Optimal Management of Anticoagulant Therapy

Educational Slides Set
American Society of Hematology 2021

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Clinical Guidelines

American Society of Hematology 2021
Guidelines for management of venous thromboembolism in Latin America

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Latin American ADOLOPMENT project

• The Latin American ADOLOPMENT project is a pilot collaborative effort of the following institutions
• Argentine Society of Hematology (SAH) Cecilia Colorio, MD
• Bolivian Society of Hematology and Hemotherapy (SBHH) Mario Luis Tejerina Valle, MD
• Brazilian Association of Hematology, Hemotherapy and Cellular Therapy (ABHH) Suely Meireles Rezende, MD PhD
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• Society of Hematology of Uruguay (SHU) Cecilia Guillermo, MD
• Venezuelan Society of Hematology (SVH) Juan Carlos Serrano, MD
• Latin American Cooperative Group of Hemostasis and Thrombosis (CLAHT) Patricia Casais, MD
• Mexican Association of Hematology Luis Meillon MD
• Colombian Association of Hematology and Oncology Guillermo Basantes MD
• American Society of Hematology
• MacGRADE Center

ASH Clinical Practice Guidelines on VTE

1. VTE prevention in hospitalized surgical patients
2. VTE prevention in hospitalized medical patients
3. Acute VTE Treatment (DVT y PE)
4. Optimum Management of Anticoagulation Therapy
5. VTE Prevention and Treatment in cancer patients
6. Heparin-induced Thrombocytopenia (HIT)
7. Thrombophilia
8. Pediatric VTE
9. VTE in the context of pregnancy
10. VTE Diagnosis
How are the ASH Guidelines developed?

PANEL CONFIRMATION
Each panel was formed based on key criteria:
• Balanced experience (including disciplines beyond hematology and patients)
• Attention to COI minimization and management

CLINICAL QUESTIONS
10 to 20 clinically relevant questions worked out on PICO format (population, intervention, comparison and outcome)

SYNTHESIS OF EVIDENCES
Analysis of evidence of each PICO question x systematic review of effects:
• Desirable and Undesirable Effects
• Use of Resources
• Feasibility
• Acceptability
• Accessibility
• Patient Values and Preferences

EXAMPLE OF PICO QUESTION
Should antithrombotic agents be administered early or late to patients submitted to surgery?

RECORD OF RECOMMENDATIONS
Recommendations made by panel members based on evidences of all factors
How should patients and physicians use these guidelines?

<table>
<thead>
<tr>
<th></th>
<th>STRONG Recommendation</th>
<th>CONDITIONAL Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(“The panel recommends ...”)</td>
<td>(“The panel suggests ...”)</td>
</tr>
<tr>
<td>For Patients</td>
<td>Most individuals will be for the intervention</td>
<td>Most individuals will be for the intervention, but several will not.</td>
</tr>
<tr>
<td>For Doctors</td>
<td>Most individuals should receive intervention.</td>
<td>Different adequate options for different patients, depending on their values and preferences. Make use of shared decisions.</td>
</tr>
</tbody>
</table>
Goals

1. By the end of this session, you will be prepared to:

2. Define the level of care and type of initial anticoagulation of VTE patients

3. Establish the anticoagulation period according to the VTE event, provoked or not provoked with or without recurrence, with new VTE events under anticoagulation

4. Determine the role of both scores of recurrence and D-dimer in provoked events.

5. Management of complications due to anticoagulation
Anticoagulants bring benefits (reduction in thrombus extension, mortal PE) and risks (potentially lethal hemorrhage).

Recognition and mitigation of the risks of damage from anticoagulants is achieved thru a management approach based on evidences.

