Family Relations and Spiritual Response to Palliative Care: a Review of Literature

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Abstract

Introduction: It is common to prioritize the critical patient care in detrainment of the caregivers, but this usual behavior must change, maintaining the family members as an extension of the care given to patients.

Results and Discussion: Spiritual wellbeing is a complement to all health strands united with physical and psychosocial status, they must be put into the routine of practicing medicine on end of life care in congruence with a good communication.

Conclusion: At all levels of assistance in palliative care the family must be included, therefore clinicians shall have information on how to deal with them and developing activities to improve communication.

Keywords
Palliative Care; Family Relations; Spirituality; Adult.

Introduction
Living with terminal illness implicates, most of the time, in dealing with existential questions, it is a state that involves the patient needs, family embrace and the health professional worries. It is common and even considered normal, to prioritize the critical patient care in detrainment of the caregivers, but there is a need to change this usual behavior and maintain the family members as an extension of the care.
Palliative care is “an approach that improves the quality of life of patients and their families facing the problem associated with life-threatening illness, through the prevention and relief of suffering by means of early identification and impeccable assessment and treatment of pain and other problems, physical, psychosocial and spiritual” by the definition of the World Health Organization (WHO, 2002) [1]. Palliative care is not just at the end of life, it must be provided anytime during a life-threatening illness [2]. It is needed to have a holistic view of the person, focusing on pain and management of symptoms as well as emotional, spiritual and social well-being, their loved ones included in it.

Spiritual caregiving meets the need of the majority of those patients. Spirituality is one of the main aspects of end of life care [3], it is necessary to put spiritual guidance into the routine of practicing medicine at special care units, intensive care units, hospices and every care of patients who integrates the group of incurable illnesses.

There are few studies reflecting the family relations and spiritual response to palliative care, our review intends to expose the family questions and needs while facing this intriguing time and how spirituality reflects on it.

Methods
Focusing on Palliative Care, a qualitative systematic review of articles about family relations and spiritual response to palliative care in hospitalized adults was realized on a basis of preselected data. A search of the literature was performed from the online database SCOPUS, BVS and PUBMED from January, 2006 to December, 2015. The search was focused from the following terms: 1. ‘Palliative Care’ [Medical Subject Headings] [MeSH term], 2. ‘Spirituality’ [Medical Subject Headings] [MeSH term], 3. ‘Family Relations’ [Medical Subject Headings] [MeSH term] and 3. ‘Adults’ [Medical Subject Headings] [MeSH term].

The choice of these terms was made from a caution conciliation, which applied to define our central matter of the article based on requested descriptors. Thus, the articles were analyzed with rigor to guarantee proper sampling.

The analysis of the articles obeyed predefined eligibility criteria. Using the following inclusion criteria: (1) Original articles with full text online access; (2) Observational, experimental or quasi-experimental studies; (3) Writings in English only; and (4) Studies which focus family relations and spiritual response to palliative care in hospitalized adults.

Exclusion criteria: (1) Other projects, such as case reports, case series, literature review and comments, (2) The non-original studies, including editorials, comments, prefaces, brief comments and letter to the editor; (3) Productions that did not address depression and the hospitalized old patients; and (4) the articles in which the objective of the study did not matched the theme purposed by the systematic review in question.

We found applicable 41 articles that, when screened, resulted in 14 articles that met the criteria of evidence and were included in this review.

Results
Results in Table 1.

Discussion
Palliative care is to improve the “quality of life of patients and their families facing the problem associated with life-threatening illness” (WHO, 2002) [1] the means to it is to have an early identification, treating and preventing pain and to support on other problems such as physical, spiritual or social. It is a specialty that requires great commitment from the health care workers, the caregivers and families, not just by the delicacy of the patient’s condition but also because of the existential and spiritual distress involved [3].

This article is available at: www.intarchmed.com and www.medbrary.com
Table 1. Distribution of the domains and facets of QoL. João Pessoa, PB, 2015.

