End-of-life care across Southern Europe: A critical review of cultural similarities and differences between Italy, Spain and Portugal

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Abstract

Evidence from a range of sources demonstrates that end-of-life (EoL) care practices and preferences vary across countries; culture is consistently one of the main explanations given for this. In order to understand how cultural factors are used to explain similarities and differences in EoL care between Spain, Italy and Portugal, database and hand searches were performed and cross-cutting core themes identified. Similarities included higher proportions of people who wished to die at home than actually died at home, a persistent trend for partial disclosure in Italy and Spain, low use of advance directives, and low incidence of all medical EoL decisions (with the exception of medical EoL decisions).

Abbreviations: ADs, advance directives; DNRs, do-not-resuscitate orders; EoL, end-of-life; ICUs, intensive care units.

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1. Introduction: different approaches to culture in EoL care

There is growing interest to integrate the cultural dimension in EoL care [1–3]. However, culture is a broad, multilayered concept [4–6] and the approach taken to understanding culture influences its potential contribution to improving EoL care. There are at least three different perspectives on the relationship between culture and medicine. These three perspectives can also be distinguished in the context of EoL care.

The most common approach links culture to the ways different groups of lay people understand and face health, illness and death: it is considered important to understand the culture of specific patient groups in order to improve their access, and adherence to medical services and treatments. This was the first approach to be used in medical anthropology and, nowadays, in western countries, it is mainly used to understand the behaviour of immigrant, minority and ethnic groups and is associated with clinical notions of ‘cultural competence’, ‘cultural sensitivity’ or ‘cultural awareness’ [7–9]. Nevertheless, it can also be associated with patients’ other characteristics: illness related practices may also differ depending on age, gender, urban or rural origin, etc.

A second perspective, or group of perspectives, recognizes medicine itself as a cultural system with a specific understanding of disease, human body, and cure [4]. It also acknowledges the differences between specialties within medicine in the way they understand disease, treatment and care [10]. This is the perspective of those who explain palliative care as a ‘new culture’ within medicine [11], whose aim is to give total care to patients approaching EoL, and thus, differs from the main curative, death-denying approach of some other specialties.

Finally, a third perspective explores the geographic, social and cultural contexts of both patients and medical systems [12]. This approach is helpful in understanding why supposedly neutral, scientific and medical practices vary from place to place. This is the approach to culture applied in this article.

Evidence from a variety of sources suggests that many EoL care related practices, such as pain management [13], medical EoL decisions [14–16], and disclosure of life threatening diseases [17,18], vary across regions of the world, countries, or regions within countries. The influence of culture is consistently given as one of the main explanations for this variability [13–18]. In these comparisons, Italy, Spain, and Portugal are usually grouped together as part of ‘Southern Europe’ [19,20], the ‘Mediterranean area’ [21,22], or as representing ‘Latin’ [23], or ‘Catholic’ [14,24] culture. Other countries, such as France and Greece, share some, but not all, of these characteristics. When grouped together, they are frequently contrasted with other sets of countries, such as those defined as ‘Northern European’, ‘Anglo-Saxon’ or ‘Protestant’.

This article aims to understand how, in the literature, cultural factors are used to explain the similarities and differences between EoL care practices and preferences in Spain, Italy and Portugal. First, the preferences and practices in several EoL care domains in Italy, Spain and Portugal are synthesised, with a focus on the similarities and differences found between the three countries. Later, the explanatory potential and limits of the cultural factors, as they are used in the literature, are discussed. Throughout, special attention is given to qualitative research as a means to understand the role of culture in EoL care in different countries.

2. Design

This critical review of the literature was undertaken as part of the EU (FP7) supported PRISMA programme which aims to inform best practice and harmonise research in EoL care for cancer patients across Europe [25]. PRISMA incorporated a work package on the influence of culture on EoL care in different European countries.

