**Sex-specific aspects of venous thromboembolism: What is new and what is next?**

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Abstract
Men seem to have a higher intrinsic risk of venous thromboembolism (VTE) than women, regardless of age. To date, this difference has not been explained. By integrating state-of-the-art research presented at the International Society on Thrombosis and Haemostasis Congress of 2021 with the available literature, we address potential explanations for this intriguing risk difference between men and women. We discuss the role of exogenous and endogenous sex hormones as the most important known sex-specific determinants of VTE risk. In addition, we highlight clues on the role of sex hormones and VTE risk from clinical scenarios such as pregnancy and the polycystic ovary syndrome. Furthermore, we address new potential sex-specific risk factors and unanswered research questions, which could provide more insight in the intrinsic risk difference between men and women, such as body height and differences in body fat distribution, leading to dysregulation of metabolism and inflammation.

**KEYWORDS**
body fat distribution, hormones, metabolism, sex, venous thromboembolism

Essentials
- Men have an, unexplained, higher intrinsic venous thromboembolism (VTE) risk than women.
- Body height does not fully explain the sex-specific VTE risk difference.
- Exogenous estrogens increase VTE risk, for endogenous sex hormones this is largely unknown.
- Body fat distribution, metabolism and inflammation are of interest for future studies.

1 | **SEX-SPECIFIC RISK OF VENOUS THROMBOEMBOLISM: IS THERE A DIFFERENCE?**

Over the past several decades, several important sex-specific aspects of risk of venous thromboembolism (VTE) have been established. Nearly universally, studies on the sex-specific incidence of first VTE across age categories report the same pattern.¹⁻⁵ Most studies report a 2- to 3-fold higher incidence of first VTE in women during their fertile lifespan compared with men of similar age groups.¹⁻⁵ After age approximately 50 years, the observed incidence is slightly higher in men than in women. After the age of 70 years, the observed incidence increases steeply in both sexes and, in some studies, the incidence in women is again higher than in men¹⁴,⁵ (see...
Figure 1 for sex-specific incidence rates from large population based studies with corresponding incidence rate ratios per age groups. As a result, on average, the risk of VTE is considered more or less equal in both sexes. This observation is remarkably in contrast to the risk of a recurrent event, which has consistently been shown to be twice as high in men than in women.\(^6\)\(^-\)\(^9\)

Different explanations have been proposed for the observed variation in incidence of both first and recurrent VTEs across the lifespan of women and men. The observed higher incidence in women during the fertile lifespan is likely explained by the presence of transient risk factors related to reproductive factors such as use of hormonal contraceptives and pregnancy. When these women-specific risk factors during premenopausal ages are taken into account, it appears that the incidence of a first VTE is also two times higher in men in the younger age group, and accordingly fits with the increased recurrence risk as observed in men.\(^5\)\(^9\) Hence, the intrinsic risk of VTE (i.e., the risk in the absence of extrinsic risk factors) appears to be higher in men than in women, at least up to the age of 70 years.\(^9\)

For the very elderly (>age 70 years), fewer data are available. In a recent simulation study, in which the competing risk of death was taken into account, the estimated lifetime risk of first VTE (i.e., at

![Figure 1](image-url)

**Figure 1** Left figures: observed incidence of first venous thromboembolism (VTE) by age group separately for women (red) and men (blue). Right figures: corresponding incidence rate ratio (IRR) of the risk of VTE comparing men with women by age groups. An IRR of 1 indicates no difference, an IRR > 1 indicates a higher incidence in women, an IRR of < 1 indicates a higher incidence in men. Based on available published data from large population-based studies that reported the incidence of first VTE separately for men and women with small (5-year) age categories available. (A) Naess et al.,\(^5\) (B) Arnesen et al.,\(^5\) (C) Kort et al\(^5\)
100 years of age) was slightly higher in women than in men (9.3% vs. 8.1%); however, 50% of all first VTE events in women occur after the age of 70 years. Hence, there seems to be a “catch-up” effect in women that is possibly explained by competing risks (e.g., death from other causes in men).5