This chapter focus on the optimal management of anticoagulants for the prevention and treatment of TVE (after choosing the anticoagulant).
Case 1: New deep venous thrombosis and acute pulmonary embolism
Male - 58 years of age

- **Previous pathological history:** Post-Operative between right kneecap replacement (TKR), AHT, chronic ischemic cardiomyopathy, overweight (BMI 29)
- **Medication:** Losartan, Carvedilol, ASA 100 mg/day
- **Clinical Profile:** Chest pain for 24 hours associates to mild moderate dyspnoea. Pain on right leg, AT 110/77 mmHg, pulse 96/min, RF 24/min SO2 92%. Increase of volume and pain over the entire right inferior limb from the last 72, lab: Dimer- D elevated, normal renal and hepatic functions, Scan Duplex showed extensive DVT of femoral and right popliteal vein, Femoral DVT in Scan duplex. AngioTAC: Subsegmental Pulmonary embolism without Right Ventricular dysfunction on Echocardiogram.
- **Diagnosis:** Proximal Deep Vein Thrombosis complicated with pattern provoked pulmonary embolism
Considering his clinical condition of low risk and hemodynamic stability; How would you consider to conduct his treatment?

• a) Fibrinolytic therapy, hospitalized
• b) Hospitalized treatment with the use of unfractionated heparin and then Warfarin
• c) Outpatient management with the use of DOACs exclusively
• d) Short hospitalization with LMWH, but outpatient management mainly and then DOAC or Warfarin according to availability

• C and D are correct
Recommendations

For patients with PE and low complication risks, the Latin American Panel suggests \textit{treatment at home or hospital treatment} (\textit{conditional recommendation, based on a very low certainty of the evidence on the effects}).

<table>
<thead>
<tr>
<th>Results (Quality of Evidence)</th>
<th>Relative Risk (95% CI)</th>
<th>Risk with Hospitalized Treatment</th>
<th>Risk with Outpatient treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mortality</td>
<td>RR 0.33 (0.01 - 0.98)</td>
<td>6 per 1000</td>
<td>4 minus per 1000 (6 minus to 42 plus)</td>
</tr>
<tr>
<td>PE</td>
<td>RR 2.95 (0.12 - 7.85)</td>
<td>0 per 1000</td>
<td>0 por 1000</td>
</tr>
<tr>
<td>Proximal Symptomatic DVT</td>
<td>Not estimable</td>
<td>0 x 1000</td>
<td>0 x 1000 (0 x 1000)</td>
</tr>
<tr>
<td>Greater Bleeding</td>
<td>RR 6.88 (0.36 - 134.1)</td>
<td>0 per 1000</td>
<td>0 x 1000</td>
</tr>
</tbody>
</table>

Evidence Quality (GRADE): Low, Moderate, Strong

Evidence of low quality, therefore benefit/damage is uncertain. The panel also considered:
- This recommendation is not applicable to patients with other major risk conditions that require hospitalization, limited support or none at home, cannot afford drugs or have a history of deficient non-compliance.
- High risk of bleeding may also need to start treatment at the hospital.
- The treatment at home may not be feasible in certain contexts due to health system limitations or insurance policy restrictions.
# Pesi score for PE severity classification

**Table 1: Original and simplified Pulmonary Embolism Severity Index (PESI).**

<table>
<thead>
<tr>
<th>Variable</th>
<th>PESI (a) Original Score</th>
<th>PESI (b) Simplified Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age &gt; 80 years</td>
<td>Age in years</td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>+10</td>
<td>-</td>
</tr>
<tr>
<td>History of Cancer</td>
<td>+30</td>
<td>1</td>
</tr>
<tr>
<td>History of Heart Failure</td>
<td>+10</td>
<td>1 (c)</td>
</tr>
<tr>
<td>History of Chronic Pulmonary Disease</td>
<td>+10</td>
<td></td>
</tr>
<tr>
<td>Pulse ≥ 110 beats /min</td>
<td>+20</td>
<td>1</td>
</tr>
<tr>
<td>Systolic Arterial pressure &lt; 100 mm Hg</td>
<td>+30</td>
<td>1</td>
</tr>
<tr>
<td>Respiratory Frequency ≥ 30 x min</td>
<td>+20</td>
<td>-</td>
</tr>
<tr>
<td>Temperature &lt; 36°C</td>
<td>+20</td>
<td>-</td>
</tr>
<tr>
<td>Altered Mental Status*</td>
<td>+60</td>
<td>-</td>
</tr>
<tr>
<td>Arterial Oxygen Saturation &lt; 90%*</td>
<td>+20</td>
<td>1</td>
</tr>
</tbody>
</table>

