Review

Fat Soluble Vitamins Role in Health Promotion

Mehri Aliasgharpour

Member of Biochemistry Laboratory Faculty, Reference Health Laboratory, Iran Ministry of Health & Medical Education, Tehran, Iran

*Corresponding Author: Mehri Aliasgharpour https://orcid.org/0000-0003-4906-2715

Abstract:

Vitamins as a class of essential nutrients in trace quantities are required for normal growth, and reproduction. They are studied in two categories; fat soluble vitamins and water soluble vitamins that are functional in many ways to maintain healthy immune system response for disease prevention, and to improve cognitive functions. The main aims of the present review are on vitamins discovery and classifications, fat soluble vitamins biological functions, conditions of deficiency/toxicity on human health promotion, their possible effect/s regarding COVID-19 infection and common neurological and genetic diseases. For this purpose many basic related literatures as well as new advances on fat soluble vitamins were assessed. Investigations indicated that malabsorption in fat-soluble vitamins is of particular significance in Cystic Fibrosis. In addition, in Parkinson's and Alzheimer's patients a diet rich in antioxidant vitamins recommended for their protective role and improvement of the cognitive functions. Furthermore, it is recognized that fat soluble vitamins use, especially vitamins A & D supplements during COVID-19 days in light of their safe and therapeutic range could be beneficial. However, their possible preventive role and/or supportive therapy against COVID-19 are yet controversial. Further clinical studies worldwide will hopefully define their role/s in reducing the severity and complications of the infection. In addition, in the absence of specific treatment for COVID-19 to date, as well as reducing the risks for other deficiency conditions, looking for alternative approaches like improving the availability, affordability and acceptability of healthy diets for all, specifically for the most vulnerable groups are important.

Keywords: Fat Soluble Vitamins- Deficiency- Toxicity- Covid-19 Infection- Immune System.

Submitted: 1 December 2022, Revised: 17 December 2022, Accepted: 23 December 2022

Introduction

The discovery of the vitamins was a major scientific achievement in understanding of health and disease. The major period of discovery began in the early nineteenth century and ended at the mid-twentieth century. In 1912, Casimir Funk originally purposed the term "vitamine". Funk believed that some human diseases, were caused by chemical deficiencies of factors of the same type. Because each of these factors had a nitrogencontaining component known as an amine, he called the compounds "vital amines," a term that he later shortened to "vitamines." The final "e" was dropped later when it was discovered that not all of the vitamins contain nitrogen and, therefore, not all are amines (1). By the last half of last century, all vitamins were identified, their chemical structures were determined and natural sources from which vitamins can be obtained were described in detail (2-3). Vitamins are a class of essential nutrients and organic compounds that are required in trace quantities (in micrograms to milligrams quantities per day) for normal health, growth, wellbuilt immune system and reproduction. Some vitamins can be synthesized in varying concentrations by humans. Generally this endogenous synthesis is not enough to cover daily needs and so dietary intake is required. If a vitamin is absent from the diet or is not properly absorbed by the body, a specific deficiency disease may develop (1-3). There are thirteen vitamins that are recognized. Based on their solubility, they are divided into fatsoluble variants (A, D, E and K) and watersoluble variants (B and C). The former mainly bind to nuclear receptors of the cells and affect the expression of specific genes (4). The latter mainly constitute a cofactor for the enzyme, the enzymatic activity affecting (5-6). Furthermore, the alphabetic nomenclature indicates the chronology of vitamins discovery; however; the subsequent observation that vitamin B consisted of multiple compounds,

gave rise to numerical nomenclature. The gaps in numbering are due to the removal of several substance that were initially described as vitamins (6-8). Water-soluble vitamins travel freely through the body and are not effectively stored. The excess amounts usually excreted by the kidneys. The body needs water-soluble vitamins in frequent, and small doses. These vitamins are not as likely as fat-soluble vitamins to reach toxic levels. Fat-soluble vitamins (A,D,E,K) need bile acid micelle solubilization to be digested .They are absorbed in intestinal tract with the help of dietary fats(7), transported, and stored in fatty tissue and the liver for longer periods of time. In addition, they are not excreted as easily as water-soluble vitamins (7-9). There are many publications involving both groups of vitamins (8-11), however, in the present review, fat soluble vitamins dietary sources, biological functions, state of deficiency/toxicity human on health promotion, their possible effect/s regarding COVID-19 infection, a two year global pandemic, and common neurological and genetic diseases are discussed.

Fat soluble Vitamins Vitamin A

Vitamin A (retinol) is an essential nutrient needed in small amounts by humans for the normal function of the visual system, growth and development, maintenance of epithelial cellular integrity, immune function, neuronal differentiation and influences on the secretion of neurotransmitters in the brain, and reproduction. It is not a single vitamin but a collection of compounds known as retinoids. Retinoids occur naturally in the human body. The most active retinoid has been found to be retinoic acid. Some foods such as meat, fish, eggs, and dairy foods provide retinols (Preformed vitamin A), which the body can use directly as vitamin A. Others provide provitamin A or carotenoids; generally in the form of alpha/beta/gamma carotene compounds. They are dietary precursors of retinol that the body converts into vitamin A. Carotenoids are dark-colored pigments and are found in vegetables such as carrots, sweet potatoes, spinach and other leafy green vegetables; and fruit such as cantaloupes and apricots (12-13). There are more than 500 known carotenoids. The most common type of them is betacarotene that is an antioxidant and protects cells from damage caused by free radicals (14-16). It is also specifically contributes to the orange color of vegetables and fruits. Vitamin A functions at two levels in the body: the first is in the visual cycle in the retina of the eye; the second is in all body tissues where it systemically maintains the growth and soundness of epithelial cells and membrane regulation (from skin to mucous to teeth and to bones) metabolism (17). Furthermore, it has anti- oxidative [13-14] properties and a major role in immune system (18). In the visual system, the major consequence of vitamin A deficiency is ocular dysfunction with abnormal dark adaptation (night blindness), conjunctival and corneal xerosis (thickening) which can lead to blindness (19-20). Night blindness in which it is difficult or impossible to see in relatively low light is usually an indicator of inadequate available retinol, but it can also be due to a deficit of other nutrients that are critical to the regeneration of rhodopsin, such as protein and zinc, and to some inherited diseases, such as retinitis pigmentosa. In the eye, the symptoms signs, together referred and to as xerophthalmia. It ranges from the milder stages of night blindness and Bitot spots to the potentially blinding stages of corneal xerosis, ulceration and necrosis (keratomalacia). The various stages of xerophthalmia are regarded both as disorders and clinical indicators of vitamin A deficiency. At the second level, the growth and differentiation of epithelial cells throughout the body are affected by vitamin A deficiency (VAD). In VAD case, in epithelial

tissues, goblet cell numbers are reduced and causes mucous secretions to diminish. As a consequence, cells lining protective tissue surfaces fail to regenerate and differentiate, hence they flatten and accumulate keratin. Both factors-the decline in mucous secretions and loss of cellular integrity-reduce the body's ability to fight infections (21). Classical symptoms of xerosis (drying) and desquamation of dead surface cells as seen in ocular tissue (i.e. xerophthalmia) are the external evidence of the changes also occurring to various degrees in internal epithelial tissues. Vitamin A deficiency has also been linked to impaired mechanisms of host resistance to infection, poor growth and increased mortality in a study of mothers and children ^[19]. Furthermore, Cystic Fibrosis is a known risk factor of VAD because of liposoluble vitamin malabsorption due to pancreatic insufficiency (21). Cystic fibrosis (CF) is a genetic disorder that affects multiple organs and causes disease in the lungs, digestive systems and pancreas. In people with CF, the pancreas often does not produce enough enzymes to allow the body to absorb digested food properly. Pancreatic insufficiency affects up to 90% of people with CF, whereby fat malabsorption occurs and pancreatic enzyme replacement is required to prevent malnutrition (21). Fat soluble vitamins (A, D, E and K) are co-absorbed with fat and thus deficiency of these vitamins may occur (20). Vitamin A deficiency can be defined as a serum retinol (SROL) concentration less than 0.70 µmol/L (18). However, SROL levels may be influenced by albumin and retinol binding protein as well as acute illnesses with infection and inflammation (22). Levels of SROL should be measured while individuals are clinically stable (20,23). Furthermore, concentration of vitamin A, do not play a role in the pathogenesis of Parkinson's disease (PD) (24). Carotene deficiency has no defined serum concentrations (25). However, as long as vitamin A levels are normal, adverse clinical manifestations of β -carotene deficiency are unknown (15). Because vitamin A is fat soluble and can be stored, primarily in the liver, routine consumption of large amounts of vitamin A over a period of time can result in toxic symptoms, including hepatotoxicity, bone abnormalities and joint pain, headaches, and skin desquamation. vomiting, Furthermore, it may contribute to osteoporosis and hip fractures. Hyper-vitaminosis A appears to be due to abnormal transport and distribution of vitamin A and retinoids caused by overloading of the plasma transport mechanisms (26-27). On the other hand, excessive carotene in the diet can temporarily yellow the skin, a condition called carotenemia, a harmless, reversible yellowing of the skin (15). In general, most authors consider excessive intake of carotene as nontoxic (15-16). It is commonly seen in infants fed largely mushed carrots.VAD is most common in populations consuming most of their vitamin A needs from pro-vitamin carotenoid sources and where minimal dietary fat is available (28). About 90% of ingested preformed vitamin A is absorbed, whereas the absorption efficiency of pro-vitamin A carotenoids varies widely, depending on the type of plant source and the fat content of the accompanying meal (29). Where possible, an increased intake of dietary fat is likely to improve the absorption of vitamin A in the body. VAD can occur in individuals of any age. However, it is a disabling and potentially fatal public health problem for children under 6 years of age. VAD related blindness is most prevalent in children under 3 years of age (30). Women of reproductive age are also thought to be vulnerable to VAD during pregnancy and lactation because they often report night blindness (31-32) and because their breast milk is frequently low in vitamin A (33-34). There is no consistent, clear indication in humans of a sex differential in vitamin A requirements during childhood. Growth rates. and

