Explaining health care providers’ perceptions about the integration of palliative care with primary health care; a qualitative study

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Abstract
Background: Easy access to palliative care is one of the basic needs of cancer patients, and this can be achieved by providing such services at the community level. One approach to provide community-based palliative care is to integrate it with primary health care (PHC). Considering the antiquity and extension of the PHC system in Iran and the importance of being aware of stakeholders’ views in order to integrate a palliative care provision model into a country’s health care system, we aimed to explain health care providers’ perception of the integration of palliative care with PHC.

Methods: The present qualitative research was conducted using the conventional content analysis method in Iran from October 2016 to July 2020. The participants of the study included the stakeholders involved in providing palliative care to cancer patients, as well as PHC system experts. The participants were selected purposefully using the snowball sampling method. Data were collected through holding 21 semi-structured interviews and one focused group session and analyzed based on the method proposed by Lundman and Graneheim.

Results: Qualitative data analysis revealed three main categories and ten subcategories. The main categories included the health system’s structure as an opportunity (with the subcategories of employing the network system for providing health services, establishment of a referral system, and establishment of the family physician program and manpower diversity), requirements (with the subcategories of the position of home care centers and their relationship with PHC, opioid use management, equipment management, financial support, and legal issues), and outcomes (with the subcategories of facilitated access to services and good death).

Conclusion: Iran’s health system possesses adequate infrastructure for providing palliative care to cancer patients within the context of PHC. Beside available opportunities, there are also problems that need to be resolved so that families can meet their patients’ care needs and provide them with an easy death by having access to home-based palliative care.

Keywords: Community health care, Home health care, Advanced cancer, Palliative care Terminally ill, Qualitative study

Background
Cancer is known as the second leading cause of death worldwide [1]. The International Agency for Research on Cancer estimated a global cancer incidence rate of 19.3 million cases with 10 million deaths in 2020 [2]. Cancer is also the second leading cause of death in Iran [3].
total number of cancer patients in Iran was 110,115 in 2018, causing 55,785 deaths in the same year [4].

Palliative care, as one of cancer patients’ needs, has been recognized by the World Health Organization (WHO) as a warrant for improving the quality of life of end-stage cancer patients [5]. Palliative care is a comprehensive care method that considers the whole existence of humans, including their physical, psychological, social, and spiritual dimensions [7]. This type of care not only helps patients live their lives until the last moment, but also supports patients’ families during the disease course, at the time of the patient’s death, and even after death, helping them better and more calmly and peacefully accept the phenomenon [6].

According to a report by the WHO, palliative care and primary health care (PHC) have a number of common principles such as continuous care, social responsiveness, respecting patients’ values, and paying attention to the patient’s care needs in the presence of his/her family members. So, the WHO recommended in 2016 that in order to achieve the goals of sustainable development and universal health coverage (UHC), health systems must integrate community-based palliative care into PHC programs [6] so that these services can be provided to patients and families by the clinicians who are in close contact with them [7]. According to Astana Declaration, PHC is the foundation of a sustainable health system for UHC and health-related Sustainable Development Goals (SDG), allowing to meet health care needs of people during their lifetimes via delivering comprehensive preventive, premonitory, curative, and rehabilitative services, as well as palliative care [8].

The advantages of community-based palliative care include the facilitation of patients’ access to health care [9], promotion of the quality of life of patients and caregivers, reduction of hospitalization length [10] and the rate of referral to emergency departments [11], and finally an increase in the number of at-home peaceful deaths [12].

Iran’s health system follows a hierarchical referral network, and its services to the community are provided in the form of PHC. In this structure, comprehensive urban and rural health centers are responsible to deliver health services to the people of a specific region [13]. Nevertheless, palliative care, as a novel care approach, has no place in this structure. In fact, such services are provided to the community by private and charity organizations [14]. It is obvious that patients and their families become baffled and experience high distress, confusion, and agony when they encounter advanced cancer. Given the desire of patients to receive at-home health care [15] and the appropriate structure of PHC in Iran, it seems that the integration of palliative care into this structure can greatly help cancer patients and their families receive their required services.

