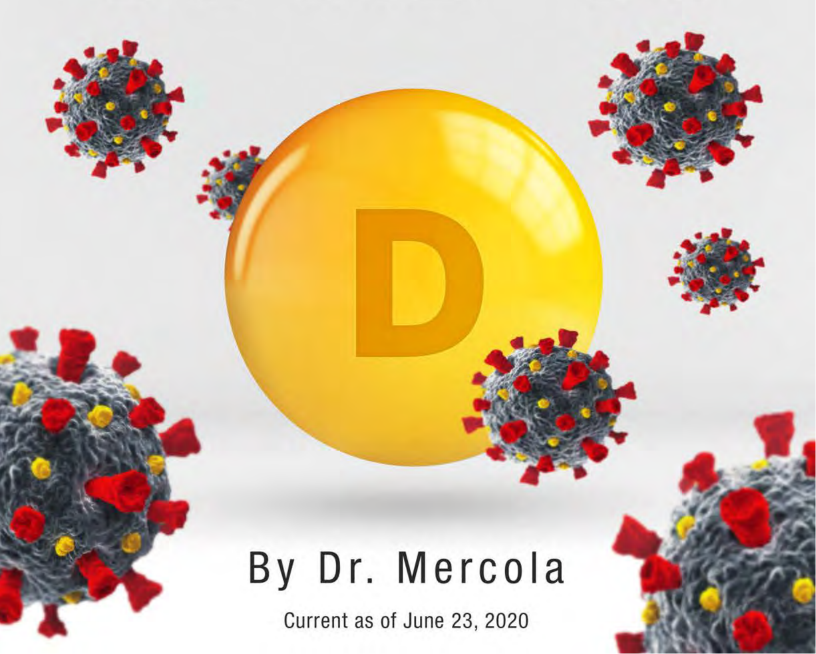
Vitamin D

In the Prevention of

COVID-19



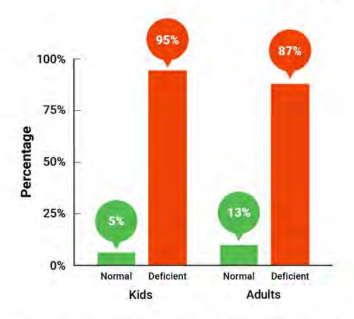
Vitamin D deficiency represents a global pandemic afflicting more than one billion individuals across all age groups worldwide¹ and it has been estimated that in excess of one billion people have vitamin D deficiency.²⁻⁴ The current pandemic of vitamin D deficiency has collided with the COVID-19 pandemic and likely radically increased the number of deaths because of vitamin D insufficiency.⁵

So just how many people are suffering from not having enough vitamin D?

A lot more than you probably thought or understood.

As you can see in Figure 1 below 95% of children and 87% of adults have less than the ideal level of vitamin D in their blood which is 40 ng/ml or 100 nmol/liter. Only 5% of children and 13% of adults have achieved ideal levels. But this is for all ethnicities. As you can see Figure 19 at the end of the document less than one percent of black children have achieved this healthy level.

Vitamin D Levels by Age



SOURCE: Centers for Disease Control and Prevention (CDC). National Center for Health Statistics (NCHS). National Health and Nutrition Examination Survey. Hyattsville, MD: U.S. Department of Health and Human Services, Centers for Disease Control and Prevention, 2013-2014. https://wwwn.cdc.gov/nchs/nhanes/ContinuousNhanes/Default.aspx?BeginYear=2013

Figure 1

Although there are currently no prospective controlled studies demonstrating vitamin D's effectiveness in COVID-19 there are many such studies underway. One can visit the clinical trials registry to review the current state of these trails.^{6 7} As of early June 2020 there were over 20 studies in progress on the use of vitamin D in COVID-19.

The purpose of this report is to help you understand why it is so important to optimize your vitamin D level in order to have healthy immune functions, and then provide you with a detailed strategy for how to do that.

This report can be an invaluable tool to share with your family and community to help prepare for a second wave of the pandemic, which is expected in the fall.

The interest in the health-promoting effects of vitamin D has increased substantially during the 21st century. There are approximately 5,500 vitamin D-related articles indexed to the US National Library of Medicine database in the past five years. The observational studies on vitamin D have received a considerable amount of attention due to a vast body of publications reporting inverse associations between vitamin D status and multiple diseases, including COVID-19.

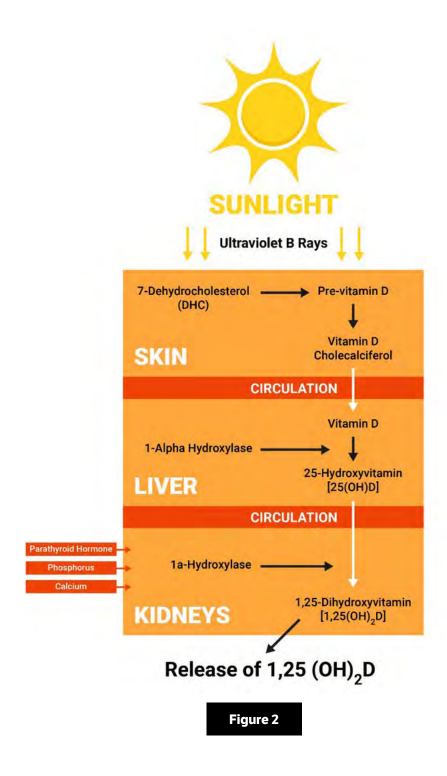
Even a former director of the Centers for Disease Control and Prevention, Dr. Tom Frieden, proposed using vitamin D to combat the COVID-19 pandemic on 23 March 2020.⁸ There have been many recent calls for widespread high-dose vitamin D supplementation in the prevention and mitigation of COVID-19.⁹⁻¹²

A recent June, 2020 article caused the editors at the BMJ Nutrition Prevention and Health to write the following supportive statement about vitamin D and COVID-19:

"Categorical general statements about the lack of benefit from vitamin D are not supported by any evidence at this time, not least because a growing number of observations and study results that point to an important role (of vitamin D)." ¹³

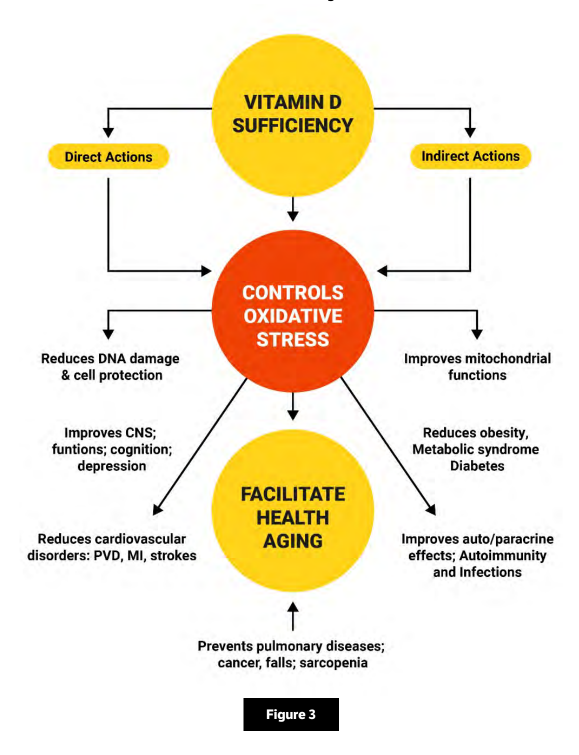
Vitamin D_3 is an ancient molecule that is produced from the direct cholesterol precursor (7-dehydrocholesterol) which is normally present in your skin using energy provided by the *UV-B* component of sunlight in a reaction that does not require any enzyme assistance.¹⁴ In its classical pathway, vitamin D_3 is converted to its single hydroxy form (25-hydroxyvitamin D) in your liver and then to its double hydroxy form (1,25-dihydroxyvitamin D) in your kidneys and even in your immune cells that fight infection.¹⁻³ ¹⁵

Vitamin D differs from most vitamins, in that your body can produce it on its own with exposure to sunlight, and that its primary active metabolite is a steroid hormone. Unlike most vitamins, which act as antioxidants or enzyme co-factors, the $1,25(OH)_2D$ form of vitamin D works by binding to the vitamin D receptor that is present in the cell membrane, or the nucleus. Once vitamin D activates the receptor it becomes a master regulator of cell function (Fig. 2).



