illness</td>
<td>18.8%</td>
<td>Very severe</td>
</tr>
<tr>
<td>Others say I have schizophrenia</td>
<td>Mental illness</td>
<td>Social activities/occupational therapy</td>
<td>4.5%</td>
<td>Variable</td>
</tr>
<tr>
<td>Non-specific illness</td>
<td>Physical illness</td>
<td>Natural remedies</td>
<td>5.4%</td>
<td></td>
</tr>
<tr>
<td>Spiritual experience</td>
<td>Personality</td>
<td>Get better</td>
<td>12.5%</td>
<td></td>
</tr>
<tr>
<td>Nothing wrong</td>
<td>Stress</td>
<td>Receiving right treatment</td>
<td>2.7%</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>Childhood events</td>
<td>None</td>
<td>4.5%</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>Other</td>
<td>Other</td>
<td>15.2%</td>
<td></td>
</tr>
</tbody>
</table>

The four ethnic groups were also compared on clinical and psychological characteristics and there were no significant differences between the groups on any of these variables (data available on request).

Explanatory models

Responses to the SEMI questions concerning concept, severity, course, aetiology and treatment from the sample as a whole were content analysed and are presented in Table 2. Although an individual may have cited more than one response, particularly with respect to aetiology and treatment preferences, the first response provided by each participant was coded for the purpose of this analysis. Four raters independently coded the raw data from the transcripts into categories. The chance-corrected proportional interrater agreement between the four raters was very good: $\kappa = 0.87$ (Altman, 1991).

Approximately 20% of the sample called their problem schizophrenia/psychosis and 18.5% called it depression/manic depression. Almost two-thirds said that their illness was either moderately or very severe. Approximately half stated that their illness had a long course and almost one-third said that it had a short course.

There was substantial variation in the primary causes of illness cited: 21.4% said they did not know; 16.2% cited interpersonal factors (e.g. ‘break-up of relationship’, ‘emotional baggage’); 15.4%, supernatural factors (e.g. ‘someone did magic to me when I was a little boy’, ‘evil forces’); 9.4%, stress mostly arising from training or employment (‘stress from work’, ‘working very hard’); 8.5%, personality factors (e.g. ‘myself for not being wiser’, ‘lack of knowledge of when I was becoming stressed’); 7.7%, drugs/alcohol (e.g. ‘heavy drinking’, ‘took crack’); 7.7%, negative childhood events (e.g. ‘physical and mental abuse in childhood’, ‘influence in life when I was young’); 5.1%, mental illness (e.g. ‘depression at a young age’, ‘breakdown’); and 5.1%, physical illness or injury (e.g. ‘heart disease’, ‘run over by a car and was in a coma’).

With respect to participants’ first treatment preference, 19.7% thought that they should receive medication; 18.8% wanted counselling, including help understanding the illness; 12.5% wanted something to make them get better but did not specify what; 5.4% wanted natural remedies (e.g. ‘natural herbal remedies’, ‘alternative herbal treatment instead of medication’); 5.4% wanted practical help (e.g. ‘help with accommodation’, ‘help with housing problems and around the house’); 4.5% wanted social/occupational activities (e.g. ‘occupational therapy’, ‘something to do like a club’); 2.7% said they were getting the right treatment; and 15.2% mentioned specific treatments that did not fit into the general categories (e.g. ‘equal opportunities as the system hates Black people’, ‘X-ray’).

Explanatory models and ethnicity

Concept of illness, or the name patients used to describe their problem, did not differ significantly between the four groups ($\chi^2 = 18.9, P = 0.09$), nor did perceived severity of illness ($\chi^2 = 16.02, P = 0.18$). Cause of illness, coded into four categories broken down by ethnic group is displayed in Table 3 and Fig. 1.

Cause of illness differed significantly according to ethnicity (see Fig. 1) in a 4 x 4 analysis ($\chi^2 = 28.7$, d.f. = 9, $P < 0.001$).