To establish health services in a specific field and to design a care provision model for these services, that field must be well-recognized.

This requires knowing the context in terms of facilities, opportunities, and barriers from the perspective of health care providers. This goal can be achieved by qualitative research. Therefore, this study was designed to explain health care providers’ perception of the integration of palliative care with PHC.

Methods

Design

This was a qualitative study performed via the conventional content analysis approach in Iran from October 2016 to July 2020.

Participants

Participants were selected through purposeful sampling and included oncologists, palliative care specialists, general practitioners, nurses, psychologists, social workers, and PHC system experts who were selected applying a snowball approach by referring to target colleges, home health care centers, comprehensive health care centers, and cancer departments. Sampling continued until reaching data saturation (i.e., the time at which no new codes were extracted from interviews) [16]. In addition to individual interviews, a focus group was held with community-based palliative care experts to enrich the data.

Policymakers should have had roles in policymaking for cancer patients and their families regarding home- and community-based palliative care services for at least three months. Inability or unwillingness to participate in the study were regarded as exclusion criteria.

Data collection

The main tool of data collection included in-depth semi-structured face-to-face interviews. The main probing question for directing the interview varied according to the interviewee’s knowledge, expertise, and experience. For data collection, cancer patients, families, and care providers were identified and selected by referring to the centers and hospitals providing home-based palliative care services. After participant identification and qualification, the time and place of the interview were specified. The duration of each interview ranged from 20 to 35 min based on the interviewee’s mood, patience, and richness of experiences. Face-to-face interviews were held at places at which the participants were comfortable.
Employees generally preferred the workplace while cancer patients and families were interviewed either in treatment centers or at their homes. All the interviews were conducted by a researcher with a Ph.D. in nursing and recorded by a voice recorder. The main questions, which varied according to the participant’s position, expertise, and knowledge, included "Would you please describe your views on the condition of home-based palliative care in Iran?", "How was your experience on home-based palliative care?", "How is it possible to integrate palliative care services into the PHC system for end-stage cancer patients?", "What are the challenges and opportunities?", and "How one can facilitate this process?". The researcher used probing questions to direct the interview towards the research objectives.

Data analysis
Data analysis was performed simultaneously with holding the interviews based on the Lundman and Graneheim approach [17]. After each interview, the interviews were immediately transcribed verbatim and read carefully several times. The initial codes were extracted from the data and compared with each other to integrative related ones and form subcategories and categories based on their differences and similarities.

Trustworthiness
Credibility, dependability, confirmability, and transferability were used according to Guba and Lincoln to ensure data rigor [18]. For acceptability, the participants were recruited in a way to obtain maximum diversity in terms of demographic variables such as age, gender, literacy level, income, and the type of cancer. After the initial analysis and extracting the primary codes, the participants were asked to confirm the accuracy of the codes and their interpretations. If the extracted codes were contradictory with the participants’ opinions, discussions and exchange of views were conducted to reach consensus on the initial codes. For transversality, the researchers tried to prepare an exact report of the details of the research process. For conformability, the researchers consulted with two faculty members who were experts in qualitative research to reach an agreement on the codes and their classifications.

Ethical considerations
All the methods were performed in accordance with the relevant guidelines and regulations of the declaration of Helsinki (ethics approval and consent to participate). The aims and methods of the study were explained to all the participants, and necessary assurance was given to them for the anonymity and confidentiality of their information and audio files. Informed consent was obtained from all subjects. The participants had the right to withdraw during the study or at any other time. The Ethics Committee of Lorestan University of Medical Sciences approved the study protocol (ethical codes: LUMS.REC.1394.57 and IR.LUMS.REC.1398.217).