Taken together, the observations from these studies demonstrate there is a substantial lifetime risk of VTE in both women and men and suggest that men have an intrinsic increased risk of VTE compared with women (see Figure 2 for a schematic illustration of this hypothesis). Intriguingly, to date, the mechanism for this higher intrinsic risk in men (or protective mechanism responsible for the lower intrinsic risk in women) has not been identified. In this International Society on Thrombosis and Haemostasis (ISTH) Congress of 2021 state-of-the-art review we will discuss the role of exogenous and endogenous sex hormones as the most important known sex-specific determinants of VTE risk. In addition, we will address new potential sex-specific risk factors and unanswered research questions, which may explain the intrinsic risk difference between men and women, such as body height and body fat distribution with subsequent changes in metabolism and inflammation. A summary of the hypothetical mechanisms that could contribute to the higher intrinsic VTE risk in men is provided in Table 1.

1.1 | Role of genes and environment

The role of genes and environmental factors as a potential explanation for the higher VTE risk in men has been discussed in detail previously.10 The most important (based on their prevalence and risk size) genetic VTE risk factors that have been identified include the Factor V Leiden and prothrombin G20201A gene mutation, non-O blood group and an inherited deficiency of protein C, protein S, and antithrombin.11,12 However, these mutations are not located on the sex chromosomes and are thus likely not distributed differently between men and women. This was confirmed in the MEGA case control study, in which there was no difference in VTE risk size of these mutations between men and women.10 The genes that code for coagulation factor (F) VIII and IX, of which higher levels are both associated with VTE risk, are located on the X chromosome and thus could result in sex-specific differences of VTE risk.13,14 However, in a large population-based study (2533 men and 2440 women), FVIII levels were slightly higher in women than in men.15 In a smaller study among healthy controls (272 women, 201 men) there was also no difference in factor IX levels.16 Hence, sex-specific differences in factors VIII and IX levels do not seem to contribute the intrinsic sex-specific VTE risk differences. Also, Y chromosomal haplogroup variation in men was not associated with differences in first or recurrent VTE risk, and it was also considered unlikely that chromosome Y variation can explain the difference in VTE risk between women and men.17

Regarding the role of the environment, the incidence of VTE is continuously subject to changes in prevalence of risk factors in a population. Environmental factors could also play a role in the observed sex-specific incidence of VTE; for example, if men would more frequently suffer from trauma and subsequent immobilization than women, this would influence the incidence of VTE. However, in previous studies, no differences were found for the prevalence of acquired strong VTE risk factors between men and women such as surgery, immobilization, malignancy, and chronic disease.10,19 However, more subtle cultural and/or lifestyle differences between men and women could still contribute. For example, on average, women may tend to choose professions associated with more physical activity such as in education or health care, whereas men on average may have more seated professions, such as driver or office worker.20 Even within similar job occupations, men were found to have more seated hours.21 On a population level, more seated hours on a daily basis may result in a higher VTE risk, which might play a role in the underlying mechanism for the observed VTE risk difference between men and women.

2 | INFLUENCE OF SEX HORMONES

2.1 | Exogenous use of sex hormones

The role of hormonal contraceptives and hormone therapy as VTE risk factors in women has been firmly established.22 The associated VTE risk is dependent on both dose and type of estrogen and progestogen and also on mode of delivery.23,24 In men, however, studies on exogenous testosterone use and VTE risk have reported conflicting results, and testosterone use may be associated with a modestly increased VTE risk.25-28 Testosterone (as opposed to hormonal contraceptives in women) is typically only used in case of concomitant disease that can be associated with an increased VTE risk in and of itself. Examples of such clinical scenarios include hypogonadism or testosterone replacement therapy after cancer treatment.25,29 In addition, use of estrogen-like therapies for management of prostate cancer are also associated with an increased risk of VTE.30

Important clues on the contribution of exogenous sex hormones to risk of VTE, independent of other morbidity and assigned sex at birth, can be derived from hormone use in the transgender population. Transwomen (persons born with male sex and female gender identity) who use gender-affirming hormone therapy (typically estrogen with antiandrogens) seem to have an increased risk of both venous and arterial thrombotic events compared with control cismen (persons born with male sex assigned at birth and a male gender identity).31-35 When comparing coagulation profiles before and after start of hormone therapy in these persons, the coagulation profiles were overall more procoagulant after the start of hormone therapy.36

