Homozygous Carrier of Prothrombin G20210A Mutation with Massive Pulmonary Embolism and His Family: Gender Differences of Susceptibility to Mutation

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INTRODUCTION
Prothrombin 20210 G>A mutation is the second most frequent inherited factor increasing the risk for developing venous thromboembolism (VTE). The risk for VTE in homozygous carriers of this mutation is not well studied because of their rarity are rare. We report a case of a homozygous carrier of prothrombin mutation: a young man with massive pulmonary embolism, and his family - an asymptomatic homozygous sister, heterozygous parents with asymptomatic mother, and father with history of deep venous thrombosis (DVT). To our knowledge, this is the first reported case of homozygous prothrombin mutation carriers in Bulgaria and the other Balkan countries. We conclude that the homozygous prothrombin mutation creates predisposition for VTE that can manifest or not depending on additional factors, one of which could be male gender.

CASE REPORT
A 25-year-old Caucasian man, never smoker, a computer specialist, presented with progressive dyspnea on exertion and at rest, palpitation, right side chest pain, and low grade fever (37.5°C). The symptoms gradually increased in a week. No external provoking factors were detected; the patient had no history of previous VTE.

Findings: tachypnea (30 per min); oral cyanosis; oxygen saturation - 93% at room air; slight pleural rubbing in the right bottom chest; heart rate - 115 beats per minute, blood pressure - 100/80.

Laboratory: prothrombin time - 14.3 sec (reference values 10 - 14 sec); D-dimer > 35 μg/ml. Transthoracic echocardiography: D-shaped left heart chamber with dyskinetic septum; dilated right chamber - 43 mm with dyskinetic free wall; 1st degree tricuspid valve regurgitation, calculated pulmonary artery systolic pressure - 41 mmHg; TAPSE - 15 mm. Computer tomographic pulmonary angiography showed massive bilateral pulmonary embolism with thrombus riding bifurcation of the truncus pulmonalis and occluding a big part of...
the cavities of the main right and left pulmonary arteries and thrombi in all segmental branches bilaterally (Fig. 1).

Figure 1. Thrombi in lobar and segmental branches of a. pulmonalis. No peripheral vascularization.

Laboratory examinations for inherited and acquired thrombophilia (thrombophilic examination) found (Fig. 2): homozgyous prothrombin 20210AA mutation; heterozygous carrier of PAI-1 4G/5G, A1298C and C677T mutations; absence of Leiden mutation, normal values of antithrombin III (AT III) - 88.5%, protein C (Pr. C) - 114.8% and protein S (Pr. S) - 130%. Antiphospholipid antibodies were not found. Homocysteine - 13.03 μmol/l (N 0 -15).

We examined the members of his family: both parents and sister. All live together, share the same lifestyle, work at computers and never smoked.

The sister: 24 years old, doesn’t take contraceptives, nulligravida, no history of operations, fully asymptomatic. The thrombophilia examination showed her to be a homozgyous carrier of prothrombin 20210AA mutation and heterozygous carrier of A1298C and C677T variants of MTHFR gene, normal carrier of PAI-1 5G/5G gene and absence of Leiden mutation (Fig. 2). ATIII, Pr. C and Pr. S were within normal ranges. Antiphospholipid antibodies were not found. Prothrombin time was 13.0 sec, homocysteine - 14.64 μmol/l.

The mother: a 52-year-old woman, multigravida (2 pregnancies) and uneventful hysterectomy at 50. The thrombophilia examination showed her to be a heterozygous carrier of prothrombin G20120 G›A and PAI-1 4G/5G mutations, homozygous carrier of mutant 1298CC and normal of 677CC variants of MTHFR gene, absence of Leiden mutation (Fig. 3). AT III, Pr. C and Pr. S levels were within normal ranges. Antiphospholipid antibodies were negative. Prothrombin time was 13.4 sec, homocysteine - 10.13 μmol/l.

The father: a 51-year-old man with history of unprovoked DVT at the age of 47. He had discontinued treatment with oral anticoagulant 3 years before. No recurrence of VTE. The thrombophilia examination showed him to be a heterozygous carrier of prothrombin G20120A, PAI-1 4G/5G, A1298C and C677T mutations and absence of Leiden mutation (Fig. 3), AT III, Pr. C and Pr. S were in normal ranges. Antiphospholipid antibodies were negative. Prothrombin time was 14.0 sec., homocysteine - 17.63 μmol/l.

DISCUSSION

The patient we report is a carrier of several inherited abnormalities. Except for the prothrombin mutation, they are all considered weak thrombophilic factors. That is why we suggest that prothrombin 20210AA mutation in this patient is the leading cause of the massive pulmonary embolism. But his sister is the same age, has similar inherited abnormalities and is
asymptomatic. Is this by chance or not? The same holds for their parents: the father and the mother are the same age and have similar abnormalities, but only the father had a VTE event, although the mother had precipitating factors such as pregnancies and operation. We hypothesize that the male gender is more susceptible to inherited thrombophilic factors compared to women in equal other circumstances. Our contention is consistent with strong evidence for twice as high risk of recurrence of VTE in men as in women.8

CONCLUSION

The homozygous prothrombin 20210G ›A mutation creates predisposition to thrombotic events that can manifest depending on additional factors. We hypothesize that the male gender is one of these factors. Our two cases can be added to the small pool of reported cases of prothrombin 20210AA mutation carriers worldwide.

REFERENCES


Гомозиготный носитель мутации гена протромбина G20210A с массивной тромбоэмболией легочной артерии и его семья: гендерные различия в восприимчивости к мутации

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Мутация гена протромбина G20210A является вторым наиболее частым наследственным фактором, увеличивающим риск развития венозной тромбоэмболии (ВТЭ). Риск ВТЭ у гомозиготных носителей мутации не очень хорошо изучен из-за их небольшого количества. В данной статье мы сообщаем о случае гомозиготного носителя мутации протромбина - молодой человек с массивной тромбоэмболией легочной артерии и его семья - бессимптомная гомозиготная сестра, гетерозиготные родители с бессимптомной матерью, а отец с историей тромбоза глубоких вен (ТГВ). Насколько нам известно, это первый случай гомозиготных носителей мутации протромбина в Болгарии и других балканских странах. Мы пришли к выводу, что гомозиготная мутация протромбина создает предрасположенность к ВТЭ, которая может проявляться или нет, в зависимости от дополнительных факторов, одним из которых может быть принадлежность к мужскому полу.

Ключевые слова: наследственные тромбофилии, мутация гена протромбина G20210A, венозная тромбоэмболия