Results
In this study, data were gathered through 22 face-to-face interviews with 21 participants (one of the participants was interviewed twice), as well as one focus group session (Tables 1 and 2). None of the participants don’t refused to participate in the study. Qualitative data analysis revealed three main categories and ten subcategories (Table 3).

The health system’s structure as an opportunity
Health care providers pointed out the capability of Iran’s health system to provide community-based health services to cancer patients. The participants acknowledged fundamental reforms in the structure of Iran’s health system after the launch of the network system, and the fact that people’s access to health services was facilitated by establishing comprehensive health service centers in residential areas. In the context of this system, patients can be referred from the first-level services to higher levels. Likewise, cancer patients can be provided with palliative care services in this structure. Three sub-categories were identified within this class, including employing the network system for providing health services, establishment of a referral system, and establishment of the family physician program and manpower diversity.

Employing the network system for providing health services
Experiences of the participants indicated that Iran’s network system possessed perfect opportunities to provide palliative care services to cancer patients. In this system, all members of the community have an electronic health record that is accessible to care providers in comprehensive health service centers. One of the experts stated: "...In health centers, the population is characterized and known..." [1]. Another participant mentioned: "...Cancer patients can receive community-based care in the context of PHC..." [18]. According to the participants’ experiences, a platform was launched in the context of comprehensive health centers in recent years, containing the health information of all people, to which caregivers can have access if needed. Therefore, known cases of cancer, as a member of the community, can be tracked and monitored using this platform. In this regard, one of the health experts noted: "...the integrated health system (i.e., SIB) has been launched in recent years, and it is possible to have access to all households’ health data...." [19]. The participants also admitted that within the context of this
network, patients’ records can be transferred from the third and second levels to the first level of referral; this was stated by one of the participants: “... the SIB system should be linked to the patient's information at hospitals, and if this is fulfilled, it would facilitate referrals and patient care programs...” [18].

### Establishment of a referral system

According to the participants, the referral system is under the attention of health policymakers in current times, and good steps have been taken towards developing and completing such a system. Actually, it has been launched in villages and suburbs for several years now. As

| Number | Age (years) | Degree of education | Work experience (years) | Position |
|--------|-------------|---------------------|-------------------------|----------|
| 1      | 62          | Specialist          | 30                      | Active in policy making and home care provision |
| 2      | 48          | Specialist          | 25                      | Active in policy making and home care provision |
| 3      | 38          | Nurse               | 16                      | In charge of coordination of home-based palliative care services |
| 4      | 52          | Ph.D. in nursing    | 26                      | Lecturer of public health nursing |
| 5      | 47          | Nurse               | 19                      | In charge of the palliative care ward |
| 6      | 46          | Theologist          | 6                       | Religious expert |
| 7      | 45          | Nurse               | 16                      | Caregiver of home-based palliative care |
| 8      | 43          | Medical doctor      | 16                      | Palliative care physician |
| 9      | 43          | Social worker       | 11                      | Social worker |
| 10     | 32          | Employee            | -                       | Caregiver |
| 11     | 25          | As assistant with a diploma | 3 | Assistant |
| 12     | 34          | General practitioner | 4                       | Palliative care physician |
| 13     | 53          | Nursing assistant   | 8                       | Nursing assistant in the palliative care ward |
| 14     | 33          | Psychologist        | 3                       | Psychologist in home-based palliative care |
| 15     | 29          | Psychologist        | 5                       | Psychologist in home-based palliative care |
| 16     | 40          | Bachelor in nursing | 17                      | Home care nurse |
| 17     | 38          | Bachelor in nursing | 10                      | Home care nurse & in charge of a home care institute |
| 18     | 38          | Ph.D. in health policymaking | 6 | Faculty member in the field of health and health promotion |
| 19     | 42          | Ph.D. in epidemiology | 17                   | Staff of the Health Deputy |
| 20     | 37          | Pharmacist          | 5                       | In charge of the drugs used in the network system |
| 21     | 54          | Employee            | 24                      | In charge of narcotics |