The patient total punctuation is obtained by adding up patient age in year and the points of each predictor when present. The score corresponds to the following risk classes: class I (≤65 points), class II (66-85 points), class III (86-105 points), class IV (106-125 points) and class V (> 125 points). Patients under risk classes I and II are defined as of low risk. (b) A patient total score is obtained by adding up the points. The score is sorted out according to the following risk classes: 0 low risk, 1 or more high risk. Empty cells indicate that variables have not been included. (c) Variables have been combined into one sole category of Chronic Cardiopulmonary Disease.

**Fibrinolysis in Acute Pulmonary Embolism**

**PATIENT CLASS I**

**Low Risk**

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Recommendation

For patients with con DVP or PE, the ASH Latin American Panel suggests the use of DOAC over AVK (conditional recommendation, based on moderate certainty of evidences about the effects).

Evidences of Research

• There are no direct comparison trials between DOAC and HBPM on this indication

• Indirect Evidences: DOAC vs HBPM have been compared only on VTE prophylaxis trials on hip and knee replacements, where DOAC reduces the risk of DVT de TVP with no increase of bleeding.

• However, in prophylaxis for hospitalized medical patients, the use of DOAC increases bleeding if compared to HBPM

Moderate Quality of evidence. The Panel also considers that:

• Patients well controlled and with no complications can remain with AVK.

• Cases de Novo may prefer DOAC with regards to safety, load of treatment, difficulties to monitor the INR.

• Increase vigilance for the risk of bleeding with DOAC, even more so when domiciled far away.
Case 1 (Continued):

- The patient was started on oral Rivaroxaban 15 mg every 12 hours for 21 days, then received 20 mg day for the following 3 and 6 months.
- On day 2 of treatment, there is substantial improvement of the respiratory condition, but with development of much pain and functional helplessness and evaluations highlight a pronounced extensive DVT throughout the femoral vein.
- Assessed for vascular surgery to decide on the approach.
The patient is assessed for Vascular Surgery, and the following proposals are discussed in the clinical team meeting; Which one do you agree with?

A. Perform immediate surgical thrombectomy
B. Keep only anticoagulation with Rivaroxaban for 3 to 6 months, individualizing recurrence risk.
C. Perform IV thrombolysis
D. Perform catheter guided thrombolysis.
## Recommendation

In patients with extensive proximal DVT, the ASH Latin American Panel suggests *against the thrombolysis in addition to anticoagulation* (*conditional* recommendation, *based on low certainty of the evidence on the effects*).

<table>
<thead>
<tr>
<th>Results (Quality of Evidence)</th>
<th>Relative Risk (95% CI)</th>
<th>Absolute Anticipated Effects (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Late Mortality</strong></td>
<td>RR 0.89 (0.46 - 1.69)</td>
<td>High 67 per 1000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Risk with anticoagulant + Thrombolysis</td>
</tr>
<tr>
<td></td>
<td></td>
<td>7 minus per 1000 (7 minus to 36 plus)</td>
</tr>
<tr>
<td><strong>PE</strong></td>
<td>RR 1.33 (0.71 to 2.46)</td>
<td>0 per 1000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Risk with anticoagulant + Thrombolysis</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0 per 1000</td>
</tr>
<tr>
<td><strong>Proximal symptomatic DVT</strong></td>
<td>RR 0.99 (0.56 to 1.76)</td>
<td>High 130 x 1000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Risk with anticoagulant + Thrombolysis</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1 minus x 1000 (57 minus to 99 plus)</td>
</tr>
<tr>
<td><strong>Post phlebitic syndrome</strong></td>
<td>RR 0.71 (0.60 to 0.85)</td>
<td>High 563 per 1,000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Risk with anticoagulant + Thrombolysis</td>
</tr>
<tr>
<td></td>
<td></td>
<td>163 minus x 1000 (225 minus to 84 less)</td>
</tr>
<tr>
<td><strong>Leg ulcer</strong></td>
<td>RR 0.75 (0.39 to 1.42)</td>
<td>High 30 per 1,000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Risk with anticoagulant + Thrombolysis</td>
</tr>
<tr>
<td></td>
<td></td>
<td>8 minus x 1000 (18 minus to 13 plus)</td>
</tr>
</tbody>
</table>