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<td>1</td>
<td>Monica Schneider, Rose Steele, Susan Cadell, David Hemsworth 2011</td>
<td>Differences on Psychosocial Outcomes Between Male and Female Caregivers of Children With Life-Limiting Illnesses</td>
<td>Journal of Pediatric Nursing</td>
<td>273 parents caring for children with life-limiting illnesses</td>
<td>Findings suggest that significant gender differences exist.</td>
<td>Women reported higher average scores compared with men for meaning in caregiving, depression, burden, and posttraumatic growth and lower average scores for optimism. Correlations also revealed some significant differences. Health care professionals need to be aware of gender differences and tailor their interventions appropriately.</td>
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<td>2</td>
<td>Timothy P. Daaleman, Barbara M. Usher, Sharon W. Williams, Jim Rawlings, Laura C. Hanson, 2008</td>
<td>An Exploratory Study of Spiritual Care at the End of Life</td>
<td>Annals of Family Medicine</td>
<td>Our study was based on qualitative research using key informant interviews and editing analysis with 12 clinicians and other health care workers nominated as spiritual caregivers by dying patients and their family members.</td>
<td>Being present was a predominant theme, marked by physical proximity and intentionality, or the deliberate ideation and purposeful action of providing care that went beyond medical treatment. Opening eyes was the process by which caregivers became aware of their patient’s life course and the individualized experience of their patient’s current illness. Participants also described another course of action, which we termed cocreating, that was a mutual and fluid activity between patients, family members, and caregivers. Cocreating began with an affirmation of the patient’s life experience and led to the generation of a wholistic care plan that focused on maintaining the patient’s humanity and dignity. Time was both a facilitator and inhibitor of effective spiritual care.</td>
<td>Clinicians and other health care workers consider spiritual care at the end of life as a series of highly fluid interpersonal processes in the context of mutually recognized human values and experiences, rather than a set of prescribed and proscribed roles.</td>
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<td>3</td>
<td>Kristin Bingen, Mary Jo Kupst and Bruce Himelstein, 2011</td>
<td>Development of the Palliative Care Parental Self-Efficacy Measure</td>
<td>Journal of Palliative Medicine</td>
<td>A list of questions were generated that asked parents their level of confidence in carrying out tasks involved in caring for a child with a potentially fatal medical condition in 6 palliative care domains: 1) medical discussion/decisions; 2) symptom management/medication; 3) daily activities; 4) feelings/concerns; 5) spirituality; and 6) end-of-life care. The PCPEM was narrowed to 58 questions after expert reviews. Then, 16 caregivers of children receiving palliative care services and 9 bereaved caregivers participated in individual focus interviews conducted by a psychologist to obtain feedback about the content and clarity of the PCPEM.</td>
<td>Results indicated that 53 of the 58 questions were rated as “comfortable being asked,” and 55 of the 58 questions were rated as “important” by the majority (&gt;80%) of the 25 caregivers.</td>
<td>This suggests that it is feasible and valuable to ask caregivers difficult questions related to end-of-life care if done in a supportive and sensitive manner. Pilot testing of the PCPEM will be conducted to determine preliminary psychometric properties.</td>
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<td>4</td>
<td>Elizabeth L. Ciemins, Jeannine Brant, Diane Kersten, Elizabeth Mullette, and Dustin Dickerson, 2015</td>
<td>A Qualitative Analysis of Patient and Family Perspectives of Palliative Care</td>
<td>Journal of Palliative medicine</td>
<td>In-depth, semi-structured patient and family interviews were conducted, transcribed, and independently reviewed using grounded theory methodology and preliminary interpretations. A combined deductive and inductive iterative qualitative approach was used to identify recurring themes. The study was conducted in a physician-led, not-for-profit, multispecialty integrated health system serving three large, western, rural states. A purposive sample of 14 individuals who received palliative care services were interviewed alone or with their families for a total of 12 interviews.</td>