Country-specific searches were performed in eight international and four Spanish databases (Table 1), using a set of search terms for the country, EoL care, and cultural factors. The search terms for cultural factors were chosen with the aim of retrieving articles concerning EoL care where cultural and social factors were sufficiently relevant to be referred to in the title, abstract, topic or key words, and articles which used qualitative, and particularly anthropological techniques (Table 1).

Reference searches were also undertaken using articles retrieved from database searches. In addition, key Spanish and Portuguese palliative care and cancer web pages, the Spanish medical anthropology bibliography compiled by Perdiguero and Comelles [26], and the abstracts from a number of Spanish and European medical anthropology conferences were also consulted (Table 1). Furthermore, a number of articles were recommended by experts participating in the PRISMA project’s network of experts in cultural issues in EoL that was set up in parallel to the literature reviews. Articles in English, Spanish, Portuguese and Italian were included.

In the final selection of articles, original studies that focused on ‘culture’, operationalised as the perceptions, opinions, understandings, knowledge, preferences, attitudes, practices and behaviours of the different actors involved in...
EoL care, were included. Reviews, overviews, theoretical proposals, reflections and opinion pieces with a cultural or societal perspective on EoL care were also included. For the purpose of this article, a cross-cultural comparison between the three countries, studies that looked at the cultural preferences and practices of specific patient groups such as migrants or ethnic minorities were excluded (14). Articles that only focused on the cultural singularities of palliative care as compared to other specialties and in the context of Western societies relation with death were also excluded (20).

Cross-cutting themes and their subcategories were identified during data extraction and used to code the articles’ content. The results for each country were systematized in data extraction tables, and integrated with qualitative meta-synthesis: the data obtained for the three countries were compared to highlight similarities and differences, and the cultural explanations given for them [27].

3. Results

One hundred and twenty-one documents for Italy, 32 for Portugal and 130 for Spain were available for full text assessment. The majority of these articles were original studies. The rest were mainly overviews, opinion pieces or critiques. Only four reviews were identified and two of them were related to EoL care (Fig. 1). Nineteen articles included data from two or three of the countries.

The main topics that arose from the search process as relevant for a cultural analysis were: setting of care and death, communication and disclosure, medical EoL decisions, knowledge and attitudes towards care and death, and informal caregivers.

3.1. Setting of care and death

In 2005, the number of healthcare centres offering palliative care services available per million inhabitants was six in Spain, four in Italy and one in Portugal, far fewer than in Belgium and Sweden (12), the UK (15), or Iceland (20) [28]. Both in Italy and Spain, home care teams constituted the main palliative care resource available. The second main palliative care resource varied across the countries. In Spain, inpatient units were most common. In Italy, hospice care had been increasingly common: the number of hospices rose from four in 1996 [29] to 90 in 2005 [28] mainly due to the influence, since 1999, of new national and regional policies on palliative care [30]. In Portugal, development of palliative care services and research began later than in Spain and Italy and palliative care services are less widespread. In all three countries, important regional inequalities in the distribution of EoL services and were found [28,31,32].

In each country, studies of the general population revealed low expectations of palliative care service provision. Only 36% of respondents to an Italian survey in 2000 knew the meaning of the term ‘palliative care’ and 85% did not know of any palliative care centre [33]. A study from Portugal in 2008, revealed greater knowledge of the meaning of palliative care; although half the interviewees were familiar with the term, they believed that palliative care was not implemented in the majority of public hospitals, and that bigger development of palliative care was needed [34]. Finally, in Spain, in 2009, less than one third of the population considered that the health system had sufficient resources to care for terminal patients [35].