presumably the need for vitamin A, from birth to 10 years for boys are consistently higher than those for girls (35). Furthermore, in light of vitamin A pulmonary and immunological roles, oral supplementation of vitamin A is currently being investigated in the treatment of severe acute respiratory syndrome coronavirus-2 (SAR- CoV-2), later named COVID-19 by the WHO. Pathology of COVID-19 involves a complex interaction between COVID-19 and the immune system. In addition, COVID-19 has been found to persuade inflammatory responses, especially involving liver, lung, and kidney, which further increase the risk of depletion of vitamin A stores (36-37).

Vitamin D

Vitamin D encompasses a number of fatsoluble secosteroids with a physiological role in mineral homeostasis, primarily calcium, magnesium, and phosphate. They are in turn needed for the normal mineralization of bone and skeletal health, muscle contraction, nerve conduction, and general cellular function in all cells of the body. Vitamin D in its natural form, cholecalciferol, is acquired through dietary sources, such as oily fish, mushrooms, and egg yolks, but is also produced through de novo synthesis in the dermis of the skin from dehydrocholesterol(cholesterol like precursor) by exposure to UVB rays from the sun or other sources (38). The version made in the skin is referred to as vitamin D3 (colecalciferol) whereas the dietary form can be vitamin D3 or a closely-related molecule of plant origin known as vitamin D2 (ergocalciferol). From a nutritional perspective, both of the above forms are inactive. They are metabolized similarly in humans, are equal in potency, and can be considered equivalent. Regardless of the route of entry, 99% of endogenous or dietary vitamin D, are transformed into the active form; [1,25dihydroxy vitamin D or1,25-(OH)2D / calcitriol] after being hydroxylated twice. First hydroxylation happens in the liver that

produces [25-hydroxyvitamin-D or 25(OH) D/calcidiol], which is the most abundant circulating metabolite of the vitamin D. The second hydroxylation occurs in the kidneys that produces [1,25-dihydroxy vitamin D or 1,25-(OH)2D / calcitriol] (39-40). This active form regulates the transcription of a number of vitamin D-dependent genes which code for calcium-transporting proteins and bone matrix proteins. Clinical assays measure 1,25-(OH)2D2 and 1,25-(OH)2D3, collectively called 1,25-(OH)2D. Similarly, calcidiol is measured as 25-OH-D but it is a mixture of 25calcium OH-D2 and 25-OH-D3. In homeostasis, 1,25-(OH)2D works in conjunction with parathyroid hormone (PTH) to produce its beneficial effects on the plasma levels of ionized calcium and phosphate (41-42). The physiologic loop starts with the calcium receptor of the parathyroid gland (43). When the level of ionized calcium in plasma falls, PTH is secreted by the parathyroid gland and stimulates the tightly regulated renal enzyme 25-OH-D-1- α -hydroxylase to make more 1,25-(OH)2D from the large circulating pool of 25-OH-D. The resulting increase in 1,25-(OH)2D (with the rise in PTH) causes an in calcium transport within the increase intestine, bone, and kidney. All these events raise plasma calcium levels back to normal, which in turn is sensed by the calcium receptor of the parathyroid gland. The further secretion of PTH is turned off not only by the feedback action of calcium, but also by a short feedback 1,25-(OH)2D loop involving directly suppressing PTH synthesis in the parathyroid gland. This model clearly demonstrates that sufficient 25-OH-D must be available to provide adequate 1,25-(OH)2D synthesis and hence an adequate level of plasma calcium. Furthermore, numerous recent studies have focused on the plasma levels of 25-OH-D and PTH to gain an insight into vitamin D status and there is a strong presumptive relationship of this variable with bone status (39,44-47).

Vitamin D deficiency is prevalent at all ages, especially in elderly. Vitamin D not only regulates the calcium homeostasis and skeletal health but also regulates the physiological and such pathological processes, as cell proliferation, differentiation, and antioxidative stress (48-50) .Children with a lack of vitamin D may suffer from rickets (bone deformity) and adults may develop osteoporosis and osteomalacia. Additionally, vitamin D deficiency is also associated with cardiovascular diseases, muscle weakness, diabetes mellitus, cancers, and multiple sclerosis (51). The relationship between vitamin D and Parkinson's disease (a multifactorial disease) has gradually attracted attention (52). Many non-interventional studies found that the high levels of serum vitamin D can reduce the risk of Parkinson's disease (PD) (53-55), and several clinical intervention trials also proposed that vitamin D supplementation attenuate the deterioration of the can Parkinson's disease and reduce the occurrence of fractures in patients with PD (56-57). Furthermore, epidemiological and clinical studies suggest that vitamin D has a positive effect on (PD). In a cohort study, over 7000 serum samples were collected for measuring 25-hydroxy vitamin D level. the and meanwhile, the occurrence of PD was instigated over a 30-year follow-up period. The results showed that individuals with higher serum vitamin D concentrations had a lower risk of PD (58). Evatt et al. also noted consistent findings (59). PD patients with lower 25-hydroxy vitamin D levels may exhibit more severe symptoms compared with normal controls (60-61). Unsurprisingly, a randomized, double-blind, placebo-controlled trial found that vitamin D3 supplementation (1200 IU/day for 12 months) significantly prevented the deterioration of PD patients (62). However, further studies are still needed to clarify the definitive correlations between vitamin D and PD. It has also been reported that vitamin D deficiency co-exists in patients with COVID-19. Dark skin color, increased age, the presence of pre-existing illnesses and vitamin D deficiency are features of severe COVID disease (63). Moreover, vitamin D has been suggested to play a role in COVID-19, as two ecological studies indicated that the rate of infection was higher in countries at higher latitudes and/or lower vitamin D status (64-65). In recommending intakes for vitamin D, it is recognized that in most locations in the world in a broad band around the equator (between latitudes 42°N and 42°S), the most physiologically relevant and efficient way of acquiring vitamin D is to synthesize it endogenously in the skin from 7dehydrocholesterol by sun (UV) light exposure. In most situations, approximately 30 minutes of skin exposure (without sunscreen) of the arms and face to sunlight can provide all the daily vitamin D needs of the body (66). However, skin synthesis of vitamin D is negatively influenced by factors which may reduce the ability of the skin to provide the total needs of the individual (66). Examples of such factors are; Latitude and season- both influence the amount of UV light reaching the skin; The ageing process-thinning of the skin reduces the efficiency of this synthetic process; Skin pigmentation-the presence of darker pigments in the skin interferes with the synthetic process because UV light cannot reach the appropriate layer of the skin; Clothing-virtually complete covering of the skin leaves insufficient skin exposed to sunlight; Sunscreen usewidespread and liberal use of sunscreen, though reducing skin damage by the sun, deleteriously affects synthesis of vitamin D. Because not all of these problems can be solved in all geographic locations, particularly during winter, it is recommended that individuals correct their vitamin D status by consuming the amounts of vitamin D appropriate for their age group (Table 1) (67). It is rare for an individual to have vitamin D toxicity and the adverse

effects of high vitamin D intakeshypercalciuria and hypercalcaemia- do not occur at the recommended intake levels as in Table 1. In fact, it is worth noting that the recommended intakes for all age groups are still well below the lowest observed adverse effect level of $50\mu g/day$ and do not reach the "no observed adverse effect level" of $20\mu g/day$ (47,68-69).

