Acceptability of low molecular weight heparin thromboprophylaxis for inpatients receiving palliative care: qualitative study

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Abstract

Objective To find out what inpatients with advanced cancer who are receiving palliative care think about the effect of thromboprophylaxis on overall quality of life.

Design Qualitative study using audiotaping of semistructured interviews.

Setting Regional cancer centre in Wales.

Participants 28 inpatients with advanced metastatic cancer receiving palliative care and low molecular weight heparin.

Main outcome measures Recurring themes on the effect of thromboprophylaxis on overall quality of life.

Results Major emerging themes showed that patients knew about the risks of venous thromboembolism and the purpose of treatment with heparin. Media coverage had raised awareness about venous thromboembolism, and many had previous experience of thromboprophylaxis. All found low molecular weight heparin an acceptable intervention, and many said that it improved their quality of life by giving them a feeling of safety and reassurance. Antiembolic stockings were considered uncomfortable and had a negative impact on quality of life. Patients were concerned that because they had advanced disease they might not be eligible for thromboprophylaxis.

Conclusion Low molecular weight heparin is acceptable to inpatients with advanced cancer receiving palliative care and has a positive impact on overall quality of life. Antiembolic stockings are an unacceptable intervention in this patient group. Guidelines on thromboprophylaxis are urgently needed for palliative care inpatient units and hospices.

Introduction

The association between venous thromboembolism and cancer is well known and seems to increase as malignancy progresses.¹⁻³ Up to 52% of patients in specialist palliative care units may have venous thromboembolism.⁴ Because one in seven inpatients with cancer dies of pulmonary embolism, venous thromboembolism probably reduces survival time in patients receiving palliative care.⁵

A consensus statement by the American College of Chest Physicians on antithrombotic and thrombolytic therapy recommends, on the basis of level 1A evidence, that hospitalised patients with cancer receive low molecular weight heparin.⁶ Recently, the House of Commons Health Committee on the prevention of venous thromboembolism for hospitalised patients reported a lack of national thromboprophylaxis guidelines within the United Kingdom.⁷

Less than 10% of palliative care units in the UK have guidelines on thromboprophylaxis, and there are concerns that daily injections of low molecular weight heparin may cause unnecessary distress.⁸ We surveyed inpatients who were receiving palliative care to find out their views on thromboprophylaxis and whether heparin was an acceptable intervention.

Methods

Sample selection

We recruited patients from the specialist palliative care unit within the regional cancer centre, which had established thromboprophylaxis guidelines. We identified patients using screening notes and drug charts.

The inclusion criteria for participants were: metastatic cancer or primary brain tumour with no curative treatment available; evidence within medical notes that the incurable nature of the disease has been discussed with the patient; the patient had received low molecular weight heparin thromboprophylaxis for at least five consecutive days.

Data collection

We audiotaped and then transcribed the semistructured interviews, which covered the following topics: cancer treatments received (such as surgery, chemotherapy, and radiotherapy); insight into prognosis; what was understood about treatment with low molecular weight heparin and thromboprophylaxis; the impact of thromboprophylaxis on overall quality of life; negative aspects of being on heparin treatment.

Analytical framework and data analysis

Qualitative research is a descriptive and interpretative approach using analytical categories to describe and explain social processes and place them within a framework of meaning relevant to the research aims. We carried out a thematic analysis, using an inductive approach to obtain categories emerging from the data that showed how the participants viewed thromboprophylaxis in this context of care.

We identified two distinct types of patient. One group comprised patients receiving rehabilitation after cord compression who previously had good performance status (median Eastern Cooperative Oncology Group score of 1; see box) but had suddenly deteriorated. The other group comprised patients admitted for symptom control who had been unwell for some time, with a gradual deterioration of performance status. Consequently, we divided the participants into two categories. Theoretical saturation (when no further recurring themes emerged from analysis) was achieved at 14 patients per group. We extracted the emerging themes from the data and constantly refined and validated them from repeated and comparative reading of the transcripts.⁹ We identified four major themes and three minor themes.
Box 2: Eastern Cooperative Oncology Group performance grades

0: Fully active; no restrictions in performance compared with before illness (Karnofsky 90-100)
1: Physically strenuous activity restricted but ambulatory and able to carry out work of a light or sedentary nature, such as light housework and office work (Karnofsky 70-80)
2: Ambulatory and capable of all self care but unable to carry out any work activities; mobile for more than 50% of waking hours (Karnofsky 50-60)
3: Capable of limited self care; confined to bed or chair more than 50% of waking hours (Karnofsky 30-40)
4: Completely disabled. Cannot carry out any self care; totally confined to bed or chair (Karnofsky 10-20)

We selected excerpts of interview text on the basis of two criteria: firstly that they illustrate the issue being discussed, and secondly that they represent a range of participants rather than rely on a few individuals, thus reducing bias. Patients are identified by an interview number in brackets and by whether they had been admitted primarily for symptom control (S) or following spinal cord compression (C).