The importance of home care in these countries could be expected to influence the place people die. Among the three countries, countrywide statistical data on place of death could only be found for Portugal: in 2005 almost one-third of all deaths occurred at home [36]. In Spain, the identified studies presented a range of percentages of home deaths: from 22% for elderly people who died in Catalonia in 1998 to 50% for terminal cancer patients in Asturias in 1995 [37–43]. Finally in Italy, the proportion of home deaths among cancer patients ranged from 36% for all 2002 cancer death certificates in Tuscany, Emilia-Romagna and Milan [44] to 58% in a mortality follow-up survey with nationally representative data from 2002 and 2003 [45]. Italian percentages for non-cancer home deaths were lower [15,44,46]. Place of death was found to be related to age, gender, socioeconomic status, cause of death, and region within the country, rural or urban environment, and the resources available in the area [38,40,43–46].
Within Italy, region was found to be a determining factor in place of death: in the south, where health services are less widely available, the percentage of home deaths increased to 94% for cancer patients [45,47].

Good practices in EoL care include people dying in their place of preference. However, both in Spain and Italy, the proportion of healthy people who would like to die at home was found to be higher than the proportion who actually died at home [35,43,46,48–52]. Only one recent Spanish survey revealed higher preferences for specific care settings for terminal patients or hospitals (50%) than for home death (45%) [35]. Patients and caregivers reported higher preferences for home care, reaching 94% in Italy [43,45,53]. Spain is the only country where doctors’ opinions regarding place of death have been frequently sought and the included studies revealed a greater consensus among healthcare professionals than among the general public that the home is the ideal place of death [54–57]. Moreover, in three of the reviewed articles, the authors (health professionals) assumed that the patient’s home is the ‘ideal’ or ‘correct’ place for EoL care [37,40,41].

3.2. Disclosure and communication

Among the reviewed articles, disclosure of information regarding diagnosis, prognosis and treatments was found to be the most frequently discussed topic (27 articles from Spain, 46 articles from Italy and 8 articles from Portugal). International comparisons describing southern European countries as examples of partial and non-disclosure [17,18,58] were the starting point of this interest, and reviews on the subject have been published both for Spain, in 1998 [59], and Italy, in 2004 [60].

The complexity and detail of measurement of disclosure and awareness has increased since the first studies in the late eighties. It is therefore important to differentiate between: the information given to the patient, his/her awareness of the situation, and the preferences of the general population, patients, relatives and/or doctors; communication about diagnosis, prognosis or treatments; and, methods, subjects and tools used to collect the information.

Spanish and Italian awareness studies published between 1994 and 2009 suggested that the partial disclosure trend persisted throughout these years. In Italy, direct assessment studies of patients’ full awareness of cancer diagnosis and prognosis showed that it remained at under 45% [61–69]. Meanwhile, the relatives interviewed in the cancer mortality follow-up survey from 2000 considered it to be higher: around 63% [70]. In Spain, full awareness of diagnosis ranged from 19% to 64%, depending on the disease, questions and design of the study. There was not however a clear trend over time [37,51,53,71–74]. In both countries, awareness was found to be higher than the information given [59,63,70,71], and more information about diagnosis was given than about prognosis [61,63,70,72]. Significant regional differences were also described in Italy [70].

Other findings, however, suggested that preferences may be evolving towards open disclosure. For example, the percentage of the Spanish population that would like to know their diagnosis and prognosis in case of terminal illness has increased over recent decades [35,49–51,55,75], and the percentage of Italian doctors who considered full disclosure ideal has also increased [76–79]. Intermediate positions were also found; for instance, the majority of Spanish doctors stated that they would inform the patient only in certain circumstances or if requested by the patient [54–57,75,80–83]. Two main obstacles to the communication of bad news were identified in Spanish health professionals’ discourse on disclosure: acceptance of the wishes of the family, hence tolerating the ‘conspiracy of silence imposed by the relatives’, and feeling uncomfortable and unprepared to give bad news. Doctors in hospitals were found to be more comfortable communicating bad news than doctors in primary healthcare settings and nurses, who often consider the communication of bad news to be doctors’ responsibility [84].

In Portugal, only two studies, both conducted in Porto, in 1999 and 2005, which addressed awareness were found. Both
studies described greater patients awareness (60–69%) and desire for information than in Spain or Italy [85,86].

