Interaction between Vitamins C and E When Scavenging the Superoxide Radical Shown by Hydrodynamic Voltammetry and DFT

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Abstract: In this study, we examine the cooperative effect between vitamins C and E that mitigates oxidative stress by using experimental and computational methods. We performed superoxide scavenging experiments on each vitamin individually and their combination using rotating ring–disk electrode voltammetry. The results indicate that vitamins E and C together produce more effective scavenging of superoxide as evaluated by a steeper slope in the efficiency graph, $-7.2 \times 10^4$, compared to that of vitamin E alone, $-1.8 \times 10^3$, or vitamin C alone, $-1.3 \times 10^4$. Density Functional Theory calculations agree with our experimental results, and we describe a mechanism for the antioxidant action of individual vitamins E and C, plus the synergistic action when both vitamins interact. This process involves the restoration of vitamin E by vitamin C and includes π-π interactions between superoxide and scavengers. The overall result produces an increase in scavenging superoxide radicals when both vitamins act together.

Keywords: vitamin C; vitamin E; nutrient synergy; antioxidant; superoxide

1. Introduction

The in vivo and in vitro relationship between vitamin C and vitamin E has been known and studied for some time [1–4]. The in vivo results of a pioneering study on healthy young adults by Jeng et al. [5] showed that combined supplementation with vitamins C and E resulted in an enhanced immune response over supplementation by either vitamin alone. Similarly, beneficial results of in vivo interaction between vitamins C and E were reported after sequential antioxidant vitamin C and E supplementation to the diet of human subjects [6]. More recently, elderly adult immune function was improved with supplementation of vitamins C and E [7]. Also, the fertility of male rats under conditions of high oxidative stress was restored by a diet containing tocopherols and vitamin C [8].

Vitamin E’s cooperative role with vitamin C as an antioxidant species in diseases associated with oxidative stress is much studied [9]. These essential nutrients play a major role in reducing and controlling oxidative stress by protecting the immune system and enhancing resistance against infectious microbes such as bacteria, viruses, and parasites. The beneficial effects of these two antioxidant vitamins on the immune system are described in several reviews [10–23]. Relevant to this work, supplementation of vitamins C and E decreased O$_2^•$− concentration in the bloodstream and improved antioxidant activity by enhancing NADPH oxidase and superoxide dismutase activity [4].

Many literature studies on the individual action of these two antioxidant vitamins exist. Vitamin C (ascorbic acid) supplementation, for example, has been shown to reduce the duration and severity of upper respiratory infections, including colds [24] and other respiratory infections, including COVID-19 [25]. Water-soluble vitamin C is an important non-enzymatic antioxidant found in human tissue that is involved in many biological
functions by acting as a reducing agent through the donation of two electrons. This redox ability is crucial to its role (1) as a cofactor in either monooxygenase or dioxygenase enzymatic reactions, including collagen synthesis, and (2) as an antioxidant against cellular oxidative stress [26]. Vitamin C also restores vitamin E through redox recycling and facilitates the absorption of iron by enhancing its intestinal absorption [27]. One of its biological roles is as a normal skin constituent, both dermis and epidermis, and it plays an important role in tissue repair and wound and burn healing. Scurvy was one of the first examples of a serious disease caused by vitamin C deficiency [28]. Current reviews describe the helpful effects of vitamin C on several other infections [13,29]. However, the relationship between insufficient vitamin C and altered inflammatory and immune responses is not yet clear. As such, the biochemical mechanisms by which vitamin C influences NF-κβ activation to produce pro-inflammatory cytokines are not obvious, and this topic is the subject of many research studies [11,30].

Vitamin E is chemically more complex and consists of a family of eight different compounds: α-, β-, γ-, and δ-tocopherols and four α-, β-, γ-, and δ-tocotrienols (Figure 1). α-Tocopherol is the predominant form in the body and has been given the greatest attention. It is a fat-soluble antioxidant, and its role in human health, particularly as it interacts with peroxyl radicals to prevent lipid peroxidation and subsequent membrane damage, is much investigated and not yet completely explained [31,32]. All naturally occurring vitamin E forms are strong antioxidants with anti-inflammatory activities [33]. A recent review describes the capability of vitamin E to behave as an antioxidant and its effects on diseases caused by oxidative stress [34]. However, despite many studies that have reported on vitamin E’s relationship with the immune system, the mechanistic details are lacking [35,36].

