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Vitamin E Functions by Association with a Novel Binding Site on the 67 kDa Laminin Receptor Activating Diacylglycerol Kinase

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Running Title: 67LR function as a vitamin E receptor

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Abstract

It is generally recognized that the main function of α -tocopherol (α Toc), which is the most active form of vitamin E, is its antioxidant effect, while non-antioxidant effects have also been reported. We previously found that α Toc ameliorates diabetic nephropathy via diacylglycerol kinase α (DGK α) activation in vivo, and the activation was not related to the antioxidant effect. However, the underlying mechanism of how α Toc activates DGK α have been enigmatic. We report that the membrane-bound 67 kDa laminin receptor (67LR), which has previously been shown to serve as a receptor for epigallocatechin gallate (EGCG), also contains a novel binding site for vitamin E, and its association with Vitamin E mediates DGK α activation by α Toc. We employed hydrogen-deuterium exchange mass spectrometry (HDX/MS) and molecular dynamics (MD) simulations to identify the specific binding site of α Toc on the 67LR and discovered the conformation of the specific hydrophobic pocket that accommodates α Toc. Also, HDX/MS and MD simulations demonstrated the detailed binding of EGCG to a water-exposed hydrophilic site on 67LR, while in contrast α Toc binds to a distinct hydrophobic site. We demonstrated that 67LR triggers an important signaling pathway mediating non-antioxidant effects of α Toc, such as DGK α activation. This is the first evidence demonstrating a membrane receptor for α Toc and one of the underlying mechanisms of a non-antioxidant function for α Toc.

Keywords: vitamin E, 67 kDa laminin receptor, diacylglycerol kinase, epigallocatechin gallate

INTRODUCTION

Tocopherols are hydrophobic antioxidant agents known generally as vitamin E. D- α -tocopherol (α Toc) is the most active Vitamin E compound that humans preferentially absorb and accumulate [1]. The main function of α Toc is recognized as its antioxidant effect preventing lipid peroxidation of polyunsaturated fatty acids (PUFAs) in the cell membrane [2], but many other functions, such as activation of signal transduction and gene expression, that are not related to the antioxidant effect, have been reported [3]. For example, it has been suggested that α Toc promotes drug-metabolizing gene expression via the nuclear receptor pregnane X receptor (PXR) [4]. Also, it has been reported that α Toc inhibits the activity of protein kinase C [5]. Vitamin E was discovered in 1922 as an essential food component, but the specific target for vitamin E that mediates the non-antioxidant effect is still enigmatic. As a specific α Toc binding protein, α -tocopherol transfer protein (α -TTP) plays a critical role in the transport of α Toc in the liver, and deficiency of α -TTP causes ataxia with vitamin E deficiency (AVED) [6–8]. However, a cell membrane receptor for α Toc, which mediates specific signal transduction, has not been reported.

Koya *et al.* [9] reported that α Toc ameliorated the symptoms of diabetic nephropathy (DN) by inhibiting diabetes-induced abnormal protein kinase C (PKC) activation in glomerulus through activation of a lipid kinase: diacylglycerol kinase (DGK). We have reported that α Toc activates DGK α and that α Toc ameliorated DN via DGK α in vivo [10–12]. Note that acetated α Toc, which lacks antioxidant effect induces DGK α activation indicating the DGK α activation is a non- antioxidant effect of α Toc [10]. These facts suggested that the effect of α Toc on DN involves the non-antioxidant effects of α Toc. Furthermore, the involvement of a receptor for α Toc in the α Toc-induced DGK α activation was suggested since the activation was an indirect effect [10]. However, the identity of the specific receptor which mediates α Toc signaling has not been reported.

The 67 kDa laminin receptor (67LR) is primarily found as a non-integrin laminin receptor [13–15]. It has also been reported that the Sindbis virus and pathogenic prion protein are the ligands for the 67LR [16,17]. In addition to these ligands, epigallocatechin gallate (EGCG), one of the green tea polyphenols, is a ligand for 67LR [18]. Finally, the expression level of 67LR is upregulated in various cancer cells, and EGCG shows an anti-cancer effect through 67LR [19–23]. Recently, we have found that EGCG indirectly activates DGK α and that 67LR is involved in the activation [24]. Also, we reported that EGCG ameliorated DN through DGK α in vivo, similar to α Toc [25]. Based on these facts, we hypothesized that α Toc also activates DGK α through the 67LR and that 67LR functions as an α Toc receptor mediating the α Toc signaling. Therefore, we explored whether α Toc activates DGK α directly through 67LR and the binding of α Toc to 67LR by taking advantage of hydrogen-deuterium exchange mass spectrometry (HDX/MS) and molecular dynamics (MD) simulation approaches.

HDX/MS is a powerful structural approach that can determine the specific interactions between a protein and its ligands in solution [26,27]. With HDX/MS, one can determine where various ligands bind to the protein by determining the inhibition of hydrogen-deuterium exchange in specific peptides in the protein backbone in the presence of specific ligands. In this study, we employed HDX/MS to discover a novel α Toc binding site and to confirm the EGCG binding site on 67LR. Moreover, we performed MD simulations to

investigate the detailed binding mode and confirmed the identity of the binding site using several mutants of 67LR.

METHODS

Cell culture

The *Mesocricetus auratus* leiomyosarcoma cell line (DDT1-MF-2 cells), *Cercopithecus aethiops* kidney cell line (Cos7 cells), and *homo sapiens* colorectal carcinoma cell line (HCT116 cells) were cultured in Dulbecco's modified Eagle medium (DMEM)-high glucose supplied 10% fetal bovine serum, penicillin (100 units/ml), streptomycin (100 mg/ml) and L-glutamine. The cells were maintained in a humidified atmosphere containing 5% CO₂ at 37°C.

