The IDEAL DVT study, individualized duration elastic compression therapy against long-term duration of therapy for the prevention of post thrombotic syndrome, a randomized controlled trial

<table>
<thead>
<tr>
<th>Journal:</th>
<th>BMJ Open</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manuscript ID:</td>
<td>bmjopen-2014-005265</td>
</tr>
<tr>
<td>Article Type:</td>
<td>Protocol</td>
</tr>
<tr>
<td>Date Submitted by the Author:</td>
<td>14-Mar-2014</td>
</tr>
</tbody>
</table>
| Complete List of Authors: | ten Cate, Arina; Maastricht University Medical Centre, Laboratory for Thrombosis and Hemostasis; Maastricht University Medical Center, Internal Medicine  
Bouman, Annemieke; Maastricht University Medical Centre, Laboratory for Thrombosis and Hemostasis; Maastricht University Medical Center, Internal Medicine  
Joore, Manuela; Maastricht University Medical Centre, Department of Clinical Epidemiology and Medical Technology Assessment (KEMTA)  
Prins, Martin; Maastricht University Medical Center, Epidemiology; Maastricht University Medical Centre, Department of Clinical Epidemiology and Medical Technology Assessment (KEMTA)  
ten Cate, Hugo; Maastricht University Medical Center, Internal Medicine; Maastricht University Medical Centre, Laboratory for Thrombosis and Hemostasis |
| Primary Subject Heading: | Cardiovascular medicine |
| Secondary Subject Heading: | Haematology (incl blood transfusion), Health economics |
| Keywords:          | Post thrombotic syndrome, Deep vein thrombosis, Elastic compression stocking therapy, Protocol, RCT |
Study Protocol

The IDEAL DVT study, individualized duration elastic compression therapy against long-term duration of therapy for the prevention of post thrombotic syndrome, a randomized controlled trial

Authors: Arina J ten Cate-Hoek¹², Annemieke C Bouman¹², Manuela Joore³, Martin Prins³⁴, Hugo ten Cate¹², for the IDEAL DVT trial investigators.

Corresponding author: Arina J ten Cate-Hoek MD, PhD.
Address: P Debyelaan 25
6229 HX Maastricht
E-mail: arina.tencate@maastrichtuniversity.nl
Telephone number: 0031(0)43-3871243
Fax number: 0031(0)43-3876096

¹Laboratory for Thrombosis and Hemostasis, Maastricht University Medical Centre, Maastricht, the Netherlands
²Department of Internal Medicine, Maastricht University Medical Centre, Maastricht, the Netherlands
³Department of Clinical Epidemiology and Medical Technology Assessment (KEMTA), Maastricht University Medical Center, Maastricht, The Netherlands
⁴Department of Epidemiology, Care and Public Health Research Institutes, Maastricht University, Maastricht, the Netherlands

Key words: Post thrombotic syndrome, deep vein thrombosis, elastic compression stocking therapy, protocol, RCT

Word count: 3345
Abstract

Introduction:
Post thrombotic syndrome (PTS) is a serious complication of deep vein thrombosis (DVT) of the leg that affects 20-50% of patients. Once a patient experiences PTS there is no treatment that effectively reduces the debilitating complaints. Two randomized controlled trials showed that elastic compression stocking (ECS) therapy after DVT for 24 months can reduce the incidence of PTS by 50%. However, it is unclear whether all patients benefit to the same extent from ECS therapy or what the optimal duration of therapy for individual patients should be. ECS therapy is costly, inconvenient, demanding and sometimes even debilitating. Tailoring therapy to individual needs could save substantial costs. The objective of the IDEAL DVT study therefore is to evaluate whether tailoring the duration of ECS therapy on signs and symptoms of the individual patient is a safe and effective method to prevent PTS, compared to standard ECS therapy.

Methods and analysis:
A multicentre, single-blinded, allocation concealed, randomized, non-inferiority trial. A total of 864 consecutive patients with acute objectively documented proximal DVT of the leg are randomized to either standard duration of 24 months or tailored duration of ECS therapy following an initial therapeutic period of 6 months. Signs and symptoms of PTS are recorded at regular clinic visits. Furthermore, quality of life, costs, patient preferences and compliance are measured. The primary outcome is the proportion of patients with PTS after 24 months.

Ethics and dissemination:
Based on current knowledge the broad application of ECS therapy is questioned. The IDEAL DVT study will address the central questions that remain unanswered: What individual patients do benefit from ECS therapy and what is the optimal individual treatment duration? Primary ethics approval was received from the Maastricht University Medical Centre. Results of the study will be disseminated via peer-reviewed publications and presentations at scientific conferences.

Trial Registration:
NCT01429714 and NTR 2597
Background

After deep vein thrombosis (DVT) 20-50% of patients develop post thrombotic syndrome (PTS). PTS is a chronic condition, characterized by leg complaints such as pain; heaviness; cramps; aching; tingling; and leg signs due to venous insufficiency, evolving in severe cases to venous ulceration of the leg that was affected by the DVT.PTS is a serious affliction, with substantial impact on quality of life and costs. Because an effective therapy is lacking, prevention of PTS is of major importance. Up till now, elastic compression stocking (ECS) therapy after DVT is the mainstay of PTS prevention. The evidence sustaining the value of ECS therapy following acute DVT is derived from 2 randomized clinical trials Incidences of PTS were reduced significantly (approximately 50%) by application of ECS therapy for 24 months in all patients in these studies. However, based on these trials it is still undecided whether all patients benefit to the same extent from ECS therapy or what the optimal duration of ECS therapy for individual patients should be. There have been other studies that highlighted different aspects of ECS therapy. A significant benefit of early ambulation and early compression measured as mean score on Villalta (a score containing 5 leg symptoms and 6 objective signs), was found in a follow-up study of a small trial that compared ambulation combined with inelastic compression (Unna boot) or ECS therapy to bed rest without any compression in the acute phase of 9 days, followed by ECS therapy for all patients during 2 years of follow up. ECS therapy may not be indicated for all patients as shown by a study that assessed whether ECS therapy initiated one year after the event of DVT would lower the incidence of PTS in patients without complaints but with reflux on duplex testing. No significant benefit of ECS therapy at this late time of onset was found. The incidence of PTS was low in both groups. Only one study so far assessed whether prolonged duration of ECS therapy was superior to 6 months of ECS therapy following an event of proximal DVT. In this study no significant difference in the incidence of PTS according to the CEAP classification was observed between the two treatment groups. The safety of shortened duration of ECS therapy based on individual patient clinical scores was assessed in our management study, and we showed that tailoring the duration of ECS therapy based on the signs and symptoms of the individual patient after an initial treatment period of 6 months, is a safe strategy to prevent PTS. We found that 50% of our patients did not need ECS therapy for as long as 2 years, while the overall incidence of PTS was 21.1% (95% CI 13.5- 28.7). This is comparable to published incidences after 24 months ECS therapy. In the Netherlands, each year around 25,000 patients experience a new event of DVT. In order to
prevent PTS, common practice is to prescribe custom fitted ECS therapy for 2 years for all patients.\textsuperscript{13} ECS therapy is costly, demanding and sometimes even debilitating. Elderly patients can often not apply the ECS by themselves but need help from family or they even need home care visits (7.5%).\textsuperscript{15-17} Total annual costs of ECS therapy roughly amounts to 2.5 million for stockings (25.000 patients*100 euro) and 21 million for home care (7.5%*25.000 patients*500 visits*20 euro).\textsuperscript{15-17} Moreover, the majority of patients do not experience any post thrombotic complaints.\textsuperscript{10} In the Netherlands, over 10 million euro each year could be saved if ECS treatment would be individualized. The results of a recently published trial performed by Kahn et al. refuted the routine wearing of ECS after DVT altogether. This study showed no treatment effect at all from ECS therapy in contrast to the large treatment effects found in previous non-placebo controlled trials.\textsuperscript{6,7} The unexpected lack of effectiveness cannot be dismissed solely as a placebo effect. The compliance to ECS therapy, a major determinant of effectiveness, was 55.6\% after 24 months, less than in the previous trials by Brandjes and Prandoni et al, where compliance was considerably higher, up to 90%.\textsuperscript{6,7,18} Because compliance to therapy is one of the most important features for successful ECS therapy, we will perform a discrete choice experiment in which patient preferences towards ECS therapy will be assessed and the main obstacles for compliance to therapy will be addressed.

Based on current knowledge it is understood that the broad application of ECS therapy is questioned. Especially the benefit to individual patients and the optimal duration of ECS therapy were never studied properly and as a result it is unclear which individual patients require therapy and for how long.

The IDEAL DVT study aims to address these topics. In addition quality of life, patient preference towards ECS therapy, compliance to therapy, as well as cost-effectiveness of ECS therapy will be assessed. The primary outcome is the proportion of patients with PTS at the end of follow-up.

\textbf{Methods/Design}

A multicentre, single-blinded, allocation concealed, randomized, non-inferiority trial comparing individually tailored duration of ECS therapy (intervention) with standard duration of 24 months ECS therapy (control), for the prevention of PTS. Randomization will guarantee a fair distribution of patients within each patient group. Stratification will be performed on centre and potentially confounding effects such as age, sex and body mass index. The study will be a multi center study in order to get a good
representation of patients and to achieve sufficient patient numbers. The primary outcome will be PTS at 24 months after the event; the observers will be blinded to the allocated treatment arm. The allocation will be concealed from study personnel involved in assessing the leg symptoms and rating the PTS scores. Both randomization and allocation concealment are used as strategies against bias. The study is not double blinded; no inactive stockings will be used in the comparator group. The use of sham stockings will very likely interfere with the quality of life and will complicate the assessment of perceived differences between groups. Patients will be followed for the entire study duration; incidence of recurrent DVT and VTE related death during the follow-up period will be documented.

Patients and treatment

Consecutive, consenting adults with an acute objectively documented proximal DVT of the leg, adequately treated with anticoagulant treatment are included in the study. Patients are included within 6 weeks after DVT. Exclusion criteria are: previous DVT in the affected leg, recurrent DVT in the 6 months following inclusion, pre-existent venous insufficiency (skin signs C3-C6 on CEAP score or requiring ECS therapy), contraindication for ECS therapy such as intermittent claudication or clinical signs of leg ischemia or asymptomatic arterial insufficiency (a pulse deficit or bruit at sites of narrowing at physical examination), active thrombolysis, and limited life expectancy (< 6 months).

Patients are recruited from 12 hospitals in the Netherlands and 2 hospitals in Italy. In the initial acute phase after DVT, until the acute oedema has disappeared, three strategies will be applied. The leg is either bandaged with short stretch bandages to a compression of 30-40 mmHg, worn day and night and redressed twice per week; or a bandage stocking with 35mmHg compression is prescribed (Mediven Struva 35® , Medi, Breda, the Netherlands), worn day and night; or no initial compression therapy is applied, according to the strategy of the hospital protocol. After the initial phase, a custom fitted, flat knitted, knee length graduated elastic compression stocking class III (ankle pressure 40 mmHg) is prescribed for all patients. Patients are advised to wear the stockings during the daytime.

Procedures

After obtaining written informed consent, randomization is performed centrally at the coordinating study centre, the Maastricht University Medical centre. Patients are randomized to one of the two
treatment arms. (Figure 1) A web-based randomisation program (TENALEA (Trans European Network for Clinical Trials Services)) is used that executes blocked randomisation with stratification on centre-level and on possible confounding patient characteristics such as age, sex, and Body Mass Index. Permuted blocks of randomly varying size are used to maintain balance of numbers in each treatment group and to ensure allocation concealment.

Information on allocation is only accessible to the study personnel at the coordinating study centre in Maastricht, and patients will receive information about their allocation via the coordinating study centre. The allocation will be concealed from study personnel involved in assessing the leg symptoms and rating the PTS scores. Patients are not blinded by the use of sham stockings, because they may influence quality of life, complicating the assessment of perceived differences between groups. Patients are asked not to reveal their treatment allocation to their own physician, and not to wear the elastic compression stocking at the day of their outpatient clinic visits.

