COVID, long flights, and deep vein thrombosis: What we know so far

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COVID, long flights, and deep vein thrombosis: What we know so far
Zbigniew Krasiński et al., Air travel-related VTE

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Abstract
The severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (coronavirus disease
2019 [COVID-19]) pandemic has presently stunted the growth of the airline industry. Despite
the setbacks, pre-COVID passenger numbers are forecasted to return by as early as 2024. As
the industry recovers, the number of long-distance flights will surely continue to increase like
it did before the pandemic. The incidence of venous thromboembolism (VTE) following air
travel is also likely to increase. Although not common, the unique environment of air travel
exposes individuals with particular health conditions to an elevated risk of acquiring VTEs.
Numerous factors increasing the risk of developing VTE related to air travel have been
identified, including inherited and acquired flight-related aspects. Non-pharmacological
approaches to reduce air travel-related VTEs involve simple foot movements, compression
socks and stockings, intermittent pneumatic compression devices, a novel modified airline
seat, and foot exercisers. Pharmacological methods include heparins and direct oral
anticoagulants. More than 30 reliable articles were evaluated to present the current knowledge
regarding air travel-related VTEs, their risk factors, and prophylactic methods. Issues in
research methodologies found in the literature were identified and discussed. Further research
involving international collaboration projects is recommended. The authors’ perspectives
regarding long flights in previously infected COVID-19 individuals are also included.
Key words: deep vein thrombosis, economy class syndrome, pulmonary embolism,
travel-related illness, venous thromboembolism, COVID-19
Introduction

Despite the effects of the ongoing severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (coronavirus disease 2019 [COVID-19]) pandemic, the airline industry is projected to recover by 2024. New estimates from the International Air Transport Association predict passenger numbers to double by the year 2039, compared to pre-COVID years [1]. As new technologies allow for more affordable travel over long distances, a yearly passenger growth rate of 7% since 2015 had been recorded until the onset of COVID-19 [2, 3]. The long periods of immobility and cramped conditions seen in most air travelers is reflected in the term “economy class syndrome” [4–6]. Therefore, the growing number of long-range routes and passenger numbers is likely to increase the incidence of pulmonary embolism caused by deep vein thrombosis (DVT) [7]. It is estimated that in other automotive forms of transport, the risk for venous thromboembolism (VTE) amounts to between 0.5% and 10% after travelling longer than 12 and 24 hours, respectively [7, 8]. The findings of the article are summarized in the Central illustration.

Review methods

Reviews of medical articles relating to air travel-related VTE, risk factors, and prophylaxis were conducted. Medical journals in the English language were selected for review using PubMed and Google Scholar. The keywords used included “economy class syndrome”, “pulmonary embolism”, “deep vein thrombosis”, “venous thromboembolism”, “venous thromboembolism prophylaxis”, and “air travel-related illness” either as standalone searches or in combination. Additional publications not recovered in the preliminary searches were reviewed and added if deemed suitable to the topic. Finally, 30 articles were selected for review. Additional literature not previously captured was added to supplement the review based on its relevance to the scope of interest.

Mechanism

Virchow’s triad details the conditions contributing to venous thrombosis, i.e. endothelial injury, venous hemodynamic changes (stasis and turbulence), and hypercoagulability [9, 10]. Damage to the endothelium causes platelets to bind to the injury site, forming a hemostatic plug, which may become the nidus of thrombosis [9]. Two common causes of vascular injury include surgery and major trauma. Hypobaric hypoxia, low humidity, and immobility are commonly seen in air travel [9, 11].
Hypobaric hypoxia and low humidity

At 10,800 m, airplanes are pressurized to create a livable environment, which equates to an altitude of between 1524 and 2134 m, and a cabin pressure of 75.8 kPa (101 kPa AMSL) [11]. In such conditions, the oxygen saturation in healthy individuals can decrease to 90–93%, whereas in the elderly and passengers suffering from cardiac or pulmonary disorders, it may be as low as 80% [9]. Although coagulation in hypobaric and normobaric hypoxia conditions have been the subject of a number of studies, there remains a lack of consensus regarding thrombin formation in the aforementioned conditions [12].

The humidity level in an aircraft is approximately 10% (compared to sea level 30–40%) [13]. This effect is exacerbated by decreased fluid intake, which may result in dehydration [9]. In a simulated long flight study, no evidence concerning dehydration was found, although fluid retention corresponding to an approximate increase of 1 kg in body weight was observed [14]. An increase in urinary and plasmatic osmolarity associated with a low humidity combined with the diuretic effect of beverages, such as coffee, tea, or alcohol contribute to hemoconcentration promoting VTE formation [9].