Until the 21st century, vitamin D was primarily recognized for its role in the regulation of calcium and bone health and the prevention of rickets.¹⁶ In the last 20 years, however, research has shown that vitamin D also has profound influences on immune cells and causes a general lowering of inflammation.^{17 18} It is a powerful epigenetic regulator influencing over 2,500 genes¹⁹

and impacting dozens of our most serious health challenges, like heart disease and cancer, autoimmune diseases like MS,²⁰ and others listed in Figure 3 below.

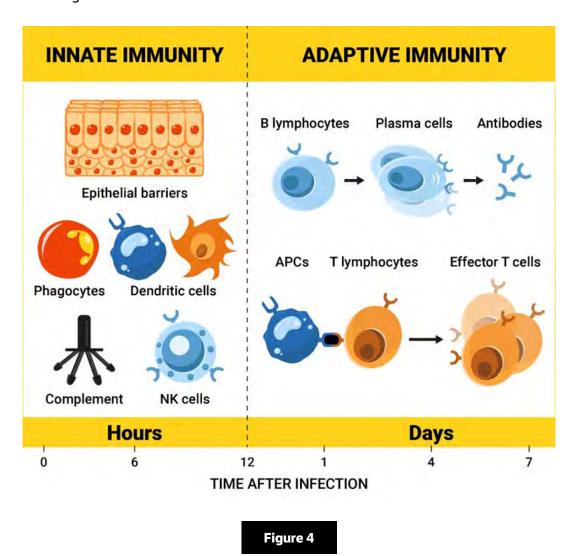


Your Innate and Adaptive Immune System

n the frame of infectious diseases, the purpose of your immune system is to recognize invading pathogens, prevent their spread, and eliminate them from your body. This extraordinarily complex system relies on billions of cells patrolling your body and a dynamic complex network.²¹

To help you understand how vitamin D impacts your immune system, it is first important to appreciate some fundamental elements of your innate and adaptive immune system. Your immune system comprises two distinct but interacting types of immunity: innate and adaptive.

Your innate immunity kicks in hours following a foreign pathogen, while your adaptive immunity takes days to react but provides long term, typically lifelong immunity, to an infection as illustrated in Figure 4 below.



Innate Immune System

Your innate immune system is your first line of defense against infections, and rapidly fights against invading pathogens. It responds in a generic way without conferring long-lasting and specific immunity.

Unlike your adaptive immune responses, innate immune responses are always general, or not specific to a particular pathogen and depend upon a group of cells and proteins that recognize conserved features of microbes that quickly promote clearance of infectious agents.

Your innate immune system includes physical barriers, like your skin and the cells lining your gut and blood vessels, and chemical barriers like your saliva, and stomach acid. These barriers help to block the entry of disease causing organisms into your body.²²

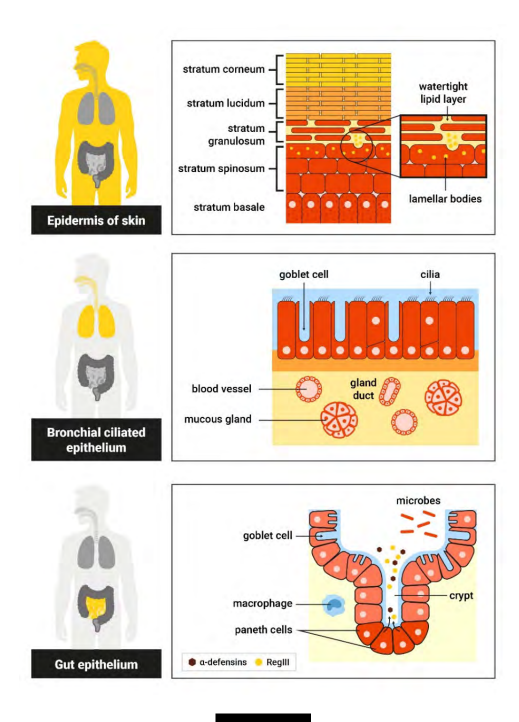


Figure 5

Vitamin D is a well-known regulator of the physical barrier portion of your innate immune system and is responsible for improving the epithelial cells that line your intestines. It also modulates your bowel's immune system. Low levels of vitamin D will increase your gut permeability and allow pathogens to sneak into your blood stream causing low-grade inflammation.

White blood cells are also part of your innate immune system and they serve as the primary initial defenders against pathogens in your body. ²³ Neutrophils are your most abundant white blood cell and contribute to your first line of defense against microbial pathogens.

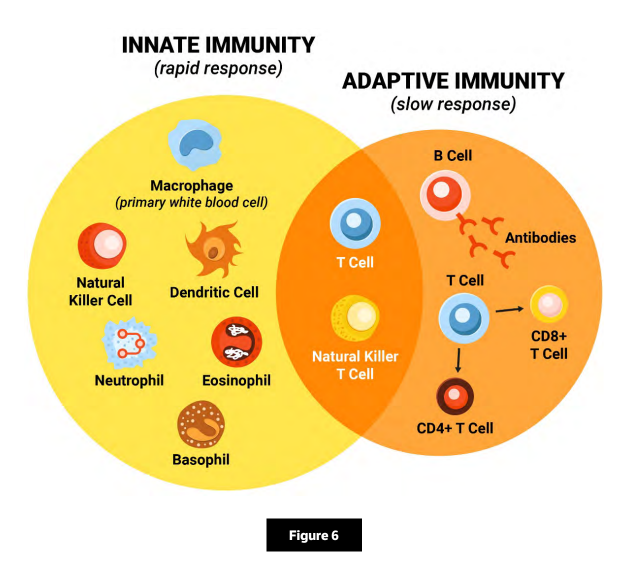
Neutrophils can clear microbes through a process called phagocytosis, or simply digesting them inside the your white blood cells where they are exposed to reactive oxygen species, which are generated in response to the pathogens, which further leads to the production of biologically active antimicrobial molecules. ²⁴

Dendritic cells play a key role in innate immune and adaptive immune responses. As the strongest antigen-presenting cells, they effectively stimulate the activation of T-lymphocytes and B-lymphocytes, thus combining innate and adaptive immunity.

Dendritic cells permanently survey your body and are specialized in absorbing antigens from pathogens. Upon exposure to inflammatory signals, they mature and migrate to your lymph nodes, and present their captured antigens to the T cells, thereby priming an antigen-specific adaptive immune response.

Macrophages are another type of white blood cell that add to the first line of your innate defense against pathogens and are important in engulfing bacteria, as white blood cells do, but also in making and secreting a whole host of inflammatory and anti-inflammatory signaling proteins.

