Social networks of patients with multimorbidity: a qualitative study of patients’ and supporters’ views

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ABSTRACT

INTRODUCTION: Multimorbidity impacts on patients’ health and wellbeing, but relationships experienced within social networks can support people to live well.

AIM: This study sought to elicit the views of New Zealanders with multimorbidity about their social networks and the views of their nominated supporters.

METHODS: Ten patients with multimorbidity and their nominated supporters each independently recorded their views of the patient’s social network on a five-concentric-circle template, indicating supporting role and importance to each patient. Sets of patients’ and nominated supporters’ templates were compared followed by comparing matched pairs of patient–supporter templates. Nominated supporters’ views about the patients’ networks and why they were nominated were collated.

RESULTS: Three patients nominated family members as supporters and seven nominated health professionals. Nominated family members identified a greater range of supporters than nominated health professionals. Nominated family members perceived that they played an integral role, whereas health professionals were less comfortable viewing relationships with patients in this way. Family members were not surprised to be nominated as supporters, and some described a considerable burden of care. Health professionals described themselves as coordinators of support and having positive relationships with patients.

DISCUSSION: Patients with multimorbidity have rich and diverse social networks. They view partners, family and health professionals as providing significant support. Family members are more aware of their role and have a deeper understanding of other network members than health professionals. Further research is needed on the use of social networks in clinical practice to support the health and wellbeing of those with multimorbidity.

KEYWORDS: Health professionals; long-term conditions; multimorbidity; patients; social networks

Introduction

Multimorbidity (multiple long-term conditions) is a rising challenge around the world and is associated with reduced quality of life, increased mortality, polypharmacy and greater health service use.1 Past and current relationships, often experienced within a social network, influence health and wellbeing. The concept of health and wellbeing is now more broadly defined as the ‘ability to adapt and to self-manage, in the face of social, physical and emotional challenges’.2 Positive relationships improve health literacy, self-efficacy, self-management and patient activation for people with multimorbidity.2–8

Social networks are ubiquitous across all countries and settings, and typically include partners, family, friends, community-based
organisations9–13 and, to a lesser extent, professionals.13,14 Informally and yet in a connected fashion, network members can act to support an individual with health and social needs.15 Irrespective of socioeconomic status12 or country,16 social networks can or have the potential to collectively enable patients to self-manage17,18 and enhance feelings of social integration and individual identity.18 Social networks yield benefits to people, health systems, health professionals and the economy.19

To date, multimorbidity research has focused on exploring individual interactions between patients and health professionals, usually within healthcare consultations, rather than exploring relationships between patients and people in their social networks.9,10,20,21 Studies exploring social networks typically use a visual template of concentric circles, indicating relative degree and importance of interaction from a patient’s perspective.12,13,17,22 In New Zealand (NZ), research with medical students has examined the use of similar templates to facilitate understanding of patients’ Communities of Clinical Practice, an approach somewhat akin to social networks,23–25 and patients’ and clinicians’ views of Communities of Clinical Practice.26 No research involving the social networks of patients with multimorbidity and that of their supporters appears to have been undertaken, although the need to do so has been recognised.27

This exploratory study sought to elicit the views of NZ patients with multimorbidity about their social networks, and the views of their nominated significant supporters.

Health system context
New Zealand has comparable multimorbidity prevalence to other Western countries,28 although people of Māori and Pacific descent have higher rates of multimorbidity and overall lower life expectancy.28–31 The care of patients with multimorbidity generally involves both primary and secondary healthcare providers, with primary care assumed to take the lead role in coordination.32,33

Context of data collection
This study was built around an existing pre-vocational registration teaching programme. A long-term conditions management module is included in an interprofessional programme for dietetic, medicine, physiotherapy and radiation therapy students.34 As part of this module, interprofessional groups of three students visited a patient with multimorbidity in their home to explore the patient’s experience of multimorbidity. A social network template comprising five concentric circles (adapted from E & B Wenger-Trayner35) was used to record each patient’s network.25 Colour-coded categories identified possible groups of network supporters. Proximity to the centre (Core, Active, Occasional, Peripheral, Transactional) indicated the relative degree of interaction or importance of individuals within patients’ social networks.

This study was approved by the University of Otago ethics committee (No. H16/007).

Methods
Researchers phoned the patients and sent written information, consent and demographic forms to patients who expressed an interest in participating in the study. Consenting patients were asked to nominate an individual who provided significant support to them. Throughout the rest of this paper, this nominated individual will be referred to as ‘the supporter’.