Immobility

The endothelium can be deprived of its non-thrombogenic state without direct injury [10]. Immobility causes stasis, which forms a chemotactic gradient across the endothelium, triggering major leukocyte migration. Furthermore, leukocyte trapping can occur between the basement membrane and the endothelium, resulting in endothelial cell separation and desquamation, which further leads to the exposure of subendothelial layers and thrombus formation [10]. Thus, the immobility of passengers in long-distance travel is referred to as “economy class syndrome” [5]. As experienced by passengers mainly in economy class, those in window seats would have a twofold increase in risk for a VTE compared to those seated by the aisle [15].

Endurance athletes

Individuals in excellent physical condition can also be at risk of a travel-related VTE. In particular, endurance athletes and long-distance runners experience repeated microtrauma, which may induce endothelial injury. Additionally, dehydration can lead to hemoconcentration while immobility during travel to and from events contribute to VTE formation [16]. In Paget-Schrötter syndrome, heavy upper extremity activities can activate coagulation in the axillo-subclavian vein [17]. Rare iliofemoral DVT known as May-Thurner
Risk factors

The most common risk factors for flight-related VTEs are thrombophilic abnormalities and the occurrence of previous DVT [11]. Others include obesity (body mass index [BMI] > 30 kg/m²), age (> 40 years old), tall and short stature (> 1.90 m or < 1.60 m), chronic disease, oestrogen administration, female gender, pregnancy, and immobility [19–21]. A new risk factor for VTE is undoubtedly related to COVID-19 infection [22]. At present, data are scarce regarding the implications for long-haul flights in individuals who are or have previously been infected with COVID-19, although the coagulopathy observed in the disease may increase the risk of VTE [23]. Because this is a developing topic within the literature, the authors offer some perspectives in the section “COVID-19: Authors’ perspectives”.

In general, although most sufferers of air travel-related VTEs are older, young and physically fit individuals can also be at risk [16, 24]. In a study, travelers’ absolute VTE risk was evaluated, and the identified risks were as follows: 1/109 for pregnant travelers, 1/140 for travelers in a plaster cast, 1/141 for travelers with malignancies, 1/164 for travelers following a recent surgery, 1/259 for travelers on contraceptives, and 1/405 for female travellers on hormone replacement therapy [25].

Inherited factors

Inherited thrombophilias predispose individuals to hypercoagulable states [26]. The prothrombin gene constitutes the most frequent cause of hereditary hypercoagulable conditions, comprising 50–60% of VTE cases, whereas factor V Leiden mutations with antithrombin, and protein C and S represent the remainder [26]. A study showed that in 72% of travel-related VTE cases thrombophilic irregularities were present [11].

**Factor V Leiden (FVL) and activated protein C (APC) resistance.** There is an elevated VTE risk in individuals with FVL (in which insensitivity to APC is observed) [9]. Moreover, APC resistance was detected in 47% of individuals with travel-related VTE [11]. Thus, even without FVL, APC resistance constitutes a risk factor for VTE, and it was observed in 15% of patients with travel-related VTE [11].

**Prothrombin G20210A.** Mutations to prothrombin constitute the second most frequently inherited thrombophilia after FVL [11]. Combined mutations of FVL and
prothrombin gene 20210A are linked with a greater risk of VTE [27]. In a travel-related VTE study, a synergistic increase in the risk for DVT was found in FVL individuals, although the VTE risk for prothrombin mutations was less pronounced [28].

**Protein C and protein S.** Protein C is a vitamin K-dependent anticoagulant protein, circulating as a zymogen. Anticoagulant effects are exerted following the activation to APC which, in turn, inactivates factors Va and VIIIa, further activating factor X for thrombin formation [27]. Protein C deficiency was identified in 4.8% of individuals with travel-related VTE [11]. Protein S is a cofactor for APC and regulates clot formation [29]. Deficiencies of Protein S were found in 7% of patients experiencing travel-related VTE. Protein C and Protein S levels are decreased by vitamin K deficiency, warfarin, and liver failure [30].

**Antithrombin.** Antithrombin is an inhibitor of thrombin, factors IXa and Xa, and other serine proteases. Deficiency of antithrombin is either inherited or acquired. Inherited deficiencies result from mutations, whereas acquired deficiencies are primarily caused by impaired production of viable antithrombin, increased utilization, or protein losses [31]. Antithrombin deficiency may result in an elevated thrombotic risk and heparin insensitivity. In an air travel-related VTE study, 3 (< 2%) subjects with VTE suffered from antithrombin deficiencies [32].