For a more complete picture of the cells involved in your innate and active immunity you can view Figure 6 below.



Adaptive Immune System

our adaptive immune system is primarily composed of your T and B lymphocytes as represented in Figure 6 above. Compared with your innate immunity, your adaptive immunity is slower to start but typically strong enough to finalize the clearance of infections that elude your innate immunity. Adaptive immunity is best characterized by its specificity to foreign antigens and its ability to generate long-lasting immune memory.

The activation of your adaptive immune system often starts with the antigen presentation by innate cells to T helper cells, which leads to their interaction with naive B cells. This then assists in activating and differentiating them into memory and antibody-secreting B cells that produce

the antibodies to protect you from future infections and that are measured to demonstrate protective immunity.

T cells, CD4+ T cells, and CD8+ T cells, particularly, play a significant antiviral role by balancing the combat against pathogens and the risk of developing autoimmunity or overwhelming inflammation.²⁵

CD4+ T cells promote the production of virus-specific antibodies by activating T-dependent B cells. However, CD8+ T cells are toxic to pathogens and can kill viral infected cells. CD8+ T cells account for about 80% of total inflammatory cells in the lungs of coronavirus infected patients and play a vital role in clearing the virus in infected cells and inducing immune injury.²⁶

The activation of naive CD4+ T cells generates different helper T-cell classes, which differ according to the type of immune response they produce. Thus, the type 1 response T helper cells supports cell-mediated immunity, whereas type 2 helper T-cell response mediates the humoral response.²⁷

This entire process is summarized in Figure 7 below.

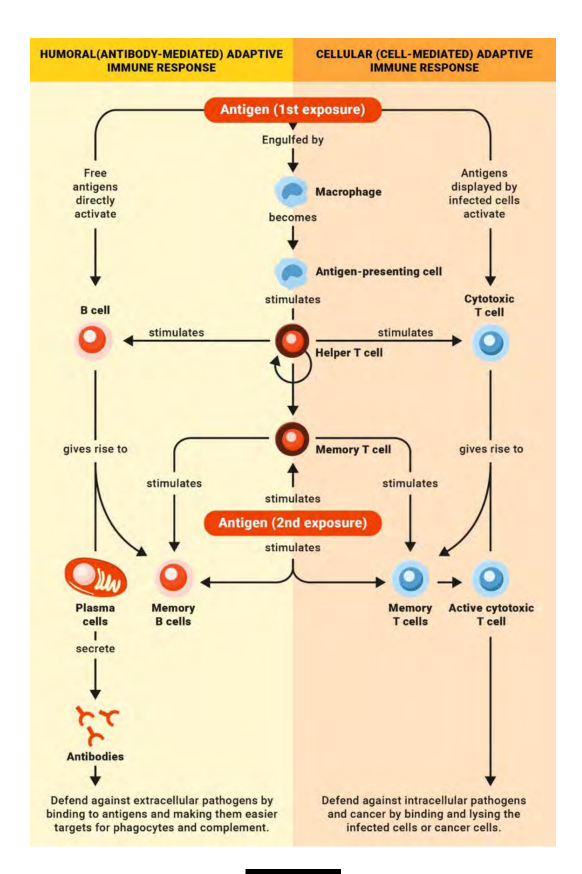
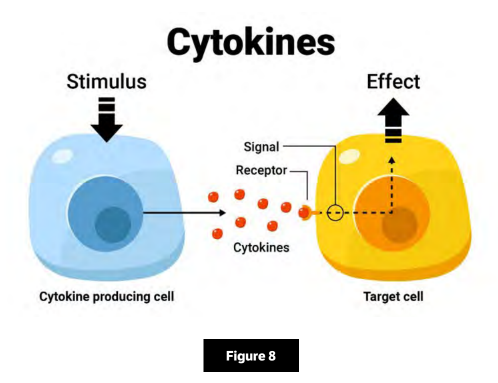


Figure 7

Vitamin D, Cytokine Storms and COVID-19

Cytokines are small proteins secreted by cells in your innate and adaptive immune systems. They serve to regulate diverse functions in your immune response. Cytokines are released by cells into your circulation or directly into your tissues. The cytokines locate target immune cells and interact with receptors on the target immune cells by binding to them. The interaction triggers or stimulates specific responses by the target cells.



In response to bacterial and viral infections such as COVID-19, your innate immune system generates both pro-inflammatory and anti-inflammatory cytokines.²⁸ The inflammatory response plays a crucial role in the clinical manifestations of COVID-19. SARS-CoV-2 triggers an immune response against the virus, which, if uncontrolled, may result in lung damage, functional impairment, and reduced lung capacity.²⁹⁻³²

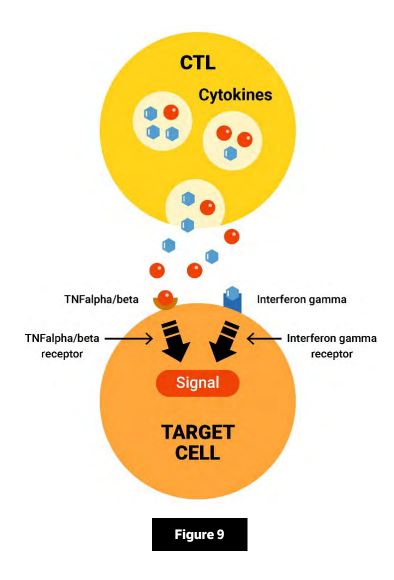
The SARS-CoV-2 viral infection-related inflammation and the subsequent cytokine storm in severe cases plays a crucial role in patient survival.³³ The extensive and uncontrolled release of proinflammatory cytokines is termed the cytokine storm. Clinically, the cytokine storm commonly presents as systemic inflammation and multiple organ failure.³⁴

The inflammatory cytokines that mediate this response are TNF α and interleukins are produced at an early stage of your innate immune response to the virus. These cytokines, among others, contribute to the recruitment and activation of cells of your adaptive immune response. ³⁵ ³⁶

Two recent reviews carefully covers the physiology of how vitamin D specifically lowers the risk of cytokine storms,^{37 38} but the process is summarized below.

There is compelling research demonstrating that vitamin D can improve endothelial stability even in cytokine storms.³⁹ This may be due to vitamin D's role in modulating your T helper cell and cytokine production, but also through promoting T regulatory cells, which are responsible for anti-infectious action, for suppressing immune responses, and for limiting inflammatory processes⁴⁰ for which vitamin D may play an important role.⁴¹

Vitamin D helps to down-regulate the immune responses mediated by your T helper cells, thus inhibiting the production of pro-inflammatory cytokines, such as type 1 interferon gamma, and interleukins like IL-6, IL-2, along with tumor necrosis factor alpha (TNF- α)⁴² ⁴³ as indicated in Figure 9 below.



