Thrombophilia or hypercoagulable state is a predisposition to form clots. Thrombophilia can be inherited or acquired, and prevalently involves venous vessels. Inherited thrombophilia consists of congenital conditions, as methylenetetrahydrofolate reductase polymorphism, Factor V Leiden and prothrombin gene mutations, natural anticoagulant deficiencies, high level of factor VIII, or dysfibrinogenemia. These congenital disorders can be responsible for venous thromboembolism, particularly deep venous thrombosis, pulmonary embolism, and, less frequently, mesenteric veins thrombosis, kidneys’ veins thrombosis or retinal vein occlusion. Acquired thrombophilia can be associated both with venous and arterial thrombosis and may be caused by antiphospholipid syndrome, aging, some malignancies, oral contraceptive use, heparin-induced thrombocytopenia, and human immunodeficiency virus. Antiplatelets’ drugs are employed in arterial thrombosis, while, heparins/oral vitamin K antagonists are indicated for acute and long-term anticoagulation. However, new oral anticoagulants can be usefully used for venous thromboembolic events. Recent experiences demonstrated that their employment is useful in some thrombophilias only, whereas other investigations are requested to evaluate their use in all hypercoagulable disorders.

Inherited thrombophilia

This condition is defined as genetic predisposition to form thrombi prevalently in the venous vessels. The leading causes responsible for the inherited hypercoagulable state are reported in Table 1.
Homocysteine

Homocysteine (Hcy) is a sulfhydryl-containing amino acid synthesized during the Methionine-metabolism. Two pathways control the serum level of Hcy: remethylation and trans-sulfuration, respectively involving the enzyme MTHFR and cystathionine-β-synthase.10,11 We only illustrate the impaired remethylation pathway, responsible for inherited mild hyperhomocysteinemia (HHcy), due to the gene mutation of the enzyme MTHFR. In homozygous trait, the MTHFR mutation reduces its activity by 50-60%, while the enzymatic activity of heterozygous trait is more modestly reduced.12

Accumulating evidences link HHcy levels to thrombosis both in arterial and venous vessels. The role of Hcy in venous thromboembolism has been studied less extensively than its role in arterial diseases, and nowadays, it seems quite controversial. It is also possible that HHcy plays a role in the pathogenesis of venous thromboembolism only as an additional risk factor in the presence of other thrombophilic disorders. Among these, Factor V Leiden, increased thrombin generation, impaired fibrinolytic potential, or proteins S-C inhibition, decreased activity of antithrombin and others, are included.13

Factor V Leiden mutation

Factor V Leiden (FVL) mutation is a moderate cause of inherited thrombophilia, accounting for 40-50% of cases. FVL mutation involves G (guanine) to A (adenine) substitution at nucleotide 1691 (G1691A). It accounts for 92% of cases of activated protein C resistance (APC-R). Therefore, the terms factor V Leiden and APC-R should be considered synonymous. APC-R caused by factor V Leiden mutation is the most common predisposition to hypercoagulability in white populations, mainly favoring deep venous thrombosis (DVT), venous thromboembolism (VTE) and pulmonary embolism (PE).14,15 This disorder also seldom causes the formation of clots in the arteries, inducing transient ischemic attack, or coronary events. Individuals with FVL heterozygous mutation have a risk of hypercoagulability increased by 3-4 fold, while homozygous mutation increases the risk approximately by 80 fold. Nevertheless, the heterozygous mutation is much more common than homozygous.

Prothrombin G20210A

Activated prothrombin is changed into thrombin, that converts fibrinogen into fibrin inducing the blood clot formation. It must be added that thrombin, apart with fibrinogen, reacts with calcium ions activating factor VIII, which stabilizes the fibrin-clots.16

Prothrombin-related thrombophilia is characterized by VTE, which manifests most commonly in adults as DVT in the legs or PE. The most common mutation of this protein is the prothrombin gene mutation (PTG) at nucleotide 20210 (G20210A), which represents the second most common inherited predisposition to hypercoagulability. People having a mutation in the prothrombin gene produces more prothrombin than usual. This mutation can be associated with both arterial and venous thromboembolism, mainly due to DVT and PE. Blood clots may also happen in unusual sites (such as the mesenteric or cerebral sinus vein), while coronary attacks or stroke represent two rare events. The mutation may be both homozygous and heterozygous.