**Other.** According to the literature, non-O blood groups have higher plasma levels of von Willebrand factor and factor VIII, which may lead to an elevated risk for thrombosis [13]. Fibrinogen gene mutation C10034T is known to produce variant fibrinogen linked with increased venous thrombosis [13]. Furthermore, lupus anticoagulant, anti-beta2-glycoprotein I, and antiprothrombin antibodies participate in prolonged coagulation in vitro [33].

Acquired factors

**Pregnancy.** Many adaptive changes occur in the hemostatic system as the body prepares for placental expulsion and vascular disruption [34]. The body enters a state of hypercoagulability and hypofibrinolysis in order to prevent excessive bleeding [34]. Although VTE risk during pregnancy is low, the postpartum risk is 5 times higher than during pregnancy [35]. However, the risk for VTE is estimated to be between 0.03% and 0.1% when the two factors, i.e. air travel and pregnancy, are combined [36]. Indeed, pregnant women on 4- to 5-hour flights have a VTE risk 5 to 10 times greater than non-pregnant women, and the risk increases to 4 and 8 times in flights longer than 8 and 12 hours, respectively [9].

Trophoblastic injury triggered by flight-related hypoxic conditions leading to premature birth and intrauterine death were reported [37]. Therefore, airlines have introduced
restrictions on pregnant women, allowing travel only up to the 36th week of pregnancy [13]. Following findings presented by the Royal College of Obstetricians and Gynecologists, it is accepted that pregnancy is at least a moderate risk factor and requires further investigation [38]. Pharmacological prevention should also be evaluated in this group of travelers [24].

**Antiphospholipid syndrome.** Antiphospholipid syndrome is an autoimmune disorder in which antibodies against proteins are bound to anionic phospholipids on plasma membranes [26]. Secondary antiphospholipid syndrome was observed in rheumatic diseases such as systemic lupus erythematosus or as a standalone disease [26]. Although VTE occurrences were reported in nearly all locales of the vascular tree in antiphospholipid syndrome patients, the most frequently reported are lower extremity DVTs and pulmonary embolisms [39].

**Chronic disease.** Numerous cases of VTE are associated with chronic disease [19]. Chronic lung or cardiovascular diseases can be exacerbated by the hypoxic conditions in air travel (i.e. induction of the coagulation system during a flight) [40]. Arthritis and inflammatory bowel disease were also identified as potential risk factors together with neoplastic diseases and chronic kidney disease [9, 21]. Indeed, the death of New Zealand international rugby icon Jonah Lomu, who had been diagnosed with nephrotic syndrome, was suspected to be caused by a VTE shortly after a long-distance flight from the United Kingdom to New Zealand [41].

Obesity (BMI > 30 kg/m²) is a widely reported risk factor in several VTE studies [9, 21]. In fact, a relative risk of 2.4 for DVT was determined when comparing non-obese and obese women [42].

Other factors

**MTHFR polymorphism and hyperhomocysteinemia.** 5,10-methylenetetrahydrofolate is reduced to 5-methyltetrahydrofolate using methylene tetrahydrofolate reductase (MTHFR). Methyl tetrahydrofolate is required in the re-methylation of homocysteine to methionine, a process that requires folate and vitamin B₁₂ [43].

Indeed, MTHFR polymorphisms were linked to an increased VTE risk [27]. MTHFR compound mutations entail a greater risk for VTE compared to heterozygous, homozygous C677T, or A1298C variants which constitute an intermediate risk [44]. It has been demonstrated that geographic and ethnic variations exist in the population. Homozygosity for C677T in North America is most prevalent in Hispanics (21–25%), followed by Whites (10–
14%), and Blacks (1–2%, particularly in the USA and Brazil), whereas homozygosity for A1298C is found more in Whites (7–12%) followed by Hispanics (4–5%) and Asians (1–4%) [45]. In contrast, heterozygosity was not considered a risk factor for VTE [11].

Hyperhomocysteinemia is another important risk factor for initial and recurring VTE, especially when fasting levels exceed 20 µmol/L [46]. The most commonly known genetic cause of hyperhomocysteinemia is MTHFR gene polymorphism [27]. Acquired hyperhomocysteinemia can stem from chronic renal failure, or it can be induced by drugs such as cyclosporine and methotrexate [11]. Additionally, folate, vitamin B<sub>6</sub>, and vitamin B<sub>12</sub> deficiencies due to a low dietary intake can also result in mild to moderate hyperhomocysteinemia [47].