It has been well established that vitamin D deficiency enhances the cytokine storm. $^{44-46}$ This is because vitamin D modulates your adaptive immunity and suppresses responses mediated by your T helper cells by repressing production of inflammatory cytokines like TNF α and interleukins like IL-2 and interferon gamma. $^{47-48}$ Furthermore, vitamin D promotes stimulates the production of your T regulatory cells that inhibit inflammatory processes. 49

There appears to be enormous value of vitamin D in COVID-19 infections as administering it reduces the expression of these pro-inflammatory cytokines and increases the expression of anti-inflammatory cytokines by macrophages.^{50 51} It has been shown that vitamin D regulates the inflammatory response, altering the pro-inflammatory/anti-inflammatory balance toward an anti-inflammatory state that controls the inflammatory burst once it is triggered.⁵²

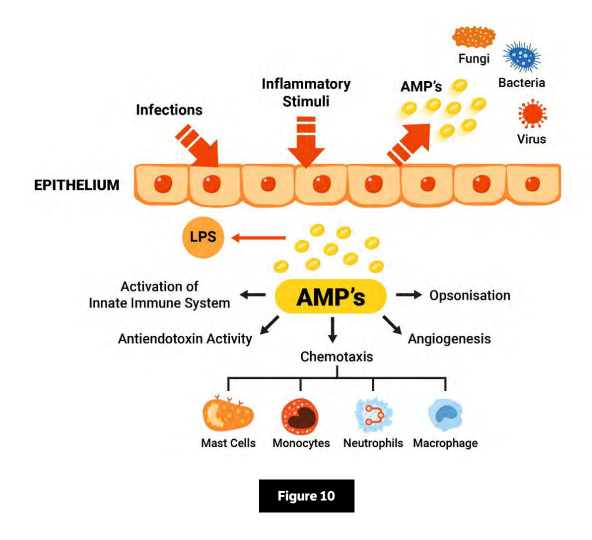
Cell culture studies have shown that vitamin D decreases the expression of pro-inflammatory cytokines, increases the production of antiviral proteins, and also has antiviral efficacy, especially facing enveloped viruses; therefore, it would likely be effective against the enveloped SARS-CoV-2 causing COVID-19.⁵³ ⁵⁴

As further proof that vitamin D reduces inflammation, a recent study showed a strong correlation with vitamin D levels and C-reactive protein (CRP). Given that CRP is a surrogate marker for cytokine storm this supports a role for vitamin D in reducing complications attributed to unregulated inflammation due to the COVID-19 cytokine storm.⁵⁵

Vitamin D Helps Your Immune Cells Create Antimicrobial Peptides

Vitamin D receptors have been identified in nearly all of your immune cells, including monocytes, B and T lymphocytes, white blood cells, macrophages and dendritic cells, as well as the epithelial cells in your lungs. ⁵⁶ This is important because if you have sufficient vitamin D in your blood it can activate these cells to create what researchers call antimicrobial peptides (AMPs). ⁵⁷

Many studies have shown that vitamin D activates your immune cells to produce AMPs which include molecules known as cathelicidins and defensins. AMPs have a broad spectrum of activity, not only microbial but also antiviral, and have been shown to inactivate the influenza virus. AMPs are the result of, among other effects, the destruction of envelope proteins done by cathelicidin.



Cathelicidins are a distinct class of proteins present in the innate immunity of mammals. In humans the primary form of cathelicidin is known as LL-37.⁶⁴ LL-37 also blocks viral entry into the cell in a similar manner to what is seen with other antimicrobial peptides.⁶⁵

Epidemiologic evidence describes a positive vitamin D related immune effect that includes many studies which feature enveloped viruses like SARS-CoV-2. This supports the notion that LL-37's anti-viral effects may be partially mediated by envelope disruption⁶⁶ as LL-37's anti-microbial effect is linked to its ability to disrupt the lipid envelopes of viruses through electrostatic interactions.⁶⁷

Vitamin D also regulates another type of AMP called beta defensin 2. Its antiviral effects result from its impact on your white blood cells like neutrophils and monocytes.⁶⁸

Vitamin D Deficiency Increases Your Risk for COVID-19

A recent retrospective analysis at the University of Chicago of over 4,000 patients⁶⁹ was designed to examine whether vitamin D deficiency and treatment are associated with testing positive for COVID-19. They found that vitamin D deficiency that was not sufficiently treated was associated with an increased risk for COVID-19 infection.

Another observational study involving 212 patients in Southeast Asia did multinomial logistic regression to predict clinical outcomes of patients infected with COVID-19 based on their vitamin D levels. Their results are summarized in the graph below which shows that of those with a COVID-19 case that was critical or severe, only 4% had normal levels, while 96% of those with mild COVID-19 had normal vitamin D levels.

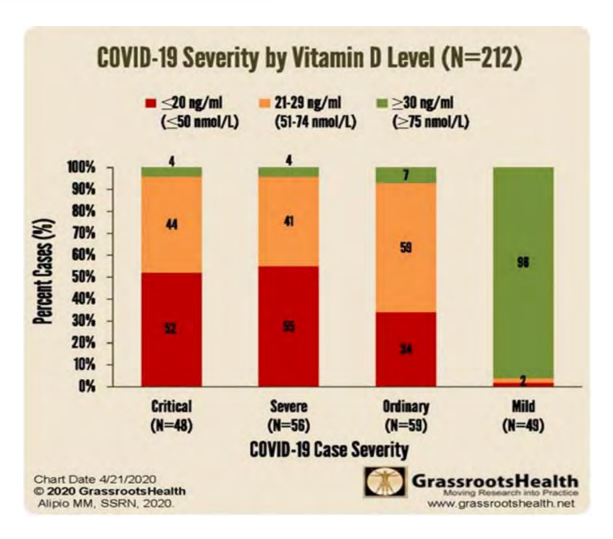


Figure 11

Another retrospective study of 780 cases with laboratory-confirmed infection of SARS-CoV-2 in Indonesia. When controlling for age, sex, and comorbidity, they found that vitamin D status was strongly associated with COVID-19 mortality outcome of cases.⁷¹ A summary of their findings are in the impressive graph below that demonstrates a radical reduction in the death rate from COVID-19 as the vitamin D level increase to over 30 ng/ml.

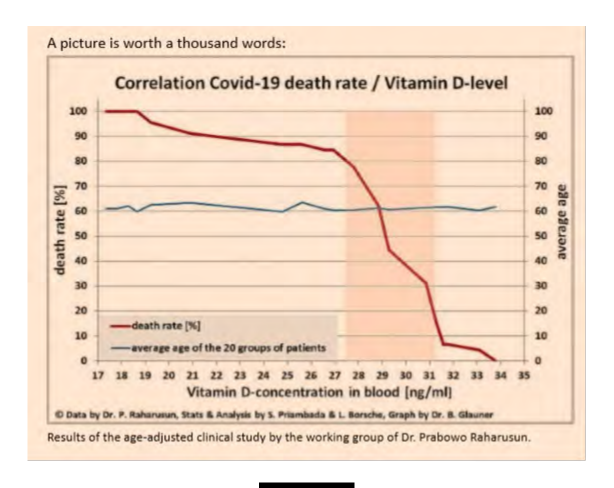


Figure 12

Similarly, a recent retrospective analysis from Sweden of 107 patients⁷² found that vitamin D concentrations were significantly lower in patients with positive PCR (polymerase chain reaction) tests for SARS-CoV-2. The researchers concluded that vitamin D3 supplementation would be useful in the treatment of COVID-19 infection, in preventing a more severe disease and/or in reducing the presence of the virus in the upper respiratory tract and making the patients less infectious.

