Cancer patients’ experiences of living with venous thromboembolism: A Systematic review and qualitative Thematic synthesis

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Abstract

Background: Cancer-Associated thrombosis is common. Recommended treatment is daily injected low-molecular-weight heparin for 6 months. Most studies focus on prophylaxis and treatment; few have explored patients’ experience.

Aims
To identify and synthesise the available literature concerning patients’ experience of cancer associated thrombosis.

Design
Systematic literature review and qualitative thematic synthesis.

Methods
MEDLINE, Embase, CINAHL, PsychINFO (until 10/2016; limited to English) were searched. Eligible papers were qualitative studies of adult patients’ experience of cancer-associated thrombosis. Two researchers screened titles/abstracts/papers against inclusion criteria with recourse to a third for disagreements. Critical Appraisal Skills Programme qualitative checklist tool was used for quality appraisal.

Results
1397 articles were identified. Five qualitative studies (total n=92; age range 32 to 84 years) met the inclusion criteria. Participants had various cancer types. Most had advanced disease and were receiving palliative care. Four major themes emerged from the data: knowledge deficit (patients and clinicians); effects of cancer associated thrombosis (physical and psychological); effects of anticoagulation; coping strategies.

Conclusion
The cancer journey is difficult in itself, but thrombosis was an additional, frightening and unexpected burden. Although the association between cancer and thromboembolism is well known, cancer patients are not educated routinely about the risk or warning symptoms/signs of thromboembolism which may otherwise be misattributed to the cancer by patient and clinician alike. This systematic review highlights the impact of cancer-associated thrombosis on the lives of cancer patients, and calls for education for patients and clinicians to be part of routine care, and further work to address this patient priority.
Key words: Venous thromboembolism, qualitative, patient experience, cancer-associated-thrombosis
INTRODUCTION

Venous thromboembolism is a life changing diagnosis. Comprising deep vein thrombosis and pulmonary embolus it is a common phenomenon worldwide. It affects one in 1,000 patients; 6.5 million people globally each year (1, 2). The risk of venous thromboembolism increases by 6-7 folds in patients with cancer compared with non-cancer patients (3, 4). Approximately 20% of all newly diagnosed cases of venous thromboembolism are cancer patients (5), and post-mortem studies have demonstrated rates of venous thromboembolism in patients with cancer to be as high as 50% (6).

Up to 20% of patients with malignancy will develop cancer-associated thrombosis (5).

Although the risk increases with late-stage and during chemotherapy, over 50% occurs during the first three months from diagnosis (3) and interferes with cancer management (6).

International guidelines for the treatment of cancer-associated thrombosis recommend anticoagulation with weight-adjusted low-molecular-weight heparin for 3–6 months. (7-9) However, even with an optimal anticoagulation, cancer-associated thrombosis is associated with a higher recurrence rate than non-cancer venous thrombosis, and a poorer prognosis than cancer patients without thrombosis (10-12).

Apart from conferring a worse prognosis, the diagnosis of venous thromboembolism is a physically and emotionally distressing phenomenon that affects patients’ experience and quality of life (13, 14). However, data available on how cancer associated thrombosis and its treatment affect the cancer patients’ experience is scarce compared with that in relation to treatment or prevention.

Systematic reviews and meta-analysis on cancer associated thrombosis are limited to biomolecular markers associated with cancer associated thrombosis (15, 16) risk assessment of venous thromboembolism in cancer patients (17, 18) or both (19), clinical outcome, thromboprophylaxis (20), management (21-23) and risk stratification (24).
In order to improve our understanding and raise awareness of cancer associated thrombosis and to stimulate improvements in the supportive care of cancer patients, we undertook a systematic literature review to answer the following question, “what is the experience of people living with cancer associated thrombosis?"