Female gender. Female gender is an independent risk factor for flight-related VTE. According to Lapostolle et al. [6], 75% (42 of 56) of confirmed VTE patients were female. Additionally, a large cohort study found the VTE risk for females is 3 times higher than for men [20]. Moreover, an increased VTE risk is also observed in menopausal women receiving hormone replacement therapy; in individuals undergoing estrogen therapy, the risk of VTE is nearly 20 times higher [21, 48]. The use of the oral contraceptive pill (OCP) in healthy women increases VTE risk fourfold [49]. According to the Centers for Disease Control and Prevention, 12.6% of women between 15 and 49 years of age take the OCP in the United States of America (USA) [50]. Worldwide, 65 million women take the OCP, which amounts to 6% of all women of reproductive age [11]. Indeed, the increased risk of VTE events when using OCP is well established in the literature [49].

Flight-related. Long-distance flights are defined as lasting 7 to 15 hours or more [51, 52]. According to the literature, the risk of DVT for such flights equals 3–12% and is 3 times higher in comparison to shorter travels [53].

Arya et al. [51] found that long-haul flights (> 8 h) were associated with DVT only if one additional risk factor was present.

Window and central seating locations are significant. Belcaro et al. [54] observed that 18 of 19 thromboses were formed in subjects sitting by the window or in central seats. In their subsequent study, all 22 DVT cases (of 422 subjects) were reported in passengers seated by the window or in central seats. As for so-called “economy class syndrome”, the risk was the same for business and economy class travelers [48, 55]. Therefore, the term “traveler’s thrombosis” has been suggested as a more appropriate term [56].
As discussed previously, alcohol contributes to the diuretic effect, thereby increasing the risk of VTE. Interestingly, 66% more alcohol is consumed in business class than in economy class [55].

**Other.** A hypercoagulable state in type I and II diabetes has been established. Chronic hyperglycemia can lead to endothelial dysfunction and is crucial for the progression of vascular complications in diabetic patients [57]. In several studies diabetes was frequently used as an indicator or as an exclusion criterion of high-risk VTE in long-haul flights [58–60].

Smoking is reportedly a risk factor for travel-related VTE because it causes hypoxia and increases blood viscosity [13, 57, 61]. In women taking OCPs, smoking acts synergistically in increasing VTE risk [62]. Interestingly, while one study demonstrated that smoking was unrelated to D-dimer development and found little evidence of its association with VTE, while another study classified smoking as a low risk for VTE [55, 63]. Nevertheless, cessation of smoking to decrease VTE risk was recommended in other studies [6, 61].

Recent surgery represents a well-described risk for VTE [9, 21, 64]. Surgery risk was divided into low (minor surgery within 3 days of a flight) and high risk (major surgery within 6 weeks of a flight) [13].

An individual with a history of previous DVT or pulmonary embolism is at high risk of developing VTE [6, 27].

The impact of race and ethnicity on VTE risk has been scarcely investigated. According to White and Kenan, African Americans had a notably higher rate of VTE, particularly following events which include surgery, illness, and trauma [65]. Pacific Islanders and Asians had between 3 and 5 times lower risk for cancer-associated VTE, and idiopathic first-time symptomatic and secondary VTE [65]. Using Caucasians as the reference ethnic group for any first-time VTE risk, the less vulnerable groups were Hispanics (50%) and Asians/Pacific Islanders (70%), whereas African Americans were 35% more vulnerable to VTE [66]. Although genetic factors are more present in some ethnic groups, the data regarding air travel-related VTEs remain insufficient.

**Prevention: Non-pharmacological**

General advice on inflight exercises for travelers is available from airline websites and on-board entertainment systems and includes stretching, foot exercises, standing up, removing bags from under the seat for more leg space, and avoiding restrictive clothing [59, 67]. Foot exercises increase the mean peak velocity in the popliteal vein and can be activated by
frequent plantarflexion and dorsiflexion [68]. However, data concerning the compliance and efficacy of such exercises are scarce.

For higher-risk individuals, compression socks/stockings, intermittent pneumatic compression devices, and active foot movements have been shown to be effective [21]. The mechanism is attributed to the high flow pulsatility induced by the vessel collapse due to distal compression (by muscle contraction) allowing deep veins to drain more readily, thereby reducing venous stasis [69]. External mechanical compression does not affect coagulation; hence, the risk of increased bleeding with this method is minimal.

Compression stockings

Passengers using compression stockings have reduced incidences of DVT and lower extremity edema [54]. In LONFLIT2, the frequency of DVT among high-risk individuals in long-haul routes was reduced 18.5 times when wearing stockings [54]. The LONFLIT4 Concorde Edema-SSL study evaluated Scholl (UK) Flight Socks (below knee, 14–17 mmHg compression at the ankles) and found a distal DVT in less than 1% of the study group compared to 6% in the controls [54, 70]. In the LONFLIT4 Concorde ECO-TRAS study, similar results were found regarding Sigvaris Traveno (Ganzoni, Switzerland) elastic stockings (below knee, 12–18 mmHg compression the ankles) [71]. Thigh-length socks were found to have equivalent effectiveness compared with knee-length although the latter has better compliance and a lower cost [72]. Similar efficacy was also reported in graded compression stockings [73].