On the risk of venous and arterial thrombotic events in transmen (persons born with female sex assigned and male gender identity) who use testosterone as gender-affirming hormone therapy, only few data are available.31,33-35 In the available studies, there does not seem to be an increased risk of thrombotic events compared with control ciswomen (persons born with female sex assigned at birth.
In line with this, the overall coagulation profiles of transmen after start of testosterone therapy did not appear more procoagulant than before start of therapy.\(^3\) Taken together, these observations suggest that use of exogenous estrogens (frequently combined with progestogens) results in procoagulant changes and increased VTE risk in both women and persons with male sex assigned at birth. Procoagulant changes in response to estrogen exposure seem of evolutionary benefit because these could result in decreased blood loss during labor.\(^3\) Interestingly, the responsible biological mechanism seems in place also in persons with male sex assigned at birth. In contrast, the use of testosterone does not appear to have a clear prothrombotic effect.

### 2.2 Influence of endogenous sex hormones

Few data are available on endogenous sex-hormone levels and risk of VTE in either women or men. In a large Danish cohort study, there was no apparent association between endogenous estrogen and testosterone levels and VTE risk in both women and persons with male sex assigned at birth. Procoagulant changes in response to estrogen exposure seem of evolutionary benefit because these could result in decreased blood loss during labor.\(^3\) Interestingly, the responsible biological mechanism seems in place also in persons with male sex assigned at birth. In contrast, the use of testosterone does not appear to have a clear prothrombotic effect.

**TABLE 1** Hypothetical mechanisms for the higher intrinsic VTE risk in men

| Risk factor | Hypothetical mechanisms | Related literature |
|-------------|-------------------------|-------------------|
| Likely contributes | | |
| Average higher body height and leg length in men | Potentially resulting in higher risk of stasis | \([19,52]\) |
| Of interest for further exploration | | |
| Cultural and/or lifestyle differences resulting in VTE risk factor differences | If men would on average have more seated hours during the day because of differences in job occupation, leading to more stasis | \([20,21]\) |
| Androgenicity | Mechanistic clues from the polycystic ovary syndrome population. Here, androgenicity and visceral and liver fat deposition seems associated with metabolic and inflammatory changes, which could result in higher VTE risk | \([44-47,66]\) |
| Body fat distribution (i.e., more visceral fat deposition) | Differences in body fat distribution (as between the sexes) are associated with metabolic and inflammatory changes, which could result in higher VTE risk | \([63,65]\) |

Abbreviation: VTE; venous thromboembolism
indicate hyperandrogenism (being one of the hallmarks of PCOS). In several studies, androgen excess in women has been associated with both increased levels of procoagulant and inflammatory markers. 46-49

On the whole, outside pregnancy and PCOS, little is known on the relevance of endogenous sex hormone levels in the context of VTE risk. The available data suggest a modestly increased VTE risk with higher levels of estrogens and especially with hyperandrogenism in young women. The latter finding is of interest in light of the intrinsic higher VTE risk in men and does not seem to be explained only by testosterone levels as discussed in previous sections. Hyperandrogenism in young women (with PCOS) is associated with dysregulation of metabolism and inflammation. 50 The question arises whether the responsible mechanism is also present in men and whether this could explain the higher intrinsic VTE risk that is observed in men.

3 | BODY HEIGHT AND SEX-SPECIFIC VTE RISK

One of the most evident physical differences between men and women is body height. Men are on average taller than women, with a global average height for men of 171 cm and 159 cm for women (aged 18 years in 2014). 51 Body height has been identified as a risk factor for VTE both in men and women, where especially the tallest persons are at higher risk. 19,52 In addition, a sedentary lifestyle and long-haul flights are risk factors that increase the risk even further in tall persons. 52,53 However, taller height does not seem to completely explain the risk difference. In men and women with a first VTE and similar body height, the recurrence risk was still 2-fold higher in men for every height (Figure 3 depicts the results as observed in the original study). 52 Accordingly, adjustment for body height only slightly attenuated the difference in recurrence risk between men and women. 52 In line with these observations, in a recent abstract reporting on a population-based study from Tromsø on the same matter, body height was found to be a risk factor for first VTE in both women and men. 54 However, in this study, adjustment for body height did attenuate the risk difference for a first VTE between men and women up to the age of 70 years, whereas this risk difference in persons older than age 70 years did not change. 54

In short, body height is a well-established VTE risk factor that could partially explain the observed higher VTE risk in men; however, this does not seem to be the full explanation.