Figure 1. Chemical structures of vitamin E components. (Top) Tocotrienols: α-Tocotrienol: R₁, R₂, R₃ = Me; β-Tocotrienol: R₁ = Me, R₂ = H, R₃ = Me; γ-Tocotrienol: R₁ = H, R₂ = Me, R₃ = Me; δ-Tocotrienol: R₁ = H, R₂ = H, R₃ = Me. (Bottom) Tocopherols have the C₁₂ long chain saturated: α-tocopherol, R₁, R₂, R₃ = Me; β-tocopherol, R₁ = Me, R₂ = H, R₃ = Me; γ-tocopherol, R₁ = H, R₂ = Me, R₃ = Me; δ-tocopherol, R₁ = H, R₂ = H, R₃ = Me.

The antioxidant role of vitamins C and E is a subject of continual investigation, and studying their interactions with reactive oxygen species (ROS), especially the superoxide radical, is of importance. The latter is biologically obtained through two main processes: (1) leakage from the electron transport chain during oxidative phosphorylation [37] and (2) the generation of the superoxide radical by NADPH oxidases in the immune system to destroy ingested microbes by neutrophils and other leukocytes [38]. However, excess superoxide is damaging and needs to be regulated. This is performed enzymatically by superoxide dismutases (SODs) and assisted through human diets that provide natural antioxidant compounds from fruits and vegetables, including vitamins C and E. Therefore, we focus our attention on trying to understand the mechanism by which vitamin C and α-tocopherol exhibit a cooperative effect to ameliorate oxidative stress, specifically due to...
the superoxide radical. Among these antioxidant mechanisms, π-π interactions between superoxide and its scavengers permit superoxide to act as a reducing agent, mimicking SOD action, as recently shown for isoflavones [39].

A recent review on superoxide chemistry [40] describes the role played by the electrodes in the electrochemical generation of $\text{O}_2^{*-}$. This is relevant to the rotating ring–disk electrode (RRDE) voltammetry technique that we developed and used in this study. In fact, we perform voltaiic experiments to measure individual superoxide scavenging by each vitamin as well as examine their combined effects. The advantage of our method is that the concentration of superoxide present in the solution can be calculated directly. Density Functional Theory (DFT) calculations supporting a synergistic effect for both vitamins are evaluated in a similar environment to the natural location of vitamin E in the cell membrane. This chemical mechanism agrees with our experimental results.

2. Materials and Methods

2.1. Hydrodynamic Voltammetry (RRDE)

Hydrodynamic Voltammetry was performed at a rotating ring–disk electrode (RRDE) using the WaveDriver 20 bipotentiostat (Pine Research, Durham, NC, USA) with the MSR Electrode Rotator, also from Pine Research, Durham, NC, USA. In RRDE, the bipotentiostat measures, at the same time, both the currents at the disk and ring electrodes (that correspond to charge movements among the ring, the disk, and the counter electrode) and the potentials of the disk and ring electrodes relating to the single reference electrode. The RRDE cell contains four electrodes: the two gold working electrodes, both the rotating ring and disk (Pine Research, Durham, NC, USA), one coiled platinum wire counter electrode, and one reference electrode consisting of a platinum wire immersed in 0.1 M dried tetrabutylammonium bromide, TBAB (Sigma-Aldrich, St. Louis, MO, USA), dissolved in 50 mL dimethyl sulfoxide, DMSO, anhydrous, ≥99.9% (Sigma-Aldrich, St. Louis, MO, USA), in a fritted glass tube. The electrodes were placed in a 5-neck electrochemical cell together with means for either bubbling or blanketing the solution with gas. The solution was subsequently bubbled with dry $\text{O}_2/\text{N}_2$ (35%/65%) for five minutes to establish the required dissolved molecular oxygen level. Careful initial cleaning of the electrodes was performed to clear potential film formation.