<td>Presence, Reassurance, and Honoring Choices emerged as central themes linked to satisfaction with palliative care services. Themes were defined as including health care professional attributes of respect, approachability, genuineness, empathy, connectedness, compassion, sensitivity, an ability to listen, good communication, provision of information, empowerment, and timeliness. Honoring Choices included those pertaining to treatment, spirituality, and family needs.</td>
<td>At end of life or during times of serious illness, patients and families identified behaviors of Presence, Reassurance, and Honoring Choices as important. According to patients/families, health care providers must be compassionate and empathetic and possess skills in listening, connecting, and interacting with patients and families.</td>
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<td>5</td>
<td>E Michiels, R Deschepper, J Bilsen, F Mortier, L Deliens, 2009</td>
<td>Information disclosure to terminally ill patients and their relatives: self-reported practice of Belgian clinical specialists and general practitioners</td>
<td>Palliative Medicine</td>
<td>A questionnaire had been sent to a random sample of 3014 Belgian physicians from different specialties frequently involved in end-of-life care. Responses were analysed using weighted percentages, Chi-square, Mann–Whitney U-tests and a multivariate ordinal logistic regression.</td>
<td>Response rate was 58%. Both clinical specialists and general practitioners (GPs) discuss most topics related to terminal illness with their patients except end-of-life hastening options, spirituality, life expectancy and options to withhold/withdraw life-sustaining treatment.</td>
<td>The topics which most physicians always discuss with relatives without informing the patient are the aim of treatment, palliative care and incurability. There is a significant difference between clinical specialists and GPs. Clinical specialists and GPs discuss most end-of-life topics with the patient but omit important issues such as end-of-life hastening options and life expectancy.</td>
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<td>6</td>
<td>Carmen G Loiselle, Michelle M Sterling 2011</td>
<td>Views on death and dying among health care workers in an Indian cancer care hospice: Balancing individual and collective perspectives</td>
<td>Palliative Medicine</td>
<td>This qualitative study explores and compares these experiences among a diverse sample of health workers (N=25) in a grassroots cancer care hospice in Bangalore, India. Our findings underscore how personal views, socio-economic status, beliefs and values, occupational experience, and workplace interventions interact to shape ‘worldviews’ about death and dying.</td>
<td>Whereas health workers report conflicting feelings of relief and sadness when confronted with the death of their patients, these mixed emotions are often lessened through open dialogue among newly trained and more experienced health workers. Moreover, experienced hospice workers wished to ensure that less experienced ones are provided with the necessary workplace support to lessen psychological ‘hardening’ that may occur with repeated exposure to death.</td>
<td>In dealing with the diverse needs of hospice workers, both individual and collective needs must be considered to ensure an optimal workplace climate. Future work should study how hospice workers’ views on death and dying evolve with time and experience.</td>
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<td>Brian Kelly, Francis T Varghese, Paul Burnett, Jane Turner, Marguerite Robertson, Patricia Kelly, Geoffrey Mitchell, Pat Treston, 2008</td>
<td>General practitioners’ experiences of the psychological aspects in the care of a dying patient</td>
<td>Palliative and Supportive Care</td>
<td>Fifteen general practitioners whose patient had been recently referred to the Mt. Olivet Palliative Home Care Services in Brisbane participated in an individual case review discussions guided by key questions within a semistructured format. These interviews focused on the psychosocial aspects of care and management of the referred patient, including aspects of the doctor/patient relationship, experience of delivering diagnosis and prognosis, addressing the psychological concerns of the patients’ family, and the doctors’ personal experiences, reactions, and responses. Qualitative analysis was conducted on the transcripts of these interviews.</td>
<td>The significant themes that emerged related to perceived barriers to exploration of emotional concerns, including spiritual issues, and the discussion of prognosis and dying, the perception of patients’ responses/coping styles, and the GP’s personal experience of the care (usually expressed in terms of identification with patient).</td>