Intermittent pneumatic compression devices

Intermittent pneumatic compression devices, calf muscle pump-facilitating devices, and simple foot movements were compared [74]. Calf muscle pump facilitating devices did not present a higher efficacy than simple foot movements, whereas the use of intermittent pneumatic compression devices was found to be justifiable for sleeping, or immobile patients [74]. The use of intermittent pneumatic compression devices on flights is restricted due to the external power source, size, and weight requirements; thus, compression stockings are preferred [75].

Modified airline seat

A modified standard airline seat (NewSit) was proposed, which elevates the feet, assisting leg mobility and allowing intermittent calf compression [76]. Improvement in
venous emptying was observed in 23 out of 25 subjects whilst sitting for 5 hours, in comparison to a conventional airline seat [76]. Currently, this is the only published paper concerning this technology.

Foot exercisers

Physical foot exercisers, such as the Airogym Exerciser (Airogym Ltd, UK) and travel footrest hammocks (various brands), are less common, although they do promote blood flow through deep veins [77].

Prevention: Pharmacological

Pharmacological methods aim to decrease coagulation and clot formation. Common drugs include low-molecular-weight heparin (LMWH), unfractionated heparin (UFH), factor Xa inhibitors, direct thrombin inhibitors, and acetylsalicylic acid [24, 48]. Such methods to decrease or prevent DVT can be employed when compression, or other physical methods are contraindicated, as in the case of severe arterial claudication, drug allergies, or high hemorrhage risk [78]. The main advantages of the pharmacological measures are the increased compliance when compared to non-pharmacological methods [78]. Crucially, for individuals undergoing long-term anticoagulant therapy with a proven prevention of recurring VTE (following an unprovoked first event), the same effect cannot be presumed when administering these medications shortly before travel [79].

Heparins

Conventional evidence-based guidelines for the treatment and prevention of DVT are with heparins [79]. Data regarding UFH use for pre-flight DVT prophylaxis is scarce because it is normally used instead in the treatment of acute VTE in controlled settings due to the intensive aPTT demands [24, 80].

Low-molecular-weight heparin has replaced UFH as the drug of choice for VTE prophylaxis [64]. The efficacy of LMWH is well documented and recommended for high-risk individuals on long-distance flights [81]. LMWH has certain advantages over heparin, such as a lower risk of heparin-induced thrombocytopenia at 0.2% vs. 2.6%, respectively, and better pharmacokinetic profile [82]. In a study, a LMWH group who were administered 1 mg/kg of enoxaparin (Clexane) between 2 and 4 hours before a long-distance flight reported 0.61% of thrombotic events in the extremities compared to 4.8% in the control group and 2.9% in the aspirin group (p = 0.002 when compared to the two other groups). Additionally,
recommendations for a single 40 mg dose of enoxaparin (Lovenox) or 5000 IU of dalteparin (Fragmin) subcutaneously prior to departure have also been made [24]. LMWH’s route of administration is not the most convenient, which decreases its compliance [83].

Direct oral anticoagulants: Factor Xa and direct thrombin inhibitors

Due to minimal food and drug interactions, direct oral anticoagulants are a safer alternative than the previous methods [61]. Furthermore, there is no evidence suggesting direct oral anticoagulants cannot be used as prophylaxis for travel-related VTE. However, as primary prophylaxis, LMWH is still preferred due to the novelty and thereby lack of data regarding direct oral anticoagulants [83].

Factor Xa transforms prothrombin to thrombin and thus is essential for coagulation. rivaroxaban (Xarelto) is a direct inhibitor of factor Xa [83]. Oral administration of 10 mg was recommended for the prevention of VTE [84]. Another direct factor Xa inhibitor is apixaban (Eliquis), but there are no data on its safety or efficacy for long-haul flight VTE prophylaxis [85].

An indirect factor Xa inhibitor is fondaparinux (Artixtra) [86]. In comparison to LMWH, fondaparinux may increase the risk for fatal hemorrhage; on the other hand, when compared to UFH, it increases all-cause mortality, simultaneously reducing VTE events [87]. An informal cost analysis of the drug indicated fondaparinux to be more expensive than LMWH [80]. A recommendation for pre-flight VTE prevention is 2.5 mg subcutaneously [24]. Contraindications for factor Xa inhibitors are renal insufficiency (creatinine clearance < 30 mL/min) and hemodialysis [24].