Aftermath software release 1.6.10523 (Pine Research, Durham, NC, USA) was used to set up the parameters needed for the experiment: the potential sweep was applied to the disk from 0.2 V to $-1.2$ V and then reversed to 0.2 V, while the potential of the ring electrode was held constant at 0.0 V. The disk voltage sweep rate was set at 25 mV/s. The rotation setting used for the rotation of the Au/Au disk electrode was chosen to be 1000 rpm at the disk electrode. The superoxide radical is generated through molecular oxygen reduction, and the peak was detected around $-0.6$ V. Meanwhile, the reverse oxidation reaction of the remaining unreacted superoxide radicals was detected at the ring electrode. An initial blank solution consisting of bubbled $\text{N}_2/\text{O}_2$, the electrolyte TBMB, and DMSO was run. The ratio of the ring/disk current was defined as “efficiency”. Next, an antioxidant aliquot of vitamin E (DL-α-tocopherol) or vitamin C (0.03 M) was introduced. The solution containing antioxidant in the voltaic cell was bubbled with the gas mixture for 5 min, a revised voltammogram was documented, and the corresponding efficiency was calculated. In this way, the rate at which the increasing concentration of the antioxidant scavenges the generated superoxide radicals during the electrochemical reaction is determined upon the addition of each antioxidant aliquot. Aftermath software was used to record the results from each run, represented as voltammograms showing the current vs. potential graphs. These were later evaluated using Microsoft Excel. The volume amount used in each of the aliquots is indicated in the related RRDE graph. Finally, the decreasing slope of the curve, describing the overall decrease in efficiency with the incremental addition of the antioxidant, serves as a quantitative measure of the antioxidant activity of the vitamins. Any decrease in the collection efficiency is anticipated to be due to the amount of superoxide consumed by the antioxidant. This method was developed in our laboratory [41].
2.2. Computational Study

Calculations were run using BIOVIA Materials Studio DMol\(^3\), implemented in Materials Studio 7.0. DMol\(^3\) is a modeling program (Dassault Systèmes, San Diego, CA, USA) that utilizes Density Functional Theory (DFT) to calculate properties of molecules such as energy, geometry, and transition state optimizations [42]. The results allow the relationships between the molecular structure with its antioxidant properties and scavenging behavior to become evident. We employed the double numerical polarized (DNP) basis set, including all the occupied atomic orbitals plus a second set of valence atomic orbitals, as well as polarized d-valence orbitals [43]. Correlation generalized gradient approximation (GGA) was used, including BLYP correlation plus BLYP-D and Becke exchange [44]. We also included Grimme’s correction when van der Waals interactions were involved [45]. The continuous model of Dmol\(^3\) was applied for solvent effects and calculation, including polar DMSO, non-polar n-hexane, or non-solvent (gas phase), showing no important variation in dimensions (bond distances or separations between isolated species) [46]. All electrons were treated explicitly, and the real space cutoff of 5 Å was set for the numerical integration of the Hamiltonian matrix elements. The self-consistent field convergence criterion was established for the root mean square variation in the electronic density to be less than \(10^{-6}\) electron/Å\(^3\). The convergence criteria applied during geometry optimization were \(2.72 \times 10^{-4}\) eV for energy and 0.054 eV/Å for force.

An attempt to use a functional of higher quality, B3LYP, which is better at estimating potential barrier heights, was made, but calculations became extremely lengthy. Ultimately, these B3LYP calculations confirmed what we saw using BLYP, e.g., some calculated potential energy minimums for our molecules are extremely shallow, which makes for extremely prolonged calculations to reach the needed energy minimum. (After almost a week for completion of the first calculation where we get convergence, we needed to refine the entire system to look for the true minimum through other very lengthy iterative refinements). Therefore, the dimensions of our molecules pose a problem, as using the strong functional (B3LYP) instead of BLYP is markedly time-consuming. From our computational studies, we are looking mainly for trends in the barriers to identify fast and slow steps for correlation with our experimental studies, which can be achieved using the simpler and faster BLYP functional. These calculations try to correlate experimental RRDE features.

