Stories of Dying and Death as told by Family Members’ of Adolescents and Young Adults (AYAs) who have Died from Cancer

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Abstract

Research of AYAs with cancer has developed significantly over the last 20 years. The research has demonstrated that AYAs with cancer are a forgotten population, who require closer study in order to understand their unique issues. This paper examines family members’ experiences of adolescents and young adults (AYAs) during the dying stage of their cancer trajectory. The results are drawn from a larger study titled ‘From Go to Woe; Family Members’ Stories of Adolescents and Young People Living with and Dying from Cancer’, which storied the family members’ experience of the diagnosis, treatment, dying and death of an AYA family member, utilizing Armstrong-Coster’s (2004) four stages of the cancer trajectory. The principal researcher’s motivation to understand and story these experiences was related to her own isolation and lack of information when her 16 year old son Anthony, was diagnosed with and eventually died of cancer at 17 years of age.

The significance of this study is the contribution made to the identification of issues that can inform health policy/ guidelines. The findings have the potential to increase understanding of, and prepare family members and AYAs with cancer, for the experience of the death and dying stage of the cancer trajectory.

Keywords: Adolescents and young adults; Cancer; Death and dying; End of life decision making

Literature Review

Research over the last 20 years has indicated that AYAs with cancer are a forgotten population who require closer study in order to understand their unique problems [1-9] prompting Michelagnoli, Pritchard and Phillip [3] to describe this group of people as a lost tribe. One factor identified to account for this disadvantage is the fragmentation of services [1,10-12].

The fragmentation of services is a result of AYAs being cared for in two systems of care, the paediatric and the adult [13] with different goals and philosophies of care.

The paediatric system has been described as mainly family focused and the adult system disease focused [14].

This has implications not only for medical treatment of AYAs with cancer, but also presents problems in providing targeted support services. O’Brien, et al., [15] identified the following issues that account for the lack of support services.

- Support services vary across adult hospitals and are fewer than those provided by specialist paediatric hospitals; support services provided both in paediatric or adult hospitals are not designed for the AYA age group, and the lack of critical mass in either system prevents optimal support service. These issues are ampliﬁed during the dying stage of the cancer journey.

- There is limited research on AYAs dying with cancer and their families. A review of literature found only four authors [16-19]. Wein, Pery and Zer [19] provided information specific to this group and their specific psychosocial needs whilst dying.

- Bisset et al. [16] identiﬁed the following needs for AYAs, their parents and siblings for palliative care services: "involvement in decision making, attention to developmental needs, an appropriate setting of care, ambivalence about taking on an adult role, a desire for independent living, the importance of school, college and employment, opportunities to do things other young people do, symptom relief and emotional support, planning of care to ensure the best end of life care possible."

- The researcher could find no research that speciﬁcally focused on AYAs and the dying trajectory, with the majority of the articles relating to children and their families. This is conﬁrmed by Mack et al [18], who emphasized that there is limited knowledge about the End of Life (EOL) care that AYAs receive.

She states: "The development of optimal tools for EOL care delivery in this population will depend on a better understanding of the care such patients receive." This paper will provide a further insight into the care of AYAs during the dying stage.

Research Aim

The aim of this article is to uncover from the stories of family members the experience of the dying and death of an adolescent or young adult (AYA) family member (aged 13-23 years).

Methodology

Narrative Inquiry was considered the most appropriate methodology for this study. Narrative inquiry uses stories to describe...
human experience [20-22]. Through story-telling meaning is constructed from the experience [21,23].

In narrative inquiry, the interview should be approached with the intent of hearing stories. Narrative has a commitment to the value of giving voice to human feelings and experience [24], through the use of “imaginative reconstruction” or “empathy” [25].

Grinyer [26] believes narrative is an appropriate method for sensitive research as it allows family members to determine the content of the information they share which she sees as strength rather than a weakness.

Narrative inquiry also includes the concept of a shared reality [20]. Bruner [20] has said that “life stories must mesh tellers and listeners must share some ‘deep structures’ about the nature of a life.” If this does not occur, both the storyteller and the listener will fail to grasp what the other is telling or hearing.