| Number | Age (Years) | Level of education | Working experience (Years) | Position |
|--------|-------------|-------------------|-----------------------------|----------|
| 1      | 62          | Specialist        | 32                          | Active in home care provision |
| 2      | 51          | Specialist        | 18                          | Active in home care provision |
| 3      | 48          | Social medicine specialist | 19                   | Representative of the Ministry of Health |
| 4      | 47          | Doctor            | 23                          | An agent of Iranian Health Insurance |
| 5      | 54          | Master degree in nursing | 30                   | Manager of the home care institute |
| 6      | 42          | Bachelor degree in Nursing | 16                   | A home care nurse |
| 7      | 44          | Social medicine specialist | 8                       | Active in home care policymaking |
| 8      | 42          | Ph.D. in nursing   | 15                          | Active in home-based palliative medicine |
mentioned by the participants, providing care to cancer patients during end-life stages can be fulfilled using this referral system in a back-referral manner. Also, the participants clarified that even though the diagnosis of cancer is generally made in clinics and hospitals, the patient should be referred to the first level of the referral system in community-based health centers. One of the participants noted: "...When cancer is diagnosed in specialized centers [tertiary level of referral] ... the patient should be referred to comprehensive health centers for further support [primary level of referral] ..." [19].

The participants reiterated the positive experience of the back-referral system for the management of other diseases such as tuberculosis and leprosy, with patients being referred from higher specialized levels to comprehensive health care centers at the community level. A faculty member who was an expert of health policymaking mentioned: "...For example, patients with tuberculosis and leprosy are usually identified at a higher level by a specialist or a general physician, who then would refer patients to us [primary level of referral] ... meaning a reverse direction...i.e., from a specialist or a subspecialist [tertiary level of referral] to the health center [primary level of referral] ..." [18].

Requirements
According to the experiences of the participants, necessary infrastructure should be available to be able to provide PHC-integrated palliative care. Data highlighted the lack of a characterized and clear link between home care centers and the PHC system, as well as a vague opioid use management approach, which was also true for equipment deliverance and social insurance support. All these can cause legal problems for palliative care provider centers. Under this category, five subcategories emerged, including the position of home care centers and their relationship with PHC, opioid use management, equipment management, financial support, and legal issues.

Establishment of the family physician program and manpower diversity
At the present time, the family physician program is running in rural and suburban areas in Iran and is also on the agenda of the Ministry of Health for large cities. So, the family physicians and care providers who are the members of the teams deployed to these areas can provide care for cancer patients at the end-stage of life. A participant said:"...Every family physician covers a limited population, so any person who needs palliative care can be under the coverage of one of these doctors..." [18].

The position of home care centers and their relationship with PHC
Experiences of the participants showed that for the better use and more benefits of patients from home-based health services, several comprehensive health centers can be brought under the coverage of a single home-care provider team. The participants described the successful experience of the network system in providing maternity services tailored according to each population's specifications and covered by several comprehensive health centers. On this issue, one of the participants stated: "...In this case, [for example] I recall maternity facilities and small

| Categories                                                                 | Subcategories                                                                 |
|---------------------------------------------------------------------------|-------------------------------------------------------------------------------|
| The health system’s structure as an opportunity                           | Employing the network system for providing health services                    |
|                                                                          | Establishment of a referral system                                             |
|                                                                          | Establishment of the family physician program and manpower diversity          |
| Requirements                                                              | The position of home care centers and their link with PHC                     |
|                                                                          | Opioid use management                                                          |
|                                                                          | Equipment management                                                           |
|                                                                          | Financial support                                                              |
|                                                                          | Legal issues                                                                   |
| Outcomes                                                                  | Facilitated access to services                                                 |
|                                                                          | Good death                                                                     |

Table 3 The categories and subcategories emerged from the data
surgery centers ... places where a population of 20,000 is under the coverage of five to six health centers ... on the natural flow of the population .... we launch maternity facilities in one of those health centers [we can follow the same pattern for providing care to cancer patients] ..." [18].