3. Results and Discussion

3.1. DFT of Vitamin E-Model and Vitamin C

3.1.1. Vitamin E-Model Scavenges Superoxide

DFT methods were applied to study the antioxidant properties of vitamins E and C. To decrease the calculation time, we used a model of vitamin E \(\alpha\)-tocopherol, which we call vitamin E-model (Figure 1), which consists of the 6-chromanol nucleus contained in the \(\alpha\)-tocopherol isomer but with the long, hydrophobic, saturated side chain replaced by a methyl group. The model then has a total of five methyl groups on the 6-chromanol nucleus (methyl groups in position R1, R2, and R3 of the aromatic ring plus the two methyl groups at the C2 position of the 6-chromanol). This model is a suitable representative of all forms of vitamin E, and a published report concluded that the antioxidant capacity of vitamin E and a model without the side chain were quite similar [47].

The superoxide radical anion, considered to be of major importance in cell biology, is primarily obtained when an oxygen molecule acquires an extra electron leaked from the mitochondria during aerobic cellular respiration that ultimately produces ATP [42,48,49]. The reactivity of the superoxide radical is controlled by endogenous enzymes, superoxide dismutases, catalase, and glutathione peroxidase, which help to cope with oxidative stress by effectively quenching free radicals [50,51]. Substantial evidence links increases in oxidative stress [52] with the decline of enzyme regulation of the superoxide anion during aging. Leakage of this radical anion during oxidative stress creates conditions for attack on DNA, proteins, and other important biological molecules, leading to disease states [53].
Due to the lipidal character of vitamin E with its long saturated hydrophobic side chain, α-tocopherol is recognized as the major antioxidant in membrane-rich fractions such as mitochondria and microsomes [54,55]. This strategic position makes vitamin E an important means of scavenging the superoxide radical inside the cell membrane [56]. It is, therefore, important to describe the initial interaction between the 6-hydroxyl in vitamin E-model and superoxide. Using computational methods, the reaction, with the minimized structures of reagent, transition state (TS), and product, shown in Figure 2, has ΔG of −0.4 Kcal/mol and an easily attainable E(barrier) of 0.8 Kcal/mol. Figure 3 shows details of the transition state search. These results demonstrate that vitamin E-model can intercept and neutralize superoxide. A potential ensuing step on the product shown in Figure 2 is described in Figure 4, where a proton is placed at van der Waals separation from the added superoxide, 2.60 Å. The minimization of Figure 4 arrangement (Figure 5) shows H2O2 formation and separated 1.654 Å from the O(polyphenol) in position 6, which is longer than the initial input conditions, 1.498 Å. Therefore, hydrogen peroxide was eliminated, and the residual semiquinone vitamin E-model was minimized. The next step in this study involves the π-π interaction with an additional molecule of superoxide. This process was shown to mimic superoxide dismutase action by some natural organic molecules and is described in previous studies and a recent review [57].

![Figure 2](image1.png)

**Figure 2.** Superoxide scavenges H(hydroxyl) in position 6 of vitamin E-model. Minimized structures of reagent (left), transition state (center), and product (bottom right) are shown. This reaction has ΔG of −0.4 Kcal/mol and E(barrier) of 0.8 Kcal/mol. The transferred H atom is equidistant from both moieties in this TS: O(superoxide)—H = 1.219 Å, O6—H = 1.209 Å. O atoms, red; C atoms, black; H atoms, light grey.

![Figure 3](image2.png)

**Figure 3.** TS search of scavenging superoxide by vitamin E-model. ΔG = −0.4 kcal/mol; E(barrier) = 0.8 kcal/mol.
Figure 4. A proton is placed near the HO₂ moiety of Figure 2 product, having van der Waals separation of 2.60 Å.

Figure 5. Minimization of Figure 4 arrangement shows H₂O₂ formation, separated 1.654 Å from the O(polyphenol) in position 6, which is longer than the initial condition, 1.498 Å, shown in Figure 4.

Therefore, the semiquinone vitamin E-model was minimized, and a superoxide radical was π-π placed above its aromatic ring (van der Waals distance, 3.50 Å). Upon geometry minimization, there is a shortening of the distance between both centroids, 2.899 Å, indicating bond formation. In the meantime, the initial superoxide bond distance of 1.373 Å shortens to 1.302 Å, which implies that some electron density becomes directed toward the ring (Figure 6). This overall arrangement is non-radical due to its even number of electrons, arising from two superoxides (each one contributing with an odd number of electrons), and it is negatively charged (−1), resulting from two negative charges of the two superoxides (from Figures 2 and 6) and one proton (from Figure 4).