Imaging of the DGK α translocation

The live imaging of the DGK α translocation was performed following our previous method [24]. In short, 1.0×10^5 DDT1-MF-2 cells were inoculated in a glass-bottom 35 mm dish and allowed to adhere for 24 h. The N-terminal GFP-tag conjugated DGK α encoding plasmid was transfected using FuGene[®] HD Transfection Reagent (Promega Corporation, Madison WI, USA). After 24 h of transfection, the medium was replaced with buffer (165 mM NaCl, 5 mM KCl, 1 mM MgCl₂, 1 mM CaCl₂, 5 mM HEPES, 10 mM Glucose) and stimulated with various concentrations of α Toc, α Toc derivatives, or EGCG. Cells were observed with a confocal scanning microscope FV-1000 (Olympus Corporation, Tokyo, Japan). When the N-terminal FLAG-tag conjugated 67LR or its mutants were overexpressed, each plasmid was mixed equally with GFP-DGK α encoding plasmid, and used the mixture for transfection. When the cells were pretreated with the 67LR antibody, 20 μ g/ μ l of the 67LR polyclonal antibody (N2C3, Cat. # GTX113485, GeneTex, Inc., Irvine CA, USA) was added into media 2 h before stimulation, and the same volume of normal rabbit serum was used as a negative control. The siRNA of 67LR (Sequence: 5'-UUCGCCCCUGAAAUUCCUCC-3' and 5'-GGAGGAAUUCAGGGCGAA-3') was transfected 24 h before transfection of GFP-DGK α . The translocation rate was calculated as a percentage of cells with translocated GFP-DGK α among at least 60 cells that express GFP-DGK α .

Binding assay of α Toc to recombinant 67LR

The FLAG-67LR and FLAG- α -TTP were overexpressed in the Cos7 cells using an electroporation method described previously [24]. The recombinant FLAG-tagged proteins were precipitated by anti-FLAG tagged antibody conjugated resin. The resin was incubated with various concentrations of α Toc contained phosphate-buffered saline (PBS) for 15 min. The resin was washed, and bound α Toc was extracted using 300 μ l of ethyl acetate/hexane (1/9 v/v). The solvent was dried with nitrogen gas, and the extract was dissolved in 50 μ l of 100% methanol. The sample was analyzed by the HPLC system using a TSKgel ODS-120T 2.0x250 mm column (Tosoh Corporation, Tokyo, Japan), and the peak of α Toc was detected by UV (excitation: 298 nm emission: 330 nm). The empty vector was used as a negative control.

Preparing tocopherol conjugated beads

The tocopherol conjugated beads were prepared using D- α -tocopherol succinate (α Tos) and NH₂ beads (TAMAGAWA SEIKI Co., Ltd., Nagano, Japan) by following the supplier recommended procedure. In brief, 10 mM α Tos, 10 mM N-hydroxysuccinimide, and 10 mM 1-(3-dimethylaminopropyl)-3-ethyl carbodiimide hydrochloride were mixed in N, N-dimethylformamide (DMF) and activated by incubating for 2 h at 25°C. The 2.5 mg of NH₂ beads was incubated with 500 μ l of activated ligand in solution for 16 h at 25°C. Finally, the beads were washed with DMF and incubated with 500 μ l of 20% acetic anhydride in DMF to mask the remaining NH₂ moiety of the beads for 2 h at 25°C. The beads were washed and stored in H₂O at 4°C until use. The control beads were prepared with the same procedures without α Tos.

Binding assay using tocopherol conjugated beads

The FLAG-67LR overexpressing Cos7 cell lysate was incubated with 0.5 mg of the tocopherol conjugated beads for 2 h at 4°C. The beads were washed with PBS containing 0.03% Triton X three times. The FLAG-67LR bound to tocopherol beads was detected by western blot using an anti-FLAG antibody for the primary antibody. The intensity of the bands was analyzed by Image J software.

Differential scanning fluorometry (DSF)

The N terminal GST and C terminal 6 \times His tag conjugated 67LR was purified using glutathione sepharose as describe following section. The mixture containing 10 or 100 μ M of EGCG and α Toc, 10 μ M of purified protein, and 2 \times SYPRO® Orange (Thermo Fisher Scientific Inc., Waltham MA, USA) in the buffer (25 mM HEPES, pH 7.5, 100 mM NaCl) was incubated for 30 min at 25°C. After the incubation, the sample was heated from 25 °C to 70 °C by 2 °C/min, and the fluorescence was scanned using CFX Connect (BioRad Laboratories, Inc., Hercules, CA, USA) every 30 sec.

Isothermal Titration Calorimetry (ITC)

The ITC experiment was performed using MicroCal iTC200. The 20 mM α Toc stock solution in DMSO was dissolved in the buffer (25 mM HEPES, pH 7.5, 100 mM NaCl) by 1/1000 dilution (the final concentration was 20 μ M) and loaded to sample cell. The 120 μ M purified recombinant GST-67LR-His protein was dialyzed against the same buffer with the α Toc sample and loaded to the injection syringe. The same amount of DMSO with the α Toc sample was added to the protein sample. The protein sample was injected into the sample cell by 19 individual injections of 1.75 μ l sample with 250 sec initial delay and 180 sec spacing time. The titration was performed at 25 °C (298K), and the sample cell was stirred by 750 rpm to ensure thorough mixing during the titration. The data were processed using Origin software, and K_d values were calculated. The experiments were performed three times using two independent batches of a protein sample, and representative data was shown as the final figure. The control experiments were performed by injecting the buffer to the α Toc sample in the cell and the protein sample to the buffer in the cell to measure the heat of dilution.

Construction of 67LR mutants

The N-terminal FLAG-tagged 67LR mutants were constructed by the long PCR method using a FLAG-67LR plasmid that was prepared previously [24] as a template. The plasmid was

amplified with respective primers shown in Table S1. The amplified products were transfected into DH5 α *E. coli*. Several single colonies were picked, and the plasmid was purified to confirm the mutation using a restriction enzyme. Finally, the mutations were confirmed by DNA sequencing.