Narrative Inquiry therefore is a suitable methodology for this research study. Within its epistemology, understanding is the development of knowledge through the telling of stories. These stories create meaning and a testament of the experience of suffering, giving voice to what might not be told.

The methodology suggests my own experience, rather than hindering the research, creates the context for me as the researcher to grasp what the family member is saying and provide a richer understanding of what is being told.

Method

Ethics approval was obtained through the Southern Cross University Human Research Ethics Committee (Approval number ECN-05-146).

The family members in this study were a purposeful sample of 26 who had experienced the death of an AYA (aged 13 to 23) to cancer. Table 1 provides a representation of the family members relationship to the AYA, the interview type and time since death. The interviews were open-ended. All interviews were recorded on an iPod and professionally transcribed.

For the purpose of this study, family members preferred not to use pseudonyms and requested that their family member’s real names be used. Additional ethical approval was obtained from Southern Cross University’s Human Research Ethics Committee for this to be possible (approval number ECN-08-029).

The study utilised narrative analytic methods to create a configured plot of core stories and a configured meta-narrative from which commonalities and differences within the core stories were identified.

Guidance was obtained from the following narrative theorists: Mishler [27], Riessman [28,29], Polkinghorne [30], Lieblich, Tuval & Zilber [31], Emden [21,32], and Frank [33], to analyse these narratives.

In addition Taylor’s [34] computer-assisted method and the qualitative research software program NVivo8 provided a framework to configure and organise the stories into a meta-narrative.

Figure 1 links the steps of the narrative analysis for this study with specific processes and relevant authors.

Results

The narrative patterns within the stories identified transitional or turning points in the family members’ stories. These turning points related to the stages of the cancer journey identified by Armstrong-Coster: [35] departure, exploration, anticipation, and destination as represented in Figure 2.
This article provides insights into the experience of family members through themes identified within the destination stage. Table 1 provides a visual representation of the themes and subthemes.

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and nobody cottoned on to actually send her for a chest x-ray. " When a months with this cough … She was told she had a lung infection … Chest X-ray was

The destination stage

Tamina who had transitioned from the paediatric to the adult system will get an x-ray just in case and that is when we found the tumours in

Table 1: The Dying Landscape.

The results are a summary of the family members' stories within the themes. Direct quotes from the participants stories are represented by quotation marks and italics.

Ending the reprieve

The death and dying stage for the AYAs and family members in this study commenced when the AYA exhibited signs and symptoms that alerted the AYA and the family members that things were not getting better or the cancer had returned. Denise was shocked when Brenton James’s melanoma reoccurred in the liver as they had been told he was fine. She said “You know we got through this shocking period of him losing his eye and we’d been given the all clear. We’d got through this, we’d escaped.” Lauren noticed that Grant was not getting any better which was frustrating "then in the April he started coughing and we called the Oncologist and they said it is probably just a virus … but we will get an x-ray just in case and that is when we found the tumours in his lungs. So that was pretty devastating.” For most, tests and investigations were quick to confirm the return of the cancer. For Tamina who had transitioned from the paediatric to the adult system there was evidence of lack of communication within these two systems, as was seen in one story that resulted in a delay in diagnosis. Pam said "we’ve been coming back and forth, or Tamina had been for a few months with this cough … She was told she had a lung infection … and nobody cottoned on to actually send her for a chest x-ray.” When a Chest X-ray was finally ordered they found she had relapsed.

The pronunciation of impending death

The way that the family members were told of the impending death was not always satisfactory. Denise spoke of the frustration she felt when a young doctor tried to tell her Brenton James was dying: "I was shocked when Brenton got melanoma … a lot of them are still finding their feet with how to deal with these things … especially the one that was trying to tell me that my son was critical and that he would die, he rambled on and on and on and on, and I wanted to shake him. To say … just say it … [I] know what you’re trying to say, but I didn’t.”