Vitamin E

Vitamin E is the major lipid-soluble antioxidant in the cell antioxidant defense system and is synthesized by plants and exclusively obtained from diet. Natural vitamin E includes two subgroups: tocopherols and tocotrienols; and they can further be divided into four lipophilic molecules, respectively: α -, β -, γ -, and δ -tocopherol and α -, β -, γ -, and δ tocotrienol. The major difference between tocopherols and tocotrienols is the side chain. Tocopherols have a saturated phytyl tail, while tocotrienols possess an unsaturated isoprenoid side chain (70). Because of this unsaturated side chain, the tocotrienol is superior to the tocopherol as an antioxidant by increasing the molecular mobility through lipid membranes and by accepting electrons readily (70). In addition to its potent antioxidant capacity, vitamin E is involved in many physiological processes such as immune function (71), cognitive function, physical performance (72-73), regulation of gene expression and skin health. Vitamin E is located primarily within the phospholipid bilayer of cell membranes and its major biological role is to protect poly unsaturated fatty acids (PUFAs) and other components of cell membranes and lowdensity lipoprotein (LDL) from oxidation by free radicals. Elevated levels of lipid peroxidation products are associated with numerous diseases and clinical conditions (74). Although vitamin E is primarily located in cell and organelle membranes where it can exert its maximum protective effect, its concentration may only be one molecule for every 2000 phospholipid molecules. This suggests that after its reaction with free radicals it is rapidly regenerated, possibly by other antioxidants (75). The requirement for vitamin E is relate to the (PUFAs) content of cellular structures and therefore, depends on the nature and quantity of dietary fat, which affects such composition. Hence, the minimum adult requirement for vitamin E is not certain and no specific recommendations regarding the intake of vitamin E have been made officially, and the optimal supplementation dosage of mixed tocopherols is still undetermined. But is probably not more than 3 to 4 mg (4.5-6 IU) for α -tocopherol per day for those who ingest a diet containing the minimum of essential fatty acids (76). Foods rich in vitamin E include canola oil, olive oil, margarine, almonds and peanuts. One can also get vitamin E from meats, dairy, leafy greens and fortified cereals. Vitamin E is absorbed in the presence of bile and from the small intestine. Most tocopherol enters the bloodstream via lymph, where it is associated with chylomicrons and very low density lipoproteins. The vitamin is stored in most tissue, with the largest amount stored in adipose tissue. Some of the tocopherol deposition is in association with lipoproteins in cellular membranes. Rapid exchange of tocopherol occurs between the erythrocyte membranes and plasma lipoproteins. When physiological amounts are administered, only a small fraction of the dose appears in urine (77-78). The assessment of the vitamin E requirement for humans is confounded by the very rare occurrence of clinical signs of deficiency because these usually only develop in infants and adults with fat malabsorption syndromes or liver disease, in individuals with genetic anomalies in transport or binding proteins, and possibly in premature infants (79-81). This suggests that diets contain sufficient vitamin E to satisfy nutritional needs. Vitamin E appears to have very low toxicity, when

obtained from food sources alone. It has no documented evidence of toxicity. However, evidence of pro-oxidant damage has been found to be associated with supplements, but usually only at very high doses (for example at >1,000 mg/day) (82-83). A few other studies suggest that tocopherols appear to inhibit platelet aggregation through the inhibition of protein kinase C (PKC) and the increased action of nitric oxide synthase (84-85). Furthermore, it has been suggested that vitamin E supplementation (200-400mg/day) may be appropriate therapeutically to moderate some aspects of degenerative diseases such as Parkinson's disease (PD) [(86). However, two population-based studies did not find the association between vitamin E intake and risk of PD (87-88). To date, there is little evidence regarding the use and/or dosage of vitamin E as a prophylactic or therapeutic agent against COVID-19 (89). Overall vitamin E supplementation improves overall immune functions, reduces respiratory tract infection incidences, severity, lowers virus load in lung tissues, and increases the antibody titers, particularly in the elderlv (90-92). Malnourished individuals should benefit from the inclusion of vitamin E supplementation in COVID-19 management. The Alzheimer's disease (AD) Cooperative Study in 1997 showed that vitamin E may slow disease progression in patients with moderately severe AD (93). High doses of vitamin E delayed the loss of the patient's ability to carry out daily activities and their consequent placement in residential care for several months. In another study, it was found that subjects with AD had reduced concentrations of plasma antioxidant micronutrients, suggesting that inadequate antioxidant activity is a factor in this disease. High plasma levels of vitamin E are associated with a reduced risk of AD in older patients and this neuroprotective effect is related to the combination of different forms of vitamin E rather than to α -atocopherol alone (94).

Alzheimer's disease occurs as a result of protein oxidation and lipid peroxidation via a free radical mechanism, where the beta amyloid protein induces cytotoxicity through a mechanism involving oxidative stress and hydrogen peroxide, leading to neuronal cell death and finally AD. Vitamin E can block the production of hydrogen peroxide and the resulting cytotoxicity (95-97). Moreover, Vitamin E is the major naturally occurring non-enzymatic lipid-soluble antioxidant protecting skin from the adverse effects of oxidative stress including photo-aging. It protects the skin from various deleterious effects due to solar radiation by acting as a free radical scavenger (98) and it is an important ingredient in many cosmetic products. Experimental studies have indicated that vitamins C and E have important protective effects in the aging process and brightening of the skin (99-100). The results have revealed significant improvements in the skin tone and increase in homogeneity. Furthermore, vitamin E is one of the treatment for yellow nail syndrome (101-102). Yellow nail syndrome includes slow growing, opaque yellow nails with exaggerated yellow curvature. In addition, the antioxidant supplementation through vitamins E and C and the mineral zinc has been seen to apparently enhance the antioxidant protection against oxidative stress and allow less time for wound healing (103).

Vitamin K

Vitamin K refers to a family of compounds with a common 2-methyl-1,4-naphthoquinone (104-105) but differ in the structures of a side chain at the 3-position. Vitamin K is needed for a unique post-translational chemical modification in a small group of proteins with properties, calcium-binding collectively known as vitamin K-dependent proteins or Gla proteins. They function in coagulation, bone development, and cardiovascular health. Vitamin K exists naturally in two bioactive

forms; as vitamin K1 (phylloquinone) and (menaquinone) (105-108).vitamin K2 Vitamin K1 contains a phytyl side chain and is synthesized by plants and algae. It is mainly found in green leafy vegetables as well as olive oil and soyabean oil. In addition, it is the main circulating form of vitamin K and is primarily provided by dietary sources. Whereas vitamin K2 consists of a group of menaquinones, which are characterized by the length of their isoprenoid side chain. Furthermore, vitamin K2 is created in the human gut by bacteria and it is found in small amounts in chicken, butter, egg yolks, cheese and fermented products (better known as natto) (109-111).Today, menaquinones are generally called MK-n, where "n "signifies the number of isoprenoid units. With regard to preventive and therapeutic aspects, menaquinone-4 (MK-4) and menaquinone-7 (MK-7) are among the most important forms of vitamin K2 with 4 and 7 isoprenoid units, respectively (106,112-113). There is a third, synthetic form K3 (menadione), the use of which has been replaced by a synthetic form of vitamin K1 due to the potential for toxicity in infants with glucose-6-phosphate dehydrogenase deficiency (114) . Although vitamin K1 (phylloquinone) in blood must have been derived exclusively from the diet, it is not known whether circulating vitamin K2 (menaquinones) such as MK-7 are derived from the diet, intestinal flora, or a combination of these sources. Vitamin K1 is the major source (>90%) in the human diet and is absorbed in the jejunum and ileum, transported by chylomicrons in circulation, and is dependent on bile, pancreatic enzymes, and dietary fat content (115). In addition, it is known to be selectively distributed in a number of hepatic and non-hepatic tissues. The biological role of vitamin K is to act as a cofactor for a specific carboxylation reaction that transforms selective glutamate (Glu) residues to g-carboxyglutamate (Gla) residues