Results

Sample characteristics

We invited 28 patients to participate and all agreed. Box 1 summarises the characteristics of the patients. We established the Eastern Cooperative Oncology Group performance grade (box 2) of each patient one week before admission. All patients had a performance status of 4 at the time of interview.

Major themes identified in both groups were insight into the prognosis; knowledge and understanding of thromboprophylaxis and guidelines on treatment of venous thromboembolism; acceptability of thromboprophylaxis with heparin; security; and optimism. Minor themes were bruising, the negative impact of antiembolic stockings on quality of life, and anger at paternalistic views towards terminally ill patients.

Major themes

Insight into prognosis

Most patients showed clear insight into the nature of their condition, describing it as “incurable” or “terminal.” These words had a similar meaning to each patient, and they were aware that the aim of treatment was the control of symptoms and not cure:

“What do I mean by terminal? Well there’s no cure. The cancer will eventually get the better of me.” (11C)

“Oh it’s incurable. They said they can’t get it better. They tried some chemotherapy but it didn’t work so now they are controlling the symptoms.” (14S)

Patients admitted for symptom control viewed their inpatient stay as “just another admission” because they experienced a slow and steady decline from a median performance status of 2. Even though worsening symptoms may suggest a change in prognosis to doctors, this was not the case for patients. Most patients admitted with spinal cord compression had a good performance status (median score of 1) before admission. The decline to performance status 4 was faster than in the symptom control group and in many cases necessitated fresh discussions about disease progression:

“You live with cancer for so long that you forget that it is eventually going to do you in. I was driving a car two weeks ago ... This has been a wake-up call.” (19C)

Knowledge and understanding

All patients understood the purpose of treatment with heparin, and many were also aware of why they may be at risk of venous thromboembolism, identifying immobility and surgery as risk factors:

“Well I understood that because I was in bed and I wasn’t moving about, mobility wise, like I usually do, they were for this deep vein thrombosis. You know well it sort of prevents it then. Much like the planes and the long trips on the planes, that’s my understanding of it.” (1S)

All patients knew that thromboembolism was undesirable. When asked about its unwanted features, all focused on the most serious potential consequence—death. No patients were aware of common symptoms of deep vein thrombosis, such as painful swollen legs, or of pulmonary embolism, such as dyspnoea:

“Well it could kill you couldn’t it. It can go to the heart or the brain. Very serious I think, a blood clot, isn’t it?” (1S)

It was clear that most patients based their knowledge of venous thromboembolism on media coverage of its association with long haul flights, with little understanding of the specific association with cancer:

“There’s been a lot about them in the news recently. They are supposed to be a bigger problem than the super bug.” (6S)

“Well it’s a bit like long airplane flights isn’t it? If you don’t move around enough like staying in bed or sat in a chair the blood will become sludgy and clot.” (13C)

Acceptability

All patients found thromboprophylaxis with low molecular weight heparin acceptable, and many could not understand why it would be considered unacceptable. Acceptability fell into three categories.
Research

- Recognition that thromboprophylaxis with heparin was part of usual practice:
  
  "It's sort of reassuring knowing that people are still doing something for me. People keep talking to me about controlling my symptoms. It seems just as important to me to prevent anything that may cause bad symptoms." (6S)

  "Just that you are getting the best care." (5C)

- Consideration that treatment with heparin was neither pleasant nor unpleasant:
  
  "The injection is of no significance and I don't feel it, and you've got to have 'nasties' to have 'nices' haven't you?" (1S)

  "... a sequence of little bruises on my tummy. I must admit that when I looked down and saw these little blue blobs I thought 'Oh that is interesting'. But no, there are no negative aspects at all." (9S)

- Balance of benefits against side effects:

  Most patients had experienced two or more treatments for cancer (surgery, chemotherapy, and radiotherapy), each with recognised side effects. In comparison, a daily injection of heparin was more acceptable than an antiembolic stocking. Patients indicated that antiembolic stockings negatively affected overall quality of life. Our study shows that low molecular weight heparin had any detrimental effects on their quality of life. Only patients receiving heparin thought it helped them live longer by preventing fatal complications of cancer. However, the acceptability of this treatment was not based purely on a "benefit versus harm" evaluation, but on the feeling that the injection was a trivial intervention with little or no impact on quality of life. Only patients receiving heparin treatment were interviewed, and their agreement to participate may have led to a positive bias. They received heparin for a relatively short time, but previous work found that long term treatment is acceptable to this patient group.