All patients are advised to wear the elastic compression stocking during the first 6 months after DVT. In the control group, all patients are instructed to wear the elastic compression stocking daily for a total duration of 2 years. In the Intervention group, duration of ECS therapy is individually tailored based on Villalta scores.\(^1\) After the first 6 months therefore, there are three scenarios:

1. The Villalta score at both the 3 and 6-month follow-up visit is ≤ 4. Then the patient is advised to discontinue ECS therapy.
2. The Villalta score at the 3-month follow-up visit is ≥ 5 but the Villalta score at the 6 month follow-up visit is ≤ 4. The patient is advised to continue the ECS therapy for another 6 months. If the Villalta score at the 12 month follow-up visit is ≤ 4, the patient is allowed to discontinue the ECS therapy,
3. The Villalta score is ≥ 5 at both the 3 and 6 months follow-up visit. The patient is advised to continue the ECS therapy for a total duration of 24 months. (Figure 2)

Once stopped, ECS treatment will not be reinstated. Using this algorithm the duration of ECS therapy is individually tailored in each patient. The decision on the duration of ECS therapy in patients in the intervention group is made centrally at the coordinating study centre, as only there the information on
treatment allocation is available. The patient is informed about the decisions on the (dis) continuation of the ECS therapy by the coordinating centre.

Follow-up

All patients are followed for 24 months after DVT. During these 24 months they visit the outpatient clinic 4 times: at 3, 6, 12, and 24 months after the DVT. At each follow-up visit the signs of PTS will be recorded and scored, by the study nurse or the treating physician, using the objective part of the Villalta clinical scale for PTS.19 In case of intermittent leg complaints or signs or symptoms suspect of recurrent VTE, patients are instructed to visit their treating physician. Study documentation will be filled out and adverse events will be recorded. The CCMO (central committee human related research) will be notified in case of serious adverse events or death.

The patient is asked to fill out 5 questionnaires: at baseline, and 3, 6, 12 and 24 months later. All questionnaires will be offered as a web based application, each patient will have a unique personal entry code. For patients who are not able or willing to use the electronic questionnaires, a paper version will be available. The first questionnaire is filled out shortly after inclusion (4-6 weeks after DVT) and the subsequent questionnaires are filled out the day before each follow-up visit. All questionnaires contain the subjective part of the Villalta clinical scale for PTS,20-23 questions on ECS compliance, a cost questionnaire, and three health related quality of life (HRQOL) questionnaires: the Dutch translated Veines-Qol,24 the SF-36,20 and the EuroQOL-5D.21 In a subset of the study population patient preferences regarding elastic compression therapy will be analysed, by conducting a discrete choice experiment (DCE). For this purpose, a DCE questionnaire will be send to a subset of the population at approximately 3 months after DVT.

Compliance to ECS therapy is monitored in two ways. Each questionnaire contains a question on compliance to the advised version of ECS therapy. Because compliance and adherence to therapy is of crucial importance the number of contact moments for the assessment of compliance with ECS therapy will be increased by adding telephone contacts to the regular clinical visits. Study-supporting staff will make computer-assisted random telephone calls during the entire follow-up period to both the patients in the group with the 2-year ECS intervention and to patients in the group with individualized ECS therapy. The telephone calls will be made randomly to create a surprise effect (patients are off
guard) and to allow for a distribution of contacts over the entire study period. Per patient 3 extra contact moments will be created. A standard questionnaire will be used to assess compliance and to address reasons for non-compliance. Crossover to another therapy arm will be discussed and discouraged.

Outcomes

The primary outcome is the proportion of patients with PTS at the end of follow-up, 24 months after DVT. PTS is defined as a Villalta score of ≥5 at two consecutive visits that are at least three months apart. Although the consensus is to diagnose PTS at the 6-month visit or later based on one single Villalta score ≥5, we modelled this study after the preceding management study in which two consecutive Villalta scores ≥5 were needed for the diagnosis PTS. We observed that a proportion of patients (about 12%) had fluctuating Villalta scores beyond 6 months after the acute event of DVT. Assigning the diagnosis based on one single score ≥5 would therefore lead to misclassification and overestimation of the PTS incidence. The Villalta scores at 6 months will be available; therefore also the proportion of patients with PTS according to the consensus scoring method will be presented.

Secondary outcomes include:

1. HRQOL, measured by questionnaires (SF-36, EuroQOL-5D, Dutch translated Veines-Qol)
2. Costs
3. Recurrent thrombosis, according to criteria as published and assessed by objective tests
4. VTE related death during follow-up, assessed by an independent and blinded adjudication committee
5. Patient preferences, assessed with a Discrete Choice Experiment (DCE)

Since both effects on costs and generic health-related quality of life are to be expected, the method of economic evaluation is a cost-utility analysis (CUA). The analysis will be from a societal perspective. The primary effect parameter is generic health-related quality of life, measured in quality adjusted life years (QALYs). We will perform two analyses: a trial based CUA with a time horizon of two years (identical to the duration of follow up in the clinical study), and a model-based CUA with a lifelong time horizon.
horizon. For the latter, we will follow the guidelines of good modelling.27 We will be able to adapt a
Markov model for diagnostic strategies in DVT, which was developed by our group earlier.28 Costs in
the economic analyses include direct health care costs (medical costs for prevention, diagnostics,
therapy, rehabilitation and care), direct non-health care costs (travel costs) and indirect costs
(productivity loss). Utilities will be calculated from the responses on the EQ-5D and SF36 classification
systems using the available multi-attribute utility functions.29 30
The results of the DCE will provide insight in the trade-offs patients make between characteristics of
the therapy. More specifically, we will investigate what duration of ECS therapy, with its drawbacks,
patients consider to be acceptable.

Sample size calculation
The proposed study is a non-inferiority trial and aims to demonstrate that the assessed alternative
therapy based on ECS for 6 months followed by individually prolonged ECS therapy based on Villalta
scores is not worse than the comparator ECS therapy for a standard duration of 24 months, by more
than the prespecified amount of 7.5% (the non-inferiority margin). The published incidence of PTS
following two years of ECS lies between 20 and 30%. (Prandoni 24.5% (95% CI 15.6- 33.4), Brandjes
20% (95% CI 12.4-29.2) As it is statistically impossible to demonstrate equivalence (prove the H0 of
no difference), Blackwelder proposed a one sided significance test to reject the null hypothesis by a
clinically acceptable amount.31 If we allow a difference of 7.5% in the outcome PTS between the group
with ECS for 2 years and the group with the alternative therapy, 70% of the effect will be preserved.
This proportion of loss in efficacy is customarily accepted in controlled randomized clinical trials. At a
one sided significance level of 0.05 and a power of 80%, with a ratio of 1, a total of 847 patients is
needed to provide sufficient patients for an adequately powered trial.31 32 Loss to follow-up of patients
is expected to be less than 2% since the intervention does not have an invasive nature. Therefore, a
total of 864 patients are needed (432 patients per treatment arm).

Statistical analysis
Descriptive statistics of the total population and of the two treatment groups separately will be
computed to provide baseline characteristics of the patients in both treatment arms.
For the primary outcome PTS, univariate analysis of all proportions will be performed with logistic regression (chi-squared) analysis. Kaplan-Meier method will be used to calculate the cumulative incidences of PTS, adjusted for centre, to compare incidence rates between the two treatment arms. Patients who die or are lost to follow-up will be censored at their last visit. ANOVA will be applied to assess the changes over time for different outcome measures.

Hazard ratio’s and 95% confidence intervals for both treatment arms will be calculated using Cox regression models. Hazard ratio’s will be adjusted for age, sex, clinical presentation of DVT, and extent of the index deep vein thrombosis.

Interim analysis (safety)
A prespecified safety analysis will be performed when 50% of the subjects have completed the 2-year follow-up. The analysis will be performed by the coordinating centre, and supervised by the principal investigators. The safety analysis will be performed to assess significant enhanced risk of PTS or excess morbidity/mortality in the intervention arm of the study population. Fisher’s exact test will be performed to compare incidence of PTS at a significance level of 0.05 (two sided). The study can be stopped in case of significant excess morbidity/mortality in the alternative treatment arm.

Ethical considerations
The medical ethical committee of the Maastricht University Medical centre approved this study. All patients are extensively informed about the study, and written informed consent is obtained from all participating patients.

Discussion
Based on current knowledge the broad application of ECS therapy after DVT is questioned. The results of a recently published trial performed by Kahn et al. refuted the routine wearing of ECS after DVT altogether (ref). This study showed no treatment effect at all from ECS therapy (graduated ECS 30-40 mmHg) compared to placebo ECS therapy (5 mmHg). This is in sharp contrast to the large treatment effects found in previous non-placebo controlled trials. The unexpected lack of effectiveness cannot be dismissed solely as a placebo effect. The compliance to ECS therapy, a major determinant of effectiveness, was 55.6% after 24 months, much less than in the previous trials by
Brandjes and Prandoni et al, where compliance was considerably higher, up to 90%. Therefore the definitive answer on the usefulness of ECS therapy for the prevention of PTS is not yet provided. The need for the IDEAL DVT remains unchanged. The benefit to individual patients and the optimal duration of ECS therapy were never studied properly and as a result it is unclear which individual patients require therapy and for how long. The IDEAL DVT study will address the central questions that remain unanswered: What individual patients do benefit from ECS therapy and what is the optimal individual treatment duration? The study that we plan to perform is not only innovative; for the first time individual tailoring of duration of ECS therapy will be investigated, but also it will provide unique additional knowledge on the safety, effectiveness and cost-effectiveness of this approach. The IDEAL DVT study started including patients March 2011 and is currently on-going.
Trial status
The IDEAL DVT study is currently in progress. Patient recruitment started March 15th 2011 and is still on-going.

Acknowledgments
The IDEAL DVT trial is funded by a grant from Zon-mW the Netherlands. (171102007)
We especially acknowledge the participating investigators.

Competing interests
The authors declare that they have no competing interests.

Funding
This work was supported by Zon-mW the Netherlands grant number 171102007.

Contributions
AtC wrote the protocol and designed the study, the other authors MJ (cost-effectiveness research), MP (study design/statistics) and HtC (study design) contributed. AB is a PhD student and coordinator of the trial.

Ethics approval
Maastricht University Medical Centre.

The IDEAL investigators:
The Netherlands:
Maastricht, Maastricht University Medical Centre
Arina J ten Cate-Hoek MD, PhD (principal investigator)
Hugo ten Cate MD, PhD
Manuela Joore PhD
Annemieke C Bouman MD
Groningen, University Medical Centre Groningen
Karina Meijer MD PhD
References


Study flow diagram
254x190mm (96 x 96 DPI)
Algorithm individually tailored ECS therapy

177x161mm (96 x 96 DPI)
The IDEAL DVT study, individualized duration elastic compression therapy against long-term duration of therapy for the prevention of post thrombotic syndrome, a randomized controlled trial

<table>
<thead>
<tr>
<th>Journal:</th>
<th>BMJ Open</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manuscript ID:</td>
<td>bmjopen-2014-005265.R1</td>
</tr>
<tr>
<td>Article Type:</td>
<td>Protocol</td>
</tr>
<tr>
<td>Date Submitted by the Author:</td>
<td>08-May-2014</td>
</tr>
</tbody>
</table>
| Complete List of Authors: | ten Cate, Arina; Maastricht University Medical Centre, Laboratory for Thrombosis and Hemostasis; Maastricht University Medical Center, Internal Medicine  
Bouman, Annemieke; Maastricht University Medical Centre, Laboratory for Thrombosis and Hemostasis; Maastricht University Medical Center, Internal Medicine  
Joore, Manuela; Maastricht University Medical Centre, Department of Clinical Epidemiology and Medical Technology Assessment (KEMTA)  
Prins, Martin; Maastricht University Medical Center, Epidemiology; Maastricht University Medical Centre, Department of Clinical Epidemiology and Medical Technology Assessment (KEMTA)  
ten Cate, Hugo; Maastricht University Medical Center, Internal Medicine; Maastricht University Medical Centre, Laboratory for Thrombosis and Hemostasis |
| Primary Subject Heading: | Cardiovascular medicine |
| Secondary Subject Heading: | Haematology (incl blood transfusion), Health economics |
| Keywords: | Post thrombotic syndrome, Deep vein thrombosis, Elastic compression stocking therapy, Protocol, RCT |
Study Protocol

The IDEAL DVT study, individualized duration elastic compression therapy against long-term duration of therapy for the prevention of post thrombotic syndrome, a randomized controlled trial

Authors: Arina J ten Cate-Hoek\textsuperscript{1,2}, Annemieke C Bouman\textsuperscript{1,2}, Manuela Joore\textsuperscript{3}, Martin Prins\textsuperscript{3,4}, Hugo ten Cate\textsuperscript{1,2}, for the IDEAL DVT trial investigators.