(104,116). The reaction is catalysed by a microsomal enzyme, g-glutamyl, or vitamin Kdependent carboxylase, which in turn is linked to a cyclic salvage pathway known as the vitamin K epoxide cycle. The four vitamin Kdependent pro-coagulants (factor II or prothrombin, and factors VII, IX, and X) are serine proteases that are synthesized in the liver and then secreted into the circulation as inactive forms (zymogens). Their biological activity depends on their normal complement of Gla residues, which are efficient chelators of calcium ions. In the presence of Gla residues and calcium ions these proteins bind to the surface membrane phospholipids of platelets and endothelial cells and together with other cofactors, form membrane-bound enzyme complexes. When coagulation is initiated, the zymogens of the four vitamin K-dependent clotting factors are cleaved to yield the active protease clotting factors and thus preventing bleeding (104,116-117). Two other vitamin K dependent proteins, protein C and protein S, play a regulatory role in the inhibition of coagulation. The function of protein C is to degrade phospholipid-bound activated factors V and VIII in the presence of calcium. Protein S acts as a synergistic cofactor to protein C by enhancing the binding of activated protein C to negatively charged phospholipids. There is evidence that protein S is synthesized by several tissues including the blood vessel wall and bone and may have other functions besides its well-established role as a coagulation inhibitor. Both vitamin K1 and vitamin K2 are required for the γ -glutamyl carboxylation of all vitamin K-dependent proteins (106) .Although mammalian bacterial intestinal flora are able to produce vitamin K2, the amount produced is thought to be negligible (106). Vitamin K is present in the other body tissues as well, including the brain, heart, pancreas, and bone (36, 118-119). It is rapidly metabolized in the liver and excreted in the urine and bile (115). This rapid metabolism accounts for its

relatively low blood levels and tissue stores compared to the other fat soluble vitamins. Furthermore, vitamin K can have a serious interaction with anticoagulants such as warfarin (Coumadin) antagonizing its activity and leading to the depletion of vitamin K dependent clotting factors. Most individuals taking warfarin are advised to avoid vitamin Kcontaining foods, such as green leafy vegetables (120). Vitamin K deficiency can be exacerbated further when warfarin is initiated (121). In addition, vitamin K deficiency can contribute to significant bleeding, poor bone development, osteoporosis, and increased cardiovascular disease. Vitamin K Deficiency Bleeding (VKDB) is rare and it is potentially life threatening bleeding disorder of early infancy. Because vitamin K stores are low at birth and its concentration in human milk is low as well. Most multi dose oral regimens provide protection for all except for a small group of infants with undetected hepatobiliary disease In adults, primary vitamin K-(122-123).deficient states that manifest as bleeding are almost unknown except when the absorption of the vitamin is impaired as a result of an underlying pathology (104,124) or during long term antibiotic or anticoagulant treatments (125). Naturally occurring vitamin K (K1 and K2) toxicity is extremely rare. However, a synthetic form of vitamin K (menadione) or K3 has been associated with neonatal haemolysis and liver damage, and therefore, no longer used therapeutically (6) .Thrombosis is a frequent manifestation of COVID-19 that contributes to poor outcomes. It has been proposed that pneumonia induced extrahepatic vitamin K depletion lead to accelerated elastic fiber damage and thrombosis in severe COVID-19 patients due to impaired activation matrix Gla Protein or (MGP) and of endothelial protein S respectively (126-127).

Discussions

Each fat soluble vitamin as well as other trace elements (128) has multiple and essential biological functions for human health. In summary; vitamin A is essential for normal vision, epithelial cell integrity, and epithelial proliferation and immunity (18). The major cause of blindness in children worldwide is xerophthalmia (19) caused by vitamin A deficiency. It is also pivotal in the maintenance of innate cell mediated and antibody mediated responses (129). In a study on vitamin A, Field et al, reported vitamin A has a capacity to resist infection (130). In this regard, several studies reported that vitamin A and related compounds have a potentially beneficial role and could be a promising option in the treating of COVID-19 pandemic by preventing lung infection (131-133). Vitamin D, promotes absorption and metabolism of calcium and phosphorus. It is required for bone health and has a role in immune function and incidence of cancer, type 1 diabetes, autoimmune disease and heart disease (3-6,134-137). Promising results vitamin D supplement is indicated that essential in treating respiratory tract infection, including COVID-19 (63-64), and pulmonary fibrosis as well (138). Furthermore, it is indicated that during the COVID-19 pandemic, it is crucial that all people in the hospital, including the patients and staff, take vitamin D supplement to raise 25(OH)D concentrations as an essential step in preventing infection spread (139). However, initial testing of vitamin D level at an interval is advocated to determine the dosing levels required. The best defined role for vitamin E is an antioxidant activity for polyunsaturated fatty acids (PUFAs) within the membranes, or preventing cell membrane and maintaining neurological oxidation. functions. The antioxidant system mainly consists of two subtypes: firstly, enzymatic antioxidant system, including superoxide dismutase (SOD), catalase (CAT), and glutathioneperoxidase (GSH-Px), and

secondly, nonenzymatic antioxidant system, including vitamin C, vitamin E, glutathione, melatonin, alpha-lipoic acid, carotenoids, and trace elements copper, zinc, and selenium (74,128). Furthermore, oxidative stress refers to the imbalance between the oxidation system and antioxidant system that results in excessive accumulation of oxidative substances. Reactive oxygen and nitrogen species and other free radicals that result from oxidative stress are potential causes of cell membrane damage (74,140). In general diseases causes an increase in oxidative stress: therefore. consumption of foods rich in antioxidants, which are potentially able to quench or neutralize excess radicals, may play an important role in modifying the development of disease. Although limited clinical data is available to establish a link between oxidative stress and viral infection due to COVID-19. many lines of evidence (141-143) still suggested that overproduction of reactive and deprived antioxidant oxygen species system play a significant role in the pathogenesis and severity of COVID-19. It is often suggested that the onset of severe lung injury in COVID- 19 patients is based on the activation of the oxidative stress mechanism coupled with an innate immune response (144-146). Furthermore, vitamin E supplementation may lead to the protection and improvement of lung tissue in Cystic Fibrosis patients. However, future studies are needed to look at this issue more specifically (147). The letters (A, B, C and so on) were assigned to the vitamins in the order of their discovery. The one exception was vitamin K which was assigned "K" from "Koagulation" by the Danish researcher Henrik Dam (2,8,148). Beyond coagulation (149), vitamin K and vitamin K -dependent proteins are essential for calcification (maintaining bone and cardiovascular health), energy metabolism and inflammation (149-150). Other important roles of vitamin K is its ability to act as a potent antioxidant reducing the lipid peroxidation in the cell by producing vitamin K hydroquinone, a robust radical scavenging species (151-152). Furthermore, vitamin K has been found to have an anti-inflammatory activity (153). Reduced vitamin K levels have also been reported in COVID-19 patients (126) and one of the most common laboratory findings in these patients is the elevation of D-dimers (154-155). This deficiency reduces the functional levels of coagulation factors II, VII, IX, and X, predisposing them to develop coagulopathy and increasing hemorrhage risk (156-157) and DIC (Disseminated Intravascular Coagulation) formation. Coagulophaty and DIC appear to be associated with high mortality rates. The other laboratory markers recommended by the International Society of Thrombosis and Hemostasis (ISTH) for monitoring DIC formation are fibrinogen, prothrombin time, and platelet count (158). Also, low vitamin K level appears to be associated with increased elastin degradation (159) preferably degrading the lung tissue and resulting in breathing difficulty in COVID-19 patients. For the treatment of COVID-19 induced coagulopathy, the use of an anticoagulant is recommended (158). Moreover, evidence for benefits of routine vitamin K supplementation for people with Cystic Fibrosis is currently weak. However, no harm was found and until further evidence is available, the present recommendation by National guidelines should be followed (124,160). Furthermore, literatures review indicated that fat soluble vitamin deficiency malabsorption is of particular significance in Cystic Fibrosis (20-21,160). In Parkinson's disease vitamins may play a protective role, however, they cannot control the progression of the disease (83) and antioxidant a diet rich in vitamins recommended to improve the cognitive functions of Alzheimer's patients (95). Table2 summarizes the main biological functions of the fat soluble vitamins and their deficiency/toxicity cases in human health.