There was no evidence of a plan as to when, where, and how the family members were told, added to this there was little support for the AYA and the family member after the pronunciation. This is seen in Jenny’s story, she said the doctor rang her at work and said: "I'm sorry, but Brenton is going to die, we can’t do anything about it. He has got metastasis everywhere”. Jenny then had the responsibility of telling her son, Brenton Duncan that he was going to die. For Pam, Tamina was told she was going to die with no support from family. Pam said "she (Tamina) was in there for a while [the doctors surgery] and he told her she was going to die … and they knew I was out in the waiting room.” Understandably the response for most was shock and devastation. Lauren recounted "the fact that that’s it, there is nothing more we can do for you was horrible.” For Fulvia the pronunciation of impending death did not occur until the AYA was near death. For example Fulvia recalled a doctor describing the dying process. She says doctor said: "Jadye’s condition has deteriorated, Jadye’s now unconscious and soon she’s going to start cheyne stoking, she’s going to start breathing different and so on and I don’t want you all to be distressed, it’s just the normal way of dying, there’s no pain and she’s not in any distress.”

Changing the focus of care

The family members noticed that with the pronunciation of impending death the focus of care changed both in how and where the care was delivered. Lauren spoke of how this quality of care changed: "whereas in the first time it was, well it doesn’t matter how sick he gets because in the end … he’ll be alive. … and so that really changed … it definitely didn’t feel that the medical profession treated him as a possibility, it was more a matter of when he died and how are we going to make from now to when he dies better than trying to prevent him dying.” Val’s story revealed the changing focus of care by deciding not to give Anthony Charles any more treatment. She said: "It was an easy decision, because why put the poor kid through so much more turmoil and hell than what he’d already been through, and who was it for, me or him? If I kept him alive it’d be for me.”

In some of the stories it is not evident if families were provided with alternative care delivery choices. There was an underlying assumption that the AYA would die in hospital. Deborah who had to insist that Matthew Anthony be allowed to go home to die, Deborah said: “… and when Matthew said he wanted to die at home and not in hospital, the doctor almost laughed at him, and said ‘well, that’s just ridiculous.’” Deborah recalled Mathew saying, “I’m not going to die in hospital. If I’ve only got a week to live, I want to be at home.” Deborah insisted he die at home saying: “You said he’s going to die, there’s nothing more you can do for him, we’ve got scripts for his morphine and … all his medications. We’ve got the stuff, we’re going home and he will die at home.” For others there appeared to be a choice. For example Thomas was cared for at home. Irene said: “they told us then there was nothing else they could do. He stayed home. … he wouldn’t stay in hospital for the last… I think it was better for him to be at home. I think if any kid’s
going to go through this, stay at home if you can get treatment at home rather than be in a hospital.”

Staying home for the last moments was not always a choice that the family member wanted. They had a sense that there was no alternative. Jenny spoke of how she felt deserted by the paediatric system because of Brenton Duncan’s age and that there was nowhere else for her to go but home. She said: “they did offer the hospice around the corner here but I went around to see it and it was just full of old people and I couldn’t put my son there.” Only one of the AYAs died in a hospice (Ben), which Sue described as a caring and compassionate place.

Going downhill

It became evident to the family members in this study that their AYA was dying when they noticed their health was deteriorating. For some this was evident when they became aware that the cancer had taken over other parts of the body. Denise noticed that Brenton James’s “stomach become bloated his legs were full of fluid the rest of him was fading away and he was having increasingly that look of someone that has not got long to live.” Fulvia spoke of how quickly the cancer spread in Jadye “Within a week it was in her liver, in her lungs, in her back, in her shoulder and in her bone marrow, horrible cancer. It’s cruel.” For others it was associated with having to deal with the symptoms associated with the cancer spread, such as blindness and deafness, seizures and behaviours associated with neurological deterioration. Cheryl spoke of Paul becoming “blind and deaf and very, very frail” whilst Helen and Geoff spoke of Mick’s epileptic seizures “and he had … 2 or 3 really bad nights … like an epileptic seizure, … [he] broke a couple of his vertebrae in his back.” While Kerry described Alinta’s neurological deterioration, “You have brain things … you might have fits … there’s agitation and fidgets.”