**METHODS**

**Search strategy**
Two independent researchers conducted the search (NB, WI). MeSH terms and text words for cancer, venous thromboembolism and quality of life (see supplementary Table 1) were combined. The following electronic databases were searched: Embase, MEDLINE, CINAHL, and PsychINFO, until October 2016 and limited to English language, according to a pre-constructed protocol. In addition, an online search was performed for the following journals: Journal of Thrombosis Hemostasis (ISTH/JTH), Thrombosis Research and Hematologica. Bibliographies from relevant articles were examined for further related studies.

**Inclusion criteria**
Studies of adult cancer patients with venous thromboembolism with or without treatment for the venous thromboembolism were included. The review included qualitative studies that assessed the quality of life or experience of this group of patients.

**Study selection**
The titles, abstracts and full studies were screened by two independent researchers (NB, WI) against the inclusion criteria. Disagreement was resolved by discussion with access to a third opinion (MJ). Studies that matched the selection criteria were retrieved and their full text version analysed.
Data extraction

Data were extracted by NB; demographics of the included papers (author, year, design, population, question, main findings) and the primary quotations presented in the results.

Quality Appraisal

All articles were assessed against the Critical Appraisal Skills Programme (CASP) checklist tool for qualitative studies by NB and WI independently (25). Studies were not excluded on the basis of quality, but the assessment of quality was taken into account during analysis.

Analysis

The primary quotation data were synthesised by NB using thematic synthesis (26) and the principles of thematic analysis to explore the understanding of long term effects of venous thromboembolism on cancer patient’s life quality (27). This allows the context of each study to be taken into account whilst aiming to produce a generalizable synthesis (28). Direct quotes from patients and the researcher comments under the headings “results, findings, or discussion” from each study were extracted for coding.

Thematic synthesis involved: line by line coding of the findings of primary articles after reading and rereading of the papers to get familiarised with the data included, then the codes were discussed with MJ, and a coding framework formed which was used to code all papers followed by development of descriptive and analytical themes from the codes, in discussion with MJ and JS (29). Both inductive (allowing themes to arise from the specific observations) and deductive (working within existing knowledge about the effect of venous thromboembolism on people without cancer, looking specifically within our data for similarities and differences) processes were involved.

RESULTS

Overview of articles

The search identified a total 13197 articles, Embase (11632); MEDLINE (1272); CINAHL (254); and PsycINFO (38) articles. One additional article was identified through searches of relevant
bibliographies. Eleven full articles were retrieved and assessed for eligibility; six articles were excluded following review. This is summarised in the PRISMA flow diagram (Figure 1).

Five qualitative studies published between 2005 and 2015, met the inclusion criteria. Four of them were conducted in the UK. The key characteristics of the studies are summarised in Table 1.

**Study populations**

A total of 92 cancer patients with venous thromboembolism were included in these studies. All were adult patients of mixed gender with mean age of 58 years (range 32-84). Participants represent a wide variety of cancer types and stage. The most cancers were: breast, colorectal, ovary, lung, prostate, pancreas, and renal.
Figure 1: (PRISMA) Flow diagram
Themes

Four major themes were identified: knowledge deficit, the effects of cancer associated thrombosis (physical and psychological effects), the effects of cancer associated thrombosis treatments, and coping strategies.

Knowledge deficit

Two studies investigated the patients’ knowledge about cancer associated thrombosis in the context of cancer journey(13, 30)

Despite the fact that cancer associated thrombosis may develop as early as the first few months of diagnosis of cancer, and that the risk is increased with cancer treatments (chemotherapy, surgery, and hospitalisation), and disease progression participants were often not aware of their increased risk or of the warning symptoms of cancer associated thrombosis.

“During my cancer treatments, I was never told that there was a risk of getting a blood clot. I didn’t know about it... I was pretty shaken up”(30)

“I have never heard of venous thromboembolism, so that’s why I was so shocked”(30)

“[…] but they don’t tell you you’re gonna get clots after chemo, that’s the one thing they haven’t, they never said but we, we just put it down to, it’s just my breathing […] just that one item of information that we weren’t aware of”. [VCC07](13)

Moreover, patients on chemotherapy usually experienced different side effects, when they develop a venous thromboembolism they associate it with chemotherapy and do not recognise that their symptoms are symptoms of a venous thromboembolism.