Molecular docking.

The 3D structure of the 67LR receptor (PDB ID: 3BCH) [28] was prepared using the Protein Preparation Wizard [29,30]. The 3D structures of α Toc and EGCG were sketched using Maestro sketcher, and they were optimized using the LigPrep [30,31]. The box center for the docking calculations was defined using the centroid of selected residues that were found to constitute the EGCG binding site [18]. The grid required for the docking procedure was generated using a scaling factor of 1.0 and a partial charge cutoff of 0.25, while the X, Y, Z dimensions of the inner box were set to 12 Å. For the ligand docking, a scaling factor of 0.8 and partial charge cutoff of 0.15 were used to allow complete flexibility of the structures. The poses were selected according to the binding mode and the XP GScore. The Glide Extra-Precision (XP) scoring function was used for the calculations [32,33].

Expression and purification of recombinant 67LR

Recombinant 67LR protein was prepared by a bacterial expression system using *E. coli*. N-terminal GST tagged and C-terminal 6 \times His tagged conjugated 67LR encoding plasmid was transfected into BL21 *E. coli*. The *E. coli* were pre-cultured for 16 h at 37°C, and 20 ml of pre-cultured *E. coli* was cultured in 1 L of LB medium until the OD₆₀₀ reached 0.7~1.0. The expression was induced with 0.2 mM of IPTG for 6 h at 25°C. The *E. coli* was harvested and lysed in lysis buffer (20 mM Tris-HCl pH 8.0, 1 mM EDTA, 1 mM DTT, 5 mM MgCl₂, 250 mM sucrose, 1 mM PMSF, 20 ng/ml leupeptin). The lysate was centrifuged, and the supernatant was subjected to purification using glutathione sepharose. The eluted protein sample was subjected to the DSF and the ITC experiment. For the HDX/MS experiment, the GST tag was removed by the following procedures. The elution fraction of the GST purification was dialyzed against buffer for HRV 3C protease (50 mM Tris-HCl pH 7.0, 150 mM NaCl, 1 mM EDTA, 1 mM DTT), and the GST tag was removed by HRV 3C protease (GE Healthcare, Chicago IL, USA). After the protease cleavage, the protein was purified with Ni Sepharose[®] (GE Healthcare, Chicago IL, USA) in accord with the supplier recommended procedure. The purified protein was dialyzed against buffer for HDX/MS (8.3 mM Tris pH7.2, 50 mM NaCl 10% glycerol) and was concentrated by Amicon[®] Ultra (Milliporesigma, Burlington MA, USA) up to 1.5 mg/ml.

Hydrogen-deuterium exchange mass spectrometry

The experiments were performed by following the methods previously published [34]. In brief, 2 μ l of protein solution was incubated with 1 μ l of 13 mM EGCG or α Toc in the buffer for HDX/MS for 20 min at 25°C. The following steps were performed at 4°C. The deuterium exchange was initiated by adding 9 μ l of D₂O buffer (8.3 mM Tris-HCl pD 7.2, 50 mM NaCl, in 98% D₂O). The exchange solution was quenched after 10, 100, 1000, and 10000 seconds with 18 μ l of quenching solution (0.08 M guanidine HCl, 0.8% formic acid, 16.6% glycerol), and the sample was immediately frozen using dry ice. The sample was kept at -80°C until subjecting to proteolysis liquid chromatography-mass spectrometry analysis using the pepsin column. The data were analyzed using Proteom Discoverer software (Thermo Fisher LLC,

San Jose CA, USA) and DXMS Explorer (Sierra Analytics Inc., Salida CA, USA) by following the previous report [34]. HDX/MS experiment was performed twice using independent batches of purified 67LR protein, and reproducible peptide regions are shown.

Receptor-membrane systems

The Membrane Builder implemented in CHARMM-GUI was employed to generate 3D models for the MD simulations [35,36]. The membrane patch consisted of phosphocholine (PC), phosphoethanolamine (PE), phosphoserine (PS), phosphoinositol (PI), and sphingomyelin (SPH). The membrane composition was selected based on the localization of 67LR on cellular membranes according to previously published data [37,38]. Each system was solvated with TIP3P water molecules and neutralized with 150 mM sodium chloride (NaCl) using the Visual Molecular Dynamics (VMD) package [39].

Equilibration and production run.

Molecular dynamics simulations were carried out using NAMD 2.12 [40]. The following minimization and equilibration protocol was performed: a minimization of 80,000 steps was initially performed by applying harmonic constraints on the receptor-ligand-membrane that were gradually turned off using a constraint scaling factor, followed by a second 120,000 steps minimization without constraints. An initial equilibration of 10,000 steps was performed by also applying harmonic constraints on the receptor-ligand-membrane that were gradually turned off using the same constraint scaling factor, followed by a second 10,000 steps equilibration without constraints. During the equilibration, each system was slowly heated and held to 310 K using a temperature reassignment with a reassignment frequency of 500-time steps (1000 fs) and a reassignment increment of 1 K. The above minimization and equilibration protocol was sufficient to induce the appropriate disorder of a fluid-like bilayer, avoid unnatural atomistic positions, and failure of the simulations by atoms moving at very high velocities. Each system was finally subjected to a 1 μ s production run. For each production run, the temperature was maintained at 310 K using the Langevin thermostat with the Langevin coupling coefficient of 1/ps [41]. The NPT ensemble was employed, and the pressure was kept constant at 1.01325 kPa using the Langevin piston method with the “useGroupPressure,” “useFlexibleCell,” and “useConstantArea” parameters turned on [42]. A time step of 2 fs was used in combination with the SHAKE algorithm to hold the bonds of hydrogen atoms similarly constrained [43]. Nonbonded interactions and full electrostatics were calculated every 1 and 2-time steps, respectively. Switching functions are used to smoothly take electrostatic and van der Waals interactions to zero with a switching distance of 10 Å and a cutoff of 12 Å. Long-range electrostatic forces in the periodic system were evaluated using the Particle Mesh Ewald (PME) Sum method with grid spacing 1/Å [44]. The CHARMM General Force Field (CGenFF) and the CHARMM36 all-atom additive force field and parameters were used for the simulations [45,46]. The simulation was run twice with the same settings. The main figures are generated from the first simulation.