*Evidence of low quality, therefore benefit / damage is uncertain. The Panel also considered:*  
- Thrombolysis to be reasonable in cases of DVT threatening limbs, with severe symptoms that do not improve with anticoagulation only and or with iliofemoral DVT with high risk of PFS and low-medium risk of  
- To take in account basal risks, patient preference and access to experimented care.
Case 1: Summary

For patients with PE and low risk of complications, suggestion of treatment either at home or hospital, according to availability of resources.

In case of low-risk PE or DVT plan the use of DOAC over AVK, even if well controlled patients can be maintained with AVK, in both cases monitor the risk of bleeding.

Thrombolysis is not recommended in extensive proximal venous thrombolysis, for the prevention of postphlebitic syndrome.
Case 2: Deep venous thrombosis not provoked with high risk of bleeding
Woman - 40 years of age

- **Pathological History:** Recurrent gastric peptic ulcer disease
- **Medication:** Esomeprazol 40 mg day

- **Clinical Profile:** As she rises from bed in the morning, she notices the development of edema and pain in the left inferior limb, difficulty to walk. Dimer-D on 1550 ug/L Scan Duplex showed left Ileofemoral DVT. Started treatment with Enoxaparin for 5 days then she is kept on Warfarin
- **Thrombophilia profile negative**

- **Diagnosis:** Proximal Deep Venous Thrombosis unprovoked
Considering her current clinical condition how long would you consider to give anticoagulation with warfarin?

A. I would give anticoagulation for 2 months
B. I would give anticoagulation for 3 to 6 months only
C. I would give anticoagulation extended to beyond 3 – 6 months and would assess for the risk of thrombotic recurrence and bleeding
D. I would give her anticoagulation indefinitely
## Recommendation

For patients with unprovoked PE or DVT, the ASH Latin American Panel suggests to **keep anticoagulation indefinitely over the interruption after a period of 3 to 6 months** (conditional recommendation based on the moderate certainty of the evidence on the effects).

### Results (Evidence Quality)

<table>
<thead>
<tr>
<th>Result</th>
<th>Relative Risk (95% CI)</th>
<th>Absolute Anticipated Effects (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mortality</td>
<td>RR 0.75 (0.49 - 1.13)</td>
<td>Risk with defined antithrombotic duration (12 months or less): 18 per 1,000 (5 minus per 1,000 (9 minus to 2 plus))</td>
</tr>
<tr>
<td>PE</td>
<td>RR 0.29 (0.15 - 0.56)</td>
<td>Risk with undefined antithrombotic duration: 29 per 1,000 (21 minus per 1,000 (25 minus to 13 minus))</td>
</tr>
<tr>
<td>Proximal symptomatic DVT.</td>
<td>RR 0.20 (0.12 to 0.34)</td>
<td>63 per 1,000 (50 minus per 1,000 (56 minus to 42 minus))</td>
</tr>
<tr>
<td>Major Bleeding</td>
<td>RR 2.24 (1.49 to 3.35)</td>
<td>5 per 1,000 (6 plus per 1,000 (2 plus to 12 plus))</td>
</tr>
</tbody>
</table>

### Evidence Quality (GRADE): Low        Moderate        Strong

Moderate evidence, so the panel considered that:

- The individual risk of DVT recurrence, risk of bleeding, costs, access to follow up and monitoring should be considered, in addition to patient values and preferences.
- This recommendation is applied to patients with average risk of bleeding.
- The risk of bleeding may change with time, benefit vs risk of anticoagulation should be reassessed periodically.
VTE – Recurrence Risk

- Acute Episode of VTE effectively treated.
- Intrinsic Risk of each case in VTE:
  - 30.3% after 8 years
  - 24.6% after 5 years
  - 17.5% after 2 years

Unprovoked VTE

Recurrence Risk after stopping anticoagulation

1) Acute Episode of VTE effectively treated.
2) Intrinsic Risk of each case in VTE

