Rylance, R, Rylance, R and Graham, P

Does the taught content of care planning live up to mental health student nurses experiences and perceptions of practice?

http://researchonline.ljmu.ac.uk/id/eprint/573/

Article

Citation (please note it is advisable to refer to the publisher’s version if you intend to cite from this work)

Rylance, R, Rylance, R and Graham, P (2014) Does the taught content of care planning live up to mental health student nurses experiences and perceptions of practice? Mental Health Practice, 18 (2). pp. 30-36. ISSN 1465-8720

LJMU has developed LJMU Research Online for users to access the research output of the University more effectively. Copyright © and Moral Rights for the papers on this site are retained by the individual authors and/or other copyright owners. Users may download and/or print one copy of any article(s) in LJMU Research Online to facilitate their private study or for non-commercial research. You may not engage in further distribution of the material or use it for any profit-making activities or any commercial gain.

The version presented here may differ from the published version or from the version of the record. Please see the repository URL above for details on accessing the published version and note that access may require a subscription.

For more information please contact researchonline@ljmu.ac.uk

http://researchonline.ljmu.ac.uk/
Does the practice of care planning live up to the theory for mental health nursing students?

Rebecca Rylance and Peter Graham examine why the person-centred care that students learn about is not always a reality on the wards

Abstract

Care planning should be a collaboration between the service user, caregivers and the relevant professionals. It is based on recovery principles, where clients identify their goals and how to work to reach them, rather than concentrating on illness, symptoms and problems. Mental health nursing students were taught the theory but observed that, in their clinical placements, this approach was often not followed in practice.

These issues were explored in two teaching sessions with six students. Subsequent focus groups were recorded, transcribed and analysed, yielding four main themes: care planning custom and practice, collaboration, organisational culture and student assumptions about their mentors. Participants detailed how care planning might not be person-centred in practice. It was suggested that this might be due to clinical customs, strains and restrictions, lack of collaboration between service users and the multidisciplinary team, and inept organisational culture. The main challenge for services has been how to manage risk as well as the person-centred approach, and the ‘competing dilemmas associated with care-versus-control issues’.

Keywords
Care planning, collaboration, recovery, student perceptions

THE CARE Programme Approach (CPA) advocates that the care planning process should be a collaborative event between the service user, caregivers and the relevant professionals (Department of Health (DH) 2008). In Making Recovery a Reality, Shepherd et al (2008) argue that recovery is about clients defining their own goals and ambitions, and suggest that this can be achieved by moving away from illness, symptoms and problem formulation.

Our study examined the perceptions of a group of mental health nursing students and their observations of care planning in theory and in practice, following a series of evidence-based teaching sessions. The aims and learning outcomes for the sessions were:

- To reinforce the fact that care planning is a shared and continuous process.
- To consider the important factors in problem identification.
- For students to be able to identify and list problems, strengths and needs with clients and caregivers.
- To introduce students to goal setting.
- For students to practise writing problem and goal statements.

Involving clients in care

The idea of mental health service users being actively involved in their own care is not new (Anthony and Crawford 2000, Tunmore 2000). The rise of the recovery movement in mental health services, with principles based on self-determination, self-management and hope (Alakeson and Perkins 2012, Hall et al 2013), has promoted the notion that clients are no longer passive recipients of care. Instead, they are actively taking charge of their own lives and being seen as ‘experts by experience’ on their own condition. The benefits of such an approach are well
attested in the literature: No Health without Mental Health (DH 2011a) describes how ‘having control over your life is associated with better physical and mental health. This also means that people with mental health problems are able to plan their own route to recovery’.

Much is written about the need for collaboration in care planning between clinicians, service users and their caregivers in mental health services. The refocusing of the CPA in 2011 (DH 2011b) recognises that clients will not engage with the care planning process unless it is meaningful to them, and their input is genuinely recognised. The Department of Health guidance states that care plans should be devised and agreed in partnership with the service users and, where appropriate, their caregivers. The therapeutic value of involving people in their own care cannot be underestimated. Stringer et al (2008) also assert that care providers should better accommodate the needs and wishes of clients in their journey of care.

