Short Report: Carers of people affected by cancer and other long-term conditions at end of life: a qualitative study of providing a bespoke package of support in a rural setting.

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ABSTRACT

Background: A UK charity, Macmillan Cancer Support has funded a local intervention whereby carers of people affected by cancer and other long-term conditions at end-of-life are offered a bespoke package of support.

Aim: This short report describes the qualitative experiences of carers in receipt of the intervention.

Design: Qualitative research utilizing in-depth interviews. Discussion were digitally recorded and transcribed verbatim. Data were analysed using thematic analysis.

Setting/participants: Participants were carers (n=10) in receipt of the intervention. Interviews were conducted between August and September 2014 in Lincolnshire (England).

Results: Five themes from the interviews were identified (1) Awareness and advertising (2) Focus of support on the carer (3) Modes of communication (4) Personal attributes and skills of the support worker (5) Streamlining and signposting.

Conclusion: The intervention was successful within a social care setting. The participants had no overtly negative opinions on the service in its current format and all held it in high regard. Carers felt a sense of reassurance from having background support and maintained that their situation would have been worse had this support not been there.
Key statements

What is already known about the topic

- The importance to care in the community of unpaid caregiving by family members and friends has been recognised progressively in UK legislation and policy over the last 20 years.
- Caring for someone with cancer can have a significant impact on the carers’ health and wellbeing.
- Tailored and specific interventions for informal caregivers in palliative care are rare.

What this paper adds

- The paper offers insight into the experiences of carers in receipt of an innovative intervention in a predominantly rural English county.

Implications for practice, theory or policy

- These issues necessitate reflection on the commissioning of carer support within palliative care service, including aspects of staff skills and modes of communication.

INTRODUCTION

The UK policy agenda around caring developed gradually over the last 25 years, with the first specific legislation arriving as the Carers (Recognition and Services) Act 1995. This Act was followed by further Carers Acts in 2000 and 2004; the UK government also published national carers strategies in 1999 and 2010. While the legislation is commonly held to have improved the recognition and visibility of carers, the impact in terms of the numbers being assessed and receiving support has been questioned. Most recently, the Care Act went beyond the previous legislation’s emphasis on the assessment of carers’ needs by specifying a duty to meet a carer’s needs for support, subject to conditions.

Many agencies are becoming increasingly interested in offering support for carers. One such example in the UK is Macmillan Cancer Support which provides information, and practical and emotional support, for carers of people with cancer and other long-term conditions. Six in ten people caring for someone with cancer experience some kind of impact on their lives and caring for someone as the illness advances and treatment becomes palliative can be physically and emotionally demanding. Despite this evidence, tailored and specific interventions for informal caregivers in palliative care are rare and the unmet needs of palliative care carers has received limited research attention.

Since March 2012, Macmillan has funded an intervention in the predominantly rural county of Lincolnshire, whereby carers of people affected by cancer and other long-term conditions at end of life are offered a bespoke package of support. The intervention involved the employment of a full-time Macmillan Carer Support Worker. The service is delivered via local carer support service Carers Connect. Referrals are received from a range of professionals including Macmillan nurses, GPs, palliative and end of life care teams.
Once a referral is received the support worker makes contact with the carer referred and conducts a comprehensive assessment of their needs. This assessment informs the nature of the bespoke package of support and carers are sign-posted to any resources available to them. The aim is to achieve a holistic and integrated package of support, to streamline lines of communication between the carer and health, social care and other services and to remove duplication of procedures across different agencies.

In order to gain insight into the experience of carers, Macmillan commissioned the University of Lincoln to undertake this research.

**METHOD**

*Design*

Given the exploratory nature of the research a qualitative approach was considered appropriate to allow for open discussion about participants’ experiences.

*Recruitment*

Twenty invitations were sent to a range of carers (demographics, locations, employment status, carer status, relationship to cared for) who had reported particularly strong positive or negative experiences of their situation and the support they received to the support worker. Invites were sent by post via the support worker on behalf of the research team; all participants were provided with an invitation and participant information sheet prior to consenting to participation. At the time of recruitment, there were approximately 70 active cases (carers currently in receipt of support) and almost 400 in total since the intervention began. Ten carers expressed an interest in taking part and so were contacted on a one-to-one basis to arrange a convenient time and location for the interview.