“[…] but um this time again first set of chemo, she felt terrible and the thing is, when we went back to hospital really desperate, the only problem we thought was that it was the chemotherapy that was causing it”. [RG02](13)
On other hand participants with prior knowledge about venous thromboembolism respond in calm
and seek medical help immediately.

“I was out of breath and I said to my partner, ‘I think we are going to hospital’ without panic because I knew that it was something that could be rectified effectively)” (30)

There was also evidence of limited awareness about venous thromboembolism and cancer amongst health care professionals (13). This is consistent with patient reports of delayed diagnosis of the venous thromboembolism; on many occasions alternative causes were considered first.

“It just got bigger and bigger and bigger, over months really […] then they doubled them (diuretics), and then they trebled them”. [RG05].(13)

Effects of cancer associated thrombosis

The effects of cancer associated thrombosis theme include three subthemes (responses to venous thromboembolism diagnosis, psychological and physical effects).

Effects of diagnosis process

Patients’ perspective on cancer associated thrombosis diagnosis varied. Some participants reacted to the diagnosis of cancer associated thrombosis as an entity distinct to the cancer, while others considered cancer associated thrombosis as a complication of their cancer. However, in both cases the diagnosis of cancer associated thrombosis had a negative impact; it led to delays in cancer treatment and added more burden to their health.

“Having the cancer and then the thrombosis on top of it, not knowing how bad it was”(13)

“The fact that there were clots meant we couldn’t operate on my leg. Not being able to operate my leg pushed back my radiation and chemotherapy. So everything was shifted in time”(30)
Psychological effects

Four studies reported that the diagnosis of cancer associated thrombosis was distressing, especially in those without prior knowledge of the symptoms, had a major impact on patients’ lives and was perceived as life threatening (13, 30-32).

“PE is not cancer but it’s dangerous too, because with both you are playing with your life”(30).

“I felt I was having a heart attack....that stress made (the symptoms) worse”(30).

“[...] having the cancer and then the thrombosis on top of it, erm, not knowing how bad it was when I went in, I know I was in terrific pain with my chest and that erm, it was frightening to be honest. [VCC01]”(13)

“It frightened the life out of me, I was more scared of that than the cancer. You know blood clots can kill you like that (clicks fingers), cancer you’ve got a little bit of chance, you know. [PT13]”(31)

The response to the diagnosis was less stressful among participants with previous experience of venous thromboembolism who reported being calm and not shocked.

“Knowing [that it is a PE] reassures you a little, nevertheless. . . I knew I need to go to the hospital as fast as possible”(30)

“Would er the main thing that did it was erm apprehension basically about er the clot and if a clot does occur it hits you like that bang” (Interview NC5)(32)

Physical Effects

The acute and chronic symptoms of cancer associated thrombosis were profound and negatively affected patients’ lives. Mockler 2012 and Seaman 2014 described the negative impact of symptoms that interfered with patients’ daily living(30, 31).

In particular participants with PE described that being short of breath prevented them from
completing even small tasks at home. Symptoms from cancer associated thrombosis prevented them from returning to normal life and activities; unable to do daily activities around the house or to mobilize unaided.

“I cannot do anything...will I always continue heading in this regression”(30)

“The lack of energy and being out of breath....it’s just so frustrating ...Frustration of not being able to be where I should be, in my mind”(30)

“I couldn’t breathe; I literally couldn’t breathe and couldn’t talk. [PT6]”(31)

“I was very breathless, even bending down to the washing machine to put a wash in I was gasping for air. [PT11](31)

Effects of venous thromboembolism treatments

This theme captured patients’ experience of anticoagulation treatment (self-injected weight-adjusted Low molecular weight heparin, warfarin and direct oral anticoagulants). Four studies assessed patients’ responses to anticoagulation focusing on the acceptability of Low molecular weight heparin (30-33) It was clear that the treatment had a positive effect on patients’ life, especially, for those who had experienced distressing symptoms.