4 | OBESITY, VISCERAL AND LIVER FAT, AND THEIR EFFECTS ON METABOLISM AND SUBSEQUENT INFLAMMATION IN RELATION TO COAGULATION AND VTE RISK

Over the past several decades, important insights have been obtained in the pathogenesis of arterial or also called atherosclerotic thrombotic events. Inflammation and dysregulation of metabolism seem intertwined, playing a key role in the pathogenesis, and have received much attention. 55 Although classically VTE and arterial thrombotic events have been considered separate diseases, it has been established that they share several risk factors such as inflammation. 56-58 Although mechanisms are not yet understood, inflammation has been well recognized as a risk factor for VTE. Especially overwhelming inflammatory states are strongly associated with an increased VTE risk. Well-known examples include sepsis-induced disseminated intravascular coagulation, the antiphospholipid syndrome, inflammatory bowel disease, and the recently described COVID-19-associated coagulopathy is a timely issue. 59,60

Whether low-grade inflammation as is observed in obesity and metabolic syndrome is of relevance in VTE pathogenesis, as in the case of arterial thrombotic disease, is largely unknown and has not been widely studied. High body mass index and obesity are well-established VTE risk factors. 61 Its important components, total body fat and amount of visceral fat, have been associated with increasing procoagulant levels. 62,63 In addition, higher liver triglyceride content (a marker of fatty liver disease) has been found to be associated with increases of coagulation factor IX, beyond body fat and visceral adipose tissue. 63 In another study, the prevalence of nonalcoholic fatty liver disease was approximately three times higher among VTE cases than in controls. 64 Moreover, important differences in body fat distribution between men and women have been described, where women have more body fat overall, yet in men the proportion of visceral and liver fat is higher, 65 which also seems the case in women with PCOS. 66-68 However, it is not known whether these differences contribute to the difference in VTE risk.

Further studies are needed investigating obesity, visceral and liver fat, and their effects on metabolism and subsequent inflammation in relation to coagulation and VTE risk. Investigations assessing these mechanisms and the significance of sex differences, in light

![FIGURE 3](image-url) Incidence rate of recurrent venous thromboembolism per body height category separately for women (n = 2315) and men (n = 1949) in the study by Flinterman et al. 52 Bars indicate 95% confidence intervals. This is from the figure in the original study, printed with permission from John Wiley and Sons.
of the higher risk in men are of interest. In addition, given the high prevalence of PCOS and increasing prevalence of obesity and disorders of fatty liver, these mechanisms are definitely of interest, and a better understanding could result in improved prevention or management strategies of VTE.

5 | ISTH CONGRESS REPORT

During the ISTH Congress of 2021, several interesting abstracts were presented that relate to the focus of this state-of-the-art review. In line with previous studies, Oakes et al. analyzed data from a large study population of transgender persons using testosterone (n = 923) and showed that although erythrocytosis as a consequence was common (up to one in five persons), the risk of thromboembolic events seemed low.\(^6\)

Chulkov and colleagues highlighted the important relationship between obesity and coagulation in young adults. Higher levels of leptin were associated with higher levels of fibrinogen and plasminogen activator inhibitor-1 in persons with an unhealthy metabolic profile.\(^7\) In a similar scope, the close relation between coagulation and inflammation was further evaluated by Pallares Robles et al., who showed an interesting interaction between FXI activity and thrombo-inflammation and lipid metabolism.\(^8\)

The study by Houghton and collaborators provides further insight in the mechanism that may explain increased risk of VTE in taller persons. They showed that reduced calf pump function as a measure of stasis was associated with the risk of ipsilateral deep vein thrombosis.\(^9\)

6 | SUMMARY AND FUTURE DIRECTIONS

In summary, men have a higher intrinsic VTE risk than women, regardless of age. To date, this difference has not been explained. Body height is a well-established risk factor for VTE but only seems to contribute partially to the observed higher VTE risk in men. Exogenous use of estrogens (combined with progestogens) increases the risk of VTE both in women and transwomen. Exogenous use of testosterone is associated with a slightly increased VTE risk in men and more robust data on use in transwomen are needed.