Conclusion

The importance of adequate and balance nutrition is vital in regulating the body's homeostasis. In this regard, a diet rich in fat soluble vitamins (A,D,E,K) are considered essential in small quantities for normal health, maintaining healthy immune system response disease prevention, and improving for cognitive functions (Table2). Use of these vitamins especially vitamins A & D supplements during a two year global pandemic COVID-19, in light of their safe and therapeutic range is also recommended. However, their possible preventive role and/or supportive therapy against COVID-19 are yet controversial. Further future clinical studies worldwide will hopefully define their role/s in reducing the severity and complications of the infection. In addition, in the absence of specific treatment for COVID-19 to date, as well as reducing the risks for other deficiency conditions, looking for alternative approaches like improving the availability, affordability and acceptability of healthy diets for all, specifically for the most vulnerable groups are important.

Conflict of Interest

None.

References

1-Funk C. The etiology of the deficiency diseases. Beri-beri, polyneuritis in birds, epidemic dropsy, scurvy, experimental scurvy in animals, infantile scurvy, ship beri-beri, pellagra. J State Med 1912; 20: 341.

2-Rosenfeld L. Vitamine- vitamin. The early years of discovery. Clin Chem 1997; 43(4):680-685.

3-Semba RD. Bloem MW. The anemia of vitamin A deficiency: Epidemiology and

pathogenesis .Euro J Clin Nutr 2002; 56(4):271-281.

4-Sánchez-Hernández D. G. Anderson H. Poon AN. *et al.* "Maternal fat-soluble vitamins, brain development, and regulation of feeding behavior: an overview of research". Nutr Res 2016;36(10): pp 1045–1054, 2016.

5-Chawlac J. Kvarnberg D. "Hydro soluble vitamins," Handbook of Clinical Neurology 2014; 120: pp 891–914.

6-Bender DA. Nutritional biochemistry of the vitamins. 2nd ed. United Kingdom: Cambridge University Press; 2003; 470.

7-Combs GF. The Vitamins: Fundamental aAspects in nutrition and health. 4th ed. London: Academic Press; 2012: pp 33-70.

8-Maltz A. Funk C. Nonconformist nomenclature, and networks surrounding the discovery of vitamins. J Nutr. 2013;143(7):1013-1020.

9-Fidanza A. Audisio M. Vitamins and lipid metabolism. Acta Vitaminologica Et Enzymologica 1982;4(1-2):105-114

10-Eggersdorfer M. Laudert D. Létinois U. McClymont T. *et al.* One hundred years of vitamins—A success story of the natural sciences. Angew Chem Int Ed Engl. 2012; 51:pp 12960–12990.

11-Ball G.F.M. Vitamins in Foods: Analysis, Bioavailability, and Stability; CRC Press: Boca Raton, FL, USA, 2005.

12-Blaner WS. Li Y. Brun, P.-J. Yuen J J. Lee S.A. Clugston RD. Vitamin A Absorption, Storage and Mobilization. Subcell. Biochem. 2016; 81: 95–125.

13-Parker RS. Absorption, metabolism, and transport of carotenoids. FASEB J 1996; 10:542–551.

14-Fiedor J. Burda K. Potential role of carotenoids as antioxidants in human health and disease. Nutr 2014; 6: 466–488.

15-Cantin AM. White TB, Cross CE. Forman_HJ.*et al.* Antioxidants in cystic fibrosis. Conclusions from the CF Antioxidant Workshop, Bethesda, Maryland, November 11-12, 2003. Free Radical Biology and Medicine 2007; 42(1):15-31.

16- Hammond BR. Renzi LM. Carotenoids. Adv Nutr 2013; 4(4):474-6.

17-Herschel Conaway H. Henning P. Lerner UH. Vitamin A metabolism, action, and role in skeletal homeostasis. Endocr. Rev 2013; 34: 766–797.

18- Huang Z. Liu Y. Qi G. Brand D. Zheng S. Role of vitamin A in the immune system. J. Clin. Med 2018; 7:258.

19-West KP. Vitamin A deficiency disorders in children and women. Food Nutr Bull 2003; 24(4 Suppl):S78-90.

20-Saxby N. Painter_C. Kench_A. King_S. *et al.* Nutrition Guidelines for Cystic Fibrosis in Australia and New Zealand. Thoracic Society of Australia and New Zealand. 2017. https://www.thoracic.org.au/documents/item/1 045.

21-Ross AC. Stephensen CB. Vitamin A and retinoids in antiviral responses. FASEB J 1996;10:979–985.

22-Napoli JL. Retinoic acid biosynthesis and metabolism. FASEB J : Official publication of the Federation of American Societies for Experimental Biology 1996; 10(9):993-1001.

23-Borowitz D. Baker_RD. Stallings_V. Consensus report on nutrition for pediatric patients with cystic fibrosis. J Pediatric Gastro Nutr 2002; 35(3):246-59.

24-Jimenez-Jimenez FJ. Fernandez-Calle P. Vazquez A. Serum levels of vitamin A in Parkinson's disease. J Neurol Sci 1992;111(1):73-6.

25-Centers for Disease Control and Prevention. Second national report on biochemical indicators of diet and nutrition in the U.S. population. 2012 April. www.cdc.gov/nutritionreport/ Report.

26-Smith FR. Goodman DS. Vitamin A transport in human vitamin A toxicity. New England J Med 1976; 294:805–808.

27-Penniston_KL. Tanumihardjo SA. The acute and chronic toxic effects of vitamin A. Am J Clin Nutr 2006; 83(2):191-201.

28-Mele L . West KP Jr. Kusdiono PA. *et al.* Nutritional and household risk factors for xerophthalmia in Aceh, Indonesia: a case– control study. Amer J Clin Nutr 1991; 53:1460–1465.

29-Erdman J Jr. The physiologic chemistry of carotenes in man. Clin Nutr 1988; 7:101–106.

30- Sommer A. Vitamin A deficiency and its consequences: a field guide to detection and control, 3rd ed. Geneva, World Health Organization, 1994.

31-.Bloem MW. Matzger H. Huq N. Vitamin A deficiency among women in the reproductive years: an ignored problem. In: Report of the XVI IVACG Meeting. Washington, DC, International Vitamin A Consultative Group, ILSI Human Nutrition Institute, 1994.

32-Christian P. West KP. Jr. Subarna KK. *et al.* Night blindness of pregnancy in rural Nepal—nutritional and health risks. Inter J Epidem 1998, 27:231–237.

33.Wallingford JC. Underwood BA. Vitamin A deficiency in pregnancy, lactation, and the nursing child. In: Baurenfeind JC, ed. Vitamin A deficiency and its control. New York, NY, Academic Press. 1986:101–152.

34.Newman V. Vitamin A and breast-feeding: a comparison of data from developed and developing countries. Food and Nutr Bulletin 1994; 15:161–176.

35-Institute of Medicine. Dietary Reference Intake for Vitamin A, Vitamin K, Arsenic, Boron, Chromium, Copper, Iodine, Iron, Manganese, Molybdenum, Nickel, Silicon, Vanadium, and Zinc. Washington, DC: National Academies Press .2001.

36-Thirumdas R. Kothakota A. Pandiselvam R. Bahrami A. Barba FJ. Role of food nutrients and supplementation in fighting against viral infections and boosting immunity: a review. Trends Food Sci Technol. 2021;110:66–77. 37-Caccialanza R. Laviano A. Lobascio F. *et al.* Early nutritional supplementation in noncritically ill patients hospitalized for the 2019 novel coronavirus disease (COVID-19): Rationale and feasibility of a shared pragmatic protocol. Nutrition. **2020**;74:110835.

38-Bikle DD. Chemistry & biology review vitamin D metabolism, mechanism of action, and clinical applications. Chem. Biol 2014; 21: 319–329.

39-Blunt JW. DeLuca HF. Schnoes HK. 25hydroxycholecalciferol. A biologically active metabolite of vitamin D3. Biochem 1968; 7:3317–22.

40-Fraser DR. Kodicek E. Unique biosynthesis by kidney of a biologically active vitamin D metabolite. Nature 1970; 228:764–766.

41-Jones G. Strugnell S. DeLuca HF. Current understanding of the molecular actions of vitamin D. Phys Rev 1998;78:1193–1231.

42-Jones G. DeLuca HF. HPLC of vitamin D and its metabolites. In: Makin HLJ. Newton R. eds. High performance liquid chromatography and its application to endocrinology. Berlin, Springer-Verlag, 1988; 95–139 (Monographs on Endocrinology, vol 30).

43-Brown EM. Pollak M. Hebert SC. The extracellular calcium-sensing receptor: its role in health and disease. Annual Rev Med 1998; 49:15–29.

44-Murad MH. Elamin KB. Abu Elnour NO. Elamin MB. *et al.* Clinical review: The effect of vitamin D on falls: A systematic review and meta-analysis. J Clin Endocri Metab 2011;96(10): 2997-3006.