Evidence was recently outlined to show that vitamin D deficiency could explain much of the reason for higher case and mortality rates for Black, Asian, and Minority Ethnic (BAME) residents in England.⁷³

There is also a preprint publication demonstrating a connection between vitamin D insufficiency and COVID-19. Louisiana State University Health Sciences Center studied 20 ICU COVID-19 patients and nearly 85%, vs. 57% in floor patients were vitamin D insufficient.⁷⁴

How Vitamin D Reduces the Risk of Viral Infections

here are many reviews that consider the ways in which vitamin D reduces the risk of viral infections.⁷⁵⁻⁸⁷

Vitamin D likely reduces the risk of viral respiratory infections because it influences several of your immune pathways, with the net effect of boosting your mucosal barrier defenses while simultaneously dampening excessive inflammation.⁸⁸ Vitamin D appears to decrease the risk of respiratory tract infections by three main mechanisms:⁸⁹

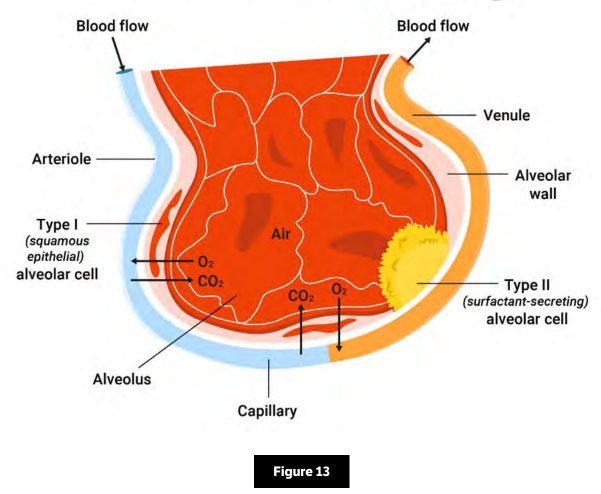
- It helps maintain tight junctions in the epithelial cells of the lungs and gut to prevent the infiltration of immune cells in lungs and other respiratory tissues,
- Inactivates some viruses through the stimulation of antiviral mechanisms such as antimicrobial peptides as discussed in the section above.
- Reduces pro-inflammatory cytokines through the modulation of the immune system as discussed in the section above.

How Vitamin D Specifically Reduces the Risk of COVID-19

he type-II pneumocytes in your lungs are the primary target for coronaviruses because the ACE2 receptors to which the virus binds are highly expressed on these cells. One of the problems with COVID-19 infections is that it impairs the function of your type-II pneumocytes which then decreases the surfactant level in your lungs.⁹⁰

This is important because surfactant prevents the collapse of the alveoli in your lungs. Surfactant allows your alveoli to stay open and compliant during both inspiration and expiration. During inspiration, your alveoli may collapse if they do not contain surfactant. If they collapse, then gas exchange across the alveoli wall cannot occur. Simply put sufficient surfactant is necessary for your alveoli to stay open and gas exchange to occur as shown in the Figure 13 below.

Alveolus – Gas Exchange



Fortunately vitamin D comes to the rescue for this problem produced by COVID-19 as it is able to stimulate the production of surfactant in alveolar type-II cells.⁹¹

Vitamin D, Angiotensin II and ACE2 Receptors

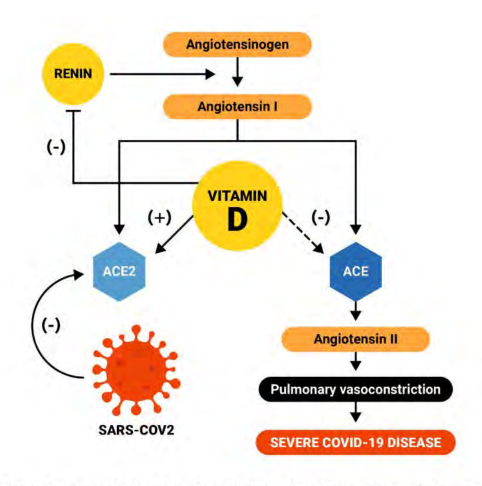
cell cultures of human alveolar type-II cells with vitamin D have shown that the SARS-CoV-2 virus interacts with the angiotensin-converting enzyme (ACE) 2 receptor expressed on the surface of your lung epithelial cells. Once the virus binds to the ACE2 receptor, it reduces its activity and, in turn, promotes ACE1 activity forming more angiotensin II which increases the severity of COVID-19.92 93 This may also be related to the vitamin D binding protein.94

Angiotensin II is a natural peptide hormone in the renin-angiotensin-aldosterone system. It is best known for increasing blood pressure through stimulating aldosterone⁹⁵ ACE2 normally consumes Angiotensin II, thereby lowering its levels. However, COVID-19 infection downregulates ACE2, which in turn leads to excessive accumulation of Angiotensin II.

High levels of Angiotensin II may cause acute respiratory distress syndrome (ARDS) or heart injury. Renin, on the other hand, is a proteolytic enzyme and a positive regulator of Angiotensin II. Vitamin D is a potent inhibitor of renin.

Vitamin D supplementation has been shown to prevent Angiotensin II accumulation and to decrease proinflammatory activity of Angiotensin II by suppressing the release of renin in patients infected with COVID, thus reducing the risk of ARDS, myocarditis, or cardiac injury. 96 Vitamin D may protect against symptoms of the COVID-19 infection by increasing the expression of ACE2 receptors on cells. 97 98 These ACE2 receptors that are expressed as a consequence of vitamin D supplementation reduce lung injury and promote binding of the virus to the lining of the lung. 99

Additionally, vitamin D may suppress renin activity. That in turn may generate less angiotensin II resulting in less lung blood vessel constriction. Although vitamin D causes the expression of ACE2, which indeed promotes the binding of the virus, it prevents the lung blood vessel constriction response in COVID-19 as illustrated in Figure 14 below.



The role of vitamin D in COVID-19. SARS-CoV2 binds to the ACE2 of alveolar cells and disturbs the ratio of ACE2/ACE activity. It increases ACE activity and, in turn, results in more angiotensin II formation causing pulmonary vasoconstriction to precipitate severe COVID-19. Vitamin D induces ACE2 expression, which limits the formation of angiotensin II via ACE and reduces lung injury. Besides, vitamin D supplementation may have a protective role against COVID-19. (Dashed line indicates indirect effect)

Figure 14

Vitamin D Seasonality and COVID-19

"Whoever wishes to investigate medicine properly should proceed thus: in the first place to consider the seasons of the year..." (Hippocrates, ca. 400 BC)." 100

It is interesting to note that there is a strong inverse correlation between sunlight exposure and the case fatality during the 1918-1919 influenza pandemic. ¹⁰¹ This strongly suggests that there is

a relationship between sunlight exposure and the risk for developing severe viral infections and secondary bacterial pneumonias.

Over 50 years ago R. Edgar Hope-Simpson, the British practitioner and self-educated epidemiologist, documented that influenza A epidemics in temperate latitudes are most intense in the months following the winter solstice.