Cell viability assay

HCT116 cells were inoculated in 60 mm dishes with 2.0×10^5 cell/dish and cultured for 48 h. The medium was changed so as to include various concentrations of EGCG or α Toc in a fresh medium. After 48 h incubation, the number of viable cells was counted. When the

endogenous 67LR was knocked down, siRNA targeting 67LR was transfected 24 h after inoculating the cells using the same method described above.

Statistical analysis

All error bars show standard error of the mean (SEM). Student's t-test was carried out when comparing two data sets. One-way ANOVA followed by Dunnet's multiple comparison test was carried out when the experiment had more than three groups using BellCurve for Excel Version 3.20. A *p*-value of less than 0.05 was considered to be significant.

RESULTS

DGK α activation by α Toc through 67LR

Vitamin E consists of four tocopherols and four tocotrienols classified α , β , γ , and δ by the position of the methyl moieties on the chroman ring structure. α Toc is a hydrophobic compound that has three methyl moieties on positions 5, 7, and 8 of the chroman ring (Fig. 1a). α Toc induces translocation of GFP-tagged DGK α to the plasma membrane, which is a hallmark of DGK α activation (Fig. 1b) [10]. To investigate if 67LR contributes to the translocation, cells were pretreated with 67LR antibody which blocks GFP-DGK α translocation by EGCG [24], and α Toc-induced translocation of GFP-DGK α was monitored. The pretreatment of the 67LR antibody significantly inhibited the translocation induced by α Toc (Fig. 1c), suggesting α Toc activates DGK α via 67LR. Indeed, the knockdown of endogenous 67LR by siRNA also significantly attenuated the translocation of DGK α (Fig. 1d). Also, we tested the effect of all tocopherols on the translocation of DGK α , and revealed that not only α Toc but also the other tocopherols induced translocation of GFP-DGK α to the plasma membrane (Fig. 1e). Then, we compared the translocation efficiency induced by all tocopherols and found that they had a similar effect on the DGK α translocation (Fig. 1f). These results indicated that all tocopherols induce DGK α activation via 67LR and the possibility that 67LR acts as a receptor for tocopherols.