Dabigatran etexilate (Pradaxa/Pradax) is a thrombin inhibitor with a similar efficacy to enoxaparin and a comparable safety profile to LMWH [24]. It has predictable pharmacokinetics, a rapid onset of action, and minimal drug and food interactions. A recommended prophylactic dosage is reported as 220 mg once per day [88].

Acetylsalicylic acid

Acetylsalicylic acid (ASA) inhibits platelet activation by the inactivation of cyclooxygenase. ASA used in combination with stockings has proven to be beneficial, although very few studies support ASA use for VTE prophylaxis [56, 89]. Subjects of the LONFLIT3 study were administered 400 mg once daily for 3 days, beginning the first dose 12 hours prior to the flight. The results indicated a small decrease of 3.6% in subjects with DVT compared with 4.8% in the control group; however, this result was not statistically significant.
The efficacy of aspirin as a standalone drug in VTE prevention is doubtful; hence, the American College of Chest Physicians have advised against its use for thromboprophylaxis [56].

Discussion

It is generally accepted that air travel is related to venous thromboembolism [53, 90, 91]. However, the issue of heterogeneity in the literature remains problematic, although explicable. Despite various definitions of a “long-distance” flight, the research performed has been extensive in methodology, variables, sampled populations, and locations, offering various opinions on the inclusion or exclusion criteria [51, 92]. Major papers and findings are summarized in Table 1.

The conundrum of air travel-related VTE is that it constitutes a multifactorial disease [9]. Distance or length travelled, individual variables, air travel conditions, passenger behaviors during travel, and recent events prior to travel such as trauma and surgery, all interact to produce different outcomes [56]. Thus, it is very challenging to identify the exact factors resulting in travel-related VTE.

The great variety of study designs in the literature is encouraging [56]. However, the variability in the study protocols has in some cases impeded subsequent meta-analyses [53]. Numerous studies do not meet the criteria for inclusion to meta-analyses (such as the MOOSE guidelines) and thus are unable to contribute to the existing literature [53, 93]. Chandra et al. [53] reported issues with study design, in particular regarding the use of control participants. The idea that control participants should be similar to the case patients is erroneous because individuals who develop VTEs or DVTs will always have more risk factors for the disease than those who remain disease-free [53]. This selection bias leading to the underrepresentation and, therefore, incorrect conclusions requires attention for future studies [53].

The literature shows a reliance on the D-dimer to determine the presence of DVT. For instance, in the New Zealand Air Traveler’s Thrombosis study, only participants with elevated D-dimer scores were included for further investigation [89]. Having a high sensitivity, D-dimer levels are well established, although the false positive rate for other conditions compromises their specificity [94]. Furthermore, a third of the 878 subjects had been administered aspirin. Hence, it was suggested that either the patients taking ASA were at high-risk for VTE, or the ASA falsely elevated D-dimers due to gastritis [55]. In another study, 12 subjects were evaluated for asymptomatic thrombosis detected by ultrasound. In 6 of these
subjects, no elevation in D-dimer was observed. However, the authors note that the short half-life of D-dimers (6 h) and the long time (up to 48 h) between the end of the journey and collecting blood samples could be contributing factors [7]. Schwarz et al. [48] noted 11 out of 27 DVTs, or isolated calf muscle vein thromboses, to have elevated D-dimers. In fact, the high negative predictive value of the D-dimer was reportedly so high that it has been clinically used to exclude DVT in low-risk patients [95]. Even though the role of D-dimers as a VTE marker has been well investigated, its continued role should be revised [28, 55].

Although using venography to detect DVTs is considered a gold standard by some researchers, others consider its use unethical in asymptomatic patients. Therefore, alternative methods, such as duplex ultrasonography, have also been used [7]. However, there is a potential to underestimate thromboses due to the specificity, which is reportedly between 79 and 99% [7]. Furthermore, duplex ultrasonography sensitivity and specificity decrease for distal DVT when compared to venography [94].

The variability in the time in which the DVT presents (from during the flight to several weeks after) poses a challenge for capturing data outside the normal ranges [53]. Optimal timing of ultrasound scans then should be considered when investigating travel-related VTE. In the BEST study, compressive ultrasonography (CUS) was used in addition to D-dimers to improve detection accuracy. However, most participants declined the CUS after the journey. Only half of the participants who presented elevated D-dimer agreed to a CUS. Additionally, because the scans were performed on arrival, it was not possible to detect developing thrombi. Interestingly, 90% of subjects with elevated D-dimers reported no VTE symptoms in the follow-up 6 months later [55].