Natural anticoagulants deficiencies

Deficiencies of protein C, protein S, or antithrombin III are important risk factors for venous thrombosis, often happening in young patients.17 The deficiencies of these natural anticoagulants are rather rare events in the general population and, combined, are found in less than 15% of all individuals. People born with a deficiency of one of the abnormal genes from either mother or father are heterozygous for this gene (more frequent). Conversely, patients can inherit the abnormal gene from both parents, and in this case, they are homozygous (more rarely occurs). As already affirmed, the prevalence of these deficiencies is associated with an increased risk of blood clots in veins, whereas that seems to play a little or no role in the blood clots in the arteries.18 Specifically, the mechanism of deficiencies of activated C/S proteins induces venous thrombosis. It consists in the inhibition of some co-factors (factor Va, factor VIIIa) acting as a risk factor in the coagulative cascade, while a deficiency in antithrombin III acts by inhibiting the serine proteases (factor III, X, XI, XII).

Elevation of factor VIII

Several studies have pointed out an association between elevated factor VIII and increased risk of
thrombophilia. This condition is due to the increased thrombin generation responsible for venous clots formation, often inducing thromboembolic events.\textsuperscript{19,20} Specifically, an excess of this factor resulted to be associated with a 6-fold increase of thrombo-embolic accidents.

**Dys-fibrinogenemia**

Defects of fibrinogenemia might be associated with increased risk for both hemorrhagic and thrombotic defects. Dysfibrinogenemia is a heterogeneous group of fibrinogen defects that may be congenital or acquired. Congenital dysfibrinogenemia is a relatively rare condition where an abnormality in the fibrin molecule results in defective clot formation. Although the exact mechanism of thrombosis is unknown, it may be related to increased fibrin formation and impaired fibrinolysis.\textsuperscript{21,22}

In Table 2, the degrees of risk of inherited thrombophilia are reported.

**Acquired thrombophilia**

Acquired thrombophilia is associated with an increased risk of VTE and arterial thrombosis. The leading causes of acquired thrombophilia are listed in Table 3.

Some of these are only described, such as antiphospholipid syndrome (APS), pregnancy, advanced age, heparin-induced thrombocytopenia (HIT), human immunodeficiency virus (HIV), some malignancies and oral contraceptive use.\textsuperscript{23}

**Antiphospholipid syndrome**

Antiphospholipid syndrome is the most common form of acquired thrombophilia, mostly present in young women. It can be defined as a hypercoagulable state characterized by recurrent by venous and/or arterial thrombosis and/or pregnancy complications of fetal loss, pre-eclampsia, or eclampsia. APS was firstly described in the setting of systemic lupus erythematosus, and subsequently recognized as an independent condition and in conjunction with a variety of other autoimmune infections, and malignant illnesses.\textsuperscript{24,25}

The spectrum of thrombosis in APS includes both venous and arterial events, even though the mechanisms of thrombosis are highly heterogeneous, multifactorial, and not yet defined.\textsuperscript{26} One hypothesis for arterial thrombosis is that antibodies associated with APS have procoagulant effects. Other proposed mechanisms include the following: activation of platelets to enhance endothelial adherence; activation of vascular endothelium, which facilitates the binding of platelets and monocytes; the reaction of antibodies to oxidized low-density lipoproteins, and interference with the coagulation cascade. The most common venous event is DVT and spontaneous abortion or unexplained fetal death in women.\textsuperscript{27}

**Pregnancy**

During pregnancy, clotting factors I, VII, VIII, IX, and X rise; protein S and fibrinolytic activity diminish, and resistance to activate protein C develops. These conditions increase the risk of venous and arterial thrombosis. The hypercoagulable state during pregnancy may increase the risk of thromboembolism.\textsuperscript{28}

**Advanced age**

Conventional risk factors in the elderly can be acquired (immobility, surgery, malignant diseases, hor-

Table 2. Risk degrees of thrombophilia related to the inherited factors.

<table>
<thead>
<tr>
<th>Degrees risk of inherited thrombophilia</th>
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<tbody>
<tr>
<td><strong>High risk of thrombophilia</strong></td>
</tr>
<tr>
<td>- Antithrombin deficiency</td>
</tr>
<tr>
<td>- Protein C deficiency</td>
</tr>
<tr>
<td>- Protein S deficiency</td>
</tr>
<tr>
<td><strong>Moderate risk of thrombophilia</strong></td>
</tr>
<tr>
<td>- Factor V Leiden</td>
</tr>
<tr>
<td>- Prothrombin gene mutation</td>
</tr>
<tr>
<td>- Factor VIII</td>
</tr>
<tr>
<td><strong>Low risk of thrombophilia</strong></td>
</tr>
<tr>
<td>- Factor IX</td>
</tr>
<tr>
<td>- Factor XI</td>
</tr>
<tr>
<td>- Hyperhomocysteinemia</td>
</tr>
</tbody>
</table>

Table 3. The most common causes favoring acquired thrombophilia.