Different interpretations of the trend for partial or non-disclosure in Italy and Spain were found in the literature. In bioethics, disclosure and the doctor/patient relationship was framed within the theoretical debate concerning the tension between the bioethical principles of ‘autonomy’ and ‘beneficence’ [18,60]. Some authors considered that an excessive preoccupation with individual autonomy does not capture the essence of the patient–doctor relationship: asymmetry of power and patient vulnerability [87]. Yet, the emphasis on beneficence could leave patients unprotected from paternalistic intrusions by family members and doctors [58,88]. In order to find the balance between autonomy and beneficence in practice, bioethicists pointed out the variability of personal situations and cultural context, and the need to act with cultural sensitivity/competence [89,90].

Other bioethical and psychological studies focused on the process of communication and the meanings of ‘truth’. Communication was described as a complex, dynamic process that accounted for differences in the awareness and wellbeing of the patient [62,91–97]. Truth was understood as a relational state [87,98,99], and in Catholic countries, communication was considered to be influenced by the Catholic tradition of gradual truth giving [24]. Overall, different communication strategies, shared between doctors and patients, were described. In countries such as the US it was considered that patients needed to be explicit if they did not wish to know this information [100], whereas in others like Spain, patients would not be given all the information if they did not ask for it explicitly [101].

Finally, non-disclosure in Italy was also understood as a way to maintain hope and continuity of social life [24,102–105]. An anthropological study described life in Tuscany as an unpredictable flow of events shared by all members of society. Amidst this flow of events, the most vulnerable were protected and cared for. Avoiding the certainty inherent in doctors’ diagnosis/prognosis was considered to avert the separation of the individual from social life and, therefore, an anticipated social death [104,105].

3.3. Advance directives (ADs)

Important legal differences were found regarding ADs between Spain, Italy and Portugal. In Italy ADs are not legally recognised. Recently a non-binding law has been proposed and there is intense debate, influenced by public opinion, concerning a number of high-profile cases [106–109]. In Portugal, the Council of Ethics has produced some guidance documents. In some instances the documents contradict one another: although ADs are considered binding in the case of a persistent vegetative state, ADs regarding blood transfusions are not binding [110]. In contrast with Italy and Portugal, in Spain, the legal and administrative development of ADs is one of the most advanced in Europe [111].

Among the three countries there were also differences in professionals’ attitudes towards ADs. In 2002, most Spanish doctors considered the implementation of ADs policies in Spain to be a positive development [112], whereas Italian oncologists and doctors had little knowledge of such documents, and at least half the oncologists and doctors were not inclined to respect patients’ previous decisions [113–115].

The general public’s knowledge and use of ADs was reported to be limited in both Spain and Italy (no data was found for Portugal). One study, carried out in Italy in 2001–2002, showed that less than 5% of patients had written ADs [15]. In Spain, the general population had very little knowledge about ADs [116–119] and in 2009 only 2% had written ADs [35].

Two issues that, as of yet, have received very little legal development, were identified in research from Spain: the idea that the process of thinking and talking about EoL decisions could be more useful than formal written living wills [116,117,120]; and the relevance of proxy decision-makers, which nowadays are usually designated verbally [119,121].

3.4. Medical EoL decisions

Most of the reviewed research on EoL decisions was carried out with doctors (37 original studies), especially in Intensive Care Units (ICUs) (9 original studies). There were fewer studies conducted with the general population (8 original studies) or patients (2 original studies) and none with patients’ family members.

International studies carried out in ICUs, showed that Do-Not-Resuscitate orders (DNRs), withholding or withdrawing treatment, and doctor-assisted death, occurred less often in Southern Europe than in Western and Northern Europe [19,122–125]. Nevertheless, national level questionnaires with more representative samples suggested that DNRs, withholding and withdrawing treatment practices in Portugal and Spain are used more often than the results of international questionnaires suggest [126–128]. Contradictions between the findings from international and local/national questionnaires regarding the involvement of family members in EoL decisions made in ICUs were also identified. International surveys reported that discussion with patients and family members regarding DNRs, withholding and withdrawing treatment occurred less frequently in southern European countries compared to other countries [123,125], whereas national and local questionnaires reported that the majority of the doctors involve or consider that family members must be involved in decision-making [79,114,127].