He hypothesized that solar radiation produces a "seasonal stimulus" that affects the risk of influenza A. He theorized that there is a seasonal steroid hormone system with an impact on the human immune system whose substrate levels are low during the influenza season, but peak when influenza is rare. 102

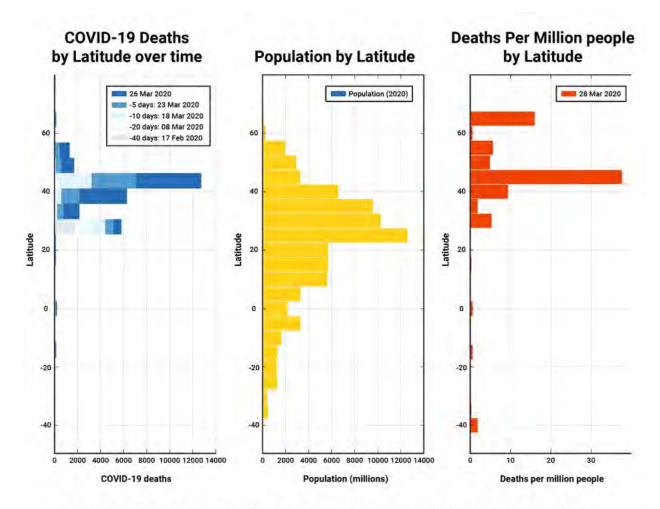
The winter incidence of influenza closely correlates with seasonal serum vitamin D levels. ¹⁰³ A British study also revealed that the prevalence of respiratory infections displayed a strong seasonal pattern in the opposite direction to the pattern for vitamin D concentrations. ¹⁰⁴ Seasonal variation in the blood levels of vitamin D, which contributes to immune function, is believed to be the underlying source of the observed influenza seasonality in temperate regions. ¹⁰⁵

More recently vitamin D deficiency was shown to be a risk factor for and/or a driver of the exaggerated and persistent inflammation that is a hallmark of ARDS. ¹⁰⁶ ¹⁰⁷ This is further evidenced with COVID -19 where the mortality from the disease has been relatively low for countries below 35 degrees latitude. ¹⁰⁸

Similarly, researchers demonstrated that the age –specific case fatality rate of COVID -19 was highest in Italy, Spain, and France, the European countries with the highest incidence severe vitamin D deficiency.¹⁰⁹

Although some have found that the rates of the cumulative COVID-19 deaths were decreased in countries with more sunshine, ¹¹⁰ Grant's analysis finds that life expectancy is the most important risk factor for those in Europe contracting COVID-19. ¹¹¹ A recent comprehensive review suggests this is likely due to the immune-senescence that occurs during aging which contributes to increased levels of inflammation and an increase in cytokine storm risk. ¹¹²

Severe COVID-19 outbreaks do indeed show a striking latitude relationship with severe outbreaks occurring exclusively in locations above the 30°N latitude line. Global reports of deaths and recoveries reveal that transmission rates and fatality rates from January to March 28, 2020 were significantly determined by latitude. Researchers aggregated world population latitude data into corresponding bins and calculated Deaths per Million as a function of latitude to come up with some powerful observations in Figure 15 shown below.¹¹³



(left) COVID-19 fatalities by latitude and over time; (middle) 2020 population by latitude; (right) COVID-19 fatalities per million people by latitude. Note: The Deaths per Million figure at -40°S is a statistical artefact due to diving two small numbers and may be ignored.

Figure 15

It is important to note that vitamin D obtained through sensible sun exposure is likely superior to oral supplementation. This may be related to other red and near-infrared light frequencies that could provide a therapeutic effect through photobiomodulation mechanisms that could elicit beneficial physiological effects such as increases in nitric oxide.¹¹⁴

Safety and Efficacy of Vitamin D Supplementation

he conventional media widely dismisses nutritional supplementation and vitamin D specifically. CNN has recently claimed vitamin D supplementation can actually "hurt you" and compared it to hydroxychloroquine whose "landmark" study the WHO used to justify stopping clinical trials was retracted by Lancet for fraudulent data use. 116

The NY Times has also recently warned about exercising caution in using vitamin D for COVID-19.¹¹⁷ ABC News cautions people that studies have yet to prove that taking a supplement will help and actively discourages vitamin D use.¹¹⁸

This differs considerably from the stance that the UK has taken. ¹¹⁹ In April 2020, Public Health England issued its advice on vitamin D, recommending that those on coronavirus lockdown (including children, pregnant and breastfeeding women and older people) should consider taking a daily supplement containing vitamin D supplement, even during the summer months, if they are not going outdoors often. ¹²⁰

Public health officials in the United Kingdom have launched an urgent review into the potential role of vitamin D in protecting people against the coronavirus. ¹²¹ The British media's Daily Mail suggested that vitamin D may be a cheap and safe way to treat the pandemic as mounting evidence supports this. ¹²² The Sun ran a story documenting how those with low vitamin D levels almost certainly die if they are hospitalized. ¹²³

The Guardian also chimed in affirmatively¹²⁴ and reporting that the public health officials are urgently reviewing the potential ability of vitamin D to reduce the risk of coronavirus.¹²⁵ Scotland also seems to be enthusiastic about adopting vitamin D strategies for COVID prevention.¹²⁶

There are two primary concerns many experts have with using vitamin D as a supplement for helping improve immune functions so your body can do its job to help mitigate the severity of infections like COVID-19. Those questions are is it safe and does it work?

First let's address the safety of vitamin D at the serum levels that are needed to achieve therapeutically meaningful blood levels, and what it takes to get there. To evaluate this, you need to understand what the existing conventional medical guidelines are for vitamin D supplementation.

The U.S. Institute of Medicine issued vitamin D and calcium guidelines nearly ten years old ago. ¹²⁷ Their guidelines are seriously dated and have not kept up with the current science as their recommendation was based solely on the effects of vitamin D for bone health and not for any of the metabolic benefits reviewed in this paper.

The institute recommended vitamin D supplementation of 600 IU/d for people younger than 70 years, 800 IU/d for those older than 70 years, and a serum 25(OH)D concentration of at least 20 ng/mL (50 nmol/L). While these doses will likely lower the risk of rickets, it will not be sufficient to decrease the risk of viral infections in those that are vitamin D deficient.

The institute did admit that no studies had reported adverse effects of supplementation with less than 10,000 IU/d of vitamin D, but still set their upper intake recommendation at 4,000 IU/d, partly out of concerns stemming from observational studies that found U-shaped 25(OH)D concentration—health outcome relationships.

However, later investigation determined that their recommendation was flawed as most reports of J- or U-shaped relationships were from observational studies that did not measure serum vitamin D blood levels and that the likely reason for those relationships was a result of enrolling some participants who had started taking vitamin D supplements shortly before enrolling. 128

It is useful to understand that significant levels of vitamin D can be produced from sun exposure during non-winter months. Approximately 10,000-25,000 IU vitamin D3 can be produced in a short time in the sun with full-body exposure, so it is obvious that your body can handle that amount easily. 129

So let's look at some of the recent studies that support a higher dose of vitamin D. One was done in a psychiatric hospital in Cincinnati, Ohio. The age range was from 18 to 90 years. Half of the patients were black, and nearly half were white.

All patients entering since 2011 were offered supplementation of 5,000 or 10,000 IU/d vitamin D3. For 36 patients who received 5000 IU/d for 12 months or longer, vitamin D levels rose from 24 to 68 ng/mL, whereas for the 78 patients who received 10,000 IU/d, mean concentrations increased from 25 to 96 ng/mL. No cases of vitamin D–induced hypercalcemia were reported. 130

Another recent study used 10,000 IU/day of vitamin D for 8–12 weeks and 93% of the subjects had vitamin D blood levels \geq 30 ng/mL after the first month and in two months the percentage increased to 100%. They also had no cases of hypercalcemia occur. ¹³¹

Although doses of 15,000 IU/day are rarely needed or recommended, they were found to be safe. ¹³² Data was collected for 3,882 participants in a community program. Blood vitamin D were measured at program entry and at follow-up within 6–18 months between 2013 and 2015.