45-Fraser DR. Kodicek E. Unique biosynthesis by kidney of a biologically active vitamin D metabolite. Nature 1970; 228:764–766.

46-Aliasgharpour M. Technical Points in Vitamin D Measurement Assays .Int J Med Invest 2019; 8(2): 1-5.

47-Food and Nutrition Board. Dietary reference intakes for calcium, phosphorus, magnesium, vitamin D, and fluoride. Washington, DC, National Academy Press 1997.

48-Samuel S . Sitrin MD. Vitamin D's role in cell proliferation and differentiation. Nutr Rev 2008; 66(10) Suppl 2: pp S116–S124.

49-Myszka M. Klinger M. The immunomodulatory role of vitamin D. Postępy Higieny i Medycyny Doświadczalnej 2014; (68) :pp 865–878.

50-Kono K. Fujii H. Nakai K.et al. Antioxidative effect of vitamin D analog on incipient vascular lesion in non-obese type 2 diabetic rats. Am J Nephr 2013; 37(2): pp 167– 174.

51- Sahota O. Understanding vitamin D deficiency. Age and Ageing 2014 ; 43(5): pp 589–591.

52-Newmark HL. Newmark j. Vitamin D and Parkinson's disease—a hypothesis. Movement Disorders 2007; 22(4): pp 461–468.

53-Wang D. Yang Y. Yu G. Shao. Wang Q. Vitamin D and sunlight exposure in newlydiagnosed Parkinson's disease. Nutrients 2016; 8(3) p 142.

54-Zhu D. Lin GY. Wen. S. Wang. WZ. Inverse associations of outdoor activity and vitamin D intake with the risk of Parkinson's disease. J Zhejiang Univer Sci B 2014; 15(10) pp 923–927.

55-Evatt ML. DeLong M. Kumari M. *et al.* High prevalence of hypovitaminosis D status in patients with early Parkinson disease. Arch Neur 2011;68(3) pp 314–319.

56-Muir SW. Montero-Odasso M. Effect of vitamin D supplementation on muscle strength, gait and balance in older adults: A systematic review and meta-analysis. J Am Geriatr Soc 2011;59: 2291-2300.

57-Sato Y. Manabe S. Kuno H. Oizumi K. Amelioration of osteopenia and hypovitaminosis D by 1α -hydroxyvitamin D3 in elderly patients with Parkinson's disease. J Neuro, Neurosur, Psychi 1999; 66(1) pp 64– 68. 58-Knekt P. Kilkkinen A. Rissanen H. Marniemi J. et al. Serum vitamin D and the risk of Parkinson disease. Archi Neuro 2010; 67(7):808–811.

59-Evatt ML. Delong MR. Khazai N. Rosen A. Prevalence of vitamin D insufficiency in patients with Parkinson disease and Alzheimer disease. Archi Neuro 2008; 65(10) 1348–1352. 60-Liu Y. Zhang BS. Serum 25hydroxyvitamin D predicts severity in Parkinson's disease patients. Neuro

Sci 2014; 35(1) 67-71.

61-Sleeman I. Aspray T. Lawson R. *et al.* The role of vitamin D in disease progression in early Parkinson's disease. J Parkinson's Dis 2017; 7(4) pp 669–675.

62-Suzuki M. Yoshioka M. Hashimoto M. et al. Randomized, double-blind, placebocontrolled trial of vitamin D supplementation in Parkinson disease. Am J Clin Nutri 2013;. 97(5) pp 1004–1013.

63-Mercola J. Grant WB. Wagner CL. Evidence Regarding Vitamin D and Risk of COVID-19 and Its Severity. Nutrients 2020;31:12(11): 3361.

64-Ilie PC . Stefanescu S. Smith L. The role of vitamin D in the prevention of coronavirus disease. 2019 infection and mortality. Aging Clin Exp Res 2020;32:1195–98.

65-Rhodes JM. Subramanian S. Laird E. Kenny RA. Editorial: Low population mortality from COVID-19 in countries south of latitude 35 degrees North supports vitamin D as a factor determining severity. Aliment

Pharmacol Ther. 2020;51: 1434–1437.

66-Holick MF. Vitamin D- new horizons for the 21st century. McCollum Award Lecture. Am J Clin Nutri 1994; 60:619–630.

67-Vitamin and mineral requirements in human nutrition : report of a joint FAO/WHO expert consultation, Bangkok, Thailand, 21–30 September 1998. ISBN :924154612 3.

68-Institute of Medicine. Dietary reference intakes for calcium and vitamin D.

Washington, D.C: National Academies Press; 2011.

69-Lachance PA. International perspective: basis, need and application of recommended dietary allowances. Nutri Rev 1998; 56:S2–S4. 70-Colombo ML. An update on vitamin E, tocopherol and tocotrienol-perspectives. Molecules 2010;. 15(4)

pp 2103–2113.

71-Beharka A. Redican S. Leka L. Meydani SN. Vitamin E status and immune function. Methods Enzy 1997; 282 pp 247–263.

72-Cesari M. Pahor M. Bartali B. *et al.* Antioxidants and physical performance in elderly persons: the Invecchiare in

Chianti (InCHIANTI) study . Am J Clin Nutri 2004; 79(2) pp 289–294.

73-Cherubini A. Andres-Lacueva MC. *et al.* Vitamin E levels, cognitive impairment and dementia in older persons: the InCHIANTI study. Neuro Aging, 2005; 26(7): pp 987–994. 74-Sies H, Berndt C. Jones DP. Oxidative stress .Annual Rev Bioch 2017; 86(1) pp 715– 748.

75-Kagan VE. Recycling and redox cycling of phenolic antioxidants. Annals New York Acad Sci 1998; 854:425–434.

76-National Research Council, Committee on Dietary Allowances: Recommended Dietary Allowances.10th revised ed. Washington, D.C, National Academy of Sciences;1989.

77-Traber MG. Regulation of human plasma vitamin E. In: Sies H, ed. Antioxidants in disease mechanisms and therapeutic strategies. San Diego, CA, Academic Press;1996:49–63.

78-Simoin EJ. Eisengart A. Sundheim L. *et al.* The metabolism of vitamin E: II. Purification and characterization of urinary metabolites of α -tocopherol. J Biol Chem 1956;.221:807.

79-McLaren DS. et al. Fat soluble vitamins. In: Garrow JS, James WPT, eds. Human nutrition and dietetics. Edinburgh, Churchill Livingstone 1993; 208–238.

80-Traber MG. Sokol RJ. Burton GW. *et al.* Impaired ability of patients with familial isolated vitamin E deficiency to incorporate atocopherol into lipoproteins secreted by the liver. J Clin Invest 1990; 85:397-407.

81-Clarke MW. Burnett JR. Croft KD. Vitamin E in human health and disease. Criti Rev Clin Lab Sci 2008; 45(5) pp 417–450.

82-Di Mascio P. Murphy ME. Sies H. Antioxidant defense systems: The role of carotenoids, tocopherols, and thiols. Am J Clin Nutr 1991; 53:194S–200S.

83.Brown KM. Morrice PC. Duthie GG. Erythrocyte vitamin E and plasma ascorbate concentrations in relation to erythrocyte peroxidation in smokers and non-smokers: dose–response of vitamin E supplementation. Am J Clin Nutr 1997; 65:496–502.

84- Li D. Saldeen T. Romeo F. Mehta JL. Different isoforms of tocopherols enhance nitric oxide synthase phosphorylation and inhibit human platelet aggregation and lipid peroxidation: Implications in therapy with vitamin E. J Cardiovasc Pharmacol Ther 2001; 6:155–61.

85-Liu M. Wallmon A. Olsson-Mortlock C. Wallin R. SaldeenT. Mixed tocopherols inhibit platelet aggregation in humans: Potential mechanisms. Am J Clin Nutr 2003; 77:700–6. 86-de Rijk MC. Breteler MM.den Breeijen J. et al. Dietary antioxidants and Parkinson disease. The Rotterdam

Study, Arch Neuro1997; 54(6) pp 762–765.

87-Zhang SM. Hernan MA. Chen H. Spiegelman D. et al. Intakes of vitamins E and C, carotenoids, vitamin supplements, and PD risk," Neuro 2002; 59(8) pp 1161–1169.

88-Logroscino G. Marder K. Cote L. Tang MX. *et al.* Dietary lipids and antioxidants in Parkinson's disease: a population-based, case-control study," Annals of Neuro1996;39(1) pp 89–94.