The variety of ways in which medical doctors were questioned makes it difficult to compare their acceptance of euthanasia across the three countries. In Italy, the findings of three studies from the nineties indicated that between 15% and 18% of doctors were in favour of euthanasia [129–131]. However, when asked about specific situations, this percentage increased to between 32% and 40% [115,130,132]. A Portuguese study from 2009 reported that up to 39% of oncol-
ogists favoured the legalisation of euthanasia [133]. Whereas a Spanish study from 2002 indicated that up to 60% of doctors believed that laws should be changed to allow patients to request and receive euthanasia and/or assisted suicide in the case of terminal illness [112]. Across the three countries, religion, sex, palliative care experience and years of practice were found to influence doctors’ opinions [115,130–133].

Europe-wide surveys of the general public found that Italy and Portugal were among the countries with the lowest acceptance of euthanasia, whereas Spain occupied an intermediate position between the low and high acceptance countries [14]. In Italy, religion played an important role and the difference of opinions between believers and non-believers was higher than in the other countries [14]. Spain and Italy were found to have experienced a greater-than-average increase in euthanasia acceptance between 1981 and 1999 compared with a group of 12 western European countries [134]. Spanish surveys showed that acceptance of euthanasia in the specific case of terminal illness has been over 60% since 1995 [35,135,136], higher among younger population groups [137], and reaching 81% in 2009 [35]. Spanish studies also demonstrated that the majority of the population was opposed to the use of life support measures in a situation of serious disease with no possibility of recovery [35,52,138].

Among the studies identified from the three countries, more research articles were found concerning euthanasia (17), than assisted suicide (6). Both the general public in Spain and doctors in Portugal were more in favour of regulation for euthanasia than assisted suicide [35,133].

Terminal sedation was more accepted than euthanasia or physician-assisted suicide [23,35,113] and more widely implemented in Italy than in other western European countries [139]. In the view of some experts, this was regarded as consistent with the Catholic tradition of opposition to euthanasia, but acceptance of pain relief to the point of death might be accelerated [23,80,126]. Moreover, terminal sedation was also considered consistent with the traditional Spanish perception that unconsciousness is the 'best way out' [23,140]. However, the debate continues: in Spain legal proceedings against the Leganés Hospital Emergency Unit have demonstrated that the boundary between euthanasia and terminal sedation is not totally clear [141], whereas some Italian authors have recently argued that sedation does not hasten death [22,142]. Part of the controversy relates to the use of terminal sedation to manage existential and family distress, which seemed more common in Spain than in other countries [143]. In Italy and Portugal, although some authors agreed that it might be difficult to distinguish pain from other kinds of suffering, terminal sedation for non-physical symptoms was rarely used [22,142,144].

3.5. Knowledge and attitudes towards care and death

The attitudes of the general public and health professionals towards death were an important topic of study. In Spain, several articles attempted to measure ‘death anxiety’ among health professionals [145–148], and among members of the general public [149–151]. Health professionals’ anxiety tended to diminish with greater experience, but no clear relationship was found in either group between ‘death anxiety’ and religion [147,151]. In Italy, one study identified, between 1989 and 1999, a progressive secularization of the values around death [48]. Italian practising Catholics and non-believers were found to share concerns about the control of pain and symptoms, and a preference to die in a calm atmosphere and in the company of close relatives and friends [46,48,152]. The concerns and fears related to pain and the process of dying were also pre-eminent among both health professionals [153–156] and the general public in Spain [50,51,149,150,157].