Corresponding author: Arina J ten Cate-Hoek MD, PhD.
Address: P Debyelaan 25
6229 HX Maastricht
E-mail: arina.tencate@maastrichtuniversity.nl
Telephone number: 0031(0)43-3871243
Fax number: 0031(0)43-3876096

\textsuperscript{1}Laboratory for Thrombosis and Hemostasis, Maastricht University Medical Centre, Maastricht, the Netherlands
\textsuperscript{2}Department of Internal Medicine, Maastricht University Medical Centre, Maastricht, the Netherlands
\textsuperscript{3}Department of Clinical Epidemiology and Medical Technology Assessment (KEMTA), Maastricht University Medical Center, Maastricht, The Netherlands
\textsuperscript{4}Department of Epidemiology, Care and Public Health Research Institutes, Maastricht University, Maastricht, the Netherlands

Key words: Post thrombotic syndrome, deep vein thrombosis, elastic compression stocking therapy, protocol, RCT

Word count: 3345
Abstract

Introduction:

Post thrombotic syndrome (PTS) is a serious complication of deep vein thrombosis (DVT) of the leg that affects 20-50% of patients. Once a patient experiences PTS there is no treatment that effectively reduces the debilitating complaints. Two randomized controlled trials showed that elastic compression stocking (ECS) therapy after DVT for 24 months can reduce the incidence of PTS by 50%. However, it is unclear whether all patients benefit to the same extent from ECS therapy or what the optimal duration of therapy for individual patients should be. ECS therapy is costly, inconvenient, demanding and sometimes even debilitating. Tailoring therapy to individual needs could save substantial costs. The objective of the IDEAL DVT study therefore is to evaluate whether tailoring the duration of ECS therapy on signs and symptoms of the individual patient is a safe and effective method to prevent PTS, compared to standard ECS therapy.

Methods and analysis:

A multicentre, single-blinded, allocation concealed, randomized, non-inferiority trial. A total of 864 consecutive patients with acute objectively documented proximal DVT of the leg are randomized to either standard duration of 24 months or tailored duration of ECS therapy following an initial therapeutic period of 6 months. Signs and symptoms of PTS are recorded at regular clinic visits. Furthermore, quality of life, costs, patient preferences and compliance are measured. The primary outcome is the proportion of patients with PTS at 24 months.

Ethics and dissemination:

Based on current knowledge the standard application of ECS therapy is questioned. The IDEAL DVT study will address the central questions that remain unanswered: What individual patients do benefit from ECS therapy and what is the optimal individual treatment duration? Primary ethics approval was received from the Maastricht University Medical Centre.

Results of the study will be disseminated via peer-reviewed publications and presentations at scientific conferences.

Trial Registration:

NCT01429714 and NTR 2597
Background

After deep vein thrombosis (DVT) 20-50% of patients develop post thrombotic syndrome (PTS). PTS is a chronic condition, characterized by leg complaints such as pain; heaviness; cramps; aching; tingling; and leg signs due to venous insufficiency, evolving in severe cases to venous ulceration of the leg that was affected by the DVT.\textsuperscript{15} PTS is a serious affliction, with substantial impact on quality of life and costs.\textsuperscript{26} Because an effective therapy is lacking, prevention of PTS is of major importance.

Up till now, elastic compression stocking (ECS) therapy after DVT has been the mainstay of PTS prevention. The evidence sustaining the value of ECS therapy following acute DVT is derived from 2 randomized clinical trials, which showed that incidences of PTS were reduced by approximately 50%, with application of ECS therapy for 24 months.\textsuperscript{7,8} However, based on these trials it is still undecided whether all patients benefit to the same extent from ECS therapy or what the optimal duration of ECS therapy for individual patients should be. We have previously assessed the safety of shortened duration of ECS therapy based on individual patient clinical scores in a management study of 125 patients, and we have shown that tailoring the duration of ECS therapy based on the signs and symptoms of the individual patient after an initial treatment period of 6 months, is a safe strategy to prevent PTS. We found that 50% of our patients did not need ECS therapy for as long as 2 years, while the overall incidence of PTS was 21.1% (95% CI 13.5-28.7).\textsuperscript{9} This incidence is comparable to published incidences after 24 months ECS therapy.\textsuperscript{7,8} While this was a prospective management cohort study with an open character and therefore prone to bias, the results of this study need to be confirmed by an adequately powered randomized controlled trial.

There have been other studies that highlighted different aspects of ECS therapy. ECS therapy may not be indicated for all patients as shown by a study that assessed whether ECS therapy initiated one year after the event of DVT would lower the incidence of PTS in patients without complaints but with reflux on duplex testing. No significant benefit of ECS therapy at this late time of onset was found. The incidence of PTS was low in both groups.\textsuperscript{10} One study so far assessed whether prolonged duration of ECS therapy was superior to 6 months of ECS therapy following an event of proximal DVT. In this study no significant difference in the incidence of PTS according to the CEAP classification was observed between the two treatment groups.\textsuperscript{11}
The proposed IDEAL (individually tailored elastic compression against (=versus) longterm therapy) DVT study is a non-inferiority trial and aims to demonstrate that the assessed alternative therapy based on ECS for 6 months followed by individually prolonged ECS therapy based on Villalta scores is non-inferior to the comparator ECS therapy for a standard duration of 24 months. In addition we are interested to study the cost-effectiveness of individual tailoring of the duration of ECS therapy via a cost-utility analysis (CUA) and we want to retrieve information on the patient’s motivations for compliance or non-compliance to therapy. Because compliance to therapy is one of the most important prerequisites for successful ECS therapy, a discrete choice experiment (DCE) will be performed in which patient preferences towards ECS therapy will be assessed, in order to reveal the main obstacles for compliance to therapy. For this purpose, a DCE questionnaire will be send to a subset of the population (300 patients) at approximately 3 months after DVT.

The results of a recently published placebo controlled trial performed by Kahn et al. refuted the routine wearing of ECS after DVT altogether. This study showed no treatment effect at all from ECS therapy in contrast to the large treatment effects found in previous non-placebo controlled trials. The unexpected lack of effectiveness cannot be dismissed solely as a placebo effect. The compliance to ECS therapy, a major determinant of effectiveness, was 55.6% after 24 months. This was less in comparison to the previous trials by Brandjes and Prandoni, where compliance was up to 90%. Although a per protocol analysis of the patients reporting regular stocking use yielded the same outcome, this sub analysis may not be adequately powered to dismiss lack of compliance as an important determinant of non-effectiveness.

Based on current knowledge it is therefore understood that the standard application of ECS therapy is questioned. Especially the benefit to individual patients and the optimal duration of ECS therapy were never studied properly and as a result it is unclear which individual patients require therapy and for how long.

The IDEAL DVT study aims to address these topics. In addition quality of life, patient preference towards ECS therapy, compliance to therapy, as well as cost-effectiveness of ECS therapy will be assessed.

The primary outcome of the IDEAL DVT study will be PTS at 24 months after the event. The secondary outcomes will be: 1. Health Related Quality Of Life, measured by questionnaires (SF-36, EuroQOL-5D, Dutch translated Veines-Qol, 2. Costs, 3. Recurrent thrombosis, according to
criteria as published\textsuperscript{17} and assessed by objective tests, 4. VTE related death during follow-up, assessed by an independent and blinded adjudication committee, 5. Patient preferences, assessed with a Discrete Choice Experiment (DCE)\textsuperscript{18}.

The scope of the problem in the Netherlands

In the Netherlands, each year around 25,000 patients experience a new event of DVT. In order to prevent PTS, common practice is to prescribe custom fitted ECS therapy for 2 years for all patients.\textsuperscript{19} ECS therapy is costly, demanding and sometimes even debilitating. Elderly patients can often not apply the ECS by themselves but need help from family or they even need home care visits (7.5%).\textsuperscript{21}

Total annual costs of ECS therapy roughly amounts to 2.5 million euro for stockings (25,000 patients*100 euro) and 21 million euro for home care (7.5%*25,000 patients*500 visits*20 euro).\textsuperscript{21-23}

Moreover, the majority of patients do not experience any post thrombotic complaints.\textsuperscript{10} In the Netherlands, over 10 million euro each year could be saved if ECS treatment would be individualized.

Methods/Design

A multicentre, single-blinded, allocation concealed, randomized, non-inferiority trial comparing individually tailored duration of ECS therapy (intervention) with standard duration of 24 months ECS therapy (control), for the prevention of PTS. Randomization will guarantee a balanced distribution of patients within each patient group. Stratification will be performed on centre and potentially confounding effects such as age, sex and body mass index. The study will be a multicenter study in order to get a good representation of patients and to achieve sufficient patient numbers. The primary outcome will be PTS at 24 months after the event; the observers will be blinded to the allocated treatment arm. The allocation will be concealed from study personnel involved in assessing the leg symptoms and rating the PTS scores. Both randomization and allocation concealment are used as strategies against bias. The study is not double blinded; no inactive stockings will be used in the comparator group. The use of sham stockings will very likely interfere with the quality of life and will complicate the assessment of perceived differences between groups. Patients will be followed for the entire study duration; incidence of recurrent DVT and VTE related death during the follow-up period will be documented.
Patients and treatment

Consecutive, consenting adults with an acute objectively documented proximal DVT of the leg, adequately treated with anticoagulant treatment and initial compression therapy according to a prespecified protocol are included in the study. Patients are included and randomized in the IDEAL DVT study within 2-6 weeks after DVT.

Exclusion criteria are:

- Previous DVT in the affected leg. Patients with a previous ipsilateral DVT might already have developed PTS after the first DVT.
- Recurrent DVT in the 6 months following inclusion, as it cannot be justified to advise these patients to discontinue ECS therapy 6 months after DVT.
- Pre-existent venous insufficiency (skin signs C3-C6 on CEAP score or requiring ECS therapy). Pre-existent venous insufficiency increases the risk of developing PTS and the majority of patients with venous insufficiency already chronically wear elastic compression stockings. In addition venous insufficiency is closely related to PTS and is therefore difficult to differentiate from PTS.
- Contraindication for ECS therapy such as intermittent claudication or clinical signs of leg ischemia or asymptomatic arterial insufficiency (a pulse deficit or bruit at sites of narrowing at physical examination).
- Active thrombolysis, as thrombolysis reduces the risk of PTS.
- Limited life expectancy (< 6 months), as the follow-up period is 2 years.

Patients are recruited from 12 hospitals in the Netherlands and 2 hospitals in Italy.

There is a prespecified protocol for the management of DVT in all participating centres.

In the initial acute phase after DVT, until the acute oedema has disappeared, one of three strategies will be applied. The leg is either bandaged with short stretch bandages to a compression of 30-40 mmHg, worn day and night and redressed twice per week; or a bandage stocking with 35mmHg compression is prescribed (Mediven Struva 35® , Medi, Breda, the Netherlands), worn day and night; or no initial compression therapy is applied, according to the prespecified strategy of the hospital protocol. After the initial phase, a custom fitted, flat knitted, knee length graduated elastic compression stocking class III (ankle pressure 40 mmHg) is prescribed for all patients. The same brand and type (Mediven 550) of compression stocking is prescribed to all patients in all participating centres.
Compression therapy with the elastic compression stocking is started immediately after the initial phase, so there is no period without compression in between, with exception of the patients who do not receive compression in the initial phase. Patients receive two new stockings every year. Patients are advised to wear the stockings during the daytime.

Procedures

After obtaining written informed consent, randomization is performed centrally at the coordinating study centre, the Maastricht University Medical centre. Patients are included and randomized within 2-6 weeks after diagnosis of DVT. Patients are randomized to one of the two treatment arms. (Figure 1) A web-based randomisation program (TENALEA (Trans European Network for Clinical Trials Services)) is used that executes blocked randomisation with stratification on centre-level and on possible confounding patient characteristics such as age, sex, and Body Mass Index (<26 kg/m² - ≥26 kg/m²). Permuted blocks of randomly varying size are used to maintain balance of numbers in each treatment group and to ensure allocation concealment.

Information on allocation is only accessible to the study personnel at the coordinating study centre in Maastricht, and patients will receive information about their allocation via the coordinating study centre. The allocation will be concealed from study personnel involved in assessing the leg symptoms and rating the PTS scores. Patients are not blinded by the use of sham stockings, because they may influence quality of life, complicating the assessment of perceived differences between groups.

Patients are asked not to reveal their treatment allocation to their own physician, and not to wear the elastic compression stocking at the day of their outpatient clinic visits.

All patients are advised to wear the elastic compression stocking during the first 6 months after DVT. In the control group, all patients are instructed to wear the elastic compression stocking daily for a total duration of 2 years. In the Intervention group, duration of ECS therapy is individually tailored based on Villalta scores. After the first 6 months therefore, there are three scenarios:

1. The Villalta score at both the 3 and 6-month follow-up visit is ≤ 4. Then the patient is advised to discontinue ECS therapy.