Student groups interviewed each patient about their experience of multimorbidity and, using
a broad list of question prompts, asked patients to complete a social network template (with or without the assistance of the students). They then prepared a PowerPoint presentation (which included the social network template) to represent this information. The presentation was retained for analysis (with the students’ consent).

The research team then approached and interviewed the nominated supporters about the patient’s social network and asked them to also complete the social network template, with or without assistance. Nominated supporters were not shown the patients completed templates so that the supporters could record their own perspective on each patient’s social network and their role in supporting the health and wellbeing of the patient, without being influenced by the patient’s view. All interviews were audio recorded, transcribed verbatim and field notes were retained.

**Data analysis**

The theoretical framework for this study drew on social network analysis focusing on relational data, including contacts, ties, connections and attachments within patients’ social networks.

The data analysis included collating information on who was in the patient’s social network. Sets of patients’ and then supporters’ templates were analysed for similarity and difference. Then, pairs of social network templates relating to each patient were independently analysed by JM and EM, who compiled and compared summary frameworks of patients and supporters.

We compared patients’ and supporters’ views about proximity and the relative importance of people named in the social network, and the sufficiency of patients’ social networks. This was undertaken through analysis of the social network templates, summary frameworks and transcribed interview data. We collated information on why supporters thought they had been nominated.

As the above steps progressed, the research team met regularly to discuss the analysis of the social network templates, to review the qualitative interview data and to contribute to the synthesis of the datasets.

**Results**

Of the 21 patients with multimorbidity involved in the teaching programme, 11 agreed to take part in the study, and nominated a supporter who provided significant support. One nominated supporter (a nurse) subsequently left her place of employment and could not be contacted for interview. Thus, 10 patients and 10 nominated others were included in the analysis (Table 1), with nine pairs of social network templates for comparison (one student group did not supply a PowerPoint presentation or a social network template).

Five females and five males aged between 35 and 86 years (five were aged 65 years or over) participated in the study, along with their nominated supporters. Supporters included family members (sister, daughter, wife) and health professionals (exercise physiologist, practice nurse, three general practitioners (GPs), a renal physician and an endocrinologist). Three patients lived alone and the remainder lived with one or more family members. Eight patients were NZ European, one Māori and one Cook Islands Māori. All patients had at least three long-term conditions.

**Who is in the patient’s social network? The views of patients and nominated supporters**

Patients’ social networks varied in number and range of supporters. Family supporters nominated more supporters and placed these supporters nearer to the Core circle, indicating greater perceived support. Health professional supporters tended to focus on other health professionals, who they placed in the Active and Occasional circle, although one had a broad understanding of the patient’s family and community-based supporters. The colour-coded categories provided a starting point, but participants could include whatever was meaningful to them. For example, a patient recorded a pet and a health professional recorded a patient’s interests.

Two paired examples were chosen as examples for this paper: the first where the supporter was a family member and the second, a health professional. In Figure 1, the social network
templates were recorded by Patient 7 and the nominated supporter, his wife. His wife recorded more people with a wider range of support roles. Compared with her husband’s view, she indicated that additional people were more closely involved nearer to the Core circle.

In Figure 2, the social networks were recorded by Patient 3 and the nominated supporter, his GP. His GP knew about this patient’s girlfriend, but very little else about him and recorded only two others involved (he and another health professional). In contrast, the patient noted more and wider supports, including his girlfriend, community supporters (gardener, cleaner), interests (bridge) and health professionals (e.g. pharmacist).

Comparison of views about proximity of participants in the social network to patients, why supporters believe they were nominated and the sufficiency of patients’ social networks

In relation to the template centre (i.e. the patient), all except one patient placed the nominated supporter in the Core circle (one placed in the Active circle), indicating a high degree of perceived importance and interaction (Table 1). The three family member supporters also placed themselves in the Core circle. Five of the seven health professional supporters placed themselves in the Active circle, while the other two positioned themselves in the Occasional circle. Supporters’