Opioid use management
Among the problems of cancer patients during their last days are pain control and access to opioids. According to the participants, cancer patients monthly refer to the National Food and Drug Administration to receive their required drugs prescribed by a specialist. However, considering the capacity of the network system and the fact that health care providers can have access to patients’ files, it is amenable to provide patients with such medications from nearby comprehensive health care centers. In this regard, one of the pharmacists involved in drug distribution management said: "...Our main objective is to ensure easy, inexpensive, and rapid access to the medications... so we can deliver these medications to comprehensive health centers so that patients can have easy access to them..." [20].

Equipment management
Findings in this study highlighted appropriate equipment management as one of the most important requirements for providing home-based palliative care. As the participants acknowledged, consumables can be provided by patients and their families; on the other hand, health care providers can entrust them with non-consumable equipment. A home-care nurse stated: "...Suction devices can be portable ...before transporting the patient to home, the patient’s bed and equipment must be prepared there ...patients can be entrusted with these equipment..." [17]. Another health expert mentioned: "...In comprehensive health centers, we also need some other essential equipment such as resuscitation kits..."[19].

Financial support
According to the participants’ experiences, one of the most essential infrastructures for providing community-based palliative care is to determine the tariffs and insurance coverage so that patients can access these services without any barrier. Regarding that comprehensive health centers are integrated into the structure of the health system, their tariffs and franchises are predetermined, and even some services in this structure are provided free of charge or at a cheap price. Likewise, palliative care services can be integrated into this structure, so that patients can benefit from these services by referring to and registering at comprehensive health centers in their places of residence. A participant noted: "...All the services provided are free of charge or are delivered at very cheap prices...within the network system, many diseases are covered by these services..." [19].

Legal issues
Data indicated the importance of addressing legal issues for patients, families, and caregivers during the provision of home-based palliative care. According to the experiences of the participants, these people should be aware of their own rights during the process and be committed to observe the rights of others. So, before the onset of the care provision process, families and comprehensive health centers should sign a contract, and all the services provided must be documented. In this regard, a participant said: "...The first thing that brings the center and the patient to feel committed to each other is to sign a contract..." [16].

Outcomes
Data revealed that by integrating palliative care services into the network system, patients’ access to health services is facilitated, leading them to experience an easy and peaceful death. In this category, two sub-categories emerged, including facilitated access to services and easy death.

Facilitated access to services
According to the participants’ experiences, the integration of palliative care into the PHC system facilitates public access to such services. As mentioned by the participants, because these services are provided at a regional scale, access to them is easy for people. On the other hand, by the integration of these services into the network structure, their costs will be adjusted to the minimum as for other services provided in the system. In this regard, one of the participants mentioned:"...When a service is provided within the framework of the PHC structure, we expect it to be either free of charge or very cheap...this means affordable, which is in line with the UHC and WHO uphanded documents...." [18].

Good death
According to the interviewees, a predetermined demise would facilitate the acceptance of death. During their last days, cancer patients are provided with the opportunity to reside at their homes and receive appropriate pharmaceutical and non-pharmacological treatments to reduce their physical and psychological complications so that they can have a better quality of life during this critical period. One of experts in this field underlined: "...patients and their family members should be endowed with peace...a psychologist can assist in improving the quality of death..." [18].
Discussion
In this study, we aimed to explain stakeholders’ perceptions about the integration of palliative care with PHC. Our findings suggested that Iran’s health system was equipped with appropriate infrastructure to facilitate the integration of palliative care services into the PHC and the easy access of cancer patients to such services. However, there are a number of obstacles in this process that need to be reevaluated and revised so that families can have access to palliative care at their homes. This not only leads to the fulfilment of their care needs, but also provides the patient a peaceful death.