Figure 6. After elimination of H₂O₂ from the previous arrangement (Figure 5), the remaining species, semiquinone vitamin E-model, was minimized, and a superoxide radical was π-π placed above the aromatic ring (van der Waals distance, 3.50 Å). Its geometry minimization shows shortening between both centroids, 2.899 Å, indicating bond formation. Meanwhile, the initial superoxide bond distance of 1.373 Å shortens to 1.302 Å, which indicates some electron density was directed toward the ring.

It can be expected that a cation will react with this arrangement, and so a proton was van der Waals placed near O6 of the anion arrangement, with an O6-proton distance of 2.60 Å. Its DFT optimization shows O6-H6 bond formation, 0.976 Å (Figure 7). This reaction
has a small energy barrier, as observed after a TS search, and the resulting structure is shown in Figure S1. The C5-O6 bond length, 1.378 Å, is characteristic of a single bond (Figure 7) and is longer than the partial double bond in the initial species, 1.295 Å (Figure 6). The effects of proton addition to O6 are also evident in the aromatic ring, with four bond lengths becoming more similar, range 1.42–1.43 Å, than those in the reacting species, range 1.42–1.46 Å, while the remaining two bond lengths in the ring continue to be shorter and unaltered (1.398–1.399 Å). These effects are related to the shorter distance between superoxide and ring centroids, 2.664 Å, compared with the initial species, 2.899 Å. Thus, the approach of a positive charge, a proton, to the reacting species makes the \( \pi-\sigma \) bound superoxide shift more electron density toward the ring. This forms a neutral and non-radical species, \( \eta-O_2 \)-vitamin E-model, and \( H_2O_2 \), resulting from the interaction of vitamin E-model with two superoxide molecules (one \( \pi \) and one \( \sigma \) added to the polyphenol), plus two protons, shown in Reaction (1).

\[
2O_2^- + \text{vitE-model} + 2H^+ \rightarrow H_2O_2 + \eta-O_2\text{-vitE-model}
\] (1)

Figure 7. DFT optimization result after a proton was placed near O6 of the previous arrangement, Figure 6, at O6-proton distance 2.60 Å. It shows O6-H6 bond formation, 0.976 Å; the C6-O6 bond length, 1.378 Å, is a typical single bond and is longer than the partial double bond in the reacting species shown above, 1.295 Å (Figure 6). This reaction has a small barrier, observed after a TS search, whose resulting structure is shown in Figure S1.

Earlier studies using UV–Vis and ESR techniques involving the scavenging of DPPH and galvinoxyl radicals by a vitamin E derivative, IRFI005, describe a “two electrons and/or 2 H-atoms” donation mechanism for each molecule of the scavenger [58].

Our mechanism is shown in Scheme 1. The scavenging mechanism here described for superoxide, a biologically relevant radical, includes partial \( \pi-\pi \) donation of an electron to the aromatic ring of vitamin E and capture of a H atom by another superoxide. In a related excellent theoretical study about vitamin E, \( \pi-\pi \) interactions have not been considered for scavenging superoxide [59].

We compare the energy barriers for the higher-quality B3LYP functional and the BLYP used in Figure 2. A TS Optimization for the TS shown in Figure 2 shows equal energy for both systems BLYP and B3LYP, suggesting no underestimation of the energy barrier by BLYP: energy of transition state: \(-806.5218885\) Ha (BLYP), \(-806.521889\) Ha (B3LYP). In addition, Figure 2 TS for the transferred proton has O(superoxide)-H = 1.219 Å, O(VitaminE)-H = 1.209 Å, and the corresponding distances are very similar to those obtained using B3LYP calculation (1.211 Å and 1.204 Å), as shown in Figure S1.
Figure 8. Initial state for vitamin C (left) transferring a H atom to the vitamin E-model semiquinone (right), which previously gave up a proton to superoxide; the total charge of this radical is 0.

Figure 9. Vitamin C transfers a hydrogen atom to vitamin E-model semiquinone. Left: Minimum of energy after approaching H(vitamin C) to vitamin E-model semiquinone (obtained after DFT minimization of Figure 8). Right: Product of reaction. Center: Transition state showing H located midway between both O atoms, 1.215 Å from vitamin C and 1.229 Å from vitamin E. This reaction is characterized by ΔG = –2.7 Kcal/mol and E(barrier) = 0.7 Kcal/mol.