89-Fernández-Quintela A . Milton-Laskibar I. Trepiana J. *et al.* Key aspects in nutritional management of COVID-19 Patients. J. Clin. Med 2020; 9: 2589. 90-Maggini S. Beveridge S. Sorbara PJ. Senatore G. Feeding the immune system: the role of micronutrients in restoring resistance to infections, CAB reviews: perspectives in agriculture, Vet. Sci. Nutr. Nat. Resour.2008; 3 : 1–21.

91-Prenticec S. They are what you eat: can nutritional factors during gestation and early infancy modulate the neonatal immune response? Front. Immunol 2017;8: 1641.

92-De la Fuente M. Hernanz A. Guayerbas N. Victor M. Arnalich F. Vitamin E ingestion improves several immune functions in elderly men and women. Free Radic. Res. 2008;42 :272–280.

93- Sano M. Ernesto C. Thomas RG. Klauber MR. *et al.* A controlled trial of selegiline, alphatocopherol, or both as treatment for Alzheimer's disease. The Alzheimer's Disease Cooperative Study. N Engl J Med 1997;336:1216–22.

94- Meydani M. Vitamin E. Lancet 1995; 345:170-5.

95-Mielech A. Puscion-Jakubik A. *et al.* Vitamins in Alzheimer Disease- Review of the latest reports. Nutrients2020;12:3458.

96-.Mangialasche F. Kivipelto M. Mecocci P. Rizzuto D.*et al.* High plasma levels of vitamin E forms and reduced Alzheimer's disease risk in advanced age. J Alzheimer's Dis 2010; 20:1029–37.

97-Mangialasche F. Westman E. Kivipelto M. Muehlboeck JS. *et al.* Classification and prediction of clinical diagnosis of Alzheimer's disease based on MRI and plasma measures of α -/ γ -tocotrienols and γ -tocopherol. J Intern Med 2013; 273:602–21.

98-keen MA. Hassan L. Vitamin E in dermatology. Indian Dermatol Online J 2016;7(4):311-5.

99-Rattanawiwatpong P. Wanitphakdeedecha R. Bumrungpert A. *et al.* Anti aging and brightening effects of a topical treatment contaninig vitamin C, vitamin E, and raspberry leaf cell culture extract; A split face randomized controlled trial. J cosmet Dermatol. 2020;19(3):671-676.

100-Lintner K. Gerstein F. Solish N. A serum containing vitamin C and E and a matrix – repair tripeptide reduces facial signs of aging as evidenced by primos analysis and frequently repeated auto perception. J Cosmet Dermatol 2020;19(12): 3262-3269.

101- Norton L. Further observations on the yellow nail syndrome with therapeutic effect of oral alpha-tocophero. Cutis. 1985;36:457–62.

102- Al Hawsawi K. Pope E. Yellow nail syndrome. Pediatr Dermatol. 2010;27:675–6.

103- Barbosa E. Faintuch J. Machado Moreira EA. Gonçalves da Silva VR. *et al.* Supplementation of vitamin E, vitamin C, and zinc attenuates oxidative stress in burned children: A randomized, double- blind, placebo-controlled pilot study. J Burn Care Res 2009;30:859–66.

104-Suttie JW. Vitamin K. In: Diplock AD. ed. Fat-soluble vitamins: their biochemistry and applications. London, Heinemann 1985:225– 311.

105-Krueger T. Westenfeld R. Schurgers L. *et al.* Coagulation meets calcification: the vitamin K system. Int J Artif Organs 2009;32:67–74.

106-Booth SL. Suttie JW. Dietary intake and adequacy of vitamin K. J Nutr 1998;128:785–8.

107- Schurgers LJ. Cranenburg EC. Vermeer C. Matrix Gla-protein: the calcification inhibitor in need of vitamin K. Thromb Haemost 2008;100:593–603.

108-Beulens JW. Booth SL. van den Heuvel EG. *et al.* The role of menaquinones (vitamin K(2)) in human health. Br J Nutr 2013;110:1357–68.

109-Thane CW. Bolton-Smith C. Coward WA. Comparative dietary intake and sources of phylloquinone (vitamin K1) among British adults in 1986–7 and 2000–1. Br J Nutr 2006;96:1105–15.

110-Schurgers LJ. Vermeer C. Determination of phylloquinone and menaquinones in food.

Effect of food matrix on circulating vitamin K concentrations. Haemostasis 2000;30:298–307.

111-Elder SJ. Haytowitz DB. Howe J. *et al.* Vitamin K contents of meat, dairy, and fast food in the US. Diet. J Agric Food Chem 2006;54:463–7.

112- Suttie JW. Vitamin K: In Health and Disease. CRC Press, 2009.

113-Olson RE. The function and metabolism of vitamin K. Annu Rev Nutr 1984; 4:281-337.

114-Mihatsch WA. Braegger C. Bronsky J. Campoy C. Domellöf M.*et al.* ESPGHAN Committee on Nutrition. Prevention of vitamin K deficiency bleeding in newborn infants: A Position Paper by the ESPGHAN Committee on Nutrition. J Pediatr Gastroenterol Nutr 2016; 63(1):123-9.

115-Shearer MJ. McBurney A. Barkhan P. Studies on the absorption and metabolism of phylloquinone (vitamin K1) in man. Vitam Horm 1974;32:513-42.

116-Furie B, Furie BC. Molecular basis of vitamin K-dependent g-carboxylation. Blood 1990; 75:1753–62.

117-Davie EW. Biochemical and molecular aspects of the coagulation cascade. Thromb Haemost 1995; 74:1–6.

118-Shearer MJ. Newman P. Metabolism and cell biology of vitamin K. Thromb Haemost 2008;100:530-47.

119-Kohlmeier M .*et al.* Transport of vitamin K to bone in humans. J Nutr 1996; 126 (Suppl.):S1192–S1196.

120-Shea MK. O'Donnell CJ. Hoffmann U. et al. Vitamin K supplementation and progression of coronary artery calcium in older men and women. Am J Clin Nutr 2009;89:1799–807.

121-Schurgers LJ. Spronk HM. Soute BA. et al. Regression of warfarin-induced medial elastocalcinosis by high intake of vitamin K in rats. Blood 2007;109:2823–31.

122-Von Kries R, Shearer MJ, Göbel U. Vitamin K in infancy. Euro J Pediat, 1988, 147:106–112. 123-Shunsuke A. Akira S. Vitamin K deficiency bleeding in infancy. Nutrients 2020; 12:780

124-Jagannath VA. Thaker V.Chang AB. Price AL. Vitamin K supplementation for Cystic Fibrosis. Cochrone Database of Systematic Reviews 2020; issue (6).Art#:CD008482.

125- Wilson A: Disorders of vitamins; Deficiency, excess, and errors of metabolism. In: Harrison's principles of internal medicine. 12th ed. Petersdorf RG. Adams RD. Braunwald E.*et al.* New York, McGraw Hill. Book Company.1991

126- Dofferhoff ASM. Piscaer L. Schurgers LJ. et al. Reduced vitamin K status as a potentially modifiable risk factor of severe Covid-19. Clin Infect Dis 2020; 27:1258.

127-Janssen R .Visser MPJ. Dofferhoff ASM. *et al.* Vitamin K metabolism as the potential missing link between lung damage and thromboembolism in coronavirus disease. Br J Nutr 2020; 7:1-8.

128- Mehri A. Trace elements in human nutrition (II) – An update. Int J Prev Med 2020;11:2.

129-Hall JA. Grainger JR. Spencer SP. *et al.* The role of retinoic acid in tolerance and immunity. Immunity 2011;35(1):13-22.

130-Field CJ. Johnson IR. Schley PD. Nutrients and their role in host resistance to infection. J Leukoc Biol. 2002;71(1):16-32.

131-AL-Sumiadai MM. Ghazzay H. Al-Dulaimy WZS. Therapeutic effect of vitamin A on severe COVID-19 patients. Eurasia J Biosci.2020;14:7347-50.

132-Michienzi SM. Badowski ME. Can vitamins and /or supplements provide hope against coronavirus? Drugs Context. 2020;9:2020-5.

133-Wu JZ P. Zha P. Treatment strategies for reducing damages to lungs in patients with coronavirus and other infections. Preprints. 2020;2020020116.11.

134-Jones G. Strugnell SA. DeLuca HF. Current understanding of the molecular actions

of vitamin D. Physiological Reviews 1998; 78:1193–1231.

135- Misra M. Pacaud D. Petryk A. *et al.* Drug and Therapeutics Committee of the Lawson Wilkins Pediatric Endocrine Society. Vitamin D deficiency in children and it's management: review of current knowledge and recommendations. Pediatrics 2008; 122:398– 417.