<table>
<thead>
<tr>
<th>Causes of acquired thrombophilia</th>
</tr>
</thead>
<tbody>
<tr>
<td>*Antiphospholipid syndrome</td>
</tr>
<tr>
<td>*Pregnancy</td>
</tr>
<tr>
<td>*Advanced age</td>
</tr>
<tr>
<td>Trauma</td>
</tr>
<tr>
<td>Immobility</td>
</tr>
<tr>
<td>*Oral contraceptive pills</td>
</tr>
<tr>
<td>Surgery</td>
</tr>
<tr>
<td>*Some malignancies</td>
</tr>
<tr>
<td>*Heparin-induced thrombocytopenia</td>
</tr>
<tr>
<td>Myeloproliferative disorders</td>
</tr>
<tr>
<td>*Human immunodeficiency virus</td>
</tr>
</tbody>
</table>

*The different forms described in the text are marked with an asterisk.
mone use), whereas the most well-known genetic risk-factors are deficiencies of natural anticoagulants protein C, protein S, antithrombin and the factor V Leiden and prothrombin 20210A mutations. An evident and common coagulative disorder with advancing age is also an elevated Factor VIII, associated with an approximately 5-to-7 fold increased risk of VTE. But, this increase also depends on more elevated levels of Factor IX and Factor X.39

**Malignancy**

Multiple prothrombotic pathogenic processes are involved in some malignancies, including activation of the coagulation cascade and fibrinolytic system, procoagulant activation of tumor cells, and tumor cell interaction with blood cells and endothelium. The incidence of thrombotic events in cancer patients increased in comparison with the normal population. Among acquired risk factors for thrombophilia in cancer, the coagulative tendency is common, especially in lymphoproliferative diseases. Polycythemia vera is a chronic myeloproliferative syndrome in which the tendency of thrombus formation is increased. The occurrence of a central venous catheter in cancer-patients is another frequent cause of thrombosis.30 However, some inherited, thrombophilic factors also may act. Mainly, factor V Leiden or G20210A prothrombin gene mutation can be present and can increase the risk of venous thromboembolism by about 2-to-4 fold.

**Oral contraceptives**

Oral contraceptive pills in women can increase the risk of thrombosis.31 Among the hereditary types of thrombophilia, a resistance to activate protein C represents nearly 50% of cases, while in 15 to 20% a deficiency of antithrombin III, protein C, or protein S is found. In addition, antiphospholipid antibodies represent a significant cause of acquired thrombophilia. They also increase during treatment with oral contraceptives and therefore represent an adjunctive thrombotic risk-factor.32

**Heparin-induced thrombocytopenia**

Heparin-induced thrombocytopenia consists in the development of thrombocytopenia due to the administration of various forms of heparin, an anticoagulant. HIT predisposes to thrombosis because platelets release microparticles that activate thrombin, leading to thrombosis. Specifically, HIT is caused by the formation of abnormal antibodies that increases in individuals the risk of both venous and arterial thromboses.33 Diagnosis of HIT is based on clinical features, including a decline in the platelet count <150,000 µL or by 50% from baseline. These antibodies are directed toward antigens present on platelets. The decline in platelet count is seen between 5 and 14 days after inhibition of heparin. HIT occurs with unfractionated heparin more commonly than with low-molecular-weight heparin. The risk of HIT increased in surgical patients (orthopedic and cardiac surgery).34

**Human immunodeficiency virus**

Another disease that induces acquired thrombophilia is human immunodeficiency virus (HIV) although the exact mechanisms for thromboembolic risk are unknown. These include the presence of antiphospholipid-anticardioliopin antibodies, decreased activity of natural anticoagulants (especially protein S), and increase platelets’ activation. A hypercoagulable state, including myocardial infarction, emerges in HIV-infected patients.