Participant attrition can affect the statistical power of the study. Because the literature largely depends on volunteer recall, many subjects do not continue with subsequent phases of the research. Dropouts due to flight connection problems or other non-medical issues have the potential to affect results, depending on the study design, e.g. in LONFLIT 1 and 2, out of the original 1663 participants, 1577 subjects did not complete the study [59, 92, 96]. Another study by Belcaro et al. [60] showed that only 198 out of 244 individuals completed the study due to logistical problems. Critically, the size of the dropout effect may be difficult to determine. Although it is conceivable that participants will be tired after the journey and dropouts may occur for a multitude of reasons, every possibility to retain subjects should be explored.

Recall bias can be managed with careful study design [28, 53]. Although difficult to eliminate, recall bias from questionnaires can be decreased with clear guidelines and timely
administration of questionnaires [32]. Particular items, such as the exact amount of water/alcohol/fluid intake and the amount of inflight exercise, are available only from the most determined research subjects.

The length and duration of flights may need to depend on the randomization of data. The unpredictable nature of flights and airborne delays or re-routing, which can be difficult to report accurately, may result in inaccuracies in the subsequent analyses.

It is vital then to conduct a series of large international collaborations where uniformity in data collection methodology and consistency in study designs can definitively capture the correct data. To date, the BEST study is an example of such a collaboration, where subjects flew directly from point A to point B, with data collection at both locations eliminating the concern regarding stopovers [6, 92]. In terms of recall biases, the aforementioned large-scale studies could even involve airlines through flight attendants, who could remind the participants to record the datapoints at the best junctures.

Pre- or post-surgery considerations

Endothelial damage can be a proponent for platelet aggregation, causing thrombus formation and increasing the risk of VTE after surgery. It is possible that the danger of VTE is enhanced when surgery and long-haul travel are combined. In fact, the risk of VTE increases nearly 20 times in passengers who had a surgery within a 3-month period [25]. Conversely, in a study involving 1465 total joint arthroplasty patients, 220 travelled by air at a mean of 2.9 days after the surgery and demonstrated no differences between flying and non-flying patients. The study concluded that air travel following total joint arthroplasty is safe [97]. Furthermore, another study also found that preoperative air travel did not influence the risk of VTE after total hip and knee arthroplasty [98].

A case report described a 37-year-old male who travelled from Europe to the USA for elective pelvic surgery. Six days postoperatively, the man died from an acute pulmonary embolism despite having heparin prophylaxis [99]. The authors suggested he had developed a DVT during travel and the symptoms appeared following the surgery. However, the patient did present additional VTE risk factors, including heavy smoking, obesity, and dehydration due to pre-operative preparations [99]. Indeed, long-haul travel prior to a major surgery increases the risk of perioperative VTE [99].

COVID-19: Authors’ perspectives
When is it safe to travel long distances by air given a previous COVID infection? The thrombotic risk in COVID-19 is well documented, and in many cases, it is a determinant of disease severity or fatality [100, 101]. However, to begin answering this question, the issue of the severity with which the individual suffered from the disease should first be evaluated. The introduction of vaccinations brings yet another set of uncertainties to the equation, which adds much complexity to the issue. This question unfortunately cannot be answered scientifically without data and large-scale studies, as described in the previous sections. Given the fluidity of the situation and the amount of new information being learned about the disease every week, the authors are hesitant and reluctant to provide any real opinions.

In any case, exercising the rule of “best judgment” and conservative management, we recommend that those previously infected with COVID-19 wait at least 6 months after resolution of the disease. This includes cases of long COVID where it is not advisable to travel at all until disease-free. Furthermore, flights in these individuals should be limited to less than 6 hours duration. Should the essential need for longer journeys arise, it is advisable to arrange multi-stop journeys with a travel break in between. Lastly, the general methods of VTE prevention as described in this review should be strictly followed:

— Take pharmacological prophylaxis under direction of the individual’s physician;
— Choose an aisle seat if possible and/or seat with more legroom;
— Strictly adhere to on-board airline guidance regarding DVT prevention strategies, i.e. calf exercises, periodic foot movements, and frequent ambulation;
— Drink plenty of water and refrain from alcohol, coffee, or other diuretics before, during, and after the flight;
— Purchase and wear recommended compressive flight stockings as directed.

Conclusions

Great interest in the literature regarding air travel-related VTE reflects global trends. The conditions in which people travel by air over long distances are likely to facilitate VTE formation, most frequently in higher-risk individuals with predisposing factors. As such, preventative measures should be evaluated to decrease the possibility of developing VTE in at-risk individuals. It is likely that as a result of lockdowns and travel restrictions imposed throughout the pandemic, the post-COVID era may well see a sharp rise in air travel as borders reopen. Because long-distance airline travel is mostly international, larger prospective research on an international level should be supported. In particular, studies relating to
individuals with post SARS-CoV-2 infection, infection severity, and vaccination effects will be of exceptional value.