According to finding, the PHC is one of the pillars of the health system structure in Iran. Iranians can seek the health services required in each of the three levels of the referral system. In line with the findings of this study, WHO has urged the integration of palliative care aids with all other levels of the community-based services provided by the health system [13].

Our data showed that one of the opportunities provided by the structure of Iran’s health system was the establishment of the family physician program. In this regard, a team of several health care providers with different expertise in comprehensive health centers can deliver community-based palliative care, which of course, this process needs empowering these centers with new manpower and necessary training [19–22].

Our data highlighted the necessity of a close contact between home-based care provision centers and comprehensive health centers so that patients can use their support if needed. Nevertheless, in Iran, there is no systematic relationship between comprehensive health centers and home-based care providers at the present time [23]. Moreover, appropriate decisions should be taken to adjust the number of available comprehensive health centers covered by home care teams in order for them to be able to cover all patients and their health care requirements.

Findings showed that one of the issues that should be addressed in the process of care provision to cancer patients was easy access to opioids. In agreement with this notion, easy access to these medications was highlighted by another study as well [24]. In Iran, opioids are generally provided to patients with incurable diseases either at hospitals or by the national Food and Drug Administration only after prescription by a doctor [25]. According to upstream laws, health centers should provide a platform for delivering opioids to patients. It seems that there is a need for revising some laws related to the distribution of these drugs in Iran. During care provision to cancer patients, the equipment required by these patients, such as beds, mattresses, oxygen generators, etc., should be given special attention and provided along with consumables [7]. Comprehensive health centers are suggested to develop an independent unit in charge of equipment management and distribution and entrust patients with these items, which are returned after the patient’s demise.

Our data revealed that one of the factors that should be considered when providing care to cancer patients was the insurance coverage of the services provided, warranting the easy and cheap access of all cancer patients to palliative care [26]. As comprehensive health centers are governmental organizations affiliated with medical universities, their services should be covered by insurance companies and have pre-determined affordable prices.

Based on our findings, to provide community-based care services to cancer patients, the issues related to their medical rights should be addressed. According to the findings of another study, the process of providing PHC-integrated palliative care to cancer patients can face ethical challenges rooted in a shortage of resources, the lack of knowledge about palliative care, and the lack of support for the cases referred due to staff’s high workload [27]. Therefore, there is a prompt need for enacting new laws and developing appropriate guidelines based on these laws to ensure that the rights of patients, families, and palliative care providers are fulfilled and quality palliative care services are delivered to cancer patients.

An important outcome of this study is the improvement of the access of cancer patients and their families to health care services, which is in line with the goals of sustainable development and UHC programs [6]. Our data showed that by providing comprehensive care, including psychological and spiritual care, patients’ quality of death and families’ preparedness for the event can be improved. Consistently, the results of another study showed that the patients who died without receiving palliative care had more unmet needs [28]. Therefore, when providing community-based palliative care, all human aspects, including spiritual and psychological dimensions, should be considered. Moreover, the patient’s family must regularly receive psychological and spiritual support from the time of disease diagnosis until after the patient’s death.

Some of the limitations of this study include the following. Palliative care and home health service are new approaches of providing care in Iran, dating back to just the recent decade. This means limited access to experts, lack of infrastructure, and scarce centers providing these services, which are the most important limitations of this study. Other limitations included the scattering of experts throughout Iran, rendering communicating with and meeting them difficult. However, this problem was largely resolved by the efforts of the research team.
Conclusion
Data analysis showed that the structure of Iran’s health system had the potential to provide palliative care to cancer patients in integration with PHC, raising numerous opportunities such as the establishment of the family physician program. Also, cancer patients can receive required services in the context of an integrated electronic health system by being referred from community-based specialized centers to comprehensive health centers around their places of residence (as the gateways of this reverse referral system). Beside these opportunities, there are also a number of obstacles for this process, such as the lack of instructions and guidelines, the lack of insurance coverage of palliative care services, and a shortage in competent human resources. These shortages must be addressed by implementing appropriate measures to finally be able to integrate palliative care services into the PHC system so that patients and their families can have easy access to such services at their homes. Consequently, patients can spend their last days beside their families and benefit from a good quality death with the preservation of human dignity along with the support of health care teams.