Most participants were started on Low molecular weight heparin and others had been on warfarin but changed to Low molecular weight heparin due to absorption difficulties, uncontrolled INR and/or venous thromboembolism recurrence. There was a general agreement by patients that Low molecular weight heparin was acceptable treatment and better than warfarin where comparison was possible. Self-injected Low molecular weight heparin allowed more self-control over their life and more freedom. However, this benefit was not without compromise, as side effects of bruising and injection-site lumps were common.

“The heparin is so much simpler than all the ****ing about with warfarin’ (22CS)” (33)
“I really don’t feel like pricking myself, but if it’s that or dying well I’d rather prick myself” (30).
“I used to spend my life travelling to hospital for a warfarin check ... sat in the car ... sat in the waiting room ... not much of a life really” (19CS) (33)

“With the warfarin, what was kind of crappy was that I had to do blood tests every two weeks. But with Low molecular weight heparin no need for draws” (30)

“I’m using the tops of my legs now so it isn’t as painful. I was using my stomach but after a while your stomach gets really hard and then you’ve got to really force them in”. [PT8](31)

Those with advanced disease felt that treatment of their cancer associated thrombosis with optimal anticoagulation meant that their doctors did not give up on them, influencing their optimism and expectation of their doctors.

“I know I’m going to die. I know that the doctors don’t have any more chemo to give ... you don’t like feeling that you’ve been put on the scrap heap ... the injection isn’t stopping the cancer but it is stopping the blood clots” (14CS)(33)

“It is important to know that people are still doing something”(15C) (33)

Some patients wish to take Low molecular weight heparin for longer than 6 months as long because of the continued risk and the peace of mind they felt with ongoing anticoagulation. This is expressed clearly by a patient in ALICAT study where views were sought on being randomised to continuation or cessation.

“The thing that bothered me at that time was that I’d already had two episodes of of a blood clot and I thought if I was in the group that didn’t have the medication um there was a good chance that I’d I’d have another one and it could could have a lot more serious repercussions than if I just continued to take this medication”. (Interview NC3)(32)

“Um I thought no I think I’ll carry on rather than, you know, spending another day in hospital being prodded and probed like I was last time”. (Interview NC2)(32)

However; some patients wanted to stop Low molecular weight heparin injection after 6 months because of the side effects of the injections and others wanted to restore their normal life without injections.

“I was just happy to get off of it to be honest with you, um it was more or less the same time every
night, um and the pain as I said eh to me was terrible, horrific and a lot of bruising and things”

(Interview NC1)(32)

“And so I was very keen I have to say, I was predisposed I don’t want any further injections once the treatments finished I just want to try to get back to as much normality as I can”. (Interview NC8)(32)

Employment of coping mechanisms

It was clear that the response to the dual diagnosis of cancer and cancer associated thrombosis was very individual, with some regarding it as a greater shock than their cancer, particularly among patients who considered cancer associated thrombosis as a setback on the road to cancer recovery, and others viewing it as less significant in comparison with their cancer (30).

“I never broke down when I was told about the cancer [...] I had the operation, went on the chemo, everything. The only time I broke down was when I went back in hospital when they told me I had blood clots [...] the cancer to a point they can treat, hold it back – blood clots they go so quick and that frightened me, it was the only time I broke down”. [PT13](31)

“During chemotherapy, I didn’t have any great nausea, and brachytherapy went well too. So I told myself, “Well, I’m going to overcome the cancer but No! Then I started to go down again”.(30)

The employment of coping mechanisms theme illustrated ways that patients developed to move on with their lives. The treatment of venous thromboembolism brings with it symptomatic relief, reassuring patients that their condition is improving. This reduces distress and allows patients, over time, to get back to ‘some sort of normality’(13). Participants described the development of strict routines and rituals to ensure Low molecular weight heparin was administered on time and without fail.(31)