<td>The findings indicate the significant challenges facing clinicians in discussions with patients and families about death, to exploring the patient’s emotional responses to terminal illness and spiritual concerns for the patient and family. These qualitative data indicate important tasks in the training and clinical support for doctors providing palliative care</td>
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<td>8</td>
<td>Jennifer Angelo &amp; Linda Wilson, 2013</td>
<td>Exploring Occupation Roles of Hospice Family Caregivers from Māori, Chinese and Tongan Ethnic Backgrounds Living in New Zealand</td>
<td>Occupational Therapy International</td>
<td>The purpose of this study was to explore palliative caregiver occupations among Māori, Chinese and Tongan ethnicities. Six informants participated, one woman and one man from each ethnic group. In each of their homes, informants were asked to discuss what it was like caring for their dying family member.</td>
<td>The occupational themes resulting from these interviews were food preparation, spirituality and family gathering. Therapists need to be aware of the differences in how people care for family members within their ethnicity.</td>
<td>Implications are that occupational therapists can help families identify activities important to them within the main occupational themes: different types of foods and their preparations, various ways to express spirituality and how families gather together members of their extended family. Further, clinicians need to take on the role of a “not-knowing” but curious health-care provider in order to meet the needs of caregivers. The limitation was the small number of participants who all lived in one geographic area. Future studies should include a wider group of ethnicities.</td>
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<td>9</td>
<td>Michiyo Ando Akira Tsuda Tatsuya Morita 2007</td>
<td>Life review interviews on the spiritual well-being of terminally ill cancer patients</td>
<td>Support Care Cancer</td>
<td>Structured life review interviews were conducted with 12 patients in a palliative care unit in Japan. They completed the SELT-M (Skalen zur Erfassung von Lebensqualität bei Tumorkranken–Modified Version) questionnaire before and after the interviews. The patients were classified into two groups: effective (patients who showed an increase in the SELT-M scores after the intervention) and noneffective groups. Meaningful spoken sentences from the patients’ life reviews were transcribed and correspondence analysis was conducted on the sentences using text mining software.</td>
<td>The mean overall QOL score and spirituality subscale score of the SELT-M significantly increased after the life reviews from 2.57±0.61 to 3.58±1.0 (P=0.013) and 2.57±0.61 to 3.14±2.25 (P=0.023), respectively. Three dimensions were extracted from the effective group based on the scores “Positive view of life,” “Pleasure in daily activities and good human relationships,” and “Balanced evaluation of life.” Similarly, three dimensions were extracted from the noneffective group: “Worries about future caused by disease,” “Conflicts in family relationship problems,” and “Confrontation of practical problems.”</td>
<td>Life review interviews may be effective in improving the spiritual well-being of terminally ill cancer patients. The potential predictors of treatment success are “positive view of life,” “pleasure in daily activities and good human relationships,” and a “balanced evaluation of life,” while those of treatment failure are “worries about future caused by disease,” “conflicts in family relationships,” and “confrontation of practical problems.” Further intervention trials on patients with predictors of treatment success are promising.</td>
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<td>Gudlaug Helga Asgeirsdottir &amp; Einar Sigurbjörnsson &amp; Rannveig Traustadottir &amp; Valgerdur Sigurdardottir &amp; Sigridur Gunnarsdotir &amp; Ewan Kelly 2013</td>
<td>“To Cherish Each Day as it Comes”: a qualitative study of spirituality among persons receiving palliative care from Palliative Care Services in Iceland.</td>
<td>Support Care Cancer</td>
<td>Qualitative interviews were conducted with ten persons receiving palliative care from Palliative Care Services in Iceland. The interviews were tape-recorded, transcribed and analysed. The study is in the field of practical theology and used the theoretical approach of hermeneutical phenomenology.</td>