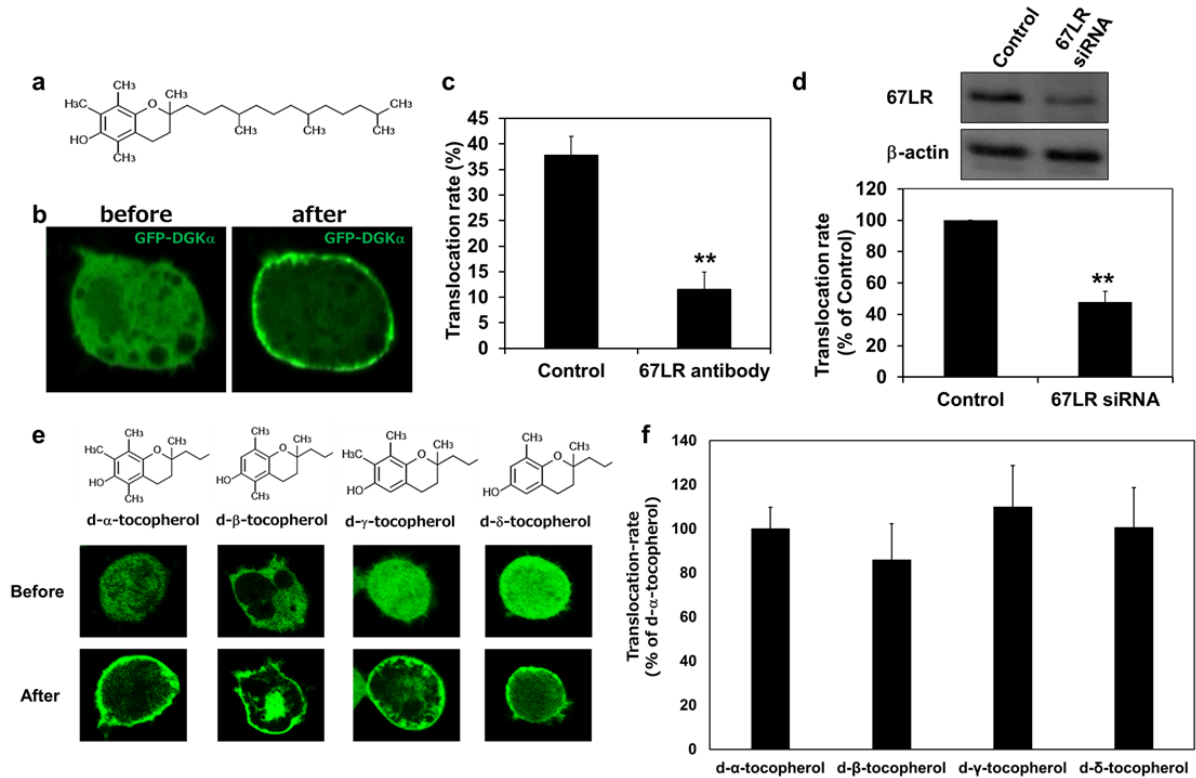


Fig. 1. Translocation of DGK α to the plasma membrane induced by tocopherols. (a) Structure of α Toc. (b) Localization of GFP-DGK α in DDT1-MF-2 cells before and 3 minutes after 200 μ M α Toc stimulation. (c) The translocation rate of GFP-DGK α in DDT1-MF-2 cells by 200 μ M α Toc in the absence or presence of anti-67LR antibody treatment. $N=4$. (d) Effect of the knockdown of 67LR on the translocation of GFP-DGK α in DDT1-MF-2 cells by 200 μ M α Toc. $N=3$. (e) Localization of GFP-DGK α in DDT1-MF-2 cells before and 3 minutes after 200 μ M tocopherol stimulation. The difference in the position of the methyl moiety in the structure of each of the tocopherols is shown above the respective pictures. (f) The translocation rate of GFP-DGK α in DDT1-MF-2 cells induced by each of the tocopherols. $N=3$. The data shows the percentage of the translocation rate relative to α Toc. The values are mean \pm SE. Asterisks indicate significance (** $p < 0.01$). Full-length blots are presented in Fig. S1.

Direct binding of α Toc to 67LR and its affinity

We investigated the direct binding of α Toc to 67LR and its affinity by various approaches. It has been reported that 67LR is a matured dimer form of the 37 kDa laminin receptor precursor (37LRP), but the detailed relationship was not understood [47]. Note that, in the present study, we employed cDNA encoded 37LRP to express recombinant 67LR. First, we precipitated FLAG-tagged recombinant 67LR by FLAG tag antibody-conjugated beads, and the beads were incubated with α Toc. The ratio of bound α Toc to precipitated 67LR was analyzed by HPLC. FLAG-tagged α -TTP was used as a positive control. Consequently, a significant amount of α Toc was detected in both FLAG-67LR and FLAG- α -TTP precipitated beads comparing to FLAG tag alone (Fig. 2a). In addition, the amount of bound α Toc

increased concentration-dependently (Fig. 2b). Next, we made tocopherol conjugated beads and analyzed the amount of FLAG-67LR bound to the tocopherol beads. The scheme for the production of tocopherol conjugated beads is shown in Fig. S3. Control beads and tocopherol conjugated beads were incubated with lysate of FLAG-67LR overexpressed cells, and the bound FLAG-67LR was detected by western blot. Although some amount of FLAG-67LR nonspecifically bound to the control beads, significantly more FLAG-67LR was detected with the tocopherol conjugated beads (Fig. 2c). While, α -TTP did not bind to the tocopherol conjugated beads (data not shown). This might be related to the binding mode of α Toc in α -TTP [48,49].

Further, to confirm the binding, we performed a differential scanning fluorometry (DSF) using 10 μ M purified recombinant GST-67LR-His protein. Two independent peaks showing the T_m value of protein, which might be delivered from 67LR and GST tag protein, were detected (Fig. S5). The first peak shifted to 1 $^{\circ}$ C and 2 $^{\circ}$ C higher temperature in the presence of 10 μ M and 100 μ M α Toc, respectively, and α Toc did not show any effect on the second peak (Fig. S5a). Importantly, although 10 μ M EGCG did not show any effect, 100 μ M EGCG increased the T_m value of the first peak without any effect on the second peak (Fig. S5b). These results indicated that the first peak shows the T_m value of 67LR, and EGCG and α Toc increase the thermal stability of 67LR by direct binding. Finally, to measure the affinity of α Toc to 67LR, we carried out the isothermal titration calorimetry (ITC). By titrating the 120 μ M purified GST-67LR-His to 20 μ M α Toc in the sample cell, absorption of heat by the interaction of α Toc to 67LR was detected, and the absorption disappeared eventually due to the saturation of the ligand (Fig. 2d). The sigmoid curve was obtained by processing the area under the curve of each peak, and the mean of K_d value was 788 ± 174 nM (Fig. 2e). Notably, the heat of dilution from α Toc and GST-67LR-His was not significant (Fig S6a, b). These approaches successfully demonstrated the direct binding of α Toc to 67LR and its affinity.