Table 3 Explanatory models of the four groups

<table>
<thead>
<tr>
<th>Primary cause of illness</th>
<th>Biological n (%)</th>
<th>Social n (%)</th>
<th>Supernatural n (%)</th>
<th>Non-specific n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A African–Caribbean (n = 30)</td>
<td>2 (6.7%)</td>
<td>18 (60%)</td>
<td>3 (10%)</td>
<td>7 (23.3%)</td>
</tr>
<tr>
<td>B Bangladeshi (n = 30)</td>
<td>0 (0%)</td>
<td>11 (42.3%)</td>
<td>7 (26.9%)</td>
<td>8 (30.8%)</td>
</tr>
<tr>
<td>C West African (n = 29)</td>
<td>3 (10.7%)</td>
<td>9 (31%)</td>
<td>8 (28.6%)</td>
<td>6 (21.4%)</td>
</tr>
<tr>
<td>D UK White (n = 30)</td>
<td>10 (34.5%)</td>
<td>9 (31%)</td>
<td>0 (0%)</td>
<td>10 (34.5%)</td>
</tr>
<tr>
<td>Total (%)</td>
<td>15 (13%)</td>
<td>49 (43%)</td>
<td>18 (16%)</td>
<td>31 (27%)</td>
</tr>
</tbody>
</table>
Because the $4 \times 4$ analysis does not show which groups differed on which measures, the effect was broken down into a series of focused $2 \times 2$ comparisons using $\chi^2$ and Fisher’s exact test in the case of low cell counts.

The sources of the overall effect were somewhat localised. Overall, there were no differences between African–Caribbeans, Bangladeshis and West Africans except when social causes were compared with supernatural causes; African–Caribbeans were more likely ($P=0.053$) to cite social causes than West Africans.

By contrast, Whites were distinguished from the other groups in four cases. First, when biological causes were compared with supernatural causes, Whites cited biological causes more frequently than African–Caribbeans ($P=0.022$), Bangladeshis ($P<0.001$) and West Africans ($P=0.001$). Second, when biological causes were compared with social causes, Whites cited biological causes more frequently than African–Caribbeans ($P=0.05$) and Bangladeshis ($P=0.003$). Given the statistical power of the study, a trend ($P=0.072$) for the same pattern when Whites were compared with West Africans is worth noting. Third, when biological causes were compared with non-specific causes, Whites cited biological causes more frequently than Bangladeshis ($P=0.014$). Finally, when supernatural causes were compared with social and non-specific causes, both Bangladeshis (cf. social $P=0.016$, cf. non-specific $P=0.013$) and West Africans (cf. social $P=0.024$, cf. non-specific $P=0.004$) cited supernatural causes more frequently than Whites.

Treatment preference also differed according to ethnicity ($\chi^2=24.7$, $P=0.054$). Bangladeshis were less likely to want treatment of any kind than Whites ($P=0.019$) and African–Caribbeans ($P=0.049$). Both Whites and African–Caribbeans were more likely to want counselling and, additionally, Whites were more likely to want medication ($P=0.051$) or to be unsure about what they wanted ($P=0.029$). Bangladeshis were also more likely to want non-conventional forms of treatment, including natural remedies and spiritual activities, than Whites ($P=0.008$), who were more likely to be unsure about what they wanted.

**Cause of illness and clinical and psychological characteristics**

Because there was substantial variation within ethnic groups, explanatory models (independent of ethnicity) were analysed with respect to the clinical and psychological characteristics assessed. With respect to concept of illness, patients who called their problem a specific or non-specific psychiatric illness or breakdown were more likely to accept that they had a mental illness than those who named it differently ($P=0.05$). For severity of illness, patients who perceived their illness to be very severe had a lower external health locus of control than those who thought that their illness was moderately severe ($P=0.05$).

**DISCUSSION**

**Explanatory models and ethnicity**

In the main there were no differences between the three non-White groups. However, there were consistent differences between the White and non-White groups. When biological causes were compared...
with supernatural causes of illness, Whites cited biological causes of illness more frequently than the three non-White groups, who cited supernatural causes more frequently. When biological causes were compared with social causes, Whites cited biological causes more frequently than African–Caribbeans and Bangladeshis. Conversely, both African–Caribbeans and Bangladeshis cited social causes more frequently. When social causes were compared with supernatural causes, Whites cited social causes more frequently than Bangladeshis and West Africans, who were more likely to cite supernatural causes.

Explanatory models, ethnicity and treatment

Bangladeshis, who cited supernatural causes more frequently, either wanted alternative forms of treatment such as religious activities or no treatment at all. Conversely, Whites, who cited biological and social causes more frequently, preferred medication and counselling. Type of explanatory model, independent of ethnicity, was related to satisfaction with treatment, suggesting that what people believe about their illness influences their experience of and benefit from mental health services (cf. Dein, 2002). There was a marked difference between a biological or social explanatory model and corresponding perceptions of treatment. People with a ‘biological’ explanatory model more often said that they were receiving the right treatment for them, were more satisfied with the treatment and had better therapeutic relationships than those with a ‘social’ explanatory model. Although the clinician’s explanatory model was not assessed in this study, it seems reasonable to suggest that because the predominant treatment model is medication-based it is primarily biological. This finding seems to support Callan & Littlewood (1998), who found that satisfaction is higher when there is concordance between the patient’s and psychiatrist’s explanatory models.