2. The Villalta score at the 3-month follow-up visit is ≥ 5 the Villalta score at the 6 month follow-up visit is ≤ 4. The patient is advised to continue the ECS therapy for another 6 months. If the
Villalta score at the 12 month follow-up visit is ≤ 4, the patient is allowed to discontinue the ECS therapy,

(3) The Villalta score is ≥ 5 at both the 3 and 6 months follow-up visit. The patient is advised to continue the ECS therapy for a total duration of 24 months. (Figure 2)

Using this algorithm the duration of ECS therapy is individually tailored in each patient. The decision on the duration of ECS therapy in patients in the intervention group is made centrally at the coordinating study centre, as only there the information on treatment allocation is available. The patient is informed about the decisions on the (dis) continuation of the ECS therapy by the coordinating centre.

When a patient develops symptoms and signs of PTS after discontinuation of ECS therapy, a predefined protocol is followed. If necessary, ECS treatment will be reinstated.

Follow-up

All patients are followed for 24 months after DVT. During these 24 months they visit the outpatient clinic 4 times: at 3, 6, 12, and 24 months after the DVT. At each follow-up visit the signs of PTS will be recorded and scored, by the study nurse or the treating physician, using the objective part of the Villalta clinical scale for PTS. In case of intermittent leg complaints or signs or symptoms suspect of recurrent VTE, patients are instructed to visit their treating physician. Study documentation will be filled out and adverse events will be recorded. The CCMO (central committee human related research) will be notified in case of serious adverse events or death.

The patient is asked to fill out 5 questionnaires: at baseline, and 3, 6, 12 and 24 months later. All questionnaires will be offered as a web based application, each patient will have a unique personal entry code. For patients who are not able or willing to use the electronic questionnaires, a paper version will be available. The first questionnaire is filled out shortly after inclusion (4-6 weeks after DVT) and the subsequent questionnaires are filled out the day before each follow-up visit. All questionnaires contain the subjective part of the Villalta clinical scale for PTS, questions on ECS compliance, a cost questionnaire, and three health related quality of life (HRQOL) questionnaires: the Dutch translated Veines-Qol, the SF-36, and the EuroQOL-5D. In a subset of the study population patient preferences regarding elastic compression therapy will be analysed, by conducting
a discrete choice experiment (DCE). For this purpose, a DCE questionnaire will be sent to a subset of the population at approximately 3 months after DVT.

Compliance to ECS therapy is monitored in two ways. Each questionnaire contains a question on compliance to the advised version of ECS therapy. Because compliance and adherence to therapy is of crucial importance the number of contact moments for the assessment of compliance with ECS therapy will be increased by adding telephone contacts to the regular clinical visits. Study-supporting staff will make computer-assisted random telephone calls during the entire follow-up period to both the patients in the group with the 2-year ECS intervention and to patients in the group with individualized ECS therapy. The telephone calls will be made randomly to create a surprise effect (patients are off guard) and to allow for a distribution of contacts over the entire study period. Per patient 3 extra contact moments will be created. A standard questionnaire will be used to assess compliance and to address reasons for non-compliance. Crossover to another therapy arm will be discussed and discouraged.

Outcomes

The primary outcome is the proportion of patients with PTS at the end of follow-up, 24 months after DVT. PTS is defined as a Villalta score of ≥5 at two consecutive visits that are at least three months apart. The time point of the second Villalta score of ≥5 will be considered the time point of PTS diagnosis. Although the consensus is to diagnose PTS at the 6-month visit or later based on one single Villalta score ≥5, we modelled this study after the preceding management study in which two consecutive Villalta scores ≥5 were needed for the diagnosis PTS. We observed that a proportion of patients (about 12%) had fluctuating Villalta scores beyond 6 months after the acute event of DVT. Assigning the diagnosis based on one single score ≥5 would therefore lead to misclassification and overestimation of the PTS incidence. The Villalta scores at 6 months will be available; therefore also the proportion of patients with PTS according to the consensus scoring method will be presented.

Secondary outcomes include:

1. HRQOL, measured by questionnaires (SF-36, EuroQOL-5D, Dutch translated Veines-Qol), 2. Costs, 3. Recurrent thrombosis, according to criteria as published and assessed by objective
tests, 4. VTE related death during follow-up, assessed by an independent and blinded adjudication committee, 5. Patient preferences, assessed with a Discrete Choice Experiment (DCE)\textsuperscript{18}.

Since both effects on costs and generic health-related quality of life are to be expected, the method of economic evaluation is a cost-utility analysis (CUA). The analysis will be from a societal perspective. The primary effect parameter is generic health-related quality of life, measured in quality adjusted life years (QALYs). We will perform two analyses: a trial based CUA with a time horizon of two years (identical to the duration of follow up in the clinical study), and a model-based CUA with a lifelong time horizon. For the latter, we will follow the guidelines of good modelling.\textsuperscript{26} We will be able to adapt a Markov model for diagnostic strategies in DVT, which was developed by our group earlier.\textsuperscript{27} Costs in the economic analyses include direct health care costs (medical costs for prevention, diagnostics, therapy, rehabilitation and care), direct non-health care costs (travel costs) and indirect costs (productivity loss). Utilities will be calculated from the responses on the EQ-5D and SF36 classification systems using the available multi-attribute utility functions.\textsuperscript{28, 29} Incremental cost-utility ratios will be calculated, and non-parametric bootstrap analyses will be used to quantify the uncertainty surrounding the cost-utility ratio of the trial-based analysis. Sensitivity analyses and subgroup analyses will be performed, to assess the impact of variation in parameters and heterogeneity of the patient population.

ECS therapy has several disadvantages for the patients (stockings are uncomfortable, ugly, and difficult to put on and off), while duration and effectiveness are uncertain. A DCE will be conducted to assess the patient preferences regarding ECS therapy, providing insight in the trade-offs patients make between characteristics of the therapy when deciding to wear the stocking or not. Duration of ECS therapy is one of the characteristics. Data will be analysed using multinomial logit models and mixed logit models (Nlogit, Econometric Software).

Sample size calculation

The proposed study is a non-inferiority trial and aims to demonstrate that the assessed alternative therapy based on ECS for 6 months followed by individually prolonged ECS therapy based on Villalta scores is not worse than the comparator ECS therapy for a standard duration of 24 months, by more than the prespecified amount of 7.5% (the non-inferiority margin). The published incidence of PTS following two years of ECS lies between 20 and 30%. (Prandoni 24.5% (95% CI 15.6- 33.4), Brandjes
20% (95% CI 12.4-29.2)\(^8\) As it is statistically impossible to demonstrate equivalence (prove the H0 of no difference), Blackwelder proposed a one sided significance test to reject the null hypothesis by a clinically acceptable amount.\(^30\) If we allow a difference of 7.5% in the outcome PTS between the group with ECS for 2 years and the group with the alternative therapy, 70% of the effect will be preserved. This proportion of loss in efficacy is customarily accepted in controlled randomized clinical trials. At a one sided significance level of 0.05 and a power of 80%, with a ratio of 1, a total of 847 patients is needed to provide sufficient patients for an adequately powered trial.\(^30\)\(^31\) Loss to follow-up of patients is expected to be less than 2% since the intervention does not have an invasive nature. Therefore, a total of 864 patients are needed (432 patients per treatment arm).

**Statistical analysis**

Descriptive statistics of the total population and of the two treatment groups separately will be computed to provide baseline characteristics of the patients in both treatment arms.

For the primary outcome PTS, univariate analysis of all proportions will be performed with logistic regression (chi-squared) analysis. Kaplan-Meier method will be used to calculate the cumulative incidences of PTS, adjusted for centre, to compare incidence rates between the two treatment arms. Patients who die or are lost to follow-up will be censored at their last visit. ANOVA will be applied to assess changes over time, by comparing different outcome measures at the different time points of follow-up.

Hazard ratio’s and 95% confidence intervals for both treatment arms will be calculated using Cox regression models. Hazard ratio’s will be stratified for centre and adjusted for age, sex, BMI, clinical presentation of DVT, and extent of the index deep vein thrombosis.

**Interim analysis (safety)**

A prespecified safety analysis will be performed when 50% of the subjects have completed the 2-year follow-up. The analysis will be performed by the coordinating centre, and supervised by the principal investigators. The safety analysis will be performed to assess significant enhanced risk of PTS or excess morbidity/mortality in the intervention arm of the study population. Fisher’s exact test will be performed to compare incidence of PTS at a significance level of 0.05 (two sided). The study can be stopped in case of significant excess morbidity/mortality in the alternative treatment arm.
Ethical considerations

The medical ethical committees of all participating hospitals in the Netherlands and Italy approved this study. All patients are extensively informed about the study, and written informed consent is obtained from all participating patients.

Discussion

Based on current knowledge the standard application of ECS therapy after DVT is questioned. The definitive answer on the usefulness of ECS therapy for the prevention of PTS is not yet provided. Therefore the need for the IDEAL DVT study remains unchanged. The benefit to individual patients and the optimal duration of ECS therapy were never studied properly and as a result it is unclear which individual patients require therapy and for how long. The IDEAL DVT study will address the central questions that remain unanswered: What individual patients do benefit from ECS therapy and what is the optimal individual treatment duration? The study that we plan to perform is not only innovative; for the first time individual tailoring of duration of ECS therapy will be investigated, but also it will provide unique additional knowledge on the safety, effectiveness and cost-effectiveness of this approach. The IDEAL DVT study started including patients March 2011 and is currently on-going. Recruitment is expected to be terminated within 1 year. As the follow-up is 2 years, the results are expected within 3 years.
Trial status

The IDEAL DVT study is currently in progress. Patient recruitment started March 15th 2011 and is still on-going.

Acknowledgments

The IDEAL DVT trial is funded by a grant from Zon-mW the Netherlands. (171102007)

We especially acknowledge the participating investigators.

Competing interests

The authors declare that they have no competing interests.

Funding

This work was supported by Zon-mW the Netherlands grant number 171102007.

Contributions

AtC wrote the protocol and designed the study, the other authors MJ (cost-effectiveness research), MP (study design/statistics) and HtC (study design) contributed. AB is a PhD student and coordinator of the trial.

Ethics approval

Maastricht University Medical Centre.
The IDEAL DVT investigators:

The Netherlands:

*Maastricht, Maastricht University Medical Centre*

Arina J ten Cate-Hoek MD, PhD (principal investigator)

Hugo ten Cate MD, PhD

Manuela Joore PhD

Annemieke C Bouman MD

Groningen, University Medical Centre Groningen

Karina Meijer MD PhD

*Amsterdam, Academic Medical Centre*

Saskia Middendorp MD, PhD

Michiel Coppens MD, PhD

Mandy N Lauw MD

Y Whitney Cheung MD

*Heerlen, Atrium Medical Centre*

Guy JM Mostard MD

Asiong Jie MD, PhD

*Hoorn, Westfriesgasthuis*

Simone M van den Heiligenberg MD

*Eindhoven, Maxima Medical Centre*

Lidwine W Tick MD, PhD

Marten R Nijziel MD, PhD

*Almere, Flevohospital*

Marije ten Wolde MD, PhD

Yuk W Cheung MD

*Amsterdam, OLVG*

Sanne van Wissen MD, PhD

Wim E Terpstra MD, PhD

*Roermond, Laurentius hospital*

 Marlène HW van de Poel MD
Amsterdam, Slotervaart hospital
Hans-Martin Otten MD, PhD
Amsterdam, VU Medical Centre
Erik H Serné MD, PhD
Nijmegen, Radboud University Nijmegen Medical Centre
Edith H Klappe MD, PhD
Mirian CH Janssen MD, PhD
Tjerk de Nijs MD
Italy:
Padua, Aziende Ospedaliera di Padova
Paolo Prandoni MD, PhD
Treviso, Aziende ULSS
Sabina Villalta MD
References


Figure 1. Study flow diagram
90x79mm (300 x 300 DPI)
Figure 2. Algorithm individually tailored ECS therapy

90x77mm (300 x 300 DPI)
The IDEAL DVT study, individualized duration elastic compression therapy against long-term duration of therapy for the prevention of post thrombotic syndrome, a randomized controlled trial

<table>
<thead>
<tr>
<th>Journal:</th>
<th>BMJ Open</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manuscript ID:</td>
<td>bmjopen-2014-005265.R2</td>
</tr>
<tr>
<td>Article Type:</td>
<td>Protocol</td>
</tr>
<tr>
<td>Date Submitted by the Author:</td>
<td>10-Jul-2014</td>
</tr>
</tbody>
</table>
| Complete List of Authors: | ten Cate, Arina; Maastricht University Medical Centre, Laboratory for Thrombosis and Hemostasis; Maastricht University Medical Center, Internal Medicine
Bouman, Annemieke; Maastricht University Medical Centre, Laboratory for Thrombosis and Hemostasis; Maastricht University Medical Center, Internal Medicine
Joore, Manuela; Maastricht University Medical Centre, Department of Clinical Epidemiology and Medical Technology Assessment (KEMTA)
Prins, Martin; Maastricht University Medical Center, Epidemiology; Maastricht University Medical Centre, Department of Clinical Epidemiology and Medical Technology Assessment (KEMTA)
ten Cate, Hugo; Maastricht University Medical Center, Internal Medicine; Maastricht University Medical Centre, Laboratory for Thrombosis and Hemostasis |
| <b>Primary Subject Heading</b>: | Cardiovascular medicine |
| Secondary Subject Heading: | Haematology (incl blood transfusion), Health economics |
| Keywords: | Post thrombotic syndrome, Deep vein thrombosis, Elastic compression stocking therapy, Protocol, RCT |
Study Protocol

The IDEAL DVT study, individualized duration elastic compression therapy against long-term duration of therapy for the prevention of post thrombotic syndrome, a randomized controlled trial

Authors: Arina J ten Cate-Hoek1,2, Annemieke C Bouman1,2, Manuela Joore3, Martin Prins3,4, Hugo ten Cate1,2, for the IDEAL DVT trial investigators.