Scheme 1. Vitamin E scavenges superoxide. (A) Superoxide approaches vitamin E hydroxyl in position 6; (B) H is captured by superoxide, forming [HO₂]⁻ plus vitamin E semiquinone; (C) a proton is captured by [HO₂]⁻, forming H₂O₂; (D) an additional superoxide interacts π-π with the aromatic ring; (E) an additional proton interacts with the semiquinone, forming π-O₂-vitamin E-model.

3.1.2. Vitamin C Restores Vitamin E-Model

As shown in Figure 7, a reacting proton is needed to accomplish Scheme 1. Therefore, the interaction between the semiquinone vitamin E-model and vitamin C, with an initial separation of 2.60 Å (Figure 8), was studied with DFT, resulting in a proton transfer to the semiquinone (Figure 9). This reaction is possible as shown by ΔG = –2.7 Kcal/mol and E(barrier) = 0.7 Kcal/mol. Since the total charge of the system is zero, the result of this interaction is an ascorbate radical and vitamin E-model.
3.1.3. Antioxidant Vitamin C Scavenges Superoxide

Our DFT calculations support that vitamin C is an antioxidant, as seen in Figures 10–13 and S2. Figure 10 shows the initial $\sigma$ approach of superoxide to the acidic proton of vitamin C. Figure 11 shows the result of DFT minimization using an n-hexane solvent effect, whose low dielectric constant is closely related to the lipidic environment in the membrane cell, the natural location of vitamin E. Figure S2 applies a DFT DMSO solvent effect, which is related to the experimental aprotic solvent used in RRDE, *vide infra*, and the structural results of Figures 11 and S2 are similar. The expected subsequent step for Figure 11 is a proton addition to obtain $\text{H}_2\text{O}_2$, which is then a substrate in the catalase enzymatic reaction to yield $\text{H}_2\text{O}$ plus $\frac{1}{2} \text{O}_2$. $\text{H}_2\text{O}_2$ is obtained after a second ascorbic acid approaches the most exposed O atom of superoxide in Figure 11. The separation between $\text{H}_2\text{O}_2$ and both ascorbates is 1.680 Å and 1.698 Å (Figure 12).

![Figure 10. Vitamin C scavenging superoxide. Initial state, superoxide is placed at van der Waals separation, 2.60 Å, from the acidic proton of vitamin C.](image)

![Figure 11. Superoxide forms a HO$_2$ moiety, well separated from the remaining species. DFT calculation performed in n-hexane, an organic environment that can mimic the membrane cell. Bond distances within HO$_2$, O-H = 1.027 Å, O-O = 1.398 Å; separation between both units = 1.617 Å; within the vitamin C derivative C-O = 1.277 Å.](image)

Since one of the two remaining vitamin C derivatives is a radical (Figure 12), this can easily react with an additional superoxide and yield a quinone-like vitamin C derivative (see Figure 13). This can be considered part of the quenching action of stopping lipid peroxidation. The alternative reaction involving a $\pi$-$\pi$ interaction, instead of the $\sigma$ superoxide attack on the quinone vitamin C derivative of Figure 10, was shown not to be feasible in performing DFT calculations. Therefore, a vitamin C quinone formation is suggested, along with $\text{H}_2\text{O}_2$ for superoxide scavenging of vitamin C. A detailed theoretical mechanism for vitamin C oxidation by the superoxide anion radical reveals the presence of
radical vitamin C quinone derivatives. These compounds have also been modeled with water molecules [60]. However, in the RRDE method used here, water molecules do not play a role as they react with superoxide when they arrive at the disk electrode. We use anhydrous DMSO solvent to have minimal water solvation for vitamins C and E, using the disk electrode where superoxide is generated.

![Figure 12.](image)

Figure 12. A 2nd vitamin C molecule, stick style, was placed near the most exposed O(superoxide) of the product arrangement shown in Figure 11. DFT showed formation of H$_2$O$_2$ well separated from two vitamin C species, 1.680 Å and 1.698 Å. Therefore, H$_2$O$_2$ is a product of vitamin C scavenging superoxide.