For others it was the AYA being overwhelmed by severe pain. Fulvia explains Jadye’s experience, “we sat up for night after night with her screaming… and she didn’t want to move, and she just begged me, … no leave me at home, leave me at home’ and I yelled at her and made her go to the hospital and she never came home.” As with Jade, pain was the beginning of the end for Brenton Duncan as Jenny describes, “he was screaming in agony and we got the district nurse to come around and give him morphine … they put a driver into him … and we put him to bed and we will be able to look after him.”

Keeping a vigil

Throughout the cancer trajectory the family members sought to stay with the AYA as much as possible and continued to care for the AYA as their condition deteriorated. For many this involved lack of sleep, being at the hospital 24/7 and being concerned that the AYA might die. Denise explains: “we were on a journey that neither of us wanted to go on but nevertheless we were going to go on this journey together and we were going to support each other along the way.” Deborah spoke of the relationship between Matthew Anthony and Lauren (his sister): “and just watching that develop between the two of them was just beautiful … even more, it just moved onto a much deeper level … it was just beautiful, absolutely beautiful.”

Experiencing lack of care

Throughout this experience, the family members occasionally felt unsupported which made keeping a vigil difficult. This was evident in the lack of care and acknowledgment, by some hospital staff, that the AYA was dying. Rebekah described an incident: “and the nurse [said] come on we’ve got you in bed and just grab her legs and swings her up onto the bed and Naomi was crying in pain and then she, … pulled the blankets up and left. I was sitting there thinking … all she had to do was wait a minute so Naomi could catch her breath and get ready because she knew it was going to hurt.” For others there was the sense that there were not enough resources to support the family members in their home. Jenny said “he wanted to die at home so I wanted him to be with me but I just needed a respite every so often because it was very hard work when you don’t know how long it is going to take, it’s just all really really hard.” Kerry reinforced this: “so as far as I could see, … someone had given me a 24 hour watch, with no help … it’s just so hard and you get more and more isolated.”

An aspect of the lack of care was the lack of coordination, particularly if the AYA was at home. Jenny described the care as “higgledy-piggledy” whilst Kerry believed it would “have helped to have some-one oversee our whole care.” For Sue this lack of support required the AYA to be admitted to a hospice. She said: “Ben came home and he was home I think, two days, and he had a fall, his legs gave way, and I knew then he had to go back to hospital because I couldn’t lift him.”

Feeling overwhelmed

The lack of support whilst the family members maintained a vigil resulted in the family member at times feeling overwhelmed with the burden of care. The extent of the burden is exemplified by Deborah’s personal battle with herself about ending the pain. She said: “he actually begged me to kill him … but that kept playing over and over in my head and I thought my God, what sort of a person am I, he’s begging me to end his suffering and I’m saying no.” The majority spoke of how exhausting and hard this vigil was, and they just wanted the AYA to die to end the suffering. As expressed by Josephine, “this sounds really awful, but I’d had enough. I couldn’t cope with it any more, I could not cope with watching him going through what he was going through, … and it was just too much, and I just wanted him to die.”

The hard work and witnessing the suffering caused some family members to become overwhelmed by emotions. For example, Lauren described a panic attack the day Grant died. She recalled “everything was just sort of blurry and hazy … I think that really it was just made by me needing to release that stress … It wasn’t real easy … I guess we were just surviving.”

Despite all the suffering and all the hard work that the family members had to endure, the journey was made easier by the courage and strength of the AYA who, rather than engage in self-pity, displayed heroic behaviours that made the journey easier for the family members. The family members spoke of the AYAs accepting the reality of their death, taking control of the dying, being grateful, protecting others and enjoying ordinary behaviours. Arlene told how Mathew James “made the decision that he would stay in hospital, keep the chest.
tube in, say his goodbyes and then we would take the tube out and he would be gone within 3 or 4 hours.” Deborah spoke of how Matthew Anthony planned his own funeral “picked his own casket, told me what he was going to wear, chose his own songs … He chose the funeral director chose the flowers that he wanted on his casket … we actually sat down and figured out the wording on the plaque, together.” The gratitude was expressed even at the end stage of dying. Lauren remembered Grant saying thank you to the nurse on the day he died “the nurse was giving him some morphine and he looked at her and he said thank you for everything.”