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| Authors | Year | Study type | Findings |
|---------|------|------------|----------|
| Ferrari et al. [90] | 1999 | Case control | Travel is a risk factor for VTE. |
| Kraaijenhagen et al. [91] | 2000 | Case control | No association between VTE and long-distance travel. |
| Arya et al. [51] | 2002 | Case control | DVT risk only increased in long-haul travelers if additional risk factors are present — prophylaxis recommended. |
| Martinelli et al. [29] | 2003 | Case control | Air travel doubles the risk for VTE, and the presence of thrombophilia or oral contraceptives increases the risk 16 and 14 times, respectively. |
| Schwarz et al. [102] | 2002 | Cohort pilot | Passengers with isolated calf vein thrombosis reported other risk factors for thrombosis. |
| Schwarz et al. [48] | 2003 | Cohort | Flights longer than 8 hours double the risk for isolated calf muscle venous thrombosis. |
| Lapostolle et al. [6] | 2001 | Retrospective | A greater distance travelled is a significant contributor to air travel-related PE. |
| Pérez-Rodríguez et al. [52] | 2003 | Retrospective | Air travel is a risk factor for VTE, and its incidence increases with the journey duration. |
| Scurr et al. [7] | 2001 | Randomized trial | Asymptomatic DVT in up to 10% of long-haul air travelers. Wearing compression stockings associated with a reduction in asymptomatic DVT. |
| Belcaro et al. LONFLIT1 [59] | 2001 | Cross-sectional | Flight related DVTs were found in individuals who presented a high risk or sitting in the window and central seats. |
| Belcaro et al. LONFLIT2 [59] | 2001 | Randomized trial | Compression therapy (stockings) decreased DVT incidence in long-haul flights. |
| Cesarone et al. LONFLIT3 [81] | 2002 | Randomized trial | LMWH use almost eradicated thrombotic events. |
| Belcaro et al. LONFLIT4 Concorde Edema-SSL [103] | 2002 | Randomized trial | Scholl Flight socks are effective in controlling edema and reducing DVT incidence in low to medium risk subjects on long-haul flights. |
| Cesarone et al. LONFLIT4 ECO-TRAS [71] | 2003 | Randomized trial | Sigvaris Traveno Stockings are effective in controlling edema in long-haul flights. |
| Cesarone et al. LONFLIT4 Concorde DVT Edema [104] | 2003 | Randomized trial | Kendall Travel Socks are effective in controlling edema and reducing DVT incidence in low- and medium-risk subjects. |
Belcaro et al.  
LONFLIT 5 JAP [70]  
2003  
Randomized trial  
Scholl Flights Socks are effective in reducing DVT incidence in high-risk subjects.

DVT — deep vein thrombosis; LMWH — low-molecular-weight heparin
Central Illustration: Graphical summary of the article's findings on air travel-related VTE.

**Air Travel-Related Venous Thromboembolism**

Krasinski, Chou & Stepak, 2021

**Issues in the literature**
- Many study designs do not conform to MOOSE guidelines
- Challenges in understanding a multifactorial disease
- Variability in time which DVT presents
- Reliance on D-dimers to detect DVTs
- Ethics of using venography on asymptomatic individuals
- Recall bias
- Participant attrition
- Variable length and duration of flights

**DVT MECHANISM AT 10,800 METRES**
- Vessel trauma
- Prior surgery
- Endothelial damage
- Hypercoagulable state
- Dehydration
- Alcohol consumption
- Low cabin humidity
- Immobility
- Sitting
- Haemostasis and turbulence

**ISSUES IDENTIFIED IN THE CURRENT LITERATURE ON AIR TRAVEL-RELATED VTE**

**Major VTE Risk Factors**
- Inherited
  - Factor V Leiden
  - Protein C Resistance
  - Prothrombin G20210A
  - Protein C
  - Protein S
  - Antithrombin
  - Pregnancy
  - Antiphospholipid syndrome
  - Chronic disease
- Acquired
  - MTHFR polymorphism
  - Hyperhomocysteinaemia
  - Female gender
  - Flight-related
  - Smoking
  - Diabetes
  - Ethnicity
  - Covid-19 infection
- Other

**VTE Prophylaxis**
- Compression socks/stockings
- Foot exercisers
- Airline seat: NewSit
- Unfractionated heparin
- Low molecular weight heparin
- Direct thrombin inhibitors
- Factor Xa inhibitors