“I usually take them between 8 and half past 8. And then I know it’s done, and I don’t forget for the day, then, because someone I was talking to, he was saying “You don’t do it in the night, do you?” and I said, “No, I get up, have my cup of tea then 8, half past 8 do it.” [PT13](31)
“[..] is a ritual now”. [VCC10](13)

DISCUSSION

The cancer journey is difficult in itself, but thrombosis was an additional, frightening and unexpected burden. The association between cancer and venous thromboembolism was first reported in the 19th century (34). However, cancer patients are still not routinely educated about the risk or warning symptoms/signs of venous thromboembolism which may otherwise be misattributed to the cancer by patient and clinician alike.

This systematic review highlights the impact of cancer associated thrombosis on the lives of cancer patients, and calls for education for patients and clinicians to be part of routine care, and further work to address this patient priority.

The four themes from this synthesis of primary qualitative studies (knowledge deficit, effects of cancer associated thrombosis, effects of anticoagulation, and employment of coping mechanisms) illustrate the ways in which cancer associated thrombosis affects quality of life. Thrombosis with its complex presentation, diagnosis and treatment, was seen by many patients as a significant additional, frightening and unexpected burden affecting cancer treatment and which impose psychosocial and functional limitations.

Lack of knowledge of cancer associated thrombosis

A survey of cancer patients found a better level of knowledge of cancer associated thrombosis risk than those in this review, but still half (53%) of participants were unaware of the increased risk of cancer associated thrombosis although three quarters knew that venous thromboembolism can be prevented.(35)
Eventhough the cancer associated thrombosis is not a new phenomenon, a lack of clinician awareness of cancer associated thrombosis appears to compound the lack of patient knowledge (36, 37). In a study of 18 patients who had had venous thromboembolism, (not cancer-related), misdiagnosis and diagnostic delay made patients feel angry and distrustful of their medical team.(38) However, similar to Mockler’s study of people with cancer associated thrombosis, patients with a previous history of venous thromboembolism were less distressed, having recognised the symptoms and sought medical help more quickly (30).

This lack of routine information giving is in contrast to other cancer-related complications such as malignant spinal cord compression and post chemotherapy neutropenia, where guidelines are systematically applied for patients and their family, and carers. Education includes the symptoms and signs to look for, when to seek medical help, and who to contact (39, 40).

Inadequate information-giving is not new (41). However, people with venous thromboembolism appear clear about the degree of information they need including understanding their diagnosis, what they should do/ do not from diagnosis and with treatment (42). Information needs are individual and vary by gender, age and stage of disease; some wishing for full details; others want basic information only (43).

Psychological burden
The long-term outcomes following acute venous thromboembolism extend beyond the physical burden (44, 45) and the experience of symptomatic pulmonary embolism is a life-changing, distressing and frightening event (42). Similar psychological effects were seen in this review, but in addition, cancer patients have to process this event in the context of the underlying cancer. For some, the potential of cancer associated thrombosis as a sudden killer came as a great shock, especially those who had viewed their cancer as a chronic illness.
**Effects of cancer associated thrombosis treatment**

Seaman et al found that efficacy of treatment was paramount despite the hypothetical preference of a tablet over an injection (31). This finding was also highlighted in the study by Noble et al (46), where patients were concerned about safety, efficacy, and lack of interference with anticancer therapies ahead of method of administration. Patients in this review included those with advanced disease, but despite previously stated concerns about patient burden with Low molecular weight heparin (47), the use of Low molecular weight heparin was acceptable. The daily injections of precalculated dose of Low molecular weight heparin giving more control than the blood tests and dose alterations of warfarin.

**Employment of coping mechanisms**

Patients tried to maintain a sort of normality in everyday life consistent with previous findings that cancer patients find ways to minimize the impact of the side effects of cancer in their new life situation. (45)

Education and support is important with regard to coping. One of the earliest reported educational group programmes for people with cancer was the ‘I Can Cope’ (ICC) programme (48). This has been well evaluated showing reduced anxiety, improved disease-related knowledge and sense of meaning (49).