![Figure 13.](image)

Figure 13. Interaction between superoxide and the vitamin C radical shown in Figure 12 (right). Superoxide gives its electron to the scavenger, which becomes a monoanionic quinone-like species. Meanwhile, the short O-O bond distance of the incoming radical, 1.296 Å, may be associated with a leaving O$_2$ molecule, as shown by the separation between both units, 3.713 Å, longer than the van der Waals separation, 2.80 Å.

3.2. Hydrodynamic Voltammetry (RRDE)

The superoxide radical must be generated experimentally to measure its scavenging. There are several methods to do this: (1) an enzymatic reaction with xanthine dehydrogenase [61] and (2) a non-enzymatic option using phenazine methosulphate, NADH, and molecular oxygen [62]. A third non-enzymatic option to generate superoxide is by dissolving potassium superoxide, KO$_2$, in a given solvent [63,64]. However, this method suffers from complex starting conditions since superoxide reacts readily with protons. Thus, for KO$_2$ use, it is necessary to operate either in an anhydrous solution or a strongly basic water environment [65]. Once generated, the superoxide concentration is followed using spectrophotometric, colorimetric, chemiluminescence, and fluorescence methods. Also,
the superoxide can be trapped with 5,5-dimethyl-1-pyrroline-N-oxide (DMPO), and the resultant DMPO-OH adduct is detectable by ESR. These methods all indirectly measure superoxide concentration as they consist of measuring a product generated through superoxide consumption, whose concentration is decreased with the addition of an antioxidant.

A simpler method to generate the superoxide radical is provided by classical cyclic voltammetry in a voltaic cell. Increasing the concentration of antioxidants in the voltaic cell detects the decrease in the current intensity of the superoxide signal [66]. However, the differences among several voltammograms, obtained after successive additions of scavenger, are not well distinguished, and a quantitative measure of superoxide is difficult [41]. We improved this system by splitting reduction and oxidation using a rotating ring–disk electrode (RRDE). At the disk, reduction will produce superoxide following Reaction (2) (lower part of Figure 14), while at the ring, the opposite oxidation reaction will be performed (upper part of Figure 14). This RRDE method measures superoxide scavenging directly, e.g., a real superoxide concentration is detected. A recent review describes several natural polyphenols analyzed with the RRDE technique [57].

\[
O_2 + e^- \rightarrow O_2^-
\]  

(2)

![Figure 14](image)

**Figure 14.** RRDE voltammograms of vitamin E. Each run is associated with a specific color (e.g., red) and corresponds to the red oxidation curve (top, positive current) detected at the ring electrode and the red reduction curve detected at the disk electrode (bottom, negative current). These individual voltammograms show almost no variation; that is, the superoxide destroyed at the ring electrode has a similar value for all runs, suggesting that vitamin E is not a strong antioxidant.

Our first experiment (Figure 14) shows voltammograms of vitamin E in which the different runs are not easily distinguished; that is, the measurements in the variation in superoxide concentration destroyed at the ring electrode, upper part, are almost all similar, which suggests that the vitamin E is not a strong antioxidant in this experiment. This is more clearly and quantitatively indicated in Figure 15, showing the efficiency collection. The estimated line equation, \( y = -0.0018x + 18.391 \), \( R^2 = 0.8981 \), has a slope of \(-1.8 \times 10^3\), which is associated with vitamin E antioxidant capability. Even so, the vitamin E slope is slightly better than that of the previously measured commercial antioxidant butylated hydroxytoluene, BHT \((-1.6 \times 10^3\) [57]. Vitamin E is a component of olive oil, previously analyzed by us [67], and both show closely related scavenging of superoxide.
RRDE vitamin C results (Figures 16 and S3), with slope $-2.6 \times 10^4$, indicate a stronger scavenging activity than that of vitamin E alone. The voltammograms (Figure 16) are much more separated from each other than those shown in Figure 14 for vitamin E. The vitamin C slope of the efficiency graph, $-2.6 \times 10^4$, is like eriodictyol ($-2.2 \times 10^4$), weaker than butein ($-11.2 \times 10^4$) and stronger than the commercial antioxidant BHT ($-0.16 \times 10^4$) [57].