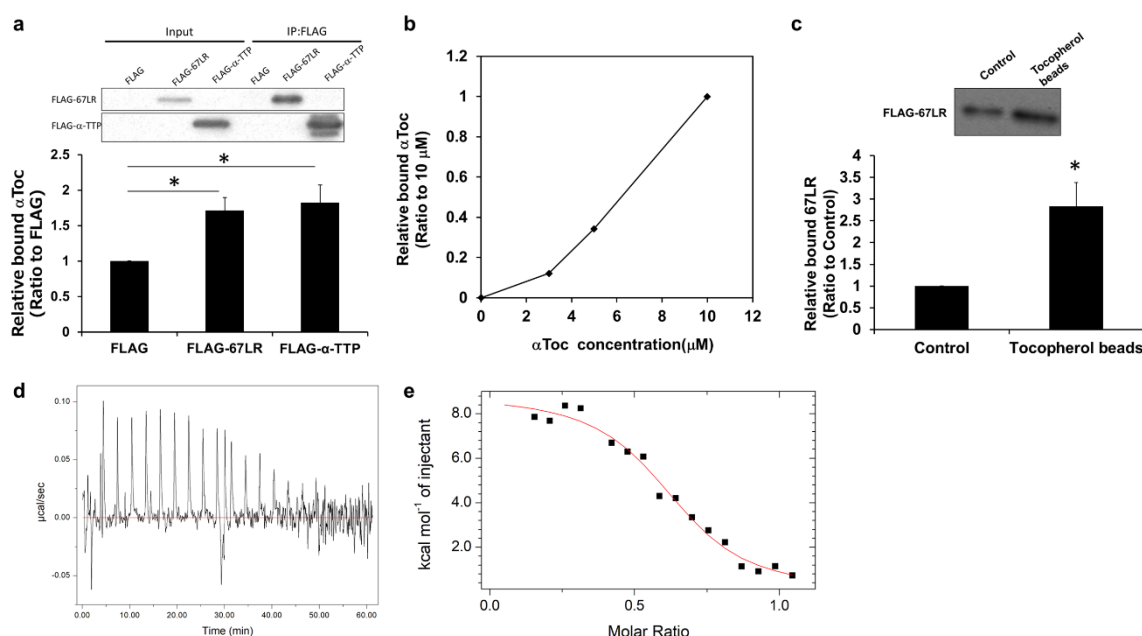


Fig. 2. Direct binding between α Toc and 67LR and its affinity. (a) The result of the binding assay between 10 μ M α Toc and precipitated FLAG-67LR or FLAG- α -TTP using HPLC is shown. The expression and precipitation of FLAG-67LR and FLAG- α -TTP were

detected by western blot. The value is the ratio of bound α Toc to FLAG alone. $N=4$. **(b)** Concentration-dependent binding of α Toc to 67LR detected by the binding assay using HPLC. The value was shown as a percentage of 10 μ M. **(c)** The result of the binding assay using tocopherol conjugated beads. The amount of precipitated FLAG-67LR by the control and tocopherol conjugated beads was detected by western blot. **(d)** The representative raw data of ITC for titration of 120 μ M GST-67LR-His to 20 μ M α Toc at 298K. **(e)** The integrated area under the curve of each peak from **(d)** was plotted by the molar ratio of GST-67LR-His and α Toc. The best fit curve was drawn by the Origin software, and the K_d value was calculated based on the curve. The intensity of bands in the western blots were analyzed by Image J. The error bars show SE. Asterisks indicate significance ($*p < 0.05$). Full-length blots are presented in Fig. S2 and S4.

α Toc binding site of 67LR

Having shown that α Toc directly binds to 67LR, we then investigated exactly how α Toc binds to 67LR. The putative EGCG binding motif of 67LR and the mutations which abolish binding to EGCG, such as the Lys166Glu mutant, have been reported, although the detailed binding model of EGCG was not known [50]. To address this question, we constructed a FLAG-67LR Lys166Glu mutant and observed its effect on the translocation of DGK α induced by EGCG and α Toc. We checked if FLAG-tagged 67LR localizes at the plasma membrane and observed membrane localization of FLAG-67LR at the plasma membrane (Fig. S7). When the mutant was overexpressed, the rate of the EGCG-induced DGK α translocation was significantly less than with wild-type overexpressing cells (Fig. 3a), confirming that Lys166 is critical for the binding to EGCG. In contrast, α Toc-induced translocation with overexpression of the mutant was the same as that with overexpression of the wild type (Fig. 3b). These results suggested that the α Toc binding site of 67LR is different from that of EGCG. Then, we performed a docking simulation of the EGCG and α Toc to the putative EGCG binding site employing Glide software and the crystal structure of 67LR [51]. The putative EGCG binding site is in a water-exposed hydrophilic region, and EGCG docks to the binding site with several hydrogen bonds (Fig. 3c, d). Indeed, the XP GScore, which represents binding stability, was decent (-6.723). However, α Toc did not dock well to the binding site based on a poor XP GScore (-0.623) (Fig. 3e), suggesting that the α Toc binding site in the 67LR is different from that of EGCG.

We then carried out HDX/MS experiments to locate the α Toc binding site on 67LR experimentally. Also, we aimed to confirm the EGCG binding site on 67LR by HDX/MS. The effect of the presence of α Toc or EGCG on the deuterium exchange ratio was analyzed by comparing the deuterium level in the absence of ligands. The peptide regions which included Lys11 to Lys17, Asn50 to Ala61, and Trp176 to Leu183 in the α -helices were inhibited after deuterium exchange by the presence of α Toc while, significant decreases in the number of deuterium were not observed in the neighboring peptides to these regions such as Phe16 to Thr28, Phe32 to Ile46, and Leu183 to Tyr202 (Fig. 3f). These residues appeared to form a reasonable pocket for α Toc, suggesting this region as a probable α Toc binding site. In the case of EGCG, significant inhibition of deuterium exchange in the regions including residues Leu16 to Met34 and Ile161 to Gly172 that are close to the putative EGCG binding

site was confirmed compared to the neighboring peptides such as Gly-5 (linker region) to Gln9 and Leu183 to Leu203 that showed no inhibition (Fig. 3g).