Corresponding author: Arina J ten Cate-Hoek MD, PhD.
Address: P Debyelaan 25
6229 HX Maastricht
E-mail: arina.tencate@maastrichtuniversity.nl
Telephone number: 0031(0)43-3871243
Fax number: 0031(0)43-3876096

1Laboratory for Thrombosis and Hemostasis, Maastricht University Medical Centre, Maastricht, the Netherlands
2Department of Internal Medicine, Maastricht University Medical Centre, Maastricht, the Netherlands
3Department of Clinical Epidemiology and Medical Technology Assessment (KEMTA), Maastricht University Medical Center, Maastricht, The Netherlands
4Department of Epidemiology, Care and Public Health Research Institutes, Maastricht University, Maastricht, the Netherlands

Key words: Post thrombotic syndrome, deep vein thrombosis, elastic compression stocking therapy, protocol, RCT

Word count: 3345
Abstract

Introduction:

Post thrombotic syndrome (PTS) is a serious complication of deep vein thrombosis (DVT) of the leg that affects 20-50% of patients. Once a patient experiences PTS there is no treatment that effectively reduces the debilitating complaints. Two randomized controlled trials showed that elastic compression stocking (ECS) therapy after DVT for 24 months can reduce the incidence of PTS by 50%. However, it is unclear whether all patients benefit to the same extent from ECS therapy or what the optimal duration of therapy for individual patients should be. ECS therapy is costly, inconvenient, demanding and sometimes even debilitating. Tailoring therapy to individual needs could save substantial costs. The objective of the IDEAL DVT study therefore is to evaluate whether tailoring the duration of ECS therapy on signs and symptoms of the individual patient is a safe and effective method to prevent PTS, compared to standard ECS therapy.

Methods and analysis:

A multicentre, single-blinded, allocation concealed, randomized, non-inferiority trial. A total of 864 consecutive patients with acute objectively documented proximal DVT of the leg are randomized to either standard duration of 24 months or tailored duration of ECS therapy following an initial therapeutic period of 6 months. Signs and symptoms of PTS are recorded at regular clinic visits. Furthermore, quality of life, costs, patient preferences and compliance are measured. The primary outcome is the proportion of patients with PTS at 24 months.

Ethics and dissemination:

Based on current knowledge the standard application of ECS therapy is questioned. The IDEAL DVT study will address the central questions that remain unanswered: What individual patients do benefit from ECS therapy and what is the optimal individual treatment duration? Primary ethics approval was received from the Maastricht University Medical Centre.

Results of the study will be disseminated via peer-reviewed publications and presentations at scientific conferences.

Trial Registration number:

NCT01429714 and NTR 2597
Background

After deep vein thrombosis (DVT) 20-50% of patients develop post thrombotic syndrome (PTS). PTS is a chronic condition, characterized by leg complaints such as pain; heaviness; cramps; aching; tingling; and leg signs due to venous insufficiency, evolving in severe cases to venous ulceration of the leg that was affected by the DVT.\(^1\)PTS is a serious affliction, with substantial impact on quality of life and costs.\(^2\) Because an effective therapy is lacking, prevention of PTS is of major importance.

Up till now, elastic compression stocking (ECS) therapy after DVT has been the mainstay of PTS prevention. The evidence sustaining the value of ECS therapy following acute DVT is derived from 2 randomized clinical trials, which showed that incidences of PTS were reduced by approximately 50%, with application of ECS therapy for 24 months.\(^3,4\) However, based on these trials it is still undecided whether all patients benefit to the same extent from ECS therapy or what the optimal duration of ECS therapy for individual patients should be. We have previously assessed the safety of shortened duration of ECS therapy based on individual patient clinical scores in a management study of 125 patients, and we have shown that tailoring the duration of ECS therapy based on the signs and symptoms of the individual patient after an initial treatment period of 6 months, is a safe strategy to prevent PTS. We found that 50% of our patients did not need ECS therapy for as long as 2 years, while the overall incidence of PTS was 21.1% (95% CI 13.5-28.7).\(^5\) This incidence is comparable to published incidences after 24 months ECS therapy.\(^6,7\) While this was a prospective management cohort study with an open character and therefore prone to bias, the results of this study need to be confirmed by an adequately powered randomized controlled trial.

There have been other studies that highlighted different aspects of ECS therapy. ECS therapy may not be indicated for all patients as shown by a study that assessed whether ECS therapy initiated one year after the event of DVT would lower the incidence of PTS in patients without complaints but with reflux on duplex testing. No significant benefit of ECS therapy at this late time of onset was found. The incidence of PTS was low in both groups.\(^8\) One study so far assessed whether prolonged duration of ECS therapy was superior to 6 months of ECS therapy following an event of proximal DVT. In this study no significant difference in the incidence of PTS according to the CEAP classification was observed between the two treatment groups.\(^9\)
The proposed IDEAL (individually tailored elastic compression against (=versus) longterm therapy) DVT study is a non-inferiority trial and aims to demonstrate that the assessed alternative therapy based on ECS for 6 months followed by individually prolonged ECS therapy based on Villalta scores is non-inferior to the comparator ECS therapy for a standard duration of 24 months. In addition we are interested to study the cost-effectiveness of individual tailoring of the duration of ECS therapy via a cost-utility analysis (CUA) and we want to retrieve information on the patient's motivations for compliance or non-compliance to therapy. Because compliance to therapy is one of the most important prerequisites for successful ECS therapy, a discrete choice experiment (DCE) will be performed in which patient preferences towards ECS therapy will be assessed, in order to reveal the main obstacles for compliance to therapy. For this purpose, a DCE questionnaire will be send to a subset of the population (300 patients) at approximately 3 months after DVT. 

The results of a recently published placebo controlled trial performed by Kahn et al. refuted the routine wearing of ECS after DVT altogether. This study showed no treatment effect at all from ECS therapy in contrast to the large treatment effects found in previous non-placebo controlled trials. The unexpected lack of effectiveness cannot be dismissed solely as a placebo effect. The compliance to ECS therapy, a major determinant of effectiveness, was 55.6% after 24 months. This was less in comparison to the previous trials by Brandjes and Prandoni, where compliance was up to 90%. Although a per protocol analysis of the patients reporting regular stocking use yielded the same outcome, this sub analysis may not be adequately powered to dismiss lack of compliance as an important determinant of non-effectiveness.

Based on current knowledge it is therefore understood that the standard application of ECS therapy is questioned. Especially the benefit to individual patients and the optimal duration of ECS therapy were never studied properly and as a result it is unclear which individual patients require therapy and for how long. 

The IDEAL DVT study aims to address these topics. In addition quality of life, patient preference towards ECS therapy, compliance to therapy, as well as cost-effectiveness of ECS therapy will be assessed.

The primary outcome of the IDEAL DVT study will be PTS at 24 months after the event. The secondary outcomes will be: 1. Health Related Quality Of Life, measured by questionnaires (SF-36, EuroQOL-5D, Dutch translated Veines-Qol, 2. Costs, 3. Recurrent thrombosis, according to
criteria as published^{17} and assessed by objective tests. 4. VTE related death during follow-up, assessed by an independent and blinded adjudication committee. 5. Patient preferences, assessed with a Discrete Choice Experiment (DCE)^{18}.

The scope of the problem in the Netherlands

In the Netherlands, each year around 25,000 patients experience a new event of DVT. In order to prevent PTS, common practice is to prescribe custom fitted ECS therapy for 2 years for all patients.^{19}^{20} ECS therapy is costly, demanding and sometimes even debilitating. Elderly patients can often not apply the ECS by themselves but need help from family or they even need home care visits (7.5%).^{21}{23}

Total annual costs of ECS therapy roughly amounts to 2.5 million euro for stockings (25,000 patients*100 euro) and 21 million euro for home care (7.5%*25,000 patients*500 visits*20 euro).^{21}{23}

Moreover, the majority of patients do not experience any post thrombotic complaints.^{10} In the Netherlands, over 10 million euro each year could be saved if ECS treatment would be individualized.

Methods/Design

A multicentre, single-blinded, allocation concealed, randomized, non-inferiority trial comparing individually tailored duration of ECS therapy (intervention) with standard duration of 24 months ECS therapy (control), for the prevention of PTS. Randomization will guarantee a balanced distribution of patients within each patient group. Stratification will be performed on centre and potentially confounding effects such as age, sex and body mass index. The study will be a multicenter study in order to get a good representation of patients and to achieve sufficient patient numbers. The primary outcome will be PTS at 24 months after the event; the observers will be blinded to the allocated treatment arm. The allocation will be concealed from study personnel involved in assessing the leg symptoms and rating the PTS scores. Both randomization and allocation concealment are used as strategies against bias. The study is not double blinded; no inactive stockings will be used in the comparator group. The use of sham stockings will very likely interfere with the quality of life and will complicate the assessment of perceived differences between groups. Patients will be followed for the entire study duration; incidence of recurrent DVT and VTE related death during the follow-up period will be documented.
Patients and treatment

Consecutive, consenting adults with an acute objectively documented proximal DVT of the leg, adequately treated with anticoagulant treatment and initial compression therapy according to a prespecified protocol are included in the study. Patients are included and randomized in the IDEAL DVT study within 2-6 weeks after DVT.

Exclusion criteria are:

- Previous DVT in the affected leg. Patients with a previous ipsilateral DVT might already have developed PTS after the first DVT.
- Recurrent DVT in the 6 months following inclusion, as it cannot be justified to advise these patients to discontinue ECS therapy 6 months after DVT.
- Pre-existent venous insufficiency (skin signs C3-C6 on CEAP score or requiring ECS therapy). Pre-existent venous insufficiency increases the risk of developing PTS and the majority of patients with venous insufficiency already chronically wear elastic compression stockings. In addition venous insufficiency is closely related to PTS and is therefore difficult to differentiate from PTS.
- Contraindication for ECS therapy such as intermittent claudication or clinical signs of leg ischemia or asymptomatic arterial insufficiency (a pulse deficit or bruit at sites of narrowing at physical examination).
- Active thrombolysis, as thrombolysis reduces the risk of PTS.
- Limited life expectancy (< 6 months), as the follow-up period is 2 years.

Patients are recruited from 12 hospitals in the Netherlands and 2 hospitals in Italy.

There is a prespecified protocol for the management of DVT in all participating centres.

In the initial acute phase after DVT, until the acute oedema has disappeared, one of three strategies will be applied. The leg is either bandaged with short stretch bandages to a compression of 30-40 mmHg, worn day and night and redressed twice per week; or a bandage stocking with 35mmHg compression is prescribed (Mediven Struva 35 ©, Medi, Breda, the Netherlands), worn day and night; or no initial compression therapy is applied, according to the prespecified strategy of the hospital protocol. After the initial phase, a custom fitted, flat knitted, knee length graduated elastic compression stocking class III (ankle pressure 40 mmHg) is prescribed for all patients. The same brand and type (Mediven 550) of compression stocking is prescribed to all patients in all participating centres.
Compression therapy with the elastic compression stocking is started immediately after the initial phase, so there is no period without compression in between, with exception of the patients who do not receive compression in the initial phase. Patients receive two new stockings every year. Patients are advised to wear the stockings during the daytime.

Procedures
After obtaining written informed consent, randomization is performed centrally at the coordinating study centre, the Maastricht University Medical centre. Patients are included and randomized within 2-6 weeks after diagnosis of DVT. Patients are randomized to one of the two treatment arms. (Figure 1) A web-based randomisation program (TENALEA (Trans European Network for Clinical Trials Services)) is used that executes blocked randomisation with stratification on centre-level and on possible confounding patient characteristics such as age, sex, and Body Mass Index (<26 kg/m$^2$ - ≥26 kg/m$^2$). Permutated blocks of randomly varying size are used to maintain balance of numbers in each treatment group and to ensure allocation concealment.