That said, to our knowledge it is not stated in the literature whether and to what extent practitioners in reality involve clients and their caregivers in care plans. The Triangle of Care (Carers Trust 2013) recommends that all care staff who undertake care planning should receive specific training on how to involve service users and their caregivers in the process.

The study
Mental health nursing students from Liverpool John Moores University undertaking the DipHE pre-registration nursing programme (March 2010 cohort) attended two dedicated teaching sessions on care planning as part of the theoretical component of the curriculum. The first session was entitled ‘Problem- and goal-centred care planning’ and was delivered by a lecturer/practitioner who leads on care planning for a local mental health trust. The teaching was recovery focused and encouraged the students to consider: the nature of collaborative practice; the use of language from the perspective of staff and client; and the issue of who ‘owns’ the care plan. A second, follow-up session explored these themes further and introduced the use of formulation to assist with the writing of the ‘problem’ statement, and was followed by focus groups to discuss what had been learned in theory and in practice. Practice settings included a range of in-patient and specialist community placements.

Method A naturalistic method of enquiry was selected to examine the relationship between care planning theory and practice as seen by mental health nursing students in the clinical practice setting. Therefore, a phenomenological approach, which seeks to consider the whole person and values their experiences (Balls 2009), was chosen as the philosophical framework for the study. Phenomenology, which is a widely accepted research tradition in qualitative enquiry (Polit and Beck 2008), is not necessarily concerned with the production of grand theory but instead seeks a description of people’s ‘lived experience’, articulated by those who did the experiencing (Balls 2009).

Phenomenology seeks to understand the ‘essence of phenomena’ by placing emphasis on the way people make sense of their world (Polit et al 2001), and by doing so it recognises and values the meanings that people ascribe to their own existence. Thus a qualitative descriptive study, informed by a phenomenological approach, was framed to explore the experiences and perceptions of this group of mental health nursing students.

To generate narrative data, the focus groups were conducted using a semi-structured interview guide. Interviews are a primary method of data collection in qualitative research (Parahoo 2006) and there is some evidence to suggest that unstructured interviews generate greater depth and detail, but that semi-structured interviews are better suited when a researcher wants to be sure that a specific set of topics are covered (Polit and Beck 2008). Thus an interview schedule (Box 1) was created with some

Clients are no longer passive recipients of care. Instead, they are actively taking charge of their own lives
preliminary questions to focus on areas of interest and to generate data.

The interview guide focused largely on the students’ observation of the care planning practices of their mentors and of other registrants. The narratives recorded in the groups were subsequently transcribed and a thematic analysis was performed using Colaizzi’s analytical framework (Colaizzi 1978) (Table 1).

Sample A total of 25 nursing students were contacted, resulting in a sample comprising six third-year mental health nursing students. Students were recruited to the study via email following a series of taught theory sessions that they had received on care planning. Participation was voluntary and written informed consent was obtained from all participants. Assurances regarding confidentiality were given and all data was anonymised in accordance with legislation and local protocols. Ethical approval was granted proportionate review by Liverpool John Moores University.

Results Ninety significant statements were extrapolated from the narrative data and 14 meanings were formulated. The meanings were then grouped into clusters of themes and subsequently sorted into four main categories: care planning custom and practice; collaboration; organisational culture; and student assumptions about mentors.

The findings and supporting narratives are shown in Figures 1 to 4 (see pages 33-35).