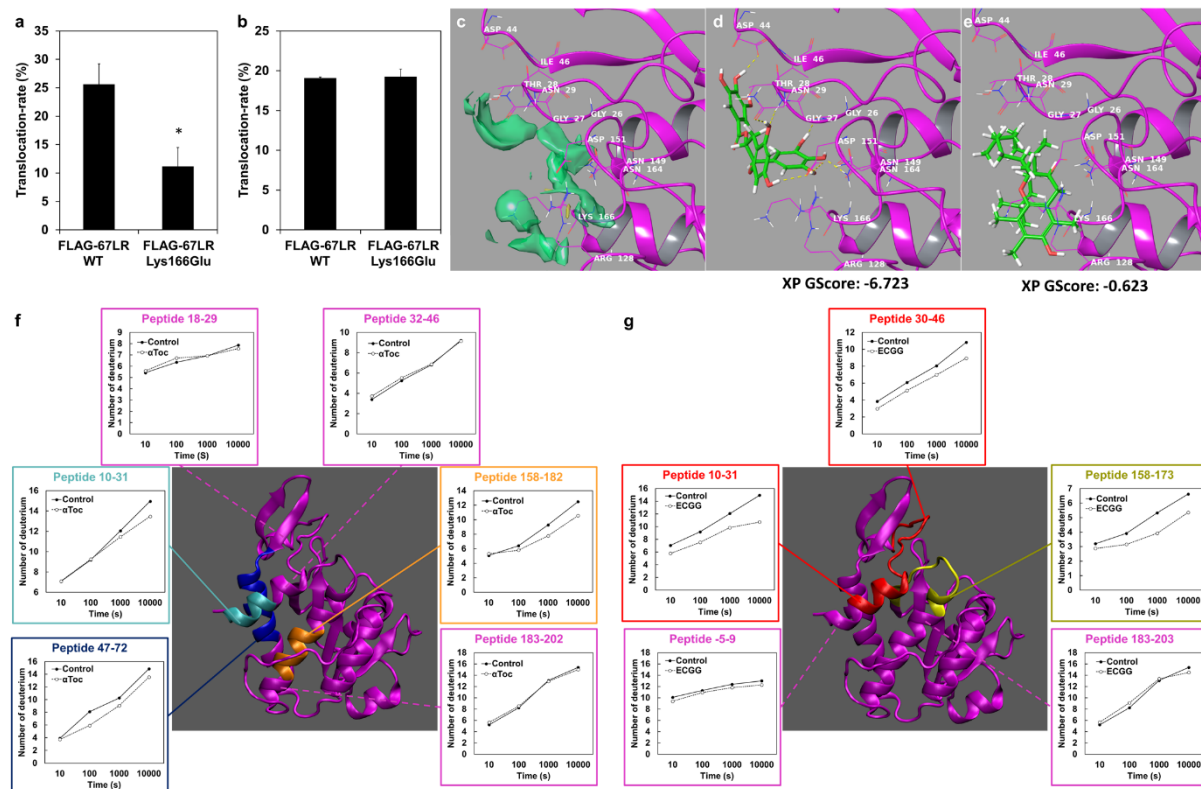


Fig. 3. The putative α Toc binding site on 67LR determined by HDX/MS. The effect of overexpression of FLAG-67LR Lys166Glu on 200 μ M (a) EGCG or (b) α Toc-induced translocation of GFP-DGK α in DDT1-MF-2 cells. $N=3$ (c) The image of the EGCG binding site of 67LR (green: hydrophilic surface). The docking simulations of (d) EGCG and (e) α Toc to the putative EGCG binding site of 67LR by Glide software. Hydrogen bonds are shown as a yellow dashed line. (f) The change in the number of deuterium in the typical reproduced regions by α Toc (Cyan: Lys11-Lys17, Blue: Asn50-Ala61, Orange: Trp176-Leu183). (g) The change in the number of deuterium in the typical reproduced regions by EGCG (Red: Leu16-Met34, Yellow: Ile161-Gly172). The dashed lines point the residues neighboring color-coded residues show no significant difference in number of deuterium. The amino acid sequence includes the remaining N-terminal linker to the GST tag and the C-terminal 6 \times His tag. The error bars show SE. Asterisks indicate significance (* $p < 0.05$).

Binding mode of α Toc and EGCG in 67LR

MD simulations were carried out to elucidate if and how α Toc docks to the region suggested by HDX/MS. We created a model of 67LR in a lipid bilayer using CHARMM-GUI Membrane Builder, with the composition of the lipid bilayer mimicking that of lipid rafts based since 67LR is present in lipid rafts [38]. In the model, 67LR penetrated the lipid bilayer, but the predicted EGCG binding site was exposed to water (Fig. S8 and S9b). Based on the model and the results of HDX/MS, the initial position of α Toc was placed in the lipid

bilayer near the region we experimentally determined (Fig. S9a), and the initial position of EGCG was placed exposed to the water near the putative binding site since EGCG is a water-soluble compound (Fig. S9b).

In the simulation of α Toc, α Toc drifted in the lipid bilayer at the beginning of the simulation but gradually approached the putative binding region (Movie S1). Finally, α Toc bound to 67LR by penetrating its phytyl moiety into the α -helix assembling region (Movie S1). The optimal binding mode showed that the phytyl moiety of α Toc completely penetrated the putative binding region of 67LR (Fig. 4a). When α Toc approached the binding site, the chroman ring structure faced to Phe203, and the phytyl moiety then got inside afterward (Movie S2). Finally, the aromatic residues such as Trp176, Phe203, and Tyr202 packed the chroman ring and would stabilize the binding (Fig. 4b). The 2D binding analysis of its optimal binding mode revealed that the backbone of Tyr202 interacts with the hydroxy group of the chroman ring structure with relatively weak hydrogen bonds (donor-acceptor distance: 2.99Å, donor-hydrogen-acceptor angle: 151.2°), and a hydrophobic pocket formed containing Phe18, Trp55, Lys17, Leu58, Trp176, and Met177 accommodate the phytyl moiety (Fig. S10a). We repeated the simulation to confirm the result, and the second simulation was well consistent with the first simulation. In brief, the phytyl moiety of α Toc got into the hydrophobic pocket with similar movement (Movie. S3). In the second run, π -H interactions were detected on Trp176 and Phe18, while hydrogen bonds between Phe202 was not detected (Fig. S10c). Importantly, the same hydrophobic pocket accommodated α Toc in both simulations.

In the EGCG binding simulation, EGCG approached its binding site and bound to the putative EGCG binding region which includes the region found by HDX/MS (Movie S4 and Fig. 4c). The simulation also confirmed an interaction between the galloyl moiety of EGCG and Lys166 in the optimal binding mode (Fig. 4d and Movie S5). In addition, it was turned out that Asp126 and Asp151 frequently interacted with the B-ring and galloyl moiety of EGCG, respectively, by hydrogen bonds (Fig. 4d and Movie S5). In the optimal binding mode, the galloyl moiety formed strong hydrogen bonds with Asp151 (donor-acceptor distance: 2.47Å, donor-hydrogen-acceptor angle: 164.3°). In addition to these residues, Asn149 and Asn165 were also contributing to the binding of EGCG by interacting with the galloyl moiety (Fig. S10b). The second simulation also showed a similar binding mode with the first run (Movie S6). Notably, although there are some differences in the hydrogen bond pair, Asp126, Asp151, Asn165, and Lys166 formed hydrogen bonds with B ring or galloyl moiety, indicating that these residues are critical for EGCG binding (Fig. 10d).