Information on allocation is only accessible to the study personnel at the coordinating study centre in Maastricht, and patients will receive information about their allocation via the coordinating study centre. The allocation will be concealed from study personnel involved in assessing the leg symptoms and rating the PTS scores. Patients are not blinded by the use of sham stockings, because they may influence quality of life, complicating the assessment of perceived differences between groups. Patients are asked not to reveal their treatment allocation to their own physician, and not to wear the elastic compression stocking at the day of their outpatient clinic visits.

All patients are advised to wear the elastic compression stocking during the first 6 months after DVT. In the control group, all patients are instructed to wear the elastic compression stocking daily for a total duration of 2 years. In the Intervention group, duration of ECS therapy is individually tailored based on Villalta scores. After the first 6 months therefore, there are three scenarios:

1. The Villalta score at both the 3 and 6-month follow-up visit is ≤ 4. Then the patient is advised to discontinue ECS therapy.

2. The Villalta score at the 3-month follow-up visit is ≥ 5 the Villalta score at the 6 month follow-up visit is ≤ 4. The patient is advised to continue the ECS therapy for another 6 months. If the...
Villalta score at the 12 month follow-up visit is ≤ 4, the patient is allowed to discontinue the ECS therapy,

(3) The Villalta score is ≥ 5 at both the 3 and 6 months follow-up visit. The patient is advised to continue the ECS therapy for a total duration of 24 months. (Figure 2)

Using this algorithm the duration of ECS therapy is individually tailored in each patient. The decision on the duration of ECS therapy in patients in the intervention group is made centrally at the coordinating study centre, as only there the information on treatment allocation is available. The patient is informed about the decisions on the (dis) continuation of the ECS therapy by the coordinating centre.

When a patient develops symptoms and signs of PTS after discontinuation of ECS therapy, a predefined protocol is followed. If necessary, ECS treatment will be reinstated.

Follow-up

All patients are followed for 24 months after DVT. During these 24 months they visit the outpatient clinic 4 times: at 3, 6, 12, and 24 months after the DVT. At each follow-up visit the signs of PTS will be recorded and scored, by the study nurse or the treating physician, using the objective part of the Villalta clinical scale for PTS. In case of intermittent leg complaints or signs or symptoms suspect of recurrent VTE, patients are instructed to visit their treating physician. Study documentation will be filled out and adverse events will be recorded. The CCMO (central committee human related research) will be notified in case of serious adverse events or death.

The patient is asked to fill out 5 questionnaires: at baseline, and 3, 6, 12 and 24 months later. All questionnaires will be offered as a web based application, each patient will have a unique personal entry code. For patients who are not able or willing to use the electronic questionnaires, a paper version will be available. The first questionnaire is filled out shortly after inclusion (4-6 weeks after DVT) and the subsequent questionnaires are filled out the day before each follow-up visit. All questionnaires contain the subjective part of the Villalta clinical scale for PTS, questions on ECS compliance, a cost questionnaire, and three health related quality of life (HRQOL) questionnaires: the Dutch translated Veines-Qol, the SF-36, and the EuroQOL-5D. In a subset of the study population patient preferences regarding elastic compression therapy will be analysed, by conducting
a discrete choice experiment (DCE). For this purpose, a DCE questionnaire will be send to a subset of the population at approximately 3 months after DVT.

Compliance to ECS therapy is monitored in two ways. Each questionnaire contains a question on compliance to the advised version of ECS therapy. Because compliance and adherence to therapy is of crucial importance the number of contact moments for the assessment of compliance with ECS therapy will be increased by adding telephone contacts to the regular clinical visits. Study-supporting staff will make computer-assisted random telephone calls during the entire follow-up period to both the patients in the group with the 2-year ECS intervention and to patients in the group with individualized ECS therapy. The telephone calls will be made randomly to create a surprise effect (patients are off guard) and to allow for a distribution of contacts over the entire study period. Per patient 3 extra contact moments will be created. A standard questionnaire will be used to assess compliance and to address reasons for non-compliance. Crossover to another therapy arm will be discussed and discouraged.

Outcomes
The primary outcome is the proportion of patients with PTS at the end of follow-up, 24 months after DVT. PTS is defined as a Villalta score of ≥5 at two consecutive visits that are at least three months apart. The time point of the second Villalta score of ≥5 will be considered the time point of PTS diagnosis. Although the consensus is to diagnose PTS at the 6-month visit or later based on one single Villalta score ≥5, we modelled this study after the preceding management study in which two consecutive Villalta scores ≥5 were needed for the diagnosis PTS. We observed that a proportion of patients (about 12%) had fluctuating Villalta scores beyond 6 months after the acute event of DVT. Assigning the diagnosis based on one single score ≥5 would therefore lead to misclassification and overestimation of the PTS incidence. The Villalta scores at 6 months will be available; therefore also the proportion of patients with PTS according to the consensus scoring method will be presented.

Secondary outcomes include:
1. HRQOL, measured by questionnaires (SF-36, EuroQOL-5D, Dutch translated Veines-Qol), 2. Costs, 3. Recurrent thrombosis, according to criteria as published and assessed by objective
tests, 4. VTE related death during follow-up, assessed by an independent and blinded adjudication committee, 5. Patient preferences, assessed with a Discrete Choice Experiment (DCE)\textsuperscript{18}.

Since both effects on costs and generic health-related quality of life are to be expected, the method of economic evaluation is a cost-utility analysis (CUA). The analysis will be from a societal perspective. The primary effect parameter is generic health-related quality of life, measured in quality adjusted life years (QALYs). We will perform two analyses: a trial based CUA with a time horizon of two years (identical to the duration of follow up in the clinical study), and a model-based CUA with a lifelong time horizon. For the latter, we will follow the guidelines of good modelling.\textsuperscript{26} We will be able to adapt a Markov model for diagnostic strategies in DVT, which was developed by our group earlier.\textsuperscript{27} Costs in the economic analyses include direct health care costs (medical costs for prevention, diagnostics, therapy, rehabilitation and care), direct non-health care costs (travel costs) and indirect costs (productivity loss). Utilities will be calculated from the responses on the EQ-5D and SF36 classification systems using the available multi-attribute utility functions.\textsuperscript{28, 29} Incremental cost-utility ratios will be calculated, and non-parametric bootstrap analyses will be used to quantify the uncertainty surrounding the cost-utility ratio of the trial-based analysis. Sensitivity analyses and subgroup analyses will be performed, to assess the impact of variation in parameters and heterogeneity of the patient population.

ECS therapy has several disadvantages for the patients (stockings are uncomfortable, ugly, and difficult to put on and off), while duration and effectiveness are uncertain. A DCE will be conducted to assess the patient preferences regarding ECS therapy, providing insight in the trade-offs patients make between characteristics of the therapy when deciding to wear the stocking or not. Duration of ECS therapy is one of the characteristics. Data will be analysed using multinomial logit models and mixed logit models.\textsuperscript{(Nlogit, Econometric Software)}

Sample size calculation

The proposed study is a non-inferiority trial and aims to demonstrate that the assessed alternative therapy based on ECS for 6 months followed by individually prolonged ECS therapy based on Villalta scores is not worse than the comparator ECS therapy for a standard duration of 24 months, by more than the prespecified amount of 7.5% (the non-inferiority margin). The published incidence of PTS following two years of ECS lies between 20 and 30%. (Prandoni 24.5% (95% CI 15.6-33.4), Brandjes...
20% (95% CI 12.4-29.2)\textsuperscript{8} As it is statistically impossible to demonstrate equivalence (prove the H0 of no difference), Blackwelder proposed a one sided significance test to reject the null hypothesis by a clinically acceptable amount.\textsuperscript{30} If we allow a difference of 7.5% in the outcome PTS between the group with ECS for 2 years and the group with the alternative therapy, 70% of the effect will be preserved. This proportion of loss in efficacy is customarily accepted in controlled randomized clinical trials. At a one sided significance level of 0.05 and a power of 80%, with a ratio of 1, a total of 847 patients is needed to provide sufficient patients for an adequately powered trial.\textsuperscript{30, 31} Loss to follow-up of patients is expected to be less than 2% since the intervention does not have an invasive nature. Therefore, a total of 864 patients are needed (432 patients per treatment arm).

Statistical analysis

Descriptive statistics of the total population and of the two treatment groups separately will be computed to provide baseline characteristics of the patients in both treatment arms.

For the primary outcome PTS, univariate analysis of all proportions will be performed with logistic regression (chi-squared) analysis. Kaplan-Meier method will be used to calculate the cumulative incidences of PTS, adjusted for centre, to compare incidence rates between the two treatment arms. Patients who die or are lost to follow-up will be censored at their last visit. ANOVA will be applied to assess changes over time, by comparing different outcome measures at the different time points of follow-up.

Hazard ratio’s and 95% confidence intervals for both treatment arms will be calculated using Cox regression models. Hazard ratio’s will be stratified for centre and adjusted for age, sex, BMI, clinical presentation of DVT, and extent of the index deep vein thrombosis.

Interim analysis (safety)

A prespecified safety analysis will be performed when 50% of the subjects have completed the 2-year follow-up. The analysis will be performed by the coordinating centre, and supervised by the principal investigators. The safety analysis will be performed to assess significant enhanced risk of PTS or excess morbidity/mortality in the intervention arm of the study population. Fisher’s exact test will be performed to compare incidence of PTS at a significance level of 0.05 (two sided). The study can be stopped in case of significant excess morbidity/mortality in the alternative treatment arm.
Ethical considerations

The medical ethical committees of all participating hospitals in the Netherlands and Italy approved this study. All patients are extensively informed about the study, and written informed consent is obtained from all participating patients.

The IDEAL DVT study started including patients at 11\textsuperscript{th} of March 2011 and inclusion is currently ongoing. Recruitment is expected to be terminated in January 2015. As the follow-up is 2 years, the results are expected within 3 years, in January 2017.

Discussion

Based on current knowledge the standard application of ECS therapy after DVT is questioned. The definitive answer on the usefulness of ECS therapy for the prevention of PTS is not yet provided. Therefore the need for the IDEAL DVT study remains unchanged. The benefit to individual patients and the optimal duration of ECS therapy were never studied properly and as a result it is unclear which individual patients require therapy and for how long. The IDEAL DVT study will address the central questions that remain unanswered: What individual patients do benefit from ECS therapy and what is the optimal individual treatment duration? The study that we plan to perform is not only innovative; for the first time individual tailoring of duration of ECS therapy will be investigated, but also it will provide unique additional knowledge on the safety, effectiveness and cost-effectiveness of this approach.
Trial status
The IDEAL DVT study is currently in progress. Patient recruitment started March 15th 2011 and is still on-going.

Acknowledgments
The IDEAL DVT trial is funded by a grant from Zon-mW the Netherlands. (171102007)
We especially acknowledge the participating investigators.

Competing interests
The authors declare that they have no competing interests.

Funding
This work was supported by Zon-mW the Netherlands grant number 171102007.

Contributions
AtC wrote the protocol and designed the study, the other authors MJ (cost-effectiveness research), MP (study design/statistics) and HtC (study design) contributed. AB is a PhD student and coordinator of the trial.

Ethics approval
Maastricht University Medical Centre.
The IDEAL DVT investigators:

The Netherlands:

*Maarstricht, Maastricht University Medical Centre*

Arina J ten Cate-Hoek MD, PhD (principal investigator)

Hugo ten Cate MD, PhD

Manuela Joore PhD

Annemieke C Bouman MD

*Groningen, University Medical Centre Groningen*

Karina Meijer MD PhD

*Amsterdam, Academic Medical Centre*

Saskia Middeldorp MD, PhD

Michiel Coppens MD, PhD

Mandy N Lauw MD

Y Whitney Cheung MD

*Heerlen, Atrium Medical Centre*

Guy JM Mostard MD

Asiong Jie MD PhD

*Hoorn, Westfriesgasthuis*

Simone M van den Heiligenberg MD

*Eindhoven, Maxima Medical Centre*

Lidwine W Tick MD, PhD

Marten R Nijziel MD, PhD

*Almere, Flevohospital*

Marije ten Wolde MD, PhD

Y Whitney Cheung MD

*Amsterdam, OLVG*

Sanne van Wissen MD, PhD

Wim E Terpstra MD, PhD

*Roermond, Laurentius hospital*

Marlène HW van de Poel MD
Amsterdam, Slotervaart hospital
Hans-Martin Otten MD, PhD

Amsterdam, VU Medical Centre
Erik H Serné MD, PhD

Nijmegen, Radboud University Nijmegen Medical Centre
Edith H Klappe MD, PhD
Mirian CH Janssen MD, PhD
Tjerk de Nijs MD

Italy:
Padua, Aziende Ospedaliera di Padova
Paolo Prandoni MD, PhD
Treviso, Aziende ULSS
Sabina Villalta MD

FIGURE LEGENDS

Figure 1. Study flow diagram

Figure 2. Algorithm individually tailored ECS therapy
References


Study Protocol

The IDEAL DVT study, individualized duration elastic compression therapy against long-term duration of therapy for the prevention of post thrombotic syndrome, a randomized controlled trial

Authors: Arina J ten Cate-Hoek\textsuperscript{1,2}, Annemieke C Bouman\textsuperscript{1,2}, Manuela Joore\textsuperscript{3}, Martin Prins\textsuperscript{3,4}, Hugo ten Cate\textsuperscript{1,2}, for the IDEAL DVT trial investigators.