Pain control strategies were mainly analyzed in articles from Italy (13 articles). The articles reported that a significant proportion of patients in studies from different parts of Italy failed to receive appropriate pain treatment [158–163]. These results from Italy are consistent with the European trend: the use of moderate doses of morphine; one third of patients assessed as in pain; and considerable variation in pain medication choices between countries [13]. In 2006, both Italy and Spain were below the European and global average of morphine consumption per capita, whereas Portugal was above the average [164]. Of the drugs used to alleviate symptoms with life-shortening as a possible side effect, Italian physicians were also found to administer opioids less often than their colleagues in Belgium, Denmark, The Netherlands, Sweden and Switzerland [165].

Different suggested explanations for low pain management included: lack of knowledge about adequate pain treatment [13]; underestimation of pain by professional caregivers [166]; health professionals’, patients’ and families’ opiophobia [13,166–169]; Catholic ideals of pain endurance [167–169]; and problems of communication related to a lack of disclosure [170]. Opioid over-regulation was not a major limitation in the three countries [164]. In Italy, health professionals’ knowledge about pain and analgesics was found to have improved between 1996 and 2003, and was better in emergency and anaesthesiology units than in the other units, and better among doctors than nurses [164,171–173]. In Spain, in the mid-nineties, nurses and nursing students did not show proper knowledge about pain control strategies [174,175].

Other aspects of palliative care were also found to be problematic for health professionals. In Spain, health professionals reportedly demanded specific training not only in pain management and other technical issues but also in psychosocial and emotional issues related to providing care to terminal patients [169,176–179]. In Italy, even though most of the doctors agreed that dealing with patients’ emotions is important, they found it difficult [76]. General practitioners and older doctors were found to deal better with patients’ anxiety and to be more emotionally protective [76].
3.6. Informal caregivers

In more than 84% of cases from Italy and Spain (no data for Portugal), patients’ relatives (mainly daughters and wives) were identified as the main informal caregiver [38,41,101,180–186]. Women made up more than two thirds of caregivers in Italy, and more than three quarters in Spain [38,41,101,180–186]. Relatives provided care not only at home, but also in the hospitals, with a continuous presence in palliative care units [74]. A European comparative study on patients’ last year of life found that the family was the main provider of support in Italy, Spain and Greece, whereas in other parts of Central and Northern Europe non-kin support was more prominent [21].

Care-giving was reported to have important consequences for relatives’ quality of life: generally they felt overburdened [72,186–190]; lacked free time for themselves or other family members [179,181–183,191]; and, experienced multiple health problems that increased with the time dedicated to home care and with patients’ worsening condition [185]. A nation-wide study in Italy showed that one-third of cancer patients’ caregivers left paid employment, and that care-giving was a significant economic burden for a quarter of them [182,183,191]. Four studies reported a lack of knowledge of available EoL care services and subsidies in Spain (3) [179,187,192] and Portugal (1) [190]. Also, Italian caregivers considered government economic subsidies insufficient and home assistance inadequate [181]. Specific problems depended on the situation of the caregivers: for young women in Portugal, care of their mothers radically altered their life and future plans [193].

Several negative portrayals of caregivers’ contribution to EoL care were found. In Italy, doctors described relatives as a major factor in preventing the ideal practice of full disclosure and limitation of non-useful treatment [77,78,114]. Two concepts with negative connotations were identified: the ‘conspiracy of silence’ in Spain and Italy [60,73,84,143], the partial or non-disclosure which is frequently attributed to family members; and, ‘claudicación familiar’ (family surrender) in Spain, when patients die in hospital and not at home (the ideal place of death according to health professionals) [194].

3.7. Qualitative research

Of the 208 original studies considered relevant to understanding the role of culture in EoL care, around one fifth (44) used qualitative techniques. The number is even lower (34) if studies that just used content analysis to design or obtain quantitative data and only analyzed open questions from mainly quantitative questionnaires are excluded. Of the 34 articles one was a Delphi study, looking for consensus regarding euthanasia in Spain [195]. Another study used an ethnoscience approach to analyze patients’ perceptions of fatigue [196]. Two studies used case studies [193,197] for very specific patients groups. Twenty-four used semi-structured or in-depth interviews and group discussions, with a large variety in their topic of interest and quality of methods: three for Portugal [189,198,199], four for Italy [92,97,152,200], sixteen for Spain [81,84,117,120,140,187,192,201–209], and one international study that includes data from Spain and Italy [210]. Finally, six articles (using data from four different studies) included long-term participant observation. Only one [168,169] of these four studies was totally focused on EoL care and the other three were wider studies on cancer [101,104,105] or intensive care units [211]. Three of these studies were focused on professionals’ perspectives [101,168,169,211], and only one took a holistic view and included the perspectives of patients and relatives [104,105].