Participants supplemented with a wide range of vitamin D doses $(1,000 - 15,000 \, \text{IU/d})$. To achieve vitamin D levels >40 ng/mL on average they needed vitamin D intakes of 6,000 IU/d for normal Body Mass Index (BMI), 7,000 IU/d for overweight and 8,000 IU/d for obese. They found no evidence of elevated calcium levels in the blood or urine at any vitamin D dose.

It has been suggested that the tolerable limit could be increased to 10,000 IU/day, as hypercalcemia is rarely encountered at lower doses, and most reports of other symptoms of vitamin D toxicity such as severe fatigue, confusion, vomiting, arrhythmia, and calcium kidney stones only occurred at doses exceeding 40,000 IU/day. In confirmation of this, a 2020 Canadian trial found the safety profile of vitamin D supplementation was similar for doses of 400, 4,000 and 10,000 IU/day in nearly 400 elderly patients.

A recent trial on a high-dose vitamin D supplementation in New Zealand involving 5,110 participants reported that, over a median of 3.3 years, monthly supplementation with 100,000 IU of vitamin D3 did not affect the incidence rate of kidney stone events or hypercalcemia. However, it should be noted that doses less than once a week are not recommended as they are not as effective.

A large meta-analysis of 25 randomized controlled trials of nearly 11,000 individual patients concluded that vitamin D supplementation was safe and it protected against acute respiratory tract infection overall. However it also showed that the benefit from vitamin D was stronger for those who received daily doses of vitamin D, but not in those who received large infrequent doses more than every two weeks. ¹³⁶ ¹³⁷

The second important question is if vitamin D supplementation works. This becomes somewhat confusing because there are many vitamin D trials showing that it is ineffective with no clinical benefit.¹³⁸

This is nearly in every case due to a common methodological flaw in the study. Typically, all of these studies that fail to show a benefit of vitamin used a specific dose of vitamin D rather than adjusting the dose to achieve an optimal vitamin D blood level. Further they have failed to measure major co-factors such as nutrient intake like magnesium, calcium and vitamins K2 and C.

Randomized controlled trials evaluating the impact of vitamin D supplementation on clinical outcomes simply need to use a study a design based on serum levels of 25-hydroxyvitamin D concentrations rather than administered vitamin D doses. Once you understand this and you carefully review the methods section of the study, you will find that nearly every negative vitamin D study failed to individualize dosing based on blood levels. Further, one of the biggest omissions was any defined co-factors.

Vitamin D Levels

Researchers have shown that levels of at least 30 ng/mL are necessary for the optimal induction of the antimicrobial peptide LL-37 (cathelicidin)¹⁴² which was discussed in an earlier section. While vitamin D levels of approximately \geq 40 ng/mL seems to provide protection against acute viral respiratory infections. ¹⁴³ ¹⁴⁴

A meta-analysis of 25 trials, of over 11,000 participants, showed vitamin D supplementation to reduce the risk of acute respiratory infections, including viral, by 12% in all participants. This was most pronounced in patients with serum vitamin D levels below 20 ng/mL.¹⁴⁵

Maintenance of circulating 25-hydroxyvitamin D levels of 40-60 ng/mL would be optimal, since concentrations of 40 ng/mL represent the beginning point of the plateau where the synthesis of the active form of vitamin D becomes consistent. 146 147

Since vitamin D can be made in your skin, the term "vitamin" seems inappropriate. However, compared to the past, most of us spend far more time indoors, largely cover our skin with clothing when outdoors, and often live at latitudes where during winter UV-B radiation is inadequate for many months. Therefore, most are unable to generate healthy levels of vitamin D, which is why most people benefit from vitamin D supplementation.

It is important to understand that blood levels targeted to a specific dose of vitamin D will be highly variable between individuals due to several demographic and biological factors:

- Baseline vitamin D status
- Status of co-factors such as magnesium, calcium, vitamin K2, vitamin C, and omega 3s
- Lower levels of 7-dehydrocholesterol in the skin¹⁴⁸
- Ethnicity and skin color
- Body fat percentage
- Genetics
- Seasonal variations and time of sun exposure ¹⁴⁹
- Type of vitamin D supplements¹⁵⁰

Increased skin pigmentation reduces the efficacy of UVB because melanin functions as a natural sunblock. In addition, aging decreases the ability of the skin to produce vitamin D $3.^{151}$ During the winter months at latitudes of greater than $28^{\circ},^{152}$ little or no UVB radiation reaches the surface of the earth.

However, residence at low latitude does not guarantee adequate vitamin D levels. Social and cultural norms may limit sun exposure, ¹⁵³ particularly as we age, leading to a tendency of serum vitamin D levels to decrease with age¹⁵⁴ which is important for COVID-19 because case-fatality rates (CFRs) increase with age. ¹⁵⁵

Finally, pharmaceutical drug use typically increases with age, and drugs such as antiepileptics, antineoplastics, antibiotics, anti-inflammatory agents, antihypertensives, antiretrovirals, endocrine drugs, and some herbal medicines can decrease vitamin D levels by activating the pregnane-X receptor. 156

Vitamin D Supplementation

agnesium supplementation is recommended when taking vitamin D supplements. Magnesium helps activate vitamin D. All the enzymes that metabolize vitamin D seem to require magnesium, which acts as a cofactor in the enzymatic reactions in the liver and kidneys. The dose of magnesium should be in the range of 250–500 mg/d, along with twice that dose of calcium.

Magnesium activates more than 600 enzymes and influences extracellular calcium levels. ¹⁵⁸ It is essential for the stability of cell function, RNA and DNA synthesis, and cell repair, as well as maintaining the antioxidant status of the cell. It is an important cofactor for the activation of a wide range of transporters and enzymes. ¹⁵⁹ ¹⁶⁰

A recent review found that as many as 50% of Americans taking vitamin D supplements may not receive significant benefits. This happens when the vitamin D they take gets stored in its inactive form because they have insufficient magnesium levels. Magnesium supplementation was shown to markedly reduce the resistance to vitamin D treatment. 161-163

In a preliminary analysis, GrassrootsHealth found¹⁶⁴ that individuals who do not take supplemental magnesium need, on average, 146% more vitamin D to achieve a blood level of 40 ng/ml (100 nmol/L), compared to those who take at least 400 mg of magnesium per day.

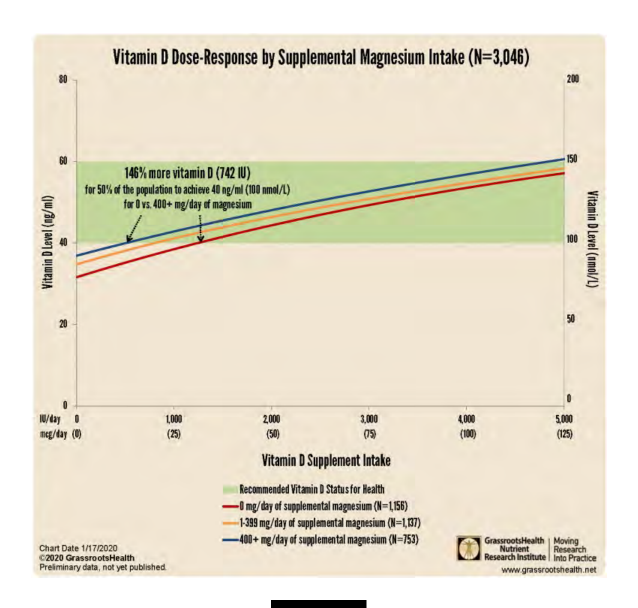


Figure 16

The interplay between magnesium and vitamin D isn't a one-way street though. It goes both ways. Interestingly, while vitamin D improves magnesium absorption, taking large doses of vitamin D can also deplete magnesium. Again, the reason for that is because magnesium is required in the conversion of vitamin D into its various forms.