*Data collection*

All interviews were conducted between 22nd August and 19th September, 2014 in Lincolnshire (England) D.N. performed all interviews with the carers; nine in their home; one at the University of Lincoln (Brayford Campus); in two cases the person they cared for was present. Interviews ranged from 45 to 90 minutes. All were digitally recorded and transcribed verbatim.

*Analysis*

Data were analysed thematically. Transcripts were independently open coded and discussed until agreement was reached. Transcripts were reviewed several times, and new codes added as appropriate and others grouped together into broader categories. Following this, themes were deduced and interpreted. Regular review and discussion contributed to data synthesis and interpretations.
Ethics

The study was approved by the National Research Ethics Service (NRES) Committee West Midlands (14/WM/0154).

RESULTS

Characteristics of carers

The sample consisted of ten participants, eight of whom were female and two male. Eight of the ten participants were aged 65 years and over and cared for their spouse or partner. Participants were also not currently in employment, with eight retired and two not employed or looking for work. At the time of interview, eight of the participants were currently caring, whilst two were no longer providing care and in receipt of bereavement support.

Five key themes from the interviews were identified as follows: (1) Awareness and advertising (2) Focus of support on the carer (3) Modes of communication (4) Personal attributes and skills of the support worker (5) Streamlining and signposting.

Awareness and advertising

Participants were most commonly referred into the service via health professionals – usually a Macmillan nurse. However, on occasion self-referrals are received. Participants explained that the timing of the intervention came at a point when they had been in receipt of little or no support and were particularly vulnerable and isolated. Of additional interest, Macmillan Cancer Support’s association with the service was not clear with some of the respondents, who were not aware that the post was funded by Macmillan.

Focus of support on the carer

It was evident that respondents were aware that the focus of the service was on their own needs as opposed to those of the individual they cared for. Responses indicated that the service was meeting a specific need of carers, not being provided elsewhere. Additionally, something that was evident was that other forms of support are patient focused and often disregard the carer.

Modes of communication

Several respondents felt it was important to meet the support worker face-to-face in their own home. All those interviewed emphasised the importance of continual contact whether this is face-to-face or via telephone. Some of the participants suggested that they may need more frequent contact in the future as the illness of the person they care for progresses. Respondents reported the sense of security
achieved simply by being registered with the service. Many of the participants acknowledged the
difficulty of delivering the service in a predominantly rural county as well as the size of the support
worker’s case load, which in some instances resulted in less frequent contact from the carer as they
were conscious of not wanting to overburden the support worker. Finally, some participants were aware
of the limited resources available to deliver the intervention.

**Personal attributes and skills of the support worker**

All of those interviewed reported that the personal attributes and specialist skills of the support worker
had influenced their satisfaction with the service. It is clear that this service offers a personal touch that
is greatly appreciated by all participants. Respondents explained how they felt that the support worker
was well skilled and suited to the effective delivery of the service.

**Streamlining and signposting**

Once carers have received an initial assessment, they are directed towards support and resources,
tailored to their specific situation and needs. In general, carers report a lack of awareness of how to
obtain relevant information and access to services. Common sources of support can include registering
with the Carers Emergency Response Service (CERS), being referred for emergency alarm systems
(telecare and lifeline), and financial assistance.

**CONCLUSION**

The study was constrained by cost in that it was funded by a small research grant from Macmillan.
Furthermore, there were time constraints on the project which prevented the recruitment of a larger
sample.

Overall this research found the Macmillan end of life support service to be a successful intervention
within a social care setting. The participants had no overtly negative opinions on the service in its current
format and all held it in very high regard. In particular, this bespoke package of support would appear
to filling a gap that is left behind by other services. The continual contact (face-to-face and by telephone)
meant that many of those interviewed felt a sense of reassurance from having background support and
consequently, maintained that their situation would have been worse had this support not been there.
Moreover, the service has streamlined lines of communication by redirecting carers to health and social
care resources that they otherwise would not have known they were entitled to.
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Declaration of conflicting interests

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