<table>
<thead>
<tr>
<th>Patient</th>
<th>Nominated supporter and relationship to the patient</th>
<th>Patient and supporter views of the proximity of the nominated supporter to the patient</th>
<th>Supporter views about why they were nominated</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Family member (sister)</td>
<td>Patient and Supporter</td>
<td>‘... as a family, we sat down (and decided who was going to be responsible), and I was the one to – because I don’t have kids, so I was the one (to) come in to help.’</td>
</tr>
<tr>
<td>2</td>
<td>Professional (nurse)</td>
<td>Patient Supporter</td>
<td>‘I do a lot of the problem-solving… If the doctor’s ‘full,’ she knows to ring me ....’</td>
</tr>
<tr>
<td>3</td>
<td>Professional (GP)</td>
<td>Patient Supporter</td>
<td>‘Yip, yip we get on very well.’</td>
</tr>
<tr>
<td>4</td>
<td>Professional (renal physician)</td>
<td>Patient Supporter</td>
<td>‘What she might be thinking about is ‘For me as a person when I look at my health care, who’s kind of coordinating that most?’</td>
</tr>
<tr>
<td>5</td>
<td>Family member (daughter)</td>
<td>Patient and Supporter</td>
<td>‘Her and my relationship will never change, until the day she dies. I’m still going to txt and call her everyday...’</td>
</tr>
<tr>
<td>6</td>
<td>Professional (exercise physiologist)</td>
<td>Patient and Supporter</td>
<td>‘Thinking of community-based support, I’d like to see myself as part of that...’</td>
</tr>
<tr>
<td>7</td>
<td>Family member (wife)</td>
<td>Patient and Supporter</td>
<td>‘But (he) does rely heavily on me for stuff and that is the way it probably should be...’</td>
</tr>
<tr>
<td>8</td>
<td>Professional (endocrinologist)</td>
<td>Supporter*</td>
<td>‘The type of relationship that we’ve evolved over time that has been very upfront and I would hope he sees (it) as supportive and not judgmental...’</td>
</tr>
<tr>
<td>9</td>
<td>Professional (GP)</td>
<td>Patient Supporter</td>
<td>‘Well there’s nobody close.... I feel quite comfortable with (him) because it’s a good communicating relationship ...’</td>
</tr>
<tr>
<td>10</td>
<td>Professional (GP)</td>
<td>Patient Supporter</td>
<td>‘I was surprised she put me as the core person.’</td>
</tr>
</tbody>
</table>

* Patient template missing.
GP (General Practitioner).

Table 1. Summary of patients and their nominated supporters, including their proximity in the patient’s social network
views on why they were nominated ranged from surprise (supporter 10); role as coordinator (supporters 2, 4); relationship (supporters 3, 5, 8, 9); family affiliation (supporters 1, 7); to community affiliation (supporter 6).

Most health professional supporters thought that patients’ social networks were sufficient (patients 2, 4, 6, 8, and 10) or had some minor shortcomings (patients 3 and 9), such as; 'No. I don't think he needs anything at the moment…. the big thing is his cognition' (supporter, patient 3); 'because it's a fairly small family network and he lives alone, the challenge will be if he gets more (unwell)' (supporter, patient 9).

The three family member supporters (patients 1, 5 and 7) directly or indirectly expressed major or significant concerns about social network sufficiency. The sister of patient 1 was under substantial pressure and described her own life as being ‘on hold’. The daughter of patient 5 described a limited network: 'It’s just about (her) going to appointments and me, which is a bit sad'. The wife of patient 7 noted her role as: 'I just have had to do everything, so I am definitely in that caring mode compared with (our previous) relationship', but noted previously, 'I think people genuinely want to help. But I suppose I always think … you’ve got to be self-sufficient'.

Discussion
This study reports the views of patients with multimorbidity regarding their social networks and the views of their nominated supporters regarding the patients’ social networks. As well as describing who is in the patients’ social
networks, it highlights the range of people supporting patients with multimorbidity. Internationally, social networks are receiving increasing attention, and this exploratory study signals a similar importance in NZ. The study builds on international research (typically asking only patients to identify their social networks) by also asking patients to nominate a significant supporter, and exploring the supporter’s view of the patient’s social network. This approach provides insight into a patient’s social network from both perspectives.

Most patients recorded diverse and rich social networks. When each pair (patient and supporter) of templates was compared, it was evident that the role and level of involvement of the supporter appeared to influence who was recorded on their template. Nominated family member supporters tended to record family, friends and community supporters, although they were also aware of the key health professionals involved. Nominated health professional supporters mainly included other health professionals, except for one who had a broader understanding of the patient’s network.