Overall, little is known on the relevance of levels of endogenous sex hormones and (sex-specific) VTE risk. Inflammation and dysregulation of metabolism are important determinants of arterial thrombotic risk and seem also of interest with respect to underlying mechanisms in VTE. Populations at higher risk of arterial thrombotic disease in which dysregulation of endogenous hormones and metabolism are key features, such as women with PCOS and subjects with differences in body fat distribution including liver fat disposition, form interesting target groups for future research. Overall better understanding of these mechanisms could result in improved (sex-specific) prevention and management strategies of VTE.

AUTHOR CONTRIBUTIONS

Luuk J. J. Scheres, Astrid van Hylckama Vlieg, and Suzanne C. Cannegieter wrote the manuscript and approved the final version.

RELATIONSHIP DISCLOSURE

The authors report no conflict of interest.

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REFERENCES

1. Næss IA, Christiansen SC, Romundstad P, Cannegieter SC, Rosendaal FR, Hammerstrøm J. Incidence and mortality of venous thrombosis: a population-based study. \textit{J Thromb Haemost}. 2007;5:692-699.
2. Alotaibi GS, Wu C, Senthilselvan A, McMurtry MS. Secular trends in incidence and mortality of acute venous thrombembolism: the AB-VTE population-based study. \textit{Am J Medicine}. 2016;129(8):879.e19-25.
3. Arshad N, Isaksen T, Hansen J-B, Brækkan SK. Time trends in incidence rates of venous thromboembolism in a large cohort recruited from the general population. \textit{Eur J Epidemiol}. 2017;32:299-305.
4. Kort D, Rein N, Meer FJM, et al. Relationship between neighborhood socioeconomic status and venous thromboembolism: results from a population-based study. \textit{J Thromb Haemost}. 2017;15:2352-2360.
5. Arnesen CAL, Veres K, Horváth-Puhó E, Hansen J-B, Sørensen HT, Brækkan SK. Estimated lifetime risk of venous thromboembolism in men and women in a Danish nationwide cohort: impact of competing risk of death. \textit{Eur J Epidemiol}. 2022;37(2):195-203.
6. Kyrie PA, Minar E, Bliauzycky C, Hirschl M, Weltermann A, Eichinger S. The risk of recurrent venous thromboembolism in men and women. \textit{N Engl J Med}. 2004;350:2558-2563.
7. Baglin T, Luddington R, Brown K, Baglin C. High risk of recurrent venous thromboembolism in men. \textit{J Thromb Haemost}. 2004;2:2152-2155.
8. Christiansen SC, Lijfering WM, Helmerhorst FM, Rosendaal FR, Cannegieter SC. Sex difference in risk of recurrent venous thrombosis and the risk profile for a second event. \textit{J Thromb Haemost}. 2010;8:2159-2168.
9. Roach REJ, Lijfering WM, Rosendaal FR, Cannegieter SC, le Cessie S. Sex difference in risk of second but not of first venous thrombosis. \textit{Circulation}. 2014;129:51-56.
10. Roach REJ, Cannegieter SC, Lijfering WM. Differential risks in men and women for first and recurrent venous thrombosis: the role of genes and environment. \textit{J Thromb Haemost}. 2014;12:1593-1600.
11. Martinelli I, Stefano VD, Mannucci PM. Inherited risk factors for venous thromboembolism. \textit{Nat Rev Cardiol}. 2014;11:140-156.
12. Zöller B, Svensson PJ, Dahlbäck B, Lind-Halldén C, Halldén C, Elf J. Genetic risk factors for venous thromboembolism. \textit{Expert Rev Hematol}. 2020;13:1-11.
13. Koster T, Vandebroucke JP, Rosendaal FR, Briët E, Rosendaal FR, Blann AD. Role of clotting factor VIII in effect of von Willebrand factor on occurrence of deep-vein thrombosis. \textit{Lancet}. 1995;345:152-155.
14. van Hylckama VA, van der Linden IK, Bertina RM, Rosendaal FR. High levels of factor IX increase the risk of venous thrombosis. \textit{Blood}. 2000;95:3678-3682.
15. Graw J, Brackmann H-H, Oldenburg J, Schnepfennig R, Spannagl M, Schwaab R. Haemophilia A: from mutation analysis to new therapies. *Nat Rev Genet.* 2005;6:488-501.