## Stratification of the Recurrence Risk of VTE

<table>
<thead>
<tr>
<th>High Risk</th>
<th>Moderate Risk</th>
<th>Low Risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>• VTE in the last 3 months</td>
<td>• VTE in the last 3-12 months</td>
<td>• VTE &gt; 12 months before</td>
</tr>
<tr>
<td>• Deficiency of protein C, protein S or antithrombin</td>
<td>• Heterozygous V Leiden Factor</td>
<td>• No other risk factors</td>
</tr>
<tr>
<td>• Antiphospholipid syndrome</td>
<td>• 20210 prothrombin mutation</td>
<td></td>
</tr>
<tr>
<td>• Multiple thrombophilic anomalies</td>
<td>• Recurrent VTE</td>
<td></td>
</tr>
<tr>
<td>• Active Cancer</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Members of the medical team treating the patient discuss the possibility of using Dimer-D or recurrence clinical scores to guide the use of anticoagulation.

Do you think this is a valid approach?

YES

NO
Recommendation

For patients with unprovoked PE of DVT, the Latin American Panel suggests to be **against** the use of Dimer - D or prognosis scores to guide the duration of the anticoagulation (conditional recommendation based on low certainty on proofs of the effects)

<table>
<thead>
<tr>
<th>Results (Evidence Quality)</th>
<th>Relative Risk (95% CI)</th>
<th>Absolute Anticipated Effects (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Mortality</strong></td>
<td>RR 1.06 (0.07 to 18.30)</td>
<td>1 per 1000</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>1 plus per 1,000</strong> (9 minus to 168 plus)</td>
</tr>
<tr>
<td><strong>PE</strong></td>
<td>RR 0.16 (0.02 to 1.33)</td>
<td>10 per 1000</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>8 minus por 1,000</strong> (10 minus to 3 plus)</td>
</tr>
<tr>
<td><strong>Proximal Symptomatic DVT</strong></td>
<td>HR 2.59 (1.90 to 3.52)</td>
<td>11 per 1000</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>17 mas por 1000</strong> (9 plus to 26 plus)</td>
</tr>
<tr>
<td><strong>Major Bleeding</strong></td>
<td>RR 3.49 (0.14 to 84.76)</td>
<td>10 per 1000</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>24 mas por 1,000</strong> (8 minus to 813 plus)</td>
</tr>
</tbody>
</table>

**Evidence of low quality, uncertain benefits:**
- Should guide towards recommendation 8, undefined anticoagulation is maintained with recurrence risk vs bleeding assessed with time
- Dimer-D only as part of a prognostic model may be useful to determine there is much indecision or difficult clinical situation.
Case 2 (continued)

The patient was kept on anticoagulation with warfarin within therapeutic INR range, but as of the 7 month shows thrombotic recurrence whilst under treatment.

What would be your anticoagulation strategy and for how long?

A. I would increase the warfarin dose (INR 3 to 4) with indefinite anticoagulation
B. I would change to DOAC within indefinite anticoagulation
C. I would change to DOAC within definite period for a year assessing the recurrence risk
D. I would recommend anticoagulation with HBPM, with re-assessment of causes of thrombosis, then defining what the most appropriate oral agent for indefinite use.
Evidence of low quality, therefore the panel has also considered:

- To appraise the vast experience in HBPM for prothrombotic conditions.
- Not to demonstrate AVK in suboptimal range. A better dose adjustment must be guaranteed.
- The need to explore the underlying causes of the recurrence under AVK.
- Final selection based on the underlying cause, patient values and preferences, cost and viability of each alternative.

Recommendation

For patients with DVT or PE during the VKA treatment, the ASH Latin American Panel suggests the use of LMWH over DOAC (conditional recommendation, based on a very low certainty of the evidence of the effects).

Research Evidences

- There are no direct comparison trials between DOAC and HBPM in this indication.
- Indirect Evidence: DOAC vs HBPM have been compared in VTE prophylaxis trials in hip and knee replacement, in which DOAC reduces the risk of DTV and there is no bleeding increase.
- However, prophylaxis in hospitalized medical patients, the use of DOAC increases bleeding when compared to HBPM.
**Recommendation**

For patients with unprovoked recurrent PE or DVT, the ASH Latin American Panel recommends to **maintain indefinite** anticoagulation over the its interruption after a period of 3 to 6 months *(strong recommendation, based on moderate certainty of the evidence on the effects)*.