<td>Thematic analysis found that the spiritual dimension was of significance for the participants who understood it as a vital element connected to seeking meaning, purpose and transcendence in life. Religious and non-religious aspects of spirituality were expressed including strong spiritual components of family relationships, the meaning of God/a higher being and spiritual practices which served as a key factor in giving strength, activating inner resources and motivating hope. Nine of the participants expressed their spirituality as faith.</td>
<td>Spirituality was experienced broadly as an important dimension of how participants lived with terminal illness. Religious and non-religious characteristics were recognized which reveals the complex nature of the phenomenon. Faith was a significant part of the participants’ spirituality indicating the importance of attending to this aspect of palliative care. The study suggests the potential contributions of theological approaches which are relevant for palliative care research and practice.</td>
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<td>11</td>
<td>Randy S. Herbert, Richard Schulz, Valire C. Copeland, and Robert M. Arnold; 2009</td>
<td>Preparing Family Caregivers for Death and Bereavement. Insights from Caregivers of Terminally Ill Patients</td>
<td>Journal of Pain and Symptom Management</td>
<td>Focus groups and ethnographic interviews were conducted with 33 family caregivers (bereaved or current) of terminally ill patients. The interviews were audiotaped, transcribed, and analyzed using the constant comparative method.</td>
<td>Life experiences such as the duration of caregiving/illness, advance care planning, previous experiences with caregiving or death, and medical sophistication all impacted preparedness, or the degree to which a caregiver is ready for the death and bereavement. Regardless of life experiences, however, all caregivers reported medical, practical, psychosocial, and religious/spiritual uncertainty. Because uncertainty was multidimensional, caregivers often needed more than prognostic information in order to prepare. Communication was the primary mechanism used to manage uncertainty. Good communication included clear, reliable information, combined with relationship-centered care from health care providers. Finally, preparedness had cognitive, affective, and behavioral dimensions.</td>
<td>To prepare, some caregivers needed information tailored to their uncertainty (cognitive), others needed to “mentally” or “emotionally” prepare (affective), and still others had important tasks to complete (behavioral). In order to better prepare family caregivers for the death of a loved one, health care providers must develop a trusting relationship with caregivers, provide them with reliable information tailored to their uncertainty, and allow time for caregivers to process the information and complete important tasks.</td>
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| 12 | Bianca Sakamoto Ribeiro Paiva & André Lopes Carvalho & Giancarlo Lucchetti & Eliane Marçon Barroso & Carlos Eduardo Paiva; 2015 | “Oh, yeah, I’m getting closer to god”: spirituality and religiousness of family caregivers of cancer patients undergoing palliative care | Support Care Cancer | This study was an exploratory and descriptive qualitative study. The qualitative approach to the data was based on Bardin’s content analysis technique. The consolidated criteria for reporting qualitative research (COREQ-32) was used in the description of the results. Thirty FCs of individuals with advanced cancer undergoing palliative care were included. | Analysis of the FCs’ narratives indicated that the FCs considered that religiousness and faith in God or a Supreme Being provide them with the strength to cope with the suffering associated with the care of relatives with advanced cancer. Many FCs emphasized that talking about God was somehow comforting and made them feel at peace with themselves. Four categories were identified in the FCs’ narratives: (1) increase in faith and closeness to God becomes stronger, (2) rethink life issues, (3) negative interference in the extrinsic religiosity, and (4) quest for religiousness to gain strength or support. A conceptual framework was developed. | The results of the present study indicated that S/R are a coping strategy frequently used by FCs of individuals with advanced cancer. The perceptions of the FCs interviewed in the present study corresponded to the four distinct categories related to spirituality and religiousness. | SCOPUS |
13. Leslie P. Scheunemann; Thomas V. Cunningham; Robert M. Arnold; Praewpannarai Buddadhumaruk; Douglas B. White; 2015
How Clinicians Discuss Critically Ill Patients’ Preferences and Values With Surrogates: An Empirical Analysis*