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Colaizzi’s analytical framework</th>
</tr>
</thead>
<tbody>
<tr>
<td>One</td>
<td>Transcribed descriptions read by investigator.</td>
</tr>
<tr>
<td>Two</td>
<td>Significant statements and phrases extracted.</td>
</tr>
<tr>
<td>Three</td>
<td>Meanings formulated from significant statements and phrases.</td>
</tr>
<tr>
<td>Four</td>
<td>Meanings clustered into themes.</td>
</tr>
<tr>
<td>Five</td>
<td>Results integrated into an exhaustive list of the phenomena.</td>
</tr>
<tr>
<td>Six</td>
<td>Descriptions of the fundamental structure of the phenomenon.</td>
</tr>
<tr>
<td>Seven</td>
<td>Descriptions returned to participants for validation.</td>
</tr>
</tbody>
</table>

(Colaizzi 1978)

Custom and practice The theme category ‘care planning custom and practice’ relates to how the students talked about their observation and experience of care planning on the wards and in community settings. A key theme cluster was that of professional jargon and, in particular, the ‘professional snobbery’ that the participants perceived as manifesting during the process. The students acknowledged that a collaborative care plan should use client-friendly language but in doing so, they perceived this to look unprofessional to other agencies.

‘You got to use all this jargon... you can’t have a care plan in simple terms because it makes you sound thick.’

Participants also commented on the practice of copying and pasting generic care plans and the use of clichés such as ‘to remain well in the community’.

‘Staff just go to copy and paste and that goes on the care plan and you just tailor it to whatever they’ve come in with.’

‘It’s scary to think people are just copying and pasting these huge kind of statements.’

In addition, it was observed that the care plans were not goal-focused but problem-focused, and tended to be a ‘paper exercise’ rather than a meaningful collaborative document.

A number of students pointed out that the assessment tools did not necessarily inform the care plans and that, when the care plan contained risk strategies, this was considered by the registered staff as a reason not to share the care plan with the service user, leading to practice misunderstanding and the potential for service user distress. This was judged by the students to be unnecessarily defensive and ethically unsound.

‘I’ve had an experience where you’re dealing with quite a high-risk individual in the community and because of the CPN’s [community psychiatric nurse’s] anxieties about his reaction to the care plan ... he can’t see it, it’s not right’.

‘In his risk assessment but it was also in his care plan so it was like he can’t sign that because he can’t see it because if he sees that he’ll know what we’re kind of doing behind closed doors.’

The students observed that a tension exists in practice: ‘The balance between risk and a person’s rights.’

Collaboration The theme category ‘collaboration’ describes the participants’ observations and perceptions of service user involvement in the care planning process. The students commented that in the inpatient environment clients were rarely involved, which was attributed to the presence of
psychosis, cognitive impairment and admissions of short duration. It was acknowledged that the community environment was more conducive to engagement and that here collaborative care planning was more likely to occur.

‘I understand why it’s [the care plan] not signed when they first get admitted to an acute ward because they might be manic or really psychotic.’

‘If it’s a first presentation you’ve got no chance of doing a proper tailor-made care plan for them, because you don’t know enough about them to get that done before they get discharged.’

Some students had noticed clients who ‘said the right thing’ in the care planning, simply to achieve leave or discharge; the students alluded to this as ‘playing the system’.
Agenda setting was perceived as being ad hoc, occasionally inappropriate and not achievable in the inpatient environment, because of lack of time and the restrictions of the ward routine.

'I've very rarely seen an agenda set unless you've got something specific you want to do with that person.'

The students perceived service user collaboration to be generally poor, despite identifying reasons why this might be the case. They all commented on the richness of information that could emerge from a collaborative approach that would benefit service users and staff.

'I've sat with people and gone through their care plans, but you don't see it enough.'

'It's also good for the practitioners.'

**Culture of organisation** The participants perceived that the culture of the organisation contributed to, or detracted from, the overall care planning process, and they alluded to the issue of time management. They suggested that the wards were less person-centred than the community setting and also that the CPA was too process-driven and task-focused. They also remarked that if the qualified staff did care plans correctly, they would actually save time.