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Authors’ contributions
HH: investigator; study design, data collection, data analysis, accrual of study participants, and writing & reviewing the manuscript. SH: Health System Research; WHO: World Health Organization, UHC: Universal Health Coverage; SDG: Sustainable Development Goals.

Supplementary Information
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Additional file 1. Raw data.

References
1. Bray F, Ferlay J, Soerjomataram I, Siegel RL, Torre LA, Jemal A. Global cancer statistics 2018: GLOBOCAN estimates of incidence and mortality worldwide for 36 cancers in 185 countries. CA Cancer J Clin. 2018;68(6):394–424.
2. Sung H, Ferlay J, Siegel RL, Laversanne M, Soerjomataram I, Jemal A, et al. Global cancer statistics 2020: GLOBOCAN estimates of incidence and mortality worldwide for 36 cancers in 185 countries. CA Cancer J Clin. 2021;71(3):209–49.
3. World Health Organisation. Profile of Noncommunicable Diseases in Iran. 2018. Available from: https://www.who.int/nmh/countries/irn_top.pdf?ua=1. Accessed 7 Jul 2021.
4. Rebbeck TR. Prostate Cancer Genetics: Variation by Race, Ethnicity, and Geography. Semin Radiat Oncol. 2017;27(1):3–10. https://doi.org/10.1016/j.semradonc.2016.08.002.
5. World Health Organisation. Strengthening of palliative care as a component of comprehensive care throughout the life course. Sixty-seventh World Health Assembly, Geneva. 2014/24. Available from: https://apps.who.int/iris/handle/10665/1623863. Accessed 7 Jul 2021.
6. World Health Organisation. Why palliative care is an essential function of primary health care. Available from: https://www.who.int/docs/default‑source/primary‑health‑care‑conference/palliative.pdf?sfvrsn=ecab9b11_2. Accessed 5 Feb 2019.
7. World Health Organisation. Integrating palliative care and symptom relief into primary health care: a WHO guide for planners, implementers and managers. Available from: http://apps.who.int/medicinedocs/docu‑ments/s23558en/s23558en.pdf. Accessed 10 Jun 2018.
8. Walraven G. The 2018 Astana Declaration on Primary Health Care, is it useful? J Glob Health. 2019;9(1):010313.

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Availability of data and materials
The audio and text files of this qualitative study are available from the corresponding author if needed.

Declarations
Ethics approval and consent to participate
All methods were performed in accordance with the relevant guidelines and regulations of the declaration of Helsinki (ethical approval and consent to participate). The aims and methods of the study were explained to all participants, and necessary assurance was given to them regarding the anonymity and confidentiality of their information and audio files. Informed consent was obtained from all subjects and/or their legal guardians. The participants had the right to withdraw during the study or at any other time. The Ethics Committee of Lorestan University of Medical Sciences approved the study’s protocol (ethical codes: LUMS.REC.1394.57 and IR.LUMS.REC.1398.217).

Consent to publication
Not applicable.

Competing interests
Authors declare that they have no competing interests.

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Abbreviations
PHC: Primary Health Care; HSR: Health System Research; WHO: World Health Organisation, UHC: Universal Health Coverage; SDG: Sustainable Development Goals.