Critical Care Medicine

Fifty-four physicians and 159 surrogates for 71 patients. We audio-recorded 71 conferences in which clinicians and surrogates discussed life-sustaining treatment decisions for an incapacitated patient near the end of life. Two coders independently coded each instance in which clinicians or surrogates discussed the patient’s previously expressed treatment preferences or values. They sub-coded for values that are commonly important to patients near the end of life. They also coded treatment recommendations by clinicians that incorporated the patient’s preferences or values.

In 30% of conferences, there was no discussion about the patient’s previously expressed preferences or values. In 37%, clinicians and surrogates discussed both the patient’s treatment preferences and values. In the remaining 33%, clinicians and surrogates discussed either the patient’s treatment preferences or values, but not both. In more than 88% of conferences, there was no conversation about the patient’s values regarding autonomy and independence, emotional well-being and relationships, physical function, cognitive function, or spirituality. On average, 3.8% (sd, 4.3; range, 0-16%) of words spoken pertained to patient preferences or values.

In roughly a third of ICU family conferences for patients at high risk of death, neither clinicians nor surrogates discussed patients’ preferences or values about end-of-life decision making. In less than 12% of conferences did participants address values of high importance to most patients, such as cognitive and physical function. Interventions are needed to ensure patients’ values and preferences are elicited and integrated into end-of-life decisions in ICUs.

14. Kirsten Wentlandt, Debika Burman, Nadia Swami, Sarah Hales, Anne Rydall, Gary Rodin, Christopher Lo and Camilla Zimmermann; 2012
Preparation for the end of life in patients with advanced cancer and association with communication with professional caregivers

Psycho-Oncology

Patients with advanced cancer but with good performance status were recruited from 24 medical oncology clinics, to participate in a cluster-randomised controlled trial of early palliative care intervention. Measures included the Quality of Life at the End of Life preparation for EOL subscale, and measures of CPC, functional status, comorbidity, spiritual well-being and symptom severity. Using chi-squared tests, t-tests and multivariate regression analyses, we examined the variables associated with preparation for EOL. We also examined the frequency distributions of individual EOL preparation items and used logistic regression to examine their associations with adequacy of CPC.

In the 469 patients, characteristics associated with better EOL preparation were better CPC, older age, living alone, less symptom burden and better spiritual well-being. Thirty-one per cent agreed that they worried ‘quite a bit’ or ‘completely’ about their family’s preparation to cope with the future, and 27% agreed that they would be a burden to their family. All preparation items except regrets about life were associated with adequacy of communication.

A substantial minority of patients with advanced cancer but with good performance status are concerned about EOL preparation, particularly in relation to their families. Better CPC may help patients prepare not only practically but also personally and socially in relation to the dying process and the welfare of their families.
It is common and even considered normal, to prioritize the critical patient care in detriment of the caregivers, but there is a need to change this usual behavior and maintain the family members as an extension of the care given to patients. According to LoiselLe and Sterling (2011) [4] the relief of pain cannot suppress the necessity of attention to how the process of death affects physical, functional, psychosocial and spiritual needs of all stakeholders involved.

One of the main characteristics of palliative care is the spiritual caregiving; 67% of people die in hospitals or long-term care facilities without assured access to spiritual care in the USA [5]. Spirituality is a multidimensional state that envelops meaning of life questions; it’s a step to deal with your own mortality, linked with religious and non-religious traces. Spiritual wellbeing is a complement to all health strands, united with physical and psychosocial status. It is necessary to put spiritual guidance into the routine of practicing medicine, Angelo and Wilson (2013) [6] suggests the facilitation of activities with patients and families with bible readings, providing conditions to go outside to gardens to energizing therapies, energy conservation techniques; all that increases family ties, listening to their needs and helping to find solutions to facilitate basic life activities autonomy.

Ando et al. (2007) [7] conducted life review interviews with palliative care patients, helping them to (1) review both good and bad memories and reevaluate these memories, (2) pay attention to, not only the negative aspects and positive aspects of their life, (3) find hobbies or interests in the reminiscence that relate with their comfort, and (4) remember good relationships with others or form new relationships. The answers for the ones with improved spiritual wellbeing were ‘positive view of life’, ‘pleasure in daily activities and good human relationships’ and balanced evaluation of life’; the ones whose cores did not improve had the dimensions including ‘worries about future caused by disease’, ‘conflicts in family relationships’ and ‘confrontation of practical problems’.