Corresponding author: Arina J ten Cate-Hoek MD, PhD.
Address: P Debyelaan 25
6229 HX Maastricht

E-mail: arina.tencate@maastrichtuniversity.nl
Telephone number: 0031(0)43-3871243
Fax number: 0031(0)43-3876096

\textsuperscript{1}Laboratory for Thrombosis and Hemostasis, Maastricht University Medical Centre, Maastricht, the Netherlands
\textsuperscript{2}Department of Internal Medicine, Maastricht University Medical Centre, Maastricht, the Netherlands
\textsuperscript{3}Department of Clinical Epidemiology and Medical Technology Assessment (KEMTA), Maastricht University Medical Center, Maastricht, The Netherlands
\textsuperscript{4}Department of Epidemiology, Care and Public Health Research Institutes, Maastricht University, Maastricht, the Netherlands

Key words: Post thrombotic syndrome, deep vein thrombosis, elastic compression stocking therapy, protocol, RCT

Word count: 3345
Abstract

Introduction:
Post thrombotic syndrome (PTS) is a serious complication of deep vein thrombosis (DVT) of the leg that affects 20-50% of patients. Once a patient experiences PTS there is no treatment that effectively reduces the debilitating complaints. Two randomized controlled trials showed that elastic compression stocking (ECS) therapy after DVT for 24 months can reduce the incidence of PTS by 50%. However, it is unclear whether all patients benefit to the same extent from ECS therapy or what the optimal duration of therapy for individual patients should be. ECS therapy is costly, inconvenient, demanding and sometimes even debilitating. Tailoring therapy to individual needs could save substantial costs. The objective of the IDEAL DVT study therefore is to evaluate whether tailoring the duration of ECS therapy on signs and symptoms of the individual patient is a safe and effective method to prevent PTS, compared to standard ECS therapy.

Methods and analysis:
A multicentre, single-blinded, allocation concealed, randomized, non-inferiority trial. A total of 864 consecutive patients with acute objectively documented proximal DVT of the leg are randomized to either standard duration of 24 months or tailored duration of ECS therapy following an initial therapeutic period of 6 months. Signs and symptoms of PTS are recorded at regular clinic visits. Furthermore, quality of life, costs, patient preferences and compliance are measured. The primary outcome is the proportion of patients with PTS at 24 months.

Ethics and dissemination:
Based on current knowledge the standard application of ECS therapy is questioned. The IDEAL DVT study will address the central questions that remain unanswered: What individual patients do benefit from ECS therapy and what is the optimal individual treatment duration? Primary ethics approval was received from the Maastricht University Medical Centre. Results of the study will be disseminated via peer-reviewed publications and presentations at scientific conferences.

Trial Registration number:
NCT01429714 and NTR 2597
Background

After deep vein thrombosis (DVT) 20-50% of patients develop post thrombotic syndrome (PTS). PTS is a chronic condition, characterized by leg complaints such as pain; heaviness; cramps; aching; tingling; and leg signs due to venous insufficiency, evolving in severe cases to venous ulceration of the leg that was affected by the DVT. PTS is a serious affliction, with substantial impact on quality of life and costs. Because an effective therapy is lacking, prevention of PTS is of major importance. Up till now, elastic compression stocking (ECS) therapy after DVT has been the mainstay of PTS prevention. The evidence sustaining the value of ECS therapy following acute DVT is derived from 2 randomized clinical trials, which showed that incidences of PTS were reduced by approximately 50%, with application of ECS therapy for 24 months. However, based on these trials it is still undecided whether all patients benefit to the same extent from ECS therapy or what the optimal duration of ECS therapy for individual patients should be. We have previously assessed the safety of shortened duration of ECS therapy based on individual patient clinical scores in a management study of 125 patients, and we have shown that tailoring the duration of ECS therapy based on the signs and symptoms of the individual patient after an initial treatment period of 6 months, is a safe strategy to prevent PTS. We found that 50% of our patients did not need ECS therapy for as long as 2 years, while the overall incidence of PTS was 21.1% (95% CI 13.5- 28.7). This incidence is comparable to published incidences after 24 months ECS therapy. While this was a prospective management cohort study with an open character and therefore prone to bias, the results of this study need to be confirmed by an adequately powered randomized controlled trial.

There have been other studies that highlighted different aspects of ECS therapy. ECS therapy may not be indicated for all patients as shown by a study that assessed whether ECS therapy initiated one year after the event of DVT would lower the incidence of PTS in patients without complaints but with reflux on duplex testing. No significant benefit of ECS therapy at this late time of onset was found. The incidence of PTS was low in both groups. One study so far assessed whether prolonged duration of ECS therapy was superior to 6 months of ECS therapy following an event of proximal DVT. In this study no significant difference in the incidence of PTS according to the CEAP classification was observed between the two treatment groups.
The proposed IDEAL (individually tailored elastic compression against (=versus) longterm therapy) DVT study is a non-inferiority trial and aims to demonstrate that the assessed alternative therapy based on ECS for 6 months followed by individually prolonged ECS therapy based on Villalta scores is non-inferior to the comparator ECS therapy for a standard duration of 24 months. In addition we are interested to study the cost-effectiveness of individual tailoring of the duration of ECS therapy via a cost-utility analysis (CUA) and we want to retrieve information on the patient’s motivations for compliance or non-compliance to therapy. Because compliance to therapy is one of the most important prerequisites for successful ECS therapy, a discrete choice experiment (DCE) will be performed in which patient preferences towards ECS therapy will be assessed, in order to reveal the main obstacles for compliance to therapy. For this purpose, a DCE questionnaire will be send to a subset of the population (300 patients) at approximately 3 months after DVT.

The results of a recently published placebo controlled trial performed by Kahn et al. refuted the routine wearing of ECS after DVT altogether.\(^7\) This study showed no treatment effect at all from ECS therapy in contrast to the large treatment effects found in previous non-placebo controlled trials.\(^7\) The unexpected lack of effectiveness cannot be dismissed solely as a placebo effect. The compliance to ECS therapy, a major determinant of effectiveness, was 55.6% after 24 months. This was less in comparison to the previous trials by Brandjes and Prandoni, where compliance was up to 90%.\(^7\)\(^8\)\(^12\) Although a per protocol analysis of the patients reporting regular stocking use yielded the same outcome, this sub analysis may not be adequately powered to dismiss lack of compliance as an important determinant of non-effectiveness.

Based on current knowledge it is therefore understood that the standard application of ECS therapy is questioned. Especially the benefit to individual patients and the optimal duration of ECS therapy were never studied properly and as a result it is unclear which individual patients require therapy and for how long. The IDEAL DVT study aims to address these topics. In addition quality of life, patient preference towards ECS therapy, compliance to therapy, as well as cost-effectiveness of ECS therapy will be assessed.

The primary outcome of the IDEAL DVT study will be PTS at 24 months after the event. The secondary outcomes will be: 1. Health Related Quality Of Life, measured by questionnaires (SF-36\(^13\), EuroQOL-5D\(^14\), Dutch translated Veines-QoL\(^15\), 2. Costs\(^16\)), 3. Recurrent thrombosis, according to
criteria as published and assessed by objective tests, 4. VTE related death during follow-up, assessed by an independent and blinded adjudication committee, 5. Patient preferences, assessed with a Discrete Choice Experiment (DCE).

The scope of the problem in the Netherlands

In the Netherlands, each year around 25,000 patients experience a new event of DVT. In order to prevent PTS, common practice is to prescribe custom fitted ECS therapy for 2 years for all patients. ECS therapy is costly, demanding and sometimes even debilitating. Elderly patients can often not apply the ECS by themselves but need help from family or they even need home care visits (7.5%).

Total annual costs of ECS therapy roughly amounts to 2.5 million euro for stockings (25,000 patients*100 euro) and 21 million euro for home care (7.5%*25,000 patients*500 visits*20 euro). Moreover, the majority of patients do not experience any post thrombotic complaints. In the Netherlands, over 10 million euro each year could be saved if ECS treatment would be individualized.

Methods/Design

A multicentre, single-blinded, allocation concealed, randomized, non-inferiority trial comparing individually tailored duration of ECS therapy (intervention) with standard duration of 24 months ECS therapy (control), for the prevention of PTS. Randomization will guarantee a balanced distribution of patients within each patient group. Stratification will be performed on centre and potentially confounding effects such as age, sex and body mass index. The study will be a multicenter study in order to get a good representation of patients and to achieve sufficient patient numbers. The primary outcome will be PTS at 24 months after the event; the observers will be blinded to the allocated treatment arm. The allocation will be concealed from study personnel involved in assessing the leg symptoms and rating the PTS scores. Both randomization and allocation concealment are used as strategies against bias. The study is not double blinded; no inactive stockings will be used in the comparator group. The use of sham stockings will very likely interfere with the quality of life and will complicate the assessment of perceived differences between groups. Patients will be followed for the entire study duration; incidence of recurrent DVT and VTE related death during the follow-up period will be documented.
Patients and treatment

Consecutive, consenting adults with an acute objectively documented proximal DVT of the leg, adequately treated with anticoagulant treatment and initial compression therapy according to a prespecified protocol are included in the study. Patients are included and randomized in the IDEAL DVT study within 2-6 weeks after DVT.

Exclusion criteria are:

- Previous DVT in the affected leg. Patients with a previous ipsilateral DVT might already have developed PTS after the first DVT.
- Recurrent DVT in the 6 months following inclusion, as it cannot be justified to advise these patients to discontinue ECS therapy 6 months after DVT.
- Pre-existing venous insufficiency (skin signs C3-C6 on CEAP score or requiring ECS therapy). Pre-existing venous insufficiency increases the risk of developing PTS and the majority of patients with venous insufficiency already chronically wear elastic compression stockings. In addition venous insufficiency is closely related to PTS and is therefore difficult to differentiate from PTS.
- Contraindication for ECS therapy such as intermittent claudication or clinical signs of leg ischemia or asymptomatic arterial insufficiency (a pulse deficit or bruit at sites of narrowing at physical examination).
- Active thrombolysis, as thrombolysis reduces the risk of PTS.
- Limited life expectancy (< 6 months), as the follow-up period is 2 years.

Patients are recruited from 12 hospitals in the Netherlands and 2 hospitals in Italy.

There is a prespecified protocol for the management of DVT in all participating centres.

In the initial acute phase after DVT, until the acute oedema has disappeared, one of three strategies will be applied. The leg is either bandaged with short stretch bandages to a compression of 30-40 mmHg, worn day and night and redressed twice per week; or a bandage stocking with 35mmHg compression is prescribed (Mediven Struva 35®, Medi, Breda, the Netherlands), worn day and night; or no initial compression therapy is applied, according to the prespecified strategy of the hospital protocol. After the initial phase, a custom fitted, flat knitted, knee length graduated elastic compression stocking class III (ankle pressure 40 mmHg) is prescribed for all patients. The same brand and type (Mediven 550) of compression stocking is prescribed to all patients in all participating centres.
Compression therapy with the elastic compression stocking is started immediately after the initial phase, so there is no period without compression in between, with exception of the patients who do not receive compression in the initial phase. Patients receive two new stockings every year. Patients are advised to wear the stockings during the daytime.