16. Hermanns MI, Grossmann V, Spronk HMH, et al. Distribution, genetic and cardiovascular determinants of FVIII:c – data from the population-based Gutenberg Health Study. *Int J Cardiol.* 2015;187:166-174.

17. Haan HG, van Hylckama VA, Gaag KJ, Knijff P, Rosendaal FR. Male-specific risk of first and recurrent venous thrombosis: a phylogenetic analysis of the Y chromosome. *J Thromb Haemost.* 2016;14:1971-1977.

18. Scheres LJ, Lijfering WM, Cannegieter SC. Current and future burden of venous thrombosis: not simply predictable. *Res Pract Thromb Haemost.* 2018;2:199-208.

19. Braekkan SK, Borch KH, Mathiesen EB, Njølstad I, Wilsgaard T, Hansen J-B. Body height and risk of venous thromboembolism: the Tromsø study. *Am J Epidemiol.* 2010;171:1109-1115.

20. Eurofound and European Commission Joint Research Centre. European jobs monitor 2021: gender gaps and the employment structure. Eurofound and European Commission Joint Research Centre. *European Jobs Monitor series, Publications Office of the European Union* 2021: 2021.

21. Johansson E, Mathiassen SE, Rasmusse CL, Hallman DM. Sitting, standing and moving during work and leisure among male and female office workers of different age: a compositional data analysis. *BMC Public Health.* 2020;20:826.

22. Middeldorp S. Thrombosis in women: what are the knowledge gaps in 2013? *J Thromb Haemost.* 2013;11:180-191.

23. van Hylckama VA, Helmerhorst FM, Vandenbroucke JP, Doggen CJM, Rosendaal FR. The venous thrombotic risk of oral contraceptives, effects of oestrogen dose and progestogen type: results of the MEGA case-control study. *BMJ.* 2009;339:b2921.

24. Bistervels IM, Scheres LJ, Hamulyák EN, Middeldorp S. Sex matters: practice 5Ps when treating young women with venous thromboembolism. *J Thromb Haemost.* 2019;17:1417-1429.

25. Martinez C, Suissa S, Rietbrock S, et al. Testosterone treatment and risk of venous thromboembolism: population based case-control study. *BMJ.* 2016;355:i5968.

26. Sharma R, Oni OA, Chen G, et al. Association between testosterone replacement therapy and the incidence of DVT and pulmonary embolism A retrospective cohort study of the veterans administration database. *Chest.* 2016;150:563-571.

27. Shores MM, Walsh TJ, Korpak A, et al. Association between testosterone treatment and risk of incident cardiovascular events among US male veterans with low testosterone levels and multiple medical comorbidities. *Am Heart Assoc.* 2021;10:e020562.

28. Houghton DE, Alsawas M, Barrioneuvo P, et al. Testosterone therapy and venous thromboembolism: a systematic review and meta-analysis. *Thromb Res.* 2018;172:94-103.

29. Blondon M, Righini M. Excess risk of venous thromboembolism associated with androgen deprivation therapy. *Eur Urol.* 2016;70:62-63.

30. Kil-Krörn AJ, Ym H, Tagalakis V, Aprikian A, Azoulay L. Androgen deprivation therapy for prostate cancer and the risk of venous thromboembolism. *Eur Urol.* 2016;70:56-61.

31. Nota NM, Wiepjes CM, de Blok CJM, Gooren LJJ, Kreukels BPC, den Heijer M. The occurrence of acute cardiovascular events in transgender individuals receiving hormone therapy: results from a large cohort study. *Circulation.* 2019;139:1461-1462.

32. Connelly PJ, Free EM, Perry C, et al. Gender-affirming hormone therapy, vascular health and cardiovascular disease in transgender adults. *Hypertension.* 1979;2019(74):1266-1274.

33. Getahun D, Nash R, Flanders WD, et al. Cross-sex hormones and acute cardiovascular events in transgender persons: a cohort study. *Ann Intern Med.* 2018;169:205.

34. Streed CG, Harfouch O, Marvel F, Blumenthal RS, Martin SS, Mukherjee M. Cardiovascular disease among transgender adults receiving hormone therapy: a narrative review. *Ann Intern Med.* 2017;167:256.

35. Connors JM, Middeldorp S. Transgender patients and the role of the coagulation clinician. *J Thromb Haemost.* 2019;17:1790-1797.