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9. Liberman D, Song E, Radbill L, Pham P, Derrington S. Early introduction of palliative care and advanced care planning for children with complex chronic medical conditions: a pilot study. Child Care Health Dev. 2016;42(3):439–49.
10. Totten AM, White-Chu EF, Wasson N, Morgan E, Kangsara D, Davis-O’Reilly C, et al. Home-based primary care interventions. 2016.
11. Owens D, Eby K, Burson S, Green M, McGoodwin W, Isaac M. Primary Palliative Care Clinic Pilot Project demonstrates benefits of a nurse practitioner-directed clinic providing primary and palliative care. J Am Acad Nurse Pract. 2012;24(1):52–8.
12. Kim SL, Tarn DM. Effect of primary care involvement on end-of-life care outcomes: a systematic review. J Am Geriatr Soc. 2016;64(10):1968–74.
13. ZanganehBaygi M, Seyedin H, Salehi M, Jafari SM. Structural and Contextual Dimensions of Iranian Primary Health Care System at Local Level. Iran Red Crescent Med J. 2015;17(1):e17222.
14. Rassouli M, Khanali L, Farahani A. Palliative care perspectives and practices in the Islamic republic of Iran, and their implication on patients’ quality of life. Palliative Care Perspectives, Practices and Impact on Quality of Life. New York: Publisher In Press, Nova Scientific; 2017.
15. Heydari H. Home-based palliative care: A missing link to patients’ care in Iran. Journal of hayat. 2018;24(2):97–101.
16. The SAGE Handbook of Qualitative Research in Psychology. London2008. Available from: https://methods.sagepub.com/book/the-sage-handbook-of-qualitative-research-in-psychology.
17. Graneheim UH, Lundman B. Qualitative content analysis in nursing research: concepts, procedures and measures to achieve trustworthiness. Nurse Educ Today. 2004;24(2):105–12.
18. Nowell LS, Norris JM, White DE, Moules NJ. Thematic analysis: Striving to meet the trustworthiness criteria. Int J Qual Methods. 2017;16(1):1609406917733847. https://doi.org/10.1177/1609406917733847.
19. Eshaghian-dorcheh A, Zandi M, Rassouli M, Tahmasiebi M. The Lack of Systematic Training for Health Care Providers, A Challenge for Providing Pediatric Palliative Home Care: A Comparative Study. Int J Pediatr. 2019;7(5):9481–96.
20. Mitchell GK, Johnson CE, Thomas K, Murray SA. Palliative care beyond that for cancer in Australia. Med J Aust. 2010;193(2):124–6.
21. Heydari H, Rahnavard Z, Ghaffari F. Exploring the Position of Community-Based Nursing in Iran: A Qualitative Study. Int J Community Based Nurs Midwifery. 2017;5(4):386–96.
22. Heydari H, Hojjat-Assari S, Almasian M, Pirjani P. Exploring health care providers’ perceptions about home-based palliative care in terminally ill cancer patients. BMC Palliat Care. 2019;18(1):66.
23. Shahsavari H, Nasrabadi AN, Almasian M, Heydari H, Hazini A. Exploration of the administrative aspects of the delivery of home health care services: a qualitative study. Asia Pac Fam Med. 2018;17(1):1–7.
24. Devi B, Tang T, Corbex M. Setting up home-based palliative care in countries with limited resources: a model from Sarawak. Ann Oncol. 2008;19(12):2061–6.
25. Regulations of drug management in Iran. Available from: https://fdo.mui.ac.ir/sites/fdo.mui.ac.ir/files/field/daroo/zavabet/4.pdf. Accessed 28 Apr 2020.
26. Benatar S, Sullivan T, Brown A. Why equity in health and in access to health care are elusive: Insights from Canada and South Africa. Glob Public Health. 2018;13(1):1533–57.
27. Saito DYT, Zoboli ELCP. Palliative care and primary health care: scoping review. Revista Bioética. 2015;23(3):593–607.
28. Currow DC, Agar M, Sanderson C, Abernethy AP. Populations who die without specialist palliative care: does lower uptake equate with unmet need? Palliat Med. 2008;22(1):43–50.

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