<table>
<thead>
<tr>
<th>Results (Evidence Quality)</th>
<th>Riesgo Relativo (95% CI)</th>
<th>Anticipated Absolute Effects (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>RR 0.75 (0.49 to 1.13)</td>
<td>Risk with defined duration anticoagulation (12 months or less) 16 per 1000</td>
</tr>
<tr>
<td></td>
<td>RR 0.29 (0.15 to 0.56)</td>
<td>Risk with undefined duration anticoagulation 4 minus per 1,000 (8 minus to 2 plus)</td>
</tr>
<tr>
<td></td>
<td>RR 0.20 (0.12 to 0.34)</td>
<td>29 per 1000</td>
</tr>
<tr>
<td></td>
<td>RR 2.17 (1.40 to 3.35)</td>
<td>63 per 1000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5 per 1000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>6 plus per 1,000 (2 plus to 12 plus)</td>
</tr>
</tbody>
</table>

**Strong evidence of Good quality, moderate certainty, the Benefit is clearer:**

- This recommendation assumes the average risk of bleeding, it cannot be applied in cases with high probability of hemorrhage.
- The risk of bleeding may change with time, so the balance between desirable and undesirable consequences of indefinite anticoagulation must be often re-assessed.
### Recurrence Risk after suspending anticoagulation

**Observational Studies Data**

<table>
<thead>
<tr>
<th>Event provoked by a transitory risk factor</th>
<th>Unprovoked Event</th>
</tr>
</thead>
<tbody>
<tr>
<td>• EVT recurrent = 4,2 per 100 patient-year</td>
<td>• EVT recurrent = 7,4 per 100 patient-year</td>
</tr>
<tr>
<td>• PE recurrent = 1,9 per 100 patient-year</td>
<td>• PE recurrent = 3,3 per 100 patient-year</td>
</tr>
<tr>
<td>• DVT recurrent = 2,3 per 100 patient-year</td>
<td>• DVT recurrent = 4,1 per 100 patient-year</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Event provoked by a chronic risk factor (cancer excluded)</th>
<th>Recurrent Unprovoked Event</th>
</tr>
</thead>
<tbody>
<tr>
<td>• EVT recurrent = 9,7 per 100 patient-year</td>
<td>• EVT recurrent = 12 per 100 patient-year</td>
</tr>
<tr>
<td>• PE recurrent = 4,4 per 100 patient-year</td>
<td>• PE recurrent = 5,4 per 100 patient-year</td>
</tr>
<tr>
<td>• DVT recurrent = 5,3 per 100 patient-year</td>
<td>• DVT recurrent = 6,6 per 100 patient-year</td>
</tr>
</tbody>
</table>

* DVT and PE rates were calculated assuming that 45% de of recurrent EVT are PE.

Case 2: Summary

For patients with unprovoked or recurrent unprovoked DTV or PE, suggestion to maintain indefinite anticoagulation over its interruption after a period of 3 to 6 months.

In cases of unprovoked PE or DVT, suggestion to oppose the use of Dimer-D or prognostic scores to guide the duration of the anticoagulation, except for some very complex situations.

In patients with DTV or PE during the VKA treatment, suggestion to use LMWH over DOAC initially, while a more efficient anticoagulation is proposed.
Caso 3. Complications due to anticoagulation

**Personal History:** Hypertension, Chronic Renal Failure (no dialysis). Has not attended medical control for 3 months.

**Clinical Profile:** Female patient, 58 years of age, on Warfarin for 1 month for prevention of recurrent unprovoked PTE. Goes to hospital complaining of headache, dizziness, vomit and difficulty to move around in those 6 previous hours. CAT of skull performed Subarachnoid hemorrhage Fisher 3 and the INR is 10.