4. Discussion

The different palliative care systems and histories in Spain, Italy and Portugal are embedded in cultural contexts that share some characteristics but also have important differences. The main cultural factors, shared between the three countries and stressed in the literature, are the importance of family and community ties, and Catholicism.

4.1. Shared cultural factors: strong family ties and Catholic tradition

Given the stereotype of these countries as places where social and family links are stronger than in the rest of Europe, it is often assumed that in Portugal, Italy and Spain, people die mainly at home, surrounded by their loved ones. However, only in Italy, particularly in the southern region, does the majority of the population die at home. Preferences for home as the place of care and death are however higher, and this suggests that there may be practical barriers to providing EoL care at home. Nevertheless, four articles emphasize the relationship between the lack of health and social services and home deaths, especially in Italy [21,31,45,47], and in Spain, half of the population has recently expressed a preference for hospice or hospital care [35]. In this context, where informal caregivers are also overburdened, preference for home death should not be used as a justification for the sole prioritization of domiciliary care that may be more related to attempts to limit healthcare expenditure [23].

The importance of diagnosis and prognosis disclosure in the Spanish, Italian and Portuguese literature contrasts with the Spanish, Italian and Portuguese literature contrasts with the findings from reviews of the literature on culture and EoL care in other European countries which give far less attention to the issue [27]. Again, the role of the family and the community has been considered crucial to explaining southern European perspectives, and the traditional Catholic approach to truth has also been considered relevant. The difference between, on one hand, the opinions and preferences of both health professionals and the general public (which have increasingly moved towards full disclosure) and on the other hand,
the real practices (the development of which is not clear) demonstrates the importance of not limiting social and cultural research in EoL to preferences and opinions. It is also necessary to describe and analyze practices and explain why they vary from stated preferences and theoretical models.

In Spain, there are also differences between common practices, legal ideals and professionals’ opinions regarding ADs. In terms of disclosure and ADs, the ideals, influenced strongly by the hegemonic ‘Anglo-Saxon’ values of medicine, tend to prioritize autonomy and rational choice, whereas in practice, the logics of care [212] and beneficence seem more prevalent. Also, in both cases, analysis of and reflection on the differences between ideals and practice have had an impact on experts’ discussions. Southern European experiences, along with those from other ‘non-Western’ countries, have enriched the debate, uncovering the complexities of patient-doctor communication and decision-making processes, and the importance of the role of family in EoL care.

The principal cultural factor stressed in the articles on medical EoL decisions is religion. Catholicism has been considered a key factor explaining the more pro-life profile of Spanish, Italian and Portuguese medical EoL decisions and the preferences for terminal sedation compared to other EoL medical practices. It has also proved to be a key reason for the differences in opinions and practices of doctors and the general public in the three countries. Only a few articles questioned the role of the family in medical EoL decisions, and it was always from the point of view of the health professionals [79,114,127]. Given the active role of the family during EoL care, there is a need for research on the perspectives of relatives and informal caregivers.

For the general public, pain is the main death-related concern, but it is frequently undertreated. Various articles stress that there are differences in pain management between European countries and recognize that different cultural factors may be involved [13,165,166,170]. In most cases, all European countries share these cultural factors. Nevertheless, there are two suggested factors that may have a specific impact in Italy, Spain and Portugal: Catholic views on pain endurance, and communication problems related to lack of disclosure. Further research is however needed to demonstrate their relevance for EoL care and total pain management in specific contexts.