In a study conducted by Asgersdottir et al. (2013) [3] the patients at end of life care experienced characteristics of existential and spiritual distress and the majority expressed how their situation influenced their loved ones, being aware of the impact of the illness on the family members even to the extent that they were considered to suffer more than they themselves, picturing the family as a non-religious spiritual resource in the caring process which draws attention to the importance of attending to the spiritual needs of the family as a whole.

Considering Daaleman et al. (2008) [5] writings clinicians and other health care workers consider spiritual care at the end of life as a series of highly fluid interpersonal processes in the context of mutually recognized human values and experiences, rather than a set of prescribed and proscribed roles. To Wëntlandt et al. (2012) [8] patients worries about their family’s ability to cope, fears of death and dying; and anxiety about becoming a burden, undermining their sense of well-being and dignity.

All caregivers share similar concerns, going from how to prepare meals to maintain the family structure and being full-time present. At Heberth et al. (2009) [9] study all caregivers have reported medical, practical, psychosocial, and religious/spiritual uncertainty; many are not prepared for death and can experience psychological morbidity. The care of individuals with severe chronic diseases imposes a heavy physical and emotional load on family caregiver’s inasmuch they combine their usual activities with the ones related to caregiving, commonly resulting in absence from their professional, family, and social life.

Being necessary to improve communication about issues (medical, psychosocial and spiritual) that encircle death and dying will lead to caregivers being better prepared for the death and, as a result, improve their well-being. Micheils et al. (2009) [10] points out the value of involving patients’ relatives...
in the information process, but also respecting the patient autonomy obtaining their consent before disclosing information to their relatives. Any intervention must be congruent with the patients’ values and preferences – according to the principles of autonomy, substituted judgment and bests interests, integrated in end of life decisions in intensive/special care units and examining the impact of those efforts in the patient and family [11].

Ciemmens et al. (2015) [12] brings out that at the end of life or during times of serious illness, patients and families identified behaviors of presence, reassurance, and honoring choices as important. According to patients/families, health care providers must be compassionate and empathetic and possess skills in listening, connecting, and interacting with patients and families.

Michiels and colleagues (2009) [10] research shows that clinical specialists and general practitioners discuss most end-of-life topics with the patient, but omit important issues such as end-of-life hastening options and life expectancy. The lack of information may influence patients’ treatment preferences, favoring life-extending therapy over comfort care, increasing anxiety, depression or dissatisfaction. The reasons for this lack of information might be dual, with some patients actively avoiding finding out overly detailed information, reflecting the tension between wanting information, but not wanting bad news and with physicians providing incorrect information or none at all.

In order to expect the patients to tell their fears, clinicians must have a ‘preparation dialogue’ [8] with end-of-life patients and their relatives may be used to probe specific patient concerns and help patients to prepare practically, personally and socially in relation to dying process and the welfare of their families. Kelly et al. (2008) [13] findings indicate the significant challenges facing clinicians in discussions with patients and families about death, to exploring the patient’s emotional responses to terminal illness and spiritual concerns for the patient and family, its qualitative research indicates important tasks in the training and clinical support for doctors providing palliative care.

Conclusion

Our review concludes that it is necessary to include the family into the routine of end of life care; studies show that the holistic view of the patients involves their family wellbeing. They must have spiritual guidance, psychological and social help, and also a good and open communication with the health professionals.

To this sensitive approach clinicians shall have information on how to deal with this situation, not only in intensive care units and hospices, but also all the levels of assistance. We suggest that further studies should focus on developing activities that help improve the involvement of families, patients and the health team, and to prove at a multi central research how caring for the family members/care givers at end of life care impacts at the patients.

References


