Family Perspectives on Communication With Healthcare Providers During End-of-Life Cancer Care

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This article has been chosen as being particularly suitable for reading and discussion in a Journal Club format. The following questions are posed to stimulate thoughtful critique and exchange of opinions, possibly leading to changes on your unit. Formulate your answers as you read the article. Photocopying of this article for group discussion purposes is permitted.

1. This article presents the results of a qualitative research study. How can we assess the level of evidence that is presented?
2. How similar is our patient and family population to the participants in the study?
3. What are the four major themes identified by the participants in the study? How well do the themes represent the types of emotions that might be expressed by family members of the patients for whom we care?
4. How well or often do we offer families the types of interventions recommended in Figure 1? Where can we improve?
5. What strategies can we use to address the issues involved in family communication that were identified in the study?

At the end of the session, take time to recap the discussion and make plans to follow through with suggested strategies.

Purpose/Objectives: To assess healthcare provider communication about end-of-life (EOL) and hospice care with patients with terminal cancer and their families, from the perspective of the family members.

Design: Exploratory, qualitative study using focus group discussion.

Setting: University of Maryland Greenebaum Cancer Center.

Sample: 24 spouses and first-degree relatives of deceased patients with cancer who had been treated at the cancer center from 2000–2002.

Methods: Family members participated in one of two focus group discussions and completed a short questionnaire regarding their sociodemographic characteristics and the type of EOL care their deceased relatives had received. Qualitative data were analyzed using descriptive statistics. Qualitative data were audiotaped and analyzed by comparing, contrasting, and summarizing content themes from the focus groups using NUD*IST (NVivo) software.

Main Research Variables: Family perceptions of communication with the healthcare team in EOL cancer care.

Findings: Participants associated the information, content, style, language, and timing of communication about EOL and hospice care from healthcare professionals with patient age, attitudes, and compliance with medical decisions. Informed decisions about EOL care by patients and their family members, including the use of hospice services, appeared to be compromised by these types of communication biases.

Conclusions: Satisfaction with EOL care was associated with the perceived quality of communication among patients, family members, and the healthcare team.

Implications for Nursing: Study findings highlight the importance of training healthcare professionals in the content, timing, and potential biases associated with information delivery to facilitate informed decisions about EOL and hospice care to dying patients and their families.

Key Points . . .

Communication about hospice and end-of-life (EOL) care can be less than optimal, leaving patients and their families struggling to get the information they need to make informed decisions.

Families stress the importance of receiving timely and accurate information about EOL care options to assist in making informed decisions and obtaining the best quality of care for patients.

The training of healthcare professionals in the content, timing, and potential biases associated with information delivery about EOL and hospice care can promote a peaceful transition for dying patients and their families.

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