**Diagnosis:** Subarachnoid hemorrhage, Warfarin intoxication, recurrent PTE
Your patient is in vital emergency, with Hemorrhagic Cerebrovascular Accident (HCVA) and warfarin overdose. What would be the initial management recommendation to approach this case?

A. Discontinue warfarin
B. Give plasma fresh frozen plasma
C. Give Vitamin K 5 mg IV
D. Use of Prothrombin complex concentrate
E. B and D are correct
Recommendation

For patients with potential lethal bleeding related to AVK during the VTE treatment, the ASH Latin American Panel suggests the use of 4 factor PCC or PFC in addition to the interruption of AVK, according to local availability and clinical circumstances (conditional recommendation, based on a very low certainty of the evidence on the effects).

<table>
<thead>
<tr>
<th>Results (Evidence Quality)</th>
<th>Relative Risk (95% CI)</th>
<th>Anticipated Absolute Effects (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Risk with Frozen Fresh Plasma</td>
<td>Risk with CCP</td>
</tr>
<tr>
<td>Mortality</td>
<td>RR 0.92 (0.37 a 2.28)</td>
<td>124 per 1000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>10 minus per 1000 (78 minus to 159 plus)</td>
</tr>
<tr>
<td>DVT (all)</td>
<td>RR 1.60 (0.70 to 3.62)</td>
<td>68 per 1000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>41 plus per 1000 (20 minus a 179 plus)</td>
</tr>
<tr>
<td>Major Bleeding</td>
<td>RR 1.34 (0.78 to 2.29)</td>
<td>91 per 1000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>31 plus per 1000 (20 minus to 117 plus)</td>
</tr>
</tbody>
</table>

Low quality evidence, therefore the panel also considered, (low certainty of the evidence on the effects):

- There is no substantial evidence in safety and efficacy of results between CCP and PFC.
- To consider the use of CCP in cases of heart failure and volume overload, in addition zones with high risk of pathogen transmission.
- We should favor the fastest option according to local availability and costs in Latin America.
Case 3 (continued)

The patient received successful surgical intervention, considering life threatening bleeding she had and the unprovoked recurrent thrombosis.

What would be your treatment strategy?

A. Discontinue anticoagulant due to high risk of new bleeding

B. Re-start oral anticoagulation once she has been clinically recovered between 15 and 90 days

C. Recommend HBPM in one week

D. Recommend AAS
### Recommendation

For patients that receive treatment to VET and survive a major bleeding episode related to the anticoagulation therapy, the ASH Latin American Panel suggests to resume oral anticoagulation therapy upon interruption (conditional recommendation based on the very low certainty of the evidence on the effects).

Resume vs interrupt anticoagulant treatment for VET in the wake of a major bleeding:

<table>
<thead>
<tr>
<th>Results (Evidence Quality)</th>
<th>Relative Effect (95% CI)</th>
<th>Anticipated Absolute Effects (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mortality</td>
<td>RR 0.59 (0.45 to 0.77)</td>
<td>Risk with Discontinuation: 845 of 2,455 (34.4%), Difference of Risk with resume of anticoagulation: 141 less deaths per 1000 (79 minus to 189 plus)</td>
</tr>
<tr>
<td>PE</td>
<td>RR 0.26 (0.08 to 0.82)</td>
<td>12 of 425 (2.8%), 21 less PE per 1000 (5 minus to 26 minus)</td>
</tr>
<tr>
<td>Symptomatic Proximal DVT</td>
<td>RR 0.66 (0.25 to 1.75)</td>
<td>11 of 464 (2.4%), 8 less DVT per 1,000 (18 minus to 18 plus)</td>
</tr>
<tr>
<td>Major Bleeding</td>
<td>RR 1.54 (1.18 to 2.02)</td>
<td>230 of 3,304 (7.0%), 38 more bleeding per 1000 (13 plus to 71 plus)</td>
</tr>
</tbody>
</table>

Evidence Quality (GRADE): Low Moderate Strong

- Increased risk of recurrent hemorrhage offset by an improvement on mortality due to all causes
- Applied to patients that require anticoagulation for long or indefinitely.
The decision to resume anticoagulation may vary according to the risk of recurrent PTE and the risk and severity of bleeding.