Anger at paternalistic views towards terminally ill patients

Inevitably, face to face interviews uncovered underlying themes that have not been prompted, but are a consideration in the research sample, whose actions and perspectives are based on their broader life experience. Some patients spoke openly about their views on dying and their experiences of being, as they described it, "terminally ill." They uniformly expressed their need to be involved in decision making, particularly with respect to the withdrawal or non-administration of treatment. Some patients had experienced what they viewed as nihilistic paternalism, and they were angry that major decisions were made about their lives without their involvement:

"I would be quite cross about that. Very, very cross indeed. It's like when I had a run in about who was going to pay for what in the nursing home and I felt afterwards that I had been put to one side for two reasons, one I had a terminal illness and two I was 72 approaching 73 and was therefore not a high priority, and that hurt." (11C)

"Just because I know I'm going to die, doesn't mean I'm happy about it. I plan to hang on as long as I can. These injections are nothing. I honestly don't know what the fuss is about." (13S)

Some patients said that heparin injections were not unpleasant, whereas they had experienced some interventions that were:

"How can someone else tell me what I should feel about a treatment? I bet they haven't even tried it. The thought of an injection may sound bad but there is nothing to it. It's much better than having to wear those stockings." (5S)

**Minor themes**

**Brusing**

Patients were asked whether thromboprophylaxis with low molecular weight heparin had any detrimental effects on their quality of life. The only negative experiences reported were of bruising:

"... a sequence of little bruises on my tummy. I must admit that when I looked down and saw these little blue blobs I thought 'Oh that is interesting'. But no, there are no negative aspects at all." (9S)

Most patients had experienced two or more treatments for cancer (surgery, chemotherapy, and radiotherapy), each with recognised side effects. In comparison, a daily injection of heparin was considered trivial.

**Impact of antiembolic stockings**

Several patients had worn antiembolic stockings during previous hospital admissions. All had found them uncomfortable:

"They are so uncomfortable to wear. You get hot in them and they feel so tight." (7C)

"They are the most uncomfortable things you can ever have. They are OK for a day but not if you try wearing them for more than that." (11S)

Patients thought that low molecular weight heparin was preferable:

"Well those stockings are so uncomfortable. They're itchy. They're sweaty. They're horrible. Give me heparin any time." (5S)

**Discussion**

The reluctance to use thromboprophylaxis with heparin in people receiving palliative care seems to be motivated by the ethical principle of non-maleficence, as some palliative care doctors believe that this treatment adversely affects patients' quality of life. Our study shows that this is not so. Antiembolic stockings have been used in preference to heparin in specialist palliative care units in the belief that they are more acceptable than an injection. Our study shows that low molecular weight heparin is both an acceptable intervention and preferable to these stockings. Patients indicated that antiembolic stockings negatively affected overall quality of life.

The problems of venous thromboembolism are widely reported in the media and palliative care inpatients are knowledgeable on the subject, although their knowledge is limited to "worst case scenarios" and based on an incomplete appreciation of the risks of thromboembolism specific to cancer. Patients viewed thromboprophylaxis positively because they thought it helped them live longer by preventing fatal complications of cancer. However, the acceptability of this treatment was not based purely on a "benefit versus harm" evaluation, but on the feeling that the injection was a trivial intervention with little or no impact on quality of life. Only patients receiving heparin treatment were interviewed, and their agreement to participate may have led to a positive bias. They received heparin for a relatively short time, but previous work found that long term treatment is acceptable to this patient group.
Implications

Our study raises two important points. Firstly, it highlights the importance of involving patients in decisions related to their care. Patients' views were often different from what we expected and acting in their best interest without consultation may have a worse outcome than if they had not received paternalistic behaviour. Secondly, our findings reiterate the need for evidence-based guidelines for preventing venous thromboembolism in selected palliative care inpatients, since patients being looked after in palliative care units now have a better prognosis, as shown by a discharge rate of greater than 50%. Clearly there will be times when established measures to control symptoms are preferable.

It seems counterintuitive to start low molecular weight heparin in patients admitted for terminal care or in those entering the dying phase. Likewise, this treatment has resource implications such as drug costs and nursing time, which would require appropriate economic evaluation. Nevertheless, this form of thromboprophylaxis was acceptable in selected palliative care inpatients, and patients receiving palliative care should be considered for treatment unless clinical reasons indicate otherwise.

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What is already known on this topic

Venous thromboembolism occurs in up to 52% of palliative care inpatients with cancer and kills one in seven cancer patients

Low molecular weight heparin is recommended for thromboprophylaxis in non-ambulant inpatients with cancer

Concerns that injections of low molecular weight heparin adversely affect quality of life are reflected in the reluctance to use this treatment in specialist palliative care units

What this study adds

Low molecular weight heparin is an acceptable intervention in palliative care inpatients with cancer

Antiembolic stockings have a negative impact on overall quality of life

Patients in palliative care units wish to be involved in making decisions about thromboprophylaxis


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