We were interested in analyzing a potential interaction between both vitamins regarding superoxide scavenging. Thus, to a fixed concentration of vitamin E, 640 µL, aliquots of vitamin C were added, and measurements were taken (Figures 17, 18, S3 and S4). Both vitamins are more effective together than individually. Therefore, combining vitamins E and C produces a steeper slope, $-7.2 \times 10^4$ (Figure 18), than that of vitamin E alone, $-1.8 \times 10^3$ (Figure 15), or vitamin C alone, $-1.3 \times 10^4$ (Figure 18). This supports earlier studies describing a cooperative association between the two antioxidant vitamins [1,14,68].
Figure 17. Voltammograms for vitamin C aliquots in fixed amount of vitamin E, 640 µL.

Figure 18. Different slopes of vitamin C and vitamins C plus E. Vitamin C (red line): $y = -13,359x + 20.573 \ R^2 = 0.9947$; vitamin E + vitamin C (blue line): $y = -71,724x + 17.833 \ R^2 = 0.99$.

4. Conclusions

The antioxidant vitamins C and E are essential nutrients that participate in crucial immune system reactions, and they mitigate the results of many diseases related to oxidative stress [6].

In this work, we use the RRDE method to measure the presence of superoxide radicals in a voltaic cell directly and have shown that when administered together, vitamin C and vitamin E antioxidant activity is enhanced. The described superoxide scavenging mechanisms for vitamin C alone and together with vitamin E show marked differences. Vitamin C reacts through its acidic proton and primarily produces $\text{H}_2\text{O}_2$, while vitamin C is transformed into a quinone derivative (Figure 13). In contrast, vitamin E possesses an aromatic ring that is also susceptible to $\pi-\pi$ interaction (not feasible for vitamin C) with a second superoxide radical, and, in fact, the second superoxide remains trapped by vitamin E to form a molecular complex, $\eta\text{-O}_2\text{-vitamin E-model}$ (Figure 7). Moreover, in Scheme 1, the antioxidant action of vitamin E includes the interaction of vitamin E semiquinone with an acidic reagent. This process has been studied in depth using DFT
when the acidic compound is vitamin C (Figure 8) and involves the transfer of an H atom to vitamin E semiquinone (Figure 9) to produce an ascorbate radical plus restored vitamin E. This concurs with the results of earlier studies that ascorbic acid can donate a hydrogen atom to a tocopherol radical at the rate of $2 \times 10^5$ Mol/s because of the difference of 1-electron reduction potential between ascorbic acid (282 mV) and a tocopherol radical (480 mV) [69].

Also, an ESR study in plasma of vitamin C and vitamin E reacting with superoxide [70], in which ascorbate and tocopherol free radicals were subjected to oxidative stress, showed an immediate increase in the concentration of ascorbate radicals, which then steadily declined. Only after the virtual disappearance of the ascorbate radical was the tocopherol radical detected. This seems consistent with our study, where the role of vitamin C is to reform vitamin E, and provides an explanation as to why the ESR signal [70] for the vitamin E radical shows up only when vitamin C is exhausted. In conclusion, these results are important to enhancing our understanding of the synergistic features of these two antioxidant vitamins in different disease states and aging effects in humans.

**Supplementary Materials:** The following supporting information can be downloaded at [https://www.mdpi.com/article/10.3390/biophysica4020022/s1](https://www.mdpi.com/article/10.3390/biophysica4020022/s1), Figure S1. TS (B3LYP calculation): O(superoxide)-H = 1.211 Å, O(VitE)-H = 1.204 Å; Figure S2. The TS structural model after approaching a proton near O6 (Figure 7), which determines O6-H6 formation. E(barrier) = 1.1 Kcal/mol $\Delta G = –82.2$ Kcal/mol. We name the related product $\eta_2$-vitamin E-model; Figure S3. Same calculation of Figure 11 but including the DMSO solvent effect, used in the RRDE experimental section. Distances have similar relevant distances seen in Figure 11. They are 1.025 Å, 1.417 Å, 1.611 Å, and 1.278 Å, respectively; Figure S4. Collection efficiency for Vit C, $y = -26.555x + 21.433$, $R^2 = 0.986$, all data included; Figure S5. Collection efficiency of vitamins C and E, referred to Figure 17; all data included ($y = -63.236x + 17.502$, $R^2 = 0.9913$).

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