Ideally one should test their vitamin D blood level as this will help to understand the appropriate starting dose. If the level is about 20 ng/mL it takes about 35 days to reach 60 ng/mL with a daily dose of 10,000 IU of vitamin D and 85 days with 4,000 IU/d. 167

If one is challenged with an acute scenario it may even be wise to use a very large initial dose. A randomized controlled trial (RCT) published in 2015 showed that after a single dose of 250,000 IU of vitamin D3 given to healthy volunteers between the ages of 18 and 65 years with baseline serum levels of <17 ng/ml, serum 25(OH)D concentrations at five days increased to an average of 41 ng/ml with no adverse effects. 168

After five days it would be reasonable to start a dose of 5,000 units a day as after 90 days, vitamin D levels will drop back to near baseline values. 169

While vitamin D supplementation could stop COVID-19 from developing at the beginning of symptoms, it probably would not be very useful after lung and organ damage occurs in the acute stage. ¹⁷⁰

While vitamin D is likely the most important nutrient to optimize for COVID-19 prevention, other nutrients, micronutrients, and phytonutrients are also known to impact your immune system and infection risk. 171172

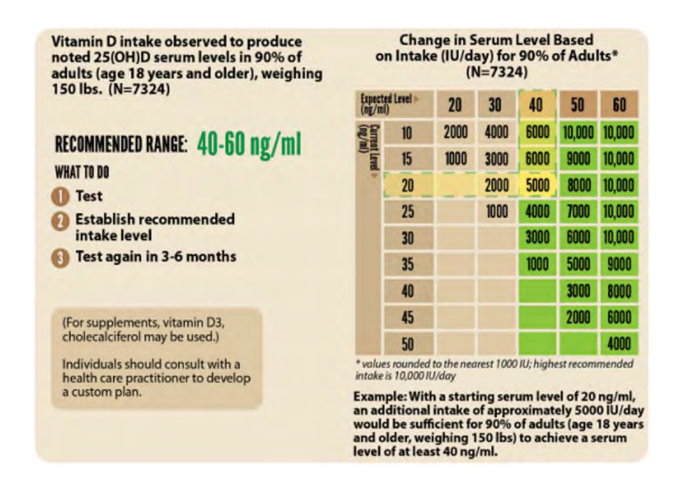


Figure 17

Target of Vitamin D Campaign

f you have ever flown you will likely recall the flight attendants take off briefing which tells you that, in the event of an emergency occurs, an oxygen mask will automatically appear in front of you. But they also tell you that if you are traveling with a child or someone who requires assistance, to secure your mask on first, and then assist the other person.

The lesson here is that it will be important to adopt the vitamin D recommendations and its cofactors in this paper for yourself and family first. But it is the intention of this document to empower an army to target the populations that are most at risk for the next wave of COVID-19 or any other respiratory infection that comes our way.

The intention of this document is to help create an army that can go out and reach these target populations that are at high risk during the next wave of the infection. The target populations are the elderly and people of color (and those with chronic diseases, and pregnant and nursing mothers). 173

It is important to know that YOU can make a difference by taking this information and sharing it with others, especially those that have influence to spread this message to these at risk populations.

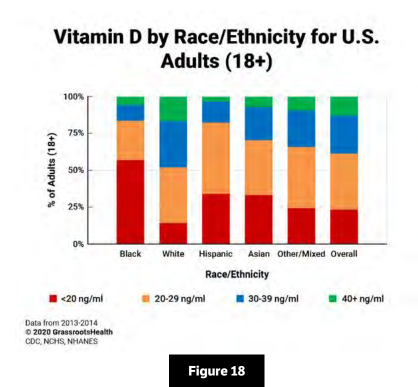
By a little investment of time you can save many lives at virtually no cost. Remember if it is the late spring, summer or early fall, you likely can get enough vitamin D for free by merely going outside around solar noon just being careful to never get burned.

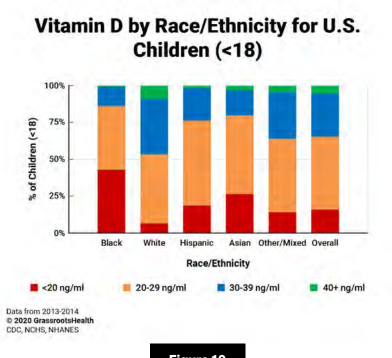
If you live below 27 degrees latitude you can get vitamin D most of the year from the sun. But if you don't live that far south or it is the winter vitamin D supplements are some of the least expensive supplements you can purchase. All you need to do is follow the dosage recommendations above.

Black Americans and People of Color

Collectively, Black Americans represent 1/8 of the population in the U.S., but they have suffered ¼ of known COVID-19 deaths. They are dying at twice their population share. ¹⁷⁴ So what could explain this dramatic difference in death rates between white and black Americans?

In the graph below that is compiled from approximately 15,000 tests done at GrassrootsHealth over the last 13 years, You will notice that the levels of vitamin D based on race in the US that only 16% of black adults have adequate vitamin D levels while over three times that number or nearly 50% of white adults have vitamin D levels over 30 ng/ml.





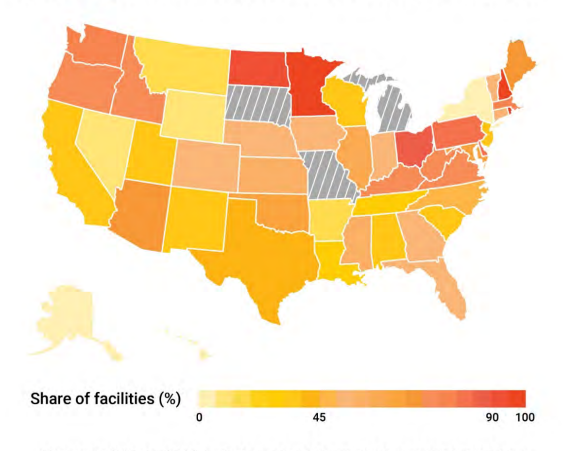
Elderly Focus

A landmark study¹⁷⁵ by Gregg Girvan and Avik Roy of the Foundation for Research of Equal Opportunity was done on long-term medical care providers to the aged and medically infirm which consist of:

- Nursing homes and skilled nursing facilities;
- Assisted living facilities, i.e., residential care communities or personal care homes;
- Adult day service centers;
- Home health Agencies; and
- Hospices

The disease caused by SARS-CoV-2 affects the elderly far more severely, on average, than younger individuals. Those living in nursing homes and assisted living facilities seem to be an extraordinarily increased risk of dying from COVID-19. As you can see in the graphic below from June 2020, 42% of deaths occurred in nursing homes and assisted living facilities.

42% of U.S. COVID-19 Deaths Occur in Nursing Homes & Assisted Living Facilities



42 percent of U.S. COVID-19 deaths have occured in nursing homes and assisted living facilities. Nursing homes are residential facilities for those needing 24/7 on-site medical supervision; assisted living facilities are for those not needing 24/7 medical supervision. The share of deaths occuring in nursing homes and assisted living facilities is highest in New Hampshire, Rhode Island, and Minnesota, using the latest data as of June 1, 2020.

SOURCE: The Foundation for Research on Equal Opportunity. G. Girvan & A. Roy / FREOPP

Figure 20

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