136-Holick MF. High prevalence of Vitamin D inadequacy and implications for health. Mayo Clin Proc. 2006; 81:353–73.

137-Holick MF. Vitamin D: importance in the prevention of cancers, type 1 diabetes, heart disease, and osteoporosis. Am J Clin Nutr 2004; 79:362–71.

138-Panfili FM. Roversi M. D'Argenio P. *et al.* Possible role of vitamin D in Covid-19 infection in pediatric population. J Endocrinal Invest. 2021;44(1):27-35.

139-Grant WB. Lahore H. McDonnell SL. *et al.* Evidence that vitamin D supplementation could reduce risk of influenza and Covid-19 infection and deaths. Nutrients 2020;12(4):988. 140-Xiuzhen Z. Ming Z. Chunxiao Li.*et al.* Benefits of vitamins in the treatment of Parkinson's disease.

Oxid Med Cell Longev 2019; 2019: 9426867 . 141-Delgado-Roche L. Mesta F. Oxidative stress as key player in severe acute respiratory syndrome coronavirus (SARS-CoV) infection. Arch Med Res. 2020;51(5):384–387.

142-Hariharan A. Hakeem AR. Radhakrishnan S. Reddy MS. Rela M. The role and therapeutic potential of NF-Kappa-B pathway in severe COVID-19 patients. Inflammopharmacology 2021;29 (1):91–100.

143-Al-Lami RA. Urban RJ. Volpi E. Algburi AMA. et al. Sex hormones and novel corona virus infectious disease (COVID- 19). Mayo Clin Proc 2020; 95(8):1710–1714.

144-de Las Heras N. Martín Giménez VM. Ferder L. Manucha W.*et al.* Implications of oxidative stress and potential role of mitochondrial dysfunction in COVID-19: therapeutic effects of Vitamin D. Antioxidants (Basel) 2020;9(9):897.

145-Kozlov EM. Ivanova E. Grechko AV. Wu WK. Starodubova AV. Orekhov AN. Involvement of oxidative stress and the innate immune system in SARS-CoV-2 infection. Diseases 2021;9 (1):17.

146-Azkur AK. Akdis M. Azkur D. *et al.* Immune response to SARS- CoV-2 and mechanisms of immunopathological changes in COVID- 19. Allergy 2020;75(7):1564– 1581.

147-Okebukola PO. Kansra S. Barrett J. Vitamin E supplementation in people with cystic fibrosis. Cochrane Data base Syst Rev.2020.

148-Dam H. Schonheyder F. Tage-Hansen E. Studies on the mode of action of vitamin K. Biochem J 1936;30:1075-9.

149-Booth SL. Roles for vitamin K beyond coagulation. Annu Rev Nutr 2009;29:89–110.

150-Maresz K. Proper calcium use: vitamin K2 as a promoter of bone and cardiovascular health. Integr Med (Encinitas). 2015;14 (1):34– 39.

151-Mukai K. Itoh S. Morimoto H. Stoppedflow kinetic study of vitamin E regeneration reaction with biological hydroquinones (reduced forms of ubiquinone, vitamin K, and tocopherolquinone) in solution. J Biol Chem. 1992;267(31):22277–81.

152-Vervoort LM. Ronden JE. Thijssen HH. The potent antioxidant activity of the vitamin K cycle in microsomal lipid peroxidation. Biochem Pharmacol. 1997;54(8):871–876.

153-Hodges SJ. Pitsillides AA. Ytrebø LM. Soper R. Anti-inflammatory actions of vitamin K. In: Vitamin K2: Vital for Health and Wellbeing. 2017;153

154.Hamblin J. Why Some People Get Sicker Than Others ? COVID- 19 is proving to be a disease of the immune system. This could, in theory, be controlled. 2020. Available from: https://www.theatlan tic.com/health/archive/2020/04/coronavirusimmune-response/ 610228.

155.Velavan TP. Meyer CG. Mild versus severe COVID-19: laboratory markers. Int J Infect Dis. 2020;95:304–307.

156-Chakraverty R. Davidson S. Peggs K. *et al*. The incidence and cause of coagulopathies in an intensive care population. Br J Haematol. 1996;93(2):460–463.

157-Crowther MA. McDonald E. Johnston M. Cook D. Vitamin K deficiency and D-dimer levels in the intensive care unit: a prospective cohort study. Blood Coagul Fibrinolysis. 2002;13(1):49–52. 158-Turshudzhyan A. Anticoagulation options for Coronavirus Disease 2019 (COVID-19)induced coagulopathy. Cureus. 2020;12(5):e8150.

159-Piscaer I. van den Ouweland JMW. Vermeersch K. *et al.* Low Vitamin K status is associated with increased elastin degradation in chronic obstructive pulmonary disease. J Clin Med. 2019;8 (8):1116.

160- Maqbool A. Stallings VA .Update on fatsoluble vitamins in cystic fibrosis. Current Opinion in Pulmonary Medicine 2008; *14* (6): 574–81.

Tables

Table 1- Recommended Nutrient Intakes (RNIs) for Vitamin D, by Group

Groups	RNI (mg/day) ^a	
Infants and children		
0–6 months	5	
7–12 months	5	
1–3 years	5	
4–6 years	5	
7–9 years	5	
Adolescents		
10–18 years	5	
Adults		
19–50 years	5	
51-65 years	10	
65+ years	15	
Pregnant women	5	
Lactating women	5	

^a Units: for vitamin D, 1 IU = 25ng, 40 IU = $1\mu g$, 200 IU = $\mu 5g$,400IU = $10\mu g$, 600 IU = $15\mu g$, 800 IU = $20\mu g$. IU-International Units

Vitamins **Biological Function Deficiency** Disease Sources Toxicity Disease Cases Cases Α Normal vision Xerophthalmia Hepatotoxicity-Bone E Eggs-(Retinol) Maintenance of epithelial cellular Night blindness. Abnormality. Meat integrity Keratinization of the central Headache &Skin Dairy Neuronal differentiation Epithelium. Dry mucous Desqamation products Antioxidant activity Membranes. Low resistance Vegetables Immune response to infection. Cystic Fibrosis is Reproduction a known risk factor of VADa. Oral supplementation is currently being investigated in the treatment of Covid-19. D Mineralization of bones Rickets in children. Rare Oily fish, (C & skeletal health Osteomalacia in adults & Mushrooms (Cholecalciferol) Egg Yolks Hom Homeostasis of calcium, Osteaporosis increased phosphorus, & magnesium risk for PDb & Covid-19 E Powerful antioxidant for Rare. Only in infants & adults Low except at very high Canola Oil (αpolvunsaturated with an inherited or acquired Supplement doses. Olive Oil Margarine (Tocopherol) fatty acids (PUFAs) condition that impairs Hemorrhagic within the membranes the vitamin absorbance Almonds toxicity Preventing cell membrane oxidation, and in those who cannot Peanuts Maintaining neurological functions. E absorb Cognitive function, physical dietary fat or have rare Meats, Dairy disorders of fat metabolism. performance, Le Regulation of gene expression and Decreased antioxidant activity Leafy skin health. in ADc . Little Greens The vitamin is stored in most tissue, evidence for prophylactic or Fortified with the largest amount stored in therapeutic agent against Cereals. adipose tissue. Covid-19 Vitamin E is one of the treatment for yellow nail syndrome Coagulation/ Blood Clotting N Naturally occurring Green leafy In Adults, primary vitamin K Κ Essential for maintaining bone and Deficient states manifest as vitamin vegetables (Phylloquinone) Olive Oil & Cardiovascular health, and bleeding are unknown except K (K1 &K2) toxicity is metabolism. when the absorption of the rare. Soyabean Oil Acting as a potent antioxidant vitamin is impaired or during Menadione (K3), Reducing the lipid per oxidation in the long term antibiotic or synthetic cell by producing vitamin K anticoagulant treatment . Form that causes liver hydroquinone. Vitamin K Deficiency Damage & neonatal It also has an anti inflammatory Bleeding (VKDB) is rare and haemolysis no activity. it is potentially Life longer used threatening bleeding disorder therapeutically . of early infancy .Thrombosis in Covid - 19 patients with elevation of D-dimers.

Table2: Fat Soluble Vitamins Biological Functions & Deficiency/ Toxicity Disease Cases in Human Health

a-VAD- Vitamin A Deficiency; b-PD-Parkinson's Disease; c-AD -Alzheimer' Disease