**Students all commented on the richness of information that could emerge from a collaborative approach**
acknowledged that the mentor-mentee relationship was central to the learning experience and all believed that the longer their mentor had been qualified, the more knowledge and experience they naturally had. Therefore those mentors who did not use clinical assessment tools were seen as not doing so because they did not need to.

‘The mentors who have been qualified for years, they’ve got so much experience that they don’t need to rely on problem-solving models to then feed into care plans. Twenty five years in, they’ve got a massive amount of knowledge and you know it’s fabulous for they are doing it all from memory.’

Related to this, was the assumption that length of time since qualifying was related to being a good mentor.

‘Mentors who have been qualified for three or four years, they’re brilliant because they still remember what it’s like to be a student.’

The value students ascribed to the length of time their mentors had been qualified was not examined further during the course of the study and undoubtedly requires more investigation.

Participants implied that mentors might behave differently when they had a student with them.

‘I think it just depends on how your mentor does things… now, whether that’s just because they’ve got a student with them…?’

Good mentors were associated with sound knowledge and application of the care planning process.

‘A really good mentor kind of understands the care planning process.’

Discussion
Our findings suggest that the principles of a recovery-focused approach to care planning are not reflected in clinical practice. The students who took part in our study noted that the inpatient areas often failed to collaborate with service users in the development of their care plans, possibly because of the person’s level of acuity. However, it may also be argued that mental health services have failed to adapt to the concept of recovery, as, historically, services had been designed around the belief that those with a severe and enduring mental illness do not recover (Anthony and Crawford 2000).

As previously stated, the push to put service users at the centre of their own care is supported by state guidance, most notably in Refocusing the Care Programme Approach (DH 2008). However, the participants in our study saw the CPA in practice to be a bureaucratic tick-box or time management exercise rather than a process for meaningful engagement and a tool to provide person-centred care.

One of the most notable observations was on assessment and management of risk in clinical practice. The narrative data suggested that the identification of risk removed any attempt by staff to collaborate with the service user in developing a care plan to manage the risk. This echoes the thoughts of Hall and Wren (2008) who state that the main challenge for services has been to manage the ‘competing dilemmas associated with care-versus-control issues’. However, the principles described in Best Practice in Managing Risk (DH 2007) clearly advocate a collaborative approach.

Our findings indicated that ‘practice’ was not necessarily practising in an evidence-based way or in a way that complemented the theory being taught in university. These phenomena may have promoted a lack of trust and confidence among the students with regard to either the theory component of care planning that is taught in university or to what they see in practice.

Whether these findings reflect a broader cultural climate that remains over-patriarchal and problem-focused is unclear and doubtless requires further exploration.

The main limitation of this study is its small scale, which means it cannot claim to be representative of wider cohorts’ perceptions of care planning as taught and in clinical practice.
Neither should it be generalised to a larger field of mental health nursing students. All students were in agreement with the discussion points.

Conclusion
Our study has provided some unique insights into the perceptions and experiences of a group of mental health nursing students and how they observe care planning in clinical practice in contrast to how it is taught on their preregistration university course. The data have unveiled a number of reasons why care planning is not person-centred in clinical practice. These include practice restrictions and customs, lack of collaboration and inept organisational culture. Our findings thus indicated a perceived marked difference between taught theory and clinical practice, which poses the question as to whether there is a theory-practice gap that should be addressed. The study also revealed some interesting and unexpected insights into the dynamics that exist between students and their mentors.

Implications for practice
■ In theory, the care planning process should be a collaboration between the service user, caregivers and the relevant professionals. However, care planning may not be collaborative, person-centred or goal-focused in clinical practice.
■ Obstacles include convention, practice restrictions, management of risk, lack of time and inept organisational culture.
■ Clients will not engage with person-centred care planning unless it uses familiar language and is meaningful, and their input is genuinely recognised.
■ The care planning approach benefits staff as well as service users and in the end saves time.

References
Department of Health (2011a) No Health without Mental Health. DH, London.

Conflict of interest
None declared