Two-thirds of the patients in this study nominated health professionals as significant supporters. All patients, except one, categorised the health professional supporters in the Core circle. In comparison, the health professionals categorised themselves as Active or Occasional supporters, possibly signalling that they perceived themselves as being central or essential in providing support. Health professional supporters typically recorded a more limited range of people in patients’ social networks, and this may be due to a lack of knowledge about patients’ family and social circumstances. Because of this, health professionals may have misjudged the degree of sufficiency of the social network. In two instances where they described the social networks as being sufficient but with minor shortcomings, this was probably not so. These patients, both elderly men, lived alone with one having no close-living family members and the other with only one close friend; one of these men was reported to have cognitive decline.

When patients nominated family members as providing significant support and placed them in the Core circle, they signalled a high level of importance and reliance on that relationship. However, it is possible that patients may have underestimated the degree of physical or emotional involvement given; the qualitative comments supported the effect on the family supporter’s quality of life. Patients may also not appreciate the reliance that family supporters have on others within their network.

The three family members expected to be involved in the patient’s care and either lived with the patient or had daily contact. However, two expressed stress, loneliness and isolation in their supporting role, accompanied by the belief that they needed to remain self-sufficient. It is possible that the expectation of family involvement in NZ is influenced by the collectivist culture of both the indigenous Māori peoples and people of Pasifika ethnicity. Despite being keen to be involved, there are likely to be limitations to the amount of work family members are able to undertake. A study by Mays notes it is unwise to regard informal care provided by social networks ‘as a straightforward source of care that can be switched on or off to meet gaps in paid care’ and cautions that people can be reluctant to accept help from family or neighbours unless they can reciprocate.

Health professionals are generally considered to be weak ties in social networks, akin to acquaintances, with relationships characterised as transient, briefer, transactional and with less connection (contrasting to the strong ties of family or close friends where trust and bonds exist). Weak ties are important in social networks, as they allow patients to seek specific, yet socially distanced support without the constraints of close relationships. This study shows it is possible for patients to categorise health professionals as being in the Core circle, yet still meet the definition of being a weak tie. It also shows that health professionals either underestimate their importance in a patient’s social network or do not want to be too important to the patient. Despite being seen as Core by patients, there was no suggestion that these patients unduly called on the health professionals’ time and care.
Health professionals are not always aware of their patients’ social networks and do not actively use them as a resource, particularly to encourage self-management. Understanding the nature of the support given by social networks is important, as not all health professionals appreciate the ‘hard work’ and effect of multimorbidity or the time, effort and self-efficacy required to self-care. Some health professionals are unrealistic about how much patients can self-manage. This study suggests that social network information is not easily recalled by health professionals, perhaps because they do not ask about it, or if elicited, it is then not specifically recorded in the patient’s clinical record. It is also not clear if and how information about social networks should be used by health professionals to support or influence care, particularly when working within a family-centred model of care. There seems considerable potential to do so. For example, when working with family, friends and others to encourage patient self-management, defining social networks would acknowledge or explore work being undertaken on behalf of the patient or help assess the burden of care on family members. Further work is needed to explore patients’ views of confidentiality and information-sharing, as well as how and when the health professionals may or might contact the social network members.

There are limitations and strengths to this study. Although students were supplied a list of question prompts to help patients complete the social network template, they may not have uniformly used these. They may have also misinterpreted the patients’ view of their social network or not recorded the information accurately. One group of students did not return the patient’s social network template. The multiple categories included in the network diagrams were useful for prompting students, but a simpler design may be easier for patients to complete. It is possible that having 10 patients and nominated supporters may limit who the study represents, but a gender balance and age range of patients was achieved, as was a balance of nominated supporters between family members and health professionals. A further methodological strength was the use of multiple sources of data, enabling data triangulation, although there may have been advantages in interviewing both health professional and family supporters about each patient.

Conclusion

Social networks are important to patients with multimorbidity and can include a diverse range of family, friends, community organisations, health professionals, interests and pets. Relationships formed are often very helpful to patients in managing their health and wellbeing. Unknown to health professionals, patients include them in their social networks and regard them as significant supporters in their care. Although health professionals know about other health professionals in patients’ social network, they have more limited knowledge about family, friends or community organisations and do not appear to actively use this information to enhance care. Patients and health professionals are not always aware of the pressure family supporters are under, how much support these family supporters also need, or the extent to which they rely on others within the patient’s social network. More research is required on how best to use information about social networks when supporting the care of people with multimorbidity.

References


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COMPETING INTERESTS
The authors have no competing interests to declare.

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