The AYAs protected and supported others as they were dying. Sue spoke of how Ben's lack of complaining enabled her to manage this part of the journey. She said: “he never complained and that’s what everyone said to me, no matter how much pain he was in, no matter what he was feeling, he never complained. He never said, why me? Never, he never complained to the nurses, the doctors, he never wallowed in self pity, and that’s I think, how I got through, I kept fighting it, even though it really wasn’t any point.” Similarly Grant said: “I didn’t realise he was going to die…right up to the end I didn’t realise”

Recalling the moment

For all the family members in this study the moment of death and being connected with the body after death were an important part of the Destination stage. For some the final moment of death was associated with waiting and watching for the last breath. Lauren recounted, “it was … kind of morbid because each breath was so far apart and so slight and we weren’t sure when he stopped… and then he kind of took this little gasp and then we were like waiting and waiting and waiting and another one didn’t come. I just started crying.”

For most waiting for the last breath was not painful it was almost like just waiting for the end of the journey. For example, Pam watching Tamina’s breathing, she said: “but I just sat on the lounge and watched her breathing and I thought … afterwards of all the wonderful people, like my friends, that I would’ve wanted to share that with me.”

Extending the living connection

It was important that after the AYA died that the family members were given the opportunity to extend the living connection with the body after death. For some this meant spending time with the body. Lauren explained, “we went back in with Grant and spent some time there … I think I had the hardest time leaving … I guess I realised it is time to go when he started not looking like himself any more I didn’t want to keep seeing him like that. So I hugged him and walked out.” Others found it important to care for the body in more physical ways such as washing and dressing the body. Deborah provided an explanation concerning the importance of caring for Matthew Anthony’s body. She said: “I was the first person to put clothes on him and I’m damn well going to be the last person. So I did, I dressed him and silly little things [is] I put 2 pairs of socks on his feet because he always got cold feet and I … put his blanket over him so he didn’t get cold … I’m glad I did it, even though technically I guess he couldn’t
feel the cold, it made me feel better that I had hugged him up and he was nice and warm.” Others viewed the body, which helped them say goodbye. Sue recalled, “so we went to see him that afternoon … he looked very peaceful and very calm, and it was very, very hard … I just wanted to pick him up.”

Discussion

This summary of the themes created in the meta-narrative of the dying stage of the cancer journey provides insights into the dying of an AYA as experienced by family members. Wein [19] identified the psychosocial developmental phase and an AYA facing death, as unique with only Mack’s 2015 [18] study to evaluate the intensity of EOL care among some of the AYAs were not given the opportunity for a good death. For ways such as maintaining their identity with friends and others. Palmer & Reason cited are “perceived lack of training, stress, no time to attend to the patients emotional needs, fear of upsetting the patients and a feeling of inadequacy or hopelessness regarding the unavailability of further curative treatment”[44]. Given these reasons for not initiating end of life discussions with adult patients, issues associated with AYAs and their families would be even more complex and confounded. The majority of family members in this study did not engage in EOL discussions and rarely was palliative care mentioned in the stories with 12 AYAs dying in the curative hospital system and only one AYA dying in a hospice. For most the pronouncement of impending death was at the end of the destination stage with little time to prepare for death or enter into EOL discussions.

The research studies have demonstrated that dying in a curative system results in EOL discussion occurring too late in the dying journey for adequate end of life decisions to occur [18,37,45,46]. Mack's 2015 [18] study to evaluate the intensity of EOL care among AYA with cancer found 68% received at least one medically intensive treatment measure within the last 30 days of life, with 11% receiving chemotherapy within the last 14 days of life. Added to this is McGrath’s [46,47] study that reported carers in the curative system were not provided with information about alternative places to care for the patient when dying. Wittenberg-Lyles, Goldsmith & Ragan [48] describe this as the isolated journey occurring within a curative, biomedical structure limiting opportunities to discuss end of life care. This is in contrast to the comfort journey in which palliative care is introduced on diagnosis enabling life prolonging treatment and comfort care. The comfort journey provides opportunities for open communication on advanced care planning and end of life decision making [48].