An approach of decisions shared that explore values given by patients to the prevention of PTE or bleeding may be a form of implementing the recommendation.

Time to resume anticoagulation remains unknown and it varies depending on particularities of each patient. It is reasonable to consider waiting for at least 2 weeks, but no longer than 90 days after the bleeding episode. However, anticoagulation should be considered to resume as early as possible if bleeding cause was identified and corrected.
Case 3 (continued)

- Patient and family members, considering the hemorrhagic risk, discuss the possibility of using antithrombotic drug of lower hemorrhagic risk, she is not a candidate to DOAC due to renal failure,
- Debate about the possibility of aspirin 100 mg day.

Do you agree with this approach?

YES  NO
**Recommendation**

Should Aspirin be used versus standard dose of anticoagulation in patients to whom an indefinite duration is preferred, after completing an initial anticoagulation course of defined duration (12 months or less). *(Conditional Recommendation, based on a moderate certainty of the evidence)*.

<table>
<thead>
<tr>
<th>Results (Evidence Quality)</th>
<th>Relative Risk (95% CI)</th>
<th>Anticipated Absolute Effects (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Risk with standard anticoagulation</td>
</tr>
<tr>
<td><strong>Mortality</strong></td>
<td>RR 0.86 (0.31 to 2.35)</td>
<td>7 per 1000</td>
</tr>
<tr>
<td><strong>PE</strong></td>
<td>RR 3.10 (1.24 to 7.73)</td>
<td>5 per 1000</td>
</tr>
<tr>
<td><strong>Proximal Symptomatic DVT.</strong></td>
<td>RR 3.15 (1.50 to 6.63)</td>
<td>8 per 1000</td>
</tr>
<tr>
<td><strong>Major Bleeding</strong></td>
<td>RR 0.49 (0.12 to 1.95)</td>
<td>5 per 1000</td>
</tr>
</tbody>
</table>

*Evidence with moderate certainty, the benefit is clearer:*
- Compared with long term anticoagulation, the treatment with Aspirin could increase the risk of VTE with all negative consequences (hospitalization costs, vs disease risks, etc.).
- The panel assumed that cost effectiveness favors long term anticoagulation.
Summary Case 3

For patients with potentially mortal bleeding related to AVK during the treatment of VTE, the use of 4 factor CCP or PFC is considered in addition to the interruption of AVK, according to local availability.

For patients with VTE who require indefinite anticoagulation, consider the possibility of resuming anticoagulation within 15 to 90 days after a major hemorrhagic episode.

For the use of long term antithrombotic agents, the use of Aspirin does not replace anticoagulants.
Other Guide Recommendations not approached in this discussion.

- Thrombolysis in submassive PE based with eco or biomarkers compatible with the right ventricle dysfunction.
- Use of compression stockings by patients with DVT and high risk of post-thrombotic syndrome.
- Use of DOAC standard dose vs lower doses in long term anticoagulation
- Use of Aspirin in cardiovascular primary prevention associated with chronic anticoagulation
- Definition of anticoagulation in recurrent provoked events and with persistent chronic factors

Those first 4 recommendations with low or very low certainty of the evidence
Summary
Back to Goals

1. Define the attention level and type of initial anticoagulation of patients with VTE
   - Cases of low risk PE and DVT can be managed as out-patients, for initial anticoagulation ACOD, thrombolysis is not indicated for extensive DVT

2. Establish anticoagulation period in VTE, provoked and unprovoked with or without recurrence, with new VTE under anticoagulation
   - In unprovoked recurrent VTE guidance towards indefinite anticoagulation

3. Determine the Role of recurrence and Dimer-D scores in unprovoked events.
   - Not appropriate the use of Dimer-D and Score System to guide routine anticoagulation

4. Manage complications caused by anticoagulation
   - Both CCP and PFC can be used to reverse Warfarin anticoagulation, in the wake of a severe hemorrhagic event, it is suitable to restart anticoagulation between 15 and 90 days
Gratitude

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For more information on the ASH Guidelines on VTE: [www.hematology.org/VTE](http://www.hematology.org/VTE)