36. Scheres LJ, Selier NLD, Nota NM, Diemen JJK, Cannegieter SC, Heijer M. Effect of gender-affirming hormone use on coagulation profiles in transmen and transwomen. *J Thromb Haemost.* 2021;19(4):1029-1037.

37. Bremme KA. Haemostatic changes in pregnancy. *Best Pract Res Clin Haematol.* 2003;16:153-168.

38. Holmeaard PN, Nordestgaard BG, Schnoor P, Tybjerg-Hansen A, Benn M. Endogenous sex hormones and risk of venous thromboembolism in women and men. *J Thromb Haemost.* 2014;12:297-305.

39. Roetker N, MacLehose R, Hoogeveen R, et al. Prospective study of endogenous hormones and incidence of venous thromboembolism: the atherosclerosis risk in communities study. *Thromb Haemost.* 2018;118:1940-1950.

40. Scheres LJ, van Hylckama VA, Ballieux BEPB, et al. Endogenous sex hormones and risk of venous thromboembolism in young women. *J Thromb Haemost.* 2019;17:1297-1304.

41. Barco S, Nijkeuter M, Middeldorp S. Pregnancy and venous thromboembolism. *Semin Thromb Hemost.* 2013;39:549-558.

42. Dumesic DA, Oberfield SE, Stener-Victorin E, Marshall JC, Laven JS, Legro RS. Scientific statement on the diagnostic criteria, epidemiology, pathophysiology, and molecular genetics of polycystic ovary syndrome. *Endocr Rev.* 2015;36:487-525.

43. McCartney CR, Marshall JC. Polycystic ovary syndrome. *N Engl J Med.* 2016;375:54-64.

44. Bird ST, Hartzema AG, Brophy JM, Etminan M, Delaney JAC. Risk of venous thromboembolism in women with polycystic ovary syndrome: a population-based matched cohort analysis. *CMAJ.* 2013;185:E115-E120.

45. Glintborg D, Rubin KH, Nybo M, Abrahamsen B, Andersen M. Morbidity and medicine prescriptions in a nationwide Danish population of patients diagnosed with polycystic ovary syndrome. *Eur J Endocrinol.* 2015;172:627-638.

46. Kelly CCJ, Lyall H, Petrie JR, Gould GW, Connell JMC, Sattar N. Low grade chronic inflammation in women with polycystic ovarian syndrome. *J Clin Endocrinol Metabolism.* 2001;86:2453-2455.

47. Toulis KA, Goulis DG, Mintziori G, et al. Meta-analysis of cardiovascular disease risk markers in women with polycystic ovary syndrome. *Hum Reprod Update.* 2011;17:741-760.

48. Peng Z, Sun Y, Lv X, Zhang H, Liu C, Dai S. Interleukin-6 levels in women with polycystic ovary syndrome: a systematic review and meta-analysis. *PLoS One.* 2016;11:e0148531.

49. Schiffer L, Kemppegowda P, Arit W, O’Reilly MW. Mechanisms in endocrinology: the sexually dimorphic role of androgens in human metabolic disease. *Eur J Endocrinol.* 2017;177:R125-R143.

50. Scicchitano P, Pentamore I, Carbonara R, et al. Cardiovascular risk in women with PCOS. *Int J Endocrinol Metabolism.* 2012;10:611-618.

51. Roser M, Appel C, Ritchie H. Human height. Published online at OurWorldInData.org. Retrieved from: https://ourworldindata.org/human-height [Online Resource].

52. Flinterman LE, van Hylckama VA, Rosendaal FR, Cannegieter SC. Body height, mobility, and risk of first and recurrent venous thrombosis. *J Thromb Haemost.* 2015;13(4):548-554.

53. Kuipers S, Cannegieter SC, Middeldorp S, Robyn L, Büller HR, Rosendaal FR. The absolute risk of venous thrombosis after air travel: a cohort study of 8,755 employees of international organisations. *PLoS Medicine.* 2007;4:e290.

54. Streed CG, Harfouch O, Marvel F, Blumenthal RS, Martin SS, Mukherjee M. Cardiovascular disease among transgender adults receiving hormone therapy: a narrative review. *Ann Intern Med.* 2017;167:256.
55. Lawler PR, Bhatt DL, Godoy LC, et al. Targeting cardiovascular inflammation: next steps in clinical translation. *Eur Heart J*. 2021;42(1):113-131.