Although cause of illness stated by patients was related to their treatment preferences and perceived benefit from and satisfaction with treatment, it was not associated with treatment compliance. A similar finding was also reported by Holzinger et al. (2002), who investigated subjective illness theory and compliance with atypical antipsychotics among people with schizophrenia.

People who cited supernatural causes of illness were less insightful (i.e. less likely to accept that they had a mental illness) but were not less compliant with treatment. Goldberg et al. (2001) found that ethnic minorities were more likely to be rated as having poor insight. It may be that insight is narrowly assessed as a difference between the perspective of the rater and the person being rated in how mental illness is conceptualised, without taking account of differences in social cultural attributions. The fact that type of explanatory model was not related to treatment compliance may highlight the lack of explanatory power of insight if assessed in this way (cf. McCabe et al., 2000; McCabe & Quayle, 2002). Indeed, other sociocultural factors, such as varying degrees of social deference, may mediate readiness to comply with treatment.

Methodological issues

In this study, individual accounts of cause of illness were not fixed but fluid and often people did not cite a single, but multiple causes. Williams & Healy (2001) suggested that ‘exploratory map’ rather than ‘model’ might reflect more accurately how people construct narratives about illness. This may be particularly relevant among people of different ethnic origin in the UK, who will vary in degree of acculturation and so may draw on multiple social and linguistic resources in narratives about illness. The fluidity of accounts of illness during the course of a single interview is compounded when test–retest reliability is considered. The stability of explanatory models over time appears to be low (McCabe & Priebe, 2004), thus limiting their value in predicting long-term outcome.

The current study used an operationalised method to assess explanatory models, as suggested by Bhui & Bhugra (2002). It highlighted the limitations as well as the potential benefits of such methods (cf. Canino et al., 1997). There was a tension in reducing explanatory models to single categories to link them with quantitative outcomes. A necessarily reductionistic approach in condensing complex accounts of illness to fixed simple categories resulted in the loss of rich contextual information, underlining the tension between the anthropological framework within which the concept of explanatory model of illness was originally developed and a reductionistic approach for quantification (McCabe & Priebe, 2004). Moreover, the categories derived from the qualitative analysis appear to be more sensitive in discriminating White from non-White groups and less sensitive in discriminating between the three non-White groups. It is possible that the analyst’s categories are less finely tuned than those members themselves (i.e. in this case the different cultural groups) use to order the social world.

The low statistical power of the small sample sizes precluded the identification of small effect sizes. Consequently, negative findings, such as the failure to detect a significant association between explanatory model and treatment compliance, must be interpreted cautiously. Also, although post hoc testing between groups for specific results was Bonferroni adjusted, there was no adjustment for multiple testing across all tested associations. This approach is justified, however, in an exploratory study investigating a number of independent research questions (Perneger, 1998).

Although it is a strength of the study that all groups were assessed with standardised instruments and that the groups were comparable in their socio-economic conditions, there may have been a selection bias. Those who did not agree to participate or were not in contact with services are likely to be less satisfied with treatment and may differ more in their explanatory models. Further research is required to shed light on the influence of explanatory models in the early phases of illness on contact with services and benefit from treatment. A related issue is that it remains unclear whether and, if so, how contact with clinicians affected patients’ explanatory models.

Although various methodological issues remain unresolved, this study is a step forward because it successfully linked qualitative data elicited in semi-structured interviews with conventional quantitative outcomes and yielded statistically significant associations that are clinically relevant and may guide further research. The differences in explanatory models between ethnic groups may be considered reliable because Fisher’s exact test is a conservative statistical test (Rosenthal & Rosnow, 1991) and so is unlikely to lead to Type I errors.

In conclusion, this study suggests that explanatory models differ reliably according to cultural background and are associated with treatment preferences and satisfaction but not with treatment compliance. Although Whites are more likely to have a biological explanatory model, African–Caribbeans, West Africans and
Bangladeshis are more likely to have a social or supernatural explanatory model. Moreover, having a biological explanatory model, especially compared with a social explanatory model, is linked with greater treatment satisfaction and better therapeutic relationships.

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