Procedures
After obtaining written informed consent, randomization is performed centrally at the coordinating study centre, the Maastricht University Medical centre. Patients are included and randomized within 2-6 weeks after diagnosis of DVT. Patients are randomized to one of the two treatment arms. (Figure 1)
A web-based randomisation program (TENALEA (Trans European Network for Clinical Trials Services)) is used that executes blocked randomisation with stratification on centre-level and on possible confounding patient characteristics such as age, sex, and Body Mass Index (<26 kg/m² - ≥26 kg/m²). Permuted blocks of randomly varying size are used to maintain balance of numbers in each treatment group and to ensure allocation concealment.
Information on allocation is only accessible to the study personnel at the coordinating study centre in Maastricht, and patients will receive information about their allocation via the coordinating study centre. The allocation will be concealed from study personnel involved in assessing the leg symptoms and rating the PTS scores. Patients are not blinded by the use of sham stockings, because they may influence quality of life, complicating the assessment of perceived differences between groups. Patients are asked not to reveal their treatment allocation to their own physician, and not to wear the elastic compression stocking at the day of their outpatient clinic visits.
All patients are advised to wear the elastic compression stocking during the first 6 months after DVT. In the control group, all patients are instructed to wear the elastic compression stocking daily for a total duration of 2 years. In the Intervention group, duration of ECS therapy is individually tailored based on Villalta scores. After the first 6 months therefore, there are three scenarios:

1. The Villalta score at both the 3 and 6-month follow-up visit is ≤ 4. Then the patient is advised to discontinue ECS therapy.
2. The Villalta score at the 3-month follow-up visit is ≥ 5 the Villalta score at the 6 month follow-up visit is ≤ 4. The patient is advised to continue the ECS therapy for another 6 months. If the
Villalta score at the 12 month follow-up visit is ≤ 4, the patient is allowed to discontinue the ECS therapy.

(3) The Villalta score is ≥ 5 at both the 3 and 6 months follow-up visit. The patient is advised to continue the ECS therapy for a total duration of 24 months. (Figure 2)

Using this algorithm the duration of ECS therapy is individually tailored in each patient. The decision on the duration of ECS therapy in patients in the intervention group is made centrally at the coordinating study centre, as only there the information on treatment allocation is available. The patient is informed about the decisions on the (dis) continuation of the ECS therapy by the coordinating centre.

When a patient develops symptoms and signs of PTS after discontinuation of ECS therapy, a predefined protocol is followed. If necessary, ECS treatment will be reinstated.

Follow-up

All patients are followed for 24 months after DVT. During these 24 months they visit the outpatient clinic 4 times: at 3, 6, 12, and 24 months after the DVT. At each follow-up visit the signs of PTS will be recorded and scored, by the study nurse or the treating physician, using the objective part of the Villalta clinical scale for PTS. In case of intermittent leg complaints or signs or symptoms suspect of recurrent VTE, patients are instructed to visit their treating physician. Study documentation will be filled out and adverse events will be recorded. The CCMO (central committee human related research) will be notified in case of serious adverse events or death.

The patient is asked to fill out 5 questionnaires: at baseline, and 3, 6, 12 and 24 months later. All questionnaires will be offered as a web based application, each patient will have a unique personal entry code. For patients who are not able or willing to use the electronic questionnaires, a paper version will be available. The first questionnaire is filled out shortly after inclusion (4-6 weeks after DVT) and the subsequent questionnaires are filled out the day before each follow-up visit. All questionnaires contain the subjective part of the Villalta clinical scale for PTS, questions on ECS compliance, a cost questionnaire, and three health related quality of life (HRQOL) questionnaires: the Dutch translated Veines-Qol, the SF-36, and the EuroQOL-5D. In a subset of the study population patient preferences regarding elastic compression therapy will be analysed, by conducting
a discrete choice experiment (DCE). For this purpose, a DCE questionnaire will be send to a subset of
the population at approximately 3 months after DVT.

Compliance to ECS therapy is monitored in two ways. Each questionnaire contains a question on
compliance to the advised version of ECS therapy. Because compliance and adherence to therapy is
of crucial importance the number of contact moments for the assessment of compliance with ECS
therapy will be increased by adding telephone contacts to the regular clinical visits. Study-supporting
staff will make computer-assisted random telephone calls during the entire follow-up period to both the
patients in the group with the 2-year ECS intervention and to patients in the group with individualized
ECS therapy. The telephone calls will be made randomly to create a surprise effect (patients are off
guard) and to allow for a distribution of contacts over the entire study period. Per patient 3 extra
contact moments will be created. A standard questionnaire will be used to assess compliance and to
address reasons for non-compliance. Crossover to another therapy arm will be discussed and
discouraged.

Outcomes
The primary outcome is the proportion of patients with PTS at the end of follow-up, 24 months after
DVT. PTS is defined as a Villalta score of ≥5 at two consecutive visits that are at least three months
apart. The time point of the second Villalta score of ≥5 will be considered the time point of PTS
diagnosis. Although the consensus is to diagnose PTS at the 6-month visit or later based on one
single Villalta score ≥5, we modelled this study after the preceding management study in which two
consecutive Villalta scores ≥5 were needed for the diagnosis PTS. We observed that a proportion of
patients (about 12%) had fluctuating Villalta scores beyond 6 months after the acute event of DVT.5
Assigning the diagnosis based on one single score ≥5 would therefore lead to misclassification and
overestimation of the PTS incidence. The Villalta scores at 6 months will be available; therefore also
the proportion of patients with PTS according to the consensus scoring method will be presented.

Secondary outcomes include:
1. HRQOL, measured by questionnaires (SF-3613, EuroQOL-5D14, Dutch translated Veines-QoL15), 2.
Costs16, 3. Recurrent thrombosis, according to criteria as published17 and assessed by objective
tests. 4. VTE related death during follow-up, assessed by an independent and blinded adjudication committee, 5. Patient preferences, assessed with a Discrete Choice Experiment (DCE). Since both effects on costs and generic health-related quality of life are to be expected, the method of economic evaluation is a cost-utility analysis (CUA). The analysis will be from a societal perspective. The primary effect parameter is generic health-related quality of life, measured in quality adjusted life years (QALYs). We will perform two analyses: a trial based CUA with a time horizon of two years (identical to the duration of follow up in the clinical study), and a model-based CUA with a lifelong time horizon. For the latter, we will follow the guidelines of good modelling. We will be able to adapt a Markov model for diagnostic strategies in DVT, which was developed by our group earlier. Costs in the economic analyses include direct health care costs (medical costs for prevention, diagnostics, therapy, rehabilitation and care), direct non-health care costs (travel costs) and indirect costs (productivity loss). Utilities will be calculated from the responses on the EQ-5D and SF36 classification systems using the available multi-attribute utility functions. Incremental cost-utility ratios will be calculated, and non-parametric bootstrap analyses will be used to quantify the uncertainty surrounding the cost-utility ratio of the trial-based analysis. Sensitivity analyses and subgroup analyses will be performed, to assess the impact of variation in parameters and heterogeneity of the patient population. ECS therapy has several disadvantages for the patients (stockings are uncomfortable, ugly, and difficult to put on and off), while duration and effectiveness are uncertain. A DCE will be conducted to assess the patient preferences regarding ECS therapy, providing insight in the trade-offs patients make between characteristics of the therapy when deciding to wear the stocking or not. Duration of ECS therapy is one of the characteristics. Data will be analysed using multinomial logit models and mixed logit models. Sample size calculation The proposed study is a non-inferiority trial and aims to demonstrate that the assessed alternative therapy based on ECS for 6 months followed by individually prolonged ECS therapy based on Villalta scores is not worse than the comparator ECS therapy for a standard duration of 24 months, by more than the prespecified amount of 7.5% (the non-inferiority margin). The published incidence of PTS following two years of ECS lies between 20 and 30%. (Prandoni 24.5% (95% CI 15.6- 33.4), Brandjes
20% (95% CI 12.4-29.2)\(^7\) As it is statistically impossible to demonstrate equivalence (prove the H0 of no difference), Blackwelder proposed a one sided significance test to reject the null hypothesis by a clinically acceptable amount.\(^30\) If we allow a difference of 7.5% in the outcome PTS between the group with ECS for 2 years and the group with the alternative therapy, 70% of the effect will be preserved. This proportion of loss in efficacy is customarily accepted in controlled randomized clinical trials. At a one sided significance level of 0.05 and a power of 80%, with a ratio of 1, a total of 847 patients is needed to provide sufficient patients for an adequately powered trial.\(^30\) \(^31\) Loss to follow-up of patients is expected to be less than 2% since the intervention does not have an invasive nature. Therefore, a total of 864 patients are needed (432 patients per treatment arm).

Statistical analysis

Descriptive statistics of the total population and of the two treatment groups separately will be computed to provide baseline characteristics of the patients in both treatment arms.

For the primary outcome PTS, univariate analysis of all proportions will be performed with logistic regression (chi-squared) analysis. Kaplan-Meier method will be used to calculate the cumulative incidences of PTS, adjusted for centre, to compare incidence rates between the two treatment arms.

Patients who die or are lost to follow-up will be censored at their last visit. ANOVA will be applied to assess changes over time, by comparing different outcome measures at the different time points of follow-up.

Hazard ratio’s and 95% confidence intervals for both treatment arms will be calculated using Cox regression models. Hazard ratio’s will be stratified for centre and adjusted for age, sex, BMI, clinical presentation of DVT, and extent of the index deep vein thrombosis.

Interim analysis (safety)

A prespecified safety analysis will be performed when 50% of the subjects have completed the 2-year follow-up. The analysis will be performed by the coordinating centre, and supervised by the principal investigators. The safety analysis will be performed to assess significant enhanced risk of PTS or excess morbidity/mortality in the intervention arm of the study population. Fisher’s exact test will be performed to compare incidence of PTS at a significance level of 0.05 (two sided). The study can be stopped in case of significant excess morbidity/mortality in the alternative treatment arm.
Ethical considerations

The medical ethical committees of all participating hospitals in the Netherlands and Italy approved this study. All patients are extensively informed about the study, and written informed consent is obtained from all participating patients.

The IDEAL DVT study started including patients at 11th of March 2011 and inclusion is currently ongoing. Recruitment is expected to be terminated in January 2015. As the follow-up is 2 years, the results are expected within 3 years, in January 2017.

Discussion

Based on current knowledge the standard application of ECS therapy after DVT is questioned. The definitive answer on the usefulness of ECS therapy for the prevention of PTS is not yet provided. Therefore the need for the IDEAL DVT study remains unchanged. The benefit to individual patients and the optimal duration of ECS therapy were never studied properly and as a result it is unclear which individual patients require therapy and for how long. The IDEAL DVT study will address the central questions that remain unanswered: What individual patients do benefit from ECS therapy and what is the optimal individual treatment duration? The study that we plan to perform is not only innovative; for the first time individual tailoring of duration of ECS therapy will be investigated, but also it will provide unique additional knowledge on the safety, effectiveness and cost-effectiveness of this approach.

The IDEAL DVT study started including patients March 2011 and is currently ongoing. Recruitment is expected to be terminated within 1 year. As the follow-up is 2 years, the results are expected within 3 years.
**Trial status**

The IDEAL DVT study is currently in progress. Patient recruitment started March 15th 2011 and is still on-going.

**Acknowledgments**

The IDEAL DVT trial is funded by a grant from Zon-mW the Netherlands. (171102007)

We especially acknowledge the participating investigators.

**Competing interests**

The authors declare that they have no competing interests.

**Funding**

This work was supported by Zon-mW the Netherlands grant number 171102007.

**Contributions**

AtC wrote the protocol and designed the study, the other authors MJ (cost-effectiveness research), MP (study design/statistics) and HtC (study design) contributed. AB is a PhD student and coordinator of the trial.

**Ethics approval**

Maastricht University Medical Centre.
The IDEAL DVT investigators:

The Netherlands:

Maastricht, Maastricht University Medical Centre
Arina J ten Cate-Hoek MD, PhD (principal investigator)
Hugo ten Cate MD, PhD
Manuela Joore PhD
Annemieke C Bouman MD
Groningen, University Medical Centre Groningen
Karina Meijer MD PhD
Amsterdam, Academic Medical Centre
Saskia Middeldorp MD, PhD
Michiel Coppens MD, PhD
Mandy N Lauw MD
Y Whitney Cheung MD
Heerlen, Atrium Medical Centre
Guy JM Mostard MD
Asiong Jie MD PhD
Hoorn, Westfriesgasthuis
Simone M van den Heiligenberg MD
Eindhoven, Maxima Medical Centre
Lidwine W Tick MD, PhD
Marten R Nijziel MD, PhD
Almere, Flevohospital
Marije ten Wolde MD, PhD
Amsterdam, OLVG
Sanne van Wissen MD, PhD
Wim E Terpstra MD, PhD
Roermond, Laurentius hospital
Marlène HW van de Poel MD
References


Figure 1. Study flow diagram
90x79mm (300 x 300 DPI)
Figure 2. Algorithm individually tailored ECS therapy

For peer review only - http://bmjopen.bmj.com/site/about/guidelines.xhtml