56. Sørensen HT, Horvath-Puho E, Pedersen L, Baron JA, Prandoni P. Venous thromboembolism and subsequent hospitalisation due to acute arterial cardiovascular events: a 20-year cohort study. *Lancet*. 2007;370:1773-1779.

57. Becattini C, Cristina Vedovati M, Ageno W, Dentali F, Agnelli G. Incidence of arterial cardiovascular events after venous thromboembolism: a systematic review and a meta-analysis. *J Thromb Haemost*. 2010;8:891-897.

58. Roach REJ, Lijfering WM, Flinterman LE, Rosendaal FR, Cannegieter SC. Increased risk of CVD after VT is determined by common etiologic factors. *Blood*. 2013;121:4948-4954.

59. Leentjens J, van Haaps TF, Wessels PF, Schutgens REG, Middeldorp S. COVID-19-associated coagulopathy and antithrombotic agents—lessons after 1 year. *Lancet Haematol*. 2021;8(7):e524-e533.

60. Colling ME, Tourdot BE, Kanthi Y. Inflammation, infection and venous thromboembolism. *Circ Res*. 2021;128:2017-2036.

61. Braekkan S, Siegerink B, Lijfering W, Hansen J-B, Cannegieter S, Rosendaal F. Role of obesity in the etiology of deep vein thrombosis and pulmonary embolism: current epidemiological insights. *Semin Thromb Hemost*. 2013;39:533-540.

62. Kotronen A, Joutsi-Korhonen L, Sevastianova K, et al. Increased coagulation factor VIII, IX, XI and XII activities in non-alcoholic liver disease. *Liver Int*. 2011;31:176-183.

63. Morelli VM, de Mutsert R, de Roos A, et al. Association between hepatic triglyceride content and coagulation factors: the Netherlands epidemiology of obesity study. *Arterioscler Thromb Vasc Biol*. 2020;40:3004-3014.

64. Minno MNDD, Tufano A, Russolillo A, Minno GD, Tarantino G. High prevalence of nonalcoholic fatty liver in patients with idiopathic venous thromboembolism. *World J Gastroenterol*. 2010;16:6119-6122.

65. Camhi SM, Bray GA, Bouchard C, et al. The relationship of waist circumference and BMI to visceral, subcutaneous, and total body fat: sex and race differences. *Obesity*. 2011;19:402-408.

66. Lord J, Thomas R, Fox B, Acharya U, Wilkin T. The central issue? Visceral fat mass is a good marker of insulin resistance and metabolic disturbance in women with polycystic ovary syndrome. *BJOG*. 2006;113:1203-1209.

67. Vassilatou E. Nonalcoholic fatty liver disease and polycystic ovary syndrome. *World J Gastroenterol*. 2014;20:8351-8363.

68. Zhu S, Li Z, Hu C, et al. Imaging-based body fat distribution in polycystic ovary syndrome: a systematic review and meta-analysis. *Front Endocrinol*. 2021;12:697223.

69. Oakes M, Arastu A, Kato C, et al. Erythrocytosis and thromboembolic events in transgender individuals undergoing masculinizing therapy with testosterone. *Res Pract Thromb Haemost*. 2021;5(suppl 2):PB1191.

70. Chulkov V, Gavrilova E, Chulkov V, et al. Assessment of cardiometabolic biomarkers in young adults with normal body weight and obesity. *Res Pract Thromb Haemost*. 2021;5(suppl 2):PB0417.

71. Pallares Robles A, ten Cate V, Schulz A, et al. Association of FXI activity with thrombo-inflammation, extracellular matrix interactions, lipid metabolism and apoptosis in venous thromboembolism. *Res Pract Thromb Haemost*. 2021;5(suppl 2):LPB0139.

72. Houghton D, Ashrani A, Liedl D, et al. Optimization of calf pump function measurements by venous plethysmography for prediction of deep vein thrombosis. *Res Pract Thromb Haemost*. 2021;5(suppl 2):OC 63.3.

73. Krishnaswamy S, Ageno W, Arabi Y, et al. Illustrated state-of-the-art capsules of the ISTH 2021 Congress. *Res Pract Thromb Haemost*. 2021;5:e12532.