**Uncertainty and information**

The varying responses to the threat of cancer associated thrombosis and its treatment seem to be related to uncertainty as to whether the cancer associated thrombosis will recur, whether it will resolve, whether the treatment will be effective and/or harmful. Uncertainty management theory (50, 51) is one theoretical framework to help the understanding of how patients encounter,
appraise levels of danger, seek information, respond to and cope with health-related threats. The differing needs for information, ways of seeking it and success in receiving it are seen within these data presented. Likewise, some patients appraised the cancer associated thrombosis as very dangerous, whereas others (often those with previous experience and better information) were able to appraise it as less dangerous because they knew what to look for and how to act. Whilst the relationship between uncertainty and danger appraisal is complex, tailored and accessible information seems to play a key part in reducing anxiety even if absolute reassurances cannot be given (52, 53). As Brashers states, uncertainty occurs when, “information is unavailable or inconsistent; and when people feel insecure in their own state of knowledge or the state of knowledge in general” (50).

Strengths and limitations
As with any systematic review it is possible to miss relevant studies. The included studies were qualitative research which is designed to give insights from the patients involved rather than to be generalizable. However through synthesis more generalizable findings can be derived (26). Only one included study came from outside the UK, however, they were from different centres, but still indicated similar concerns.

Only limited papers were found, illustrating that this area has been under-researched. The serious concerns highlighted by this review show that further work is needed.

Implications for clinical practice and policy makers
Raised clinical awareness and the provision of basic information for patients about the risk of cancer associated thrombosis is a policy priority in the UK. (54) Information about cancer associated thrombosis, both written and verbal, should be provided routinely for patients at diagnosis. cancer associated thrombosis should be part of standard training and education for all clinicians caring for people with cancer, including those in primary and palliative care. Recent initiatives such as the
International Initiative on Thrombosis and Cancer(55) should help raise awareness and help with high quality training. Streamlined clinical services for diagnosis and treatment of cancer associated thrombosis aiming to minimise time in hospital awaiting tests, especially for those with advanced disease, should improve clinical decision making.(37, 56, 57)

Conclusion

This systematic review highlights the impact of cancer associated thrombosis on the lives of cancer patients, and calls for education for patients and clinicians to be part of routine care, and further work to address this patient priority equal to that of other cancer complications such as spinal cord compression or neutropenic sepsis.

Declaration.

Authorship; NB, MJ and AM conceived the research question, NB, MJ developed the search strategy, NB, MJ, JS and AH contributed to design. NB, WI screened titles and abstracts with recourse to MJ; NB extracted data and wrote the first draft; All authors interpreted the data and contributed to drafts and approved the final manuscript.

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Conflicts of interest: Authors have the following interests to declare: AH, AM and MJ received funding to complete a qualitative substudy of anticoagulation patient experience funded by BAYER, JD, WI and NB have nothing to declare.

Ethics and consent: This was not required for this published anonymised evidence synthesis.

Data sharing: Data are available from the published papers in this review.
References