Up to now, the family and Catholicism have been described as the main explanatory cultural factors underlying characteristics of EoL care shared by Portugal, Spain and Italy. Relatives are involved during the whole EoL care process: caring for the patient at home; accompanying the patient most of the time in the hospital; worrying about the patients’ pain; and participating in disclosure, ADs and other EoL decisions. Catholic traditions have influenced medical paternalism, gradual disclosure, pro-life positions, negative attitudes towards withholding and withdrawal of treatment, preferences for sedation over other EoL options, and approaches to death and pain in these countries. However, reflection on the role of the family and religion in EoL care is not, and should not, be restricted to these countries: family and religion also influence EoL care in all countries, in different ways in every local context. Nor should reflections be restricted to simplistic univocal relations or generalizations for one or several countries. Simplified, limited understandings of culture and cultural factors have repeatedly been shown to have greater political than interpretative power, and run the risk of stereotyping [4,213].

4.2. Cultural analysis and improvement of EoL care in Italy, Spain and Portugal

There are three ways that an analysis of culture in Italy, Spain and Portugal could contribute to the improvement of EoL care. Firstly, a deeper analysis of the role of both family ties and religion is required.

In a context where family structures are changing and informal caregivers are clearly overburdened, an improved understanding of family dynamics during EoL care is crucial to improve care. Relatives cannot be excluded from the debate, as sometimes happens when Anglo-Saxon practices, such as ADs, are adopted uncritically in these countries. The negative images of caregivers’ contribution to EoL care also deserve more detailed study, as they point to a new example of ‘victim blaming’. Finally, given the preponderance of female caregivers, it is necessary to include a gender perspective in analysis.

Regarding Catholicism, at least two different ways of understanding religion have been included in this review: the Catholic cultural tradition, which involves a number of values that are clearly shared by many with a Catholic identity, but have also permeated society in general [23,24,80,126,167–169]; and the personal experience of religion, which may include a person’s self-definition as Catholic [14,115,130–133,151], and the degree to which religious practices are followed [147]. The anthropological perspective differs from the use of religion in quantitative research as an isolated variable based on self-definition and from religious and experiential views. Here, these two understandings of religion are considered to influence each other, to be interrelated and difficult to separate. With this perspective in mind, it is essential to reflect on the process of secularization: nowadays non-believers make up a significant proportion of the population in these countries. The tensions between conservative religious and progressive secular groups are evident in the debates concerning high-profile medical cases in both Italy and Spain, and it is important to analyze the complex intertwining in these countries of politics and religion.

Secondly, cultural analysis should not be restricted to the family and religion, which are topics widely cited in the literature. Anthropological analysis is useful to understand EoL care in the context of social relations, needs, values, interests and power positions of the different actors involved in the process. This analysis should be carried out in specific contexts and take into account differences and inequalities in

5. Concluding remarks

A synthesis of the cultural factors involved on EoL care in Southern Europe is always at risk of reinforcing a simplified, stereotypical portrayal of a place where most people are cared by their families and die at home without being aware of their diagnosis, where palliative care services are less developed, medical EoL decisions occur less frequently, and patients are less involved in decision making than in northern Europe. Some of these characteristics can be used appropriately to describe specific situations in particular areas of Southern Europe, but the reality is far more diverse and complex.

In this article the similarities and differences in EoL care between Italy, Spain and Portugal, and within each of these countries have been described. Gaps in research on culture and EoL care have also been identified and a cultural approach, which could be useful to avoid stereotypes and to understand and improve EoL in these countries, has been suggested. A locally-focused approach that interrelates values and practices, does not exclude any of the actors in EoL care, including the family, and dialogues with the quantitative literature can help to improve communication, decision-making and care processes in concrete and flexible social, cultural and institutional contexts.

Conflict of interest

The authors of this review have no conflict of interest.

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Busquet Durán X, Pujol Sabanés T. Los estudiantes de enfermería


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