It is not known if the AYA or the family members in this study were engaged in any EOL discussions. That there was no reference to this in the stories suggests that this did not occur. Supporting this is the fact that only one AYA died in a Hospice and of the five who died at home only Irene appeared to have seen this as a choice. Jenny and Kerry felt they had no choice and Deborah had to fight for her choice for Matthew Anthony to die at home. Jenny described the care as higgledy-piggledy with no plan or coordination. This supports the idea that palliative care be introduced early in order for patients and family members to plan their model of care [7,40,41,47,49], which is timely and integrated [50] and structured and coordinated [7].

Overlaying what appeared to be little opportunity for EOL discussions is the family members keeping a vigil with the AYA throughout the destination stage which involved many sacrifices. This keeping a vigil resulted in many feeling overwhelmed with the day-to-day care of a loved one who was dying. Added to this was the experience of feeling unsupported in the delivery of care. Despite this there was no mention of counseling being provided for either the AYA or the family members. The OnTrac Practice Framework states that access to supportive care is essential for young people facing death [7]. The National Service Delivery Framework mentions the profound effects on families when witnessing the cancer journey or death of a young person. The Framework recommends that service delivery should be mindful of these relationships and recommends grief and loss support is needed for AYAs and their families [50]. This is supported by the OnTrac Practice framework which recommends supportive care be available to family members and partners [7].

The destination stage culminates in the end stage of dying in which the AYA becomes increasingly dependent on the care of others. Palliative Care Australia stresses the importance of loved ones being informed about what to expect, how to respond and who is available to assist them. This information should include “the physical process of dying (i.e. changes in breathing patterns), psycho-education, and looking for signs of symptom relief” [7]. It is not evident in Deborah's story if she had been provided with information about dying and how to manage the pain associated with Matthew Anthony’s dying. There is the possibility that with adequate information she may not have experienced her dilemma of how to end his suffering.

It was only at the time that death was imminent that the family members were free to let the AYA go and end their suffering. It was important to say goodbye and to be there at the moment of death. After the death the family members continued their connection with the AYA with body caring rituals. The importance of this for the family members was evident in that they wished to describe in detail the moment of death and what occurred after the death. This highlights the need for health care professionals to be cognizant of the family’s
needs during and after the death and to see the person as not just a body. Whilst there was a sense of relief at the moment of death, this moment augurs the transition to another stage incorporating the loved one into their life with memories and rituals.

**Conclusion**

Barling [8] sought to understand the experience of the AYA cancer journey from the perspective of family members. This article presents selected findings from the Barling [8] study that explained family members’ experience of during the destinations stage of the cancer journey.

The summary of the meta-narrative provides acknowledgment that the destination stage is difficult for the family members and the AYA. The stories reveal a lack of support and being overwhelmed with the burden of care as the family members maintained their vigil. Despite this the family members found it important to talk about the death and how they continued their living connection with their loved one after death. In addition, the absence of EOL discussions within the majority of the meta–narratives suggests that AYAs and family members are often not provided with the opportunity to consider EOL choices during the destination stage.

This article has therefore highlighted the need to continue research into the needs of AYAs and their families during the destination stage to further increase understanding and assist in the implementation of the National and OnTrack Framework recommendations, with particular emphasis on specific issues associated with the destination stage of the cancer journey.

**Anthony’s words:**

*Mortality is a blessing taken without recognition.*

*We all walk the same tightrope.*

*Only some dare to look down, whereas others have to.*

*There is only one thing we can do: try not to walk the rope alone.*

**References**


50. (2009) CanTeen and Cancer Australia National Service Delivery Framework for Adolescents and Young Adults with Cancer.