Table 1. Overview of the articles included

<table>
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<tr>
<th>Study</th>
<th>Study design</th>
<th>Methods &amp; Sitting</th>
<th>Aim</th>
<th>Participants characteristics</th>
<th>Analysis</th>
<th>Principal findings</th>
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</table>
| Noble, S. et al. 2015 (13) | Qualitative | Semi-structured interview. | Exploring the patients experiences of cancer associated thrombosis within the context of cancer journey | N = 20 patients (10 women and 10 men) Age: 53-81 years Different primary cancers receiving Low molecular weight heparin for (2-20 months) | Framework analysis | Diagnosis and treatment of cancer associated thrombosis:  
- Lack of knowledge of venous thromboembolism in the context of cancer, patients unaware of risks of thrombosis or symptoms to look out for,  
- Limited awareness among health professionals.  
- Symptoms of cancer associated thrombosis attributed to cancer or chemotherapy and therefore delayed presentation to hospital  
- Initial reaction is shock, little information. Living with cancer associated thrombosis:  
- Treatment helps get over the initial shock, i getting on with life, ritualization of new routines |
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<tr>
<th>Study</th>
<th>Methodology</th>
<th>Participants</th>
<th>Setting</th>
<th>Themes/Findings</th>
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| Mockler, A, et al. 2012(30) | Qualitative, Semi-structured interview. | N = 10 (4 women and 6 men) aged 35-78 years | Inpatients and outpatients of a large urban university-affiliated hospital, Montreal, Canada | Exploring the experiences of patients with cancer who developed venous thromboembolism. Various cancer types diagnosed 2-18 months prior to the interview. Various stages from early with active treatment to advanced stage. Thematic analysis:
- Prior knowledge of cancer associated thrombosis risk and symptoms (or lack of knowledge) determined reaction to cancer associated thrombosis symptoms.
- For some, cancer-related concerns overshadowed those due to cancer associated thrombosis. Cancer associated thrombosis as a setback in cancer care:
  - Cancer associated thrombosis symptoms preventing a return to normal life after cancer treatment.
  - Cancer associated thrombosis treatment interfering with their cancer care. Attitudes about venous thromboembolism treatments:
  - Positive for some participants however associated with a sense of obligation.
  - Many show acceptance of self-injection of Low molecular weight heparin especially among those with previous experience with warfarin. |
| Seaman, S et al. 2014(31) | Qualitative, Semi-structured interview. | N =14 (8women and 6 men) Age 52-84 years | Palliative care and cancer associated thrombosis unit, Cardiff, UK | Exploring the acceptability of long term Low molecular weight heparin for the treatment of cancer associated thrombosis in the contexts of living with cancer and quality of life. Thematic analysis:
- Symptom burden of cancer associated thrombosis.
- Impact on activities of daily living.
- Acceptability of Low molecular weight heparin:
  - Necessary inconvenience.
  - Systematic approach to injection. Hypothetical views on New Oral Anticoagulants:
  - Efficacy paramount.
  - Willing to engage in clinical trials. |
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<tr>
<th>Noble S, et al. 2015 (32)</th>
<th>ALICAT</th>
<th>Embedded Qualitative study within a RCT</th>
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<td>Focus groups with clinicians.</td>
<td>3-11 group (3 focus group)</td>
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<td>Semistructured interviews with patients and their relatives</td>
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<td>Oncology, Haematology and Primary care.</td>
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<td>N of patients=8 (4 females)</td>
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<td>Locally advanced or metastatic cancer. Receiving Low molecular weight heparin for cancer associated thrombosis.</td>
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<td>● Patients’ perception of cancer associated thrombosis and Anticoagulation</td>
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<td>N = 40 (18 male and 22 female)</td>
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<td>● Simplicity: the majority found that daily injection of Low molecular weight heparin simpler than the frequent INR needed for warfarin.</td>
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<td>● Freedom: many patients expressed a feeling of freedom from hospitals, from being restricted to their home.</td>
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<td>● Optimism: the feeling that something active being done.</td>
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<td>● Bruising: 11 patients described bruising as a negative aspect of Low molecular weight heparin.</td>
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<th>Noble S and Finlay, I.G. 2005 (33)</th>
<th>Qualitative</th>
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<td>Semi-structured interview. Palliative care patients, both in the community and in-patient units. Cardiff. UK</td>
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<td>Assessing the appropriateness of Low molecular weight heparin in palliative care patients and the extent of daily injection burden</td>
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<td>● Simplicity: the majority found that daily injection of Low molecular weight heparin simpler than the frequent INR needed for warfarin.</td>
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<td>● Freedom: many patients expressed a feeling of freedom from hospitals, from being restricted to their home.</td>
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<td>● Optimism: the feeling that something active being done.</td>
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<tr>
<td>● Bruising: 11 patients described bruising as a negative aspect of Low molecular weight heparin.</td>
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</tbody>
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