It has been proposed that these changes are the result of disseminated intravascular coagulopathy and endothelial injury. It must be remembered that protein C and S deficiency and APS induce 6-fold increase in the risk of VTE. Other factors, such as an increased tissue factor expression may correlate with an increased risk of VTE.35,36

**Therapy**

**Antiplatelet therapy**

To prevent recurrent arterial thrombotic events related to a hypercoagulable state, predisposed individuals and patients that have already suffered from a first episode of arterial thrombosis must be treated with antiplatelet drugs indefinitely.37

**Anticoagulant therapy**

Current guidelines recommend that patients who have already suffered from a previous venous thromboembolism must receive anticoagulation therapy for a minimum of three months. Those who have a family history of venous thrombosis could receive anticoagulant treatment indefinitely. This therapy may also be continued indefinitely in patients who suffered from two or more spontaneous thrombosis, in those who had one spontaneous venous thrombosis at an unusual site, and in patients who suffered from one episode in the presence of single and/or multiple genetic coagulative defects.38,39

The anticoagulant therapy for the prevention of recurrent VTE must be started through an initial treatment with low-molecular-weight heparin followed by an oral VKA for long-term anticoagulation. Index of normalized ratio (INR) must be routinely defined, and it is crucial to perform does-adjustments in order to maintain the INR value between 2.5-3.0. INR must be maintained persistently below these values in patients...
with an increased risk of bleeding. These INR values must also be maintained in elderly patients, in patients with renal impairment, in those having low body weight, and a history of gastrointestinal bleeding.

Direct oral anticoagulants

But, in recent years, direct oral anticoagulants, and oral non-vitamin K anticoagulants, are increasingly used instead of VKAs in the prevention of recurrent VTE including that associated with thrombophilia. DOACs offer numerous advantages over VKAs, such as a predictable response, fewer drugs, and food interactions and no need for laboratory monitoring of the INR or other coagulation tests.

Dabigatran, rivaroxaban, apixaban, and edoxaban have been developed as a treatment alternative to heparins/VKAs. Several clinical trials found that new oral anticoagulants (NOACs) have non-inferior efficacy than heparins/VKA for both acute treatment and prevention of recurrent VTE. NOACs were till now approved for the treatment of acute DVT, PE, and the prevention of recurrent VTE. But, their use in these pathologic findings are still conflicting.

Little is known about the efficacy and safety of DOACs in patients with venous thromboembolism in a thrombophilic pattern. Referring to hereditary thrombophilia alone, Crowther et al. classified that into two main groups, following the risk level of thrombosis.

Group 1 (also called high-risk thrombophilias), caused by hereditary deficiencies of natural anticoagulants and include protein C, protein S, and antithrombin deficiencies.

Group 2 (also called minor or low-risk thrombophilias), associated with increased levels of the coagulation factors and include Factor V Leiden, G20210A prothrombin mutations, an elevated level of factor VIII, IX, XI and HHcy. The most common thrombophilias are induced by Factor V Leiden and G20210A prothrombin with a prevalent heterozygosity (low-risk factors). In these patients, the results were favorable. On the contrary, the results obtained in protein C and S deficiency remain uncertain. APS is an acquired thrombophilic disorder associated with increased risk for venous and arterial thrombosis. DOACs treatment in APS supplied conflicting results. In fact, in RAPS (Rivaroxaban in Anti-Phospholipid Study), no thrombotic or significant bleeding was found after 210 days of follow-up, in the group that received Rivaroxaban in comparison with the patients treated with warfarin. Contrarily, the TRAPS (Trial of Rivaroxaban in Anti-Phospholipid Syndrome) study, designed to evaluate the efficacy of Rivaroxaban versus warfarin in APS patients, was prematurely stopped because an interim analysis showed a higher rate of thromboembolic and frequent bleeding events among the patients receiving DOAC (rivaroxaban).

Finally, Elsebaie et al. in a systematic review of 10 studies performed on thrombophilic patients, affirmed that rates of VTE recurrence and bleeding events were comparable in patients with venous thrombophilias receiving both VKA or DOACs.

Conclusive remarks

In conclusion, little is still known about the efficacy and safety of DOACs in patients with different forms of inherited or acquired thrombophilia. For this reason, the unlimited use of DOACs in these patients is now controversial. However, some evidence indicates that DOACs are a valuable and promising therapeutic option in therapy of VTE and its complications (TVP, PE), as well as in the secondary prevention of thrombotic events in individuals suffering from some thrombophilias. Nevertheless, other experiences performed in a wide range are requested to demonstrate their efficacy in all patients with thrombophilia clearly. However, the observations reported seem to support that DOACs are useful in low-risk thrombophilia. Their efficacy and safety appear to be insufficient in thrombophilias by protein C, protein S, and antithrombin deficiency (high-risk thrombophilias), while the use of these drugs is uncertain in APS patients and those with a history of arterial thrombosis.

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