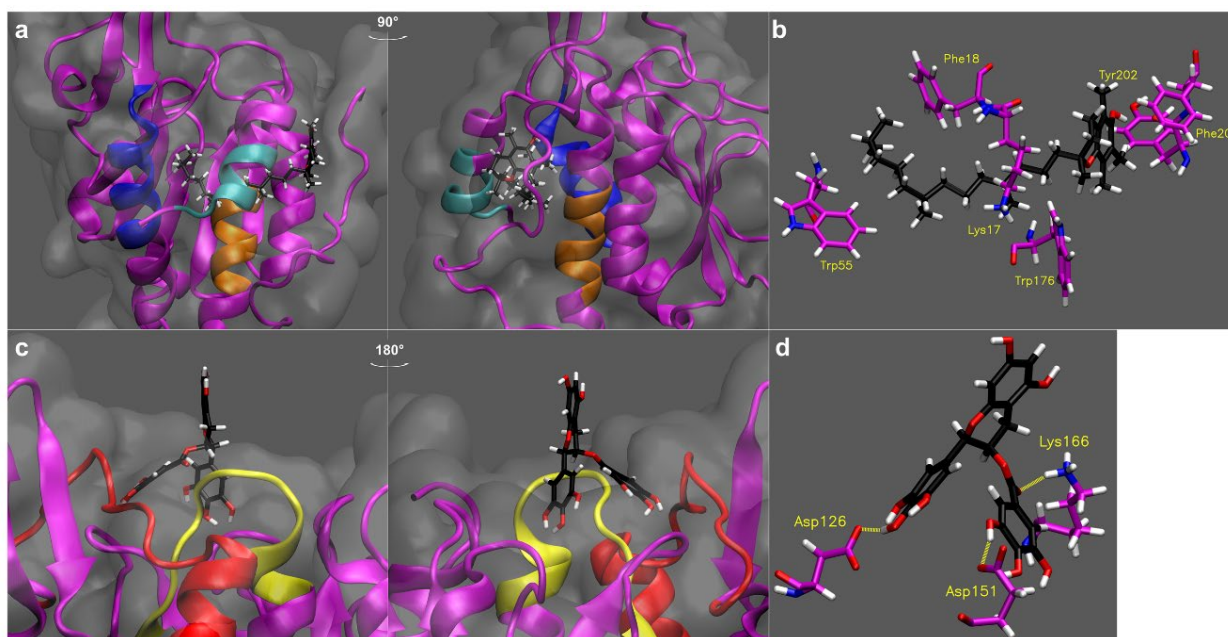


Fig. 4. Binding simulations of α Toc and EGCG to their respective putative binding sites. The binding mode of (a) α Toc and (c) EGCG in 67LR (the final frame of the simulations shown in Movie S1 and S3). The typical amino acid residues in the binding mode of (b) α Toc and (d) EGCG in 67LR. The structure of α Toc and EGCG are shown in black, and the colored regions are the same as shown in Fig. 3, respectively. The surface of the protein was shown in a gray transparent cloud. The hydrogen bonds are shown as a yellow dashed line (distance cutoff: 3.2Å, angle cutoff: 30°).

The effect of mutations in the α Toc binding pocket of 67LR on DGK α translocation

To confirm the identity of the α Toc binding site of 67LR, we constructed several point mutants of 67LR based on the optimal binding mode from MD simulation (Fig. 5a) and observed the effect of overexpression of the mutants on the α Toc and EGCG-induced translocation of GFP-DGK α . Consistent with the results in Fig. 3a and b, overexpression of Lys166Glu, as well as wild-type (WT) 67LR showed significantly enhanced α Toc-induced translocation of DGK α comparing to the control, while overexpression of Lys166Glu showed no effect on EGCG-induced translocation (Fig. 5b, c), indicating Lys166, contributes EGCG binding but not α Toc binding as HDX/MS and MD simulation revealed. First, among the regions suggested by HDX/MS and MD simulations for α Toc binding, we tested the mutants of Trp55 and Trp176. The former is in the hydrophobic pocket, and the latter is close to the chroman ring structure (Fig. 4b and Fig. 5a). These mutants did not enhance the α Toc-induced translocation of DGK α (Fig. 5b), indicating that these residues contribute to the binding of α Toc. In addition, we tested two mutants that are close to the region accommodating the chroman ring structure but seemed not to be essential for the binding, such as Phe90 and Lys89, to confirm the accuracy of the binding model from the MD simulations (Fig. 5a). Although there is no statistical significance, Phe90Ala showed a tendency to enhance α Toc-induced translocation, while Lys89Ala significantly enhanced the translocation (Fig. 5b). These results suggest that Phe90 partly contributes to the binding of

α Toc, but it is not essential and that Lys89 does not contribute to the binding of α Toc as MD simulation indicated. Indeed, the distance from the chroman ring to Lys89 is much longer (more than 6 Å) than that to Phe90 (5.63 Å) in the optimal binding mode of α Toc. Notably, these mutants, except for Lys166Glu, significantly enhanced the EGCG-induced translocation of DGK α as WT, confirming that the mutants only affected the binding to α Toc (Fig. 5c). From these results, it is clear that Trp55, Phe90, and Trp176 are in the putative α Toc binding site suggested by HDX/MS and MD simulations and are partly or critically contributing to the α Toc binding, confirming that the proposed binding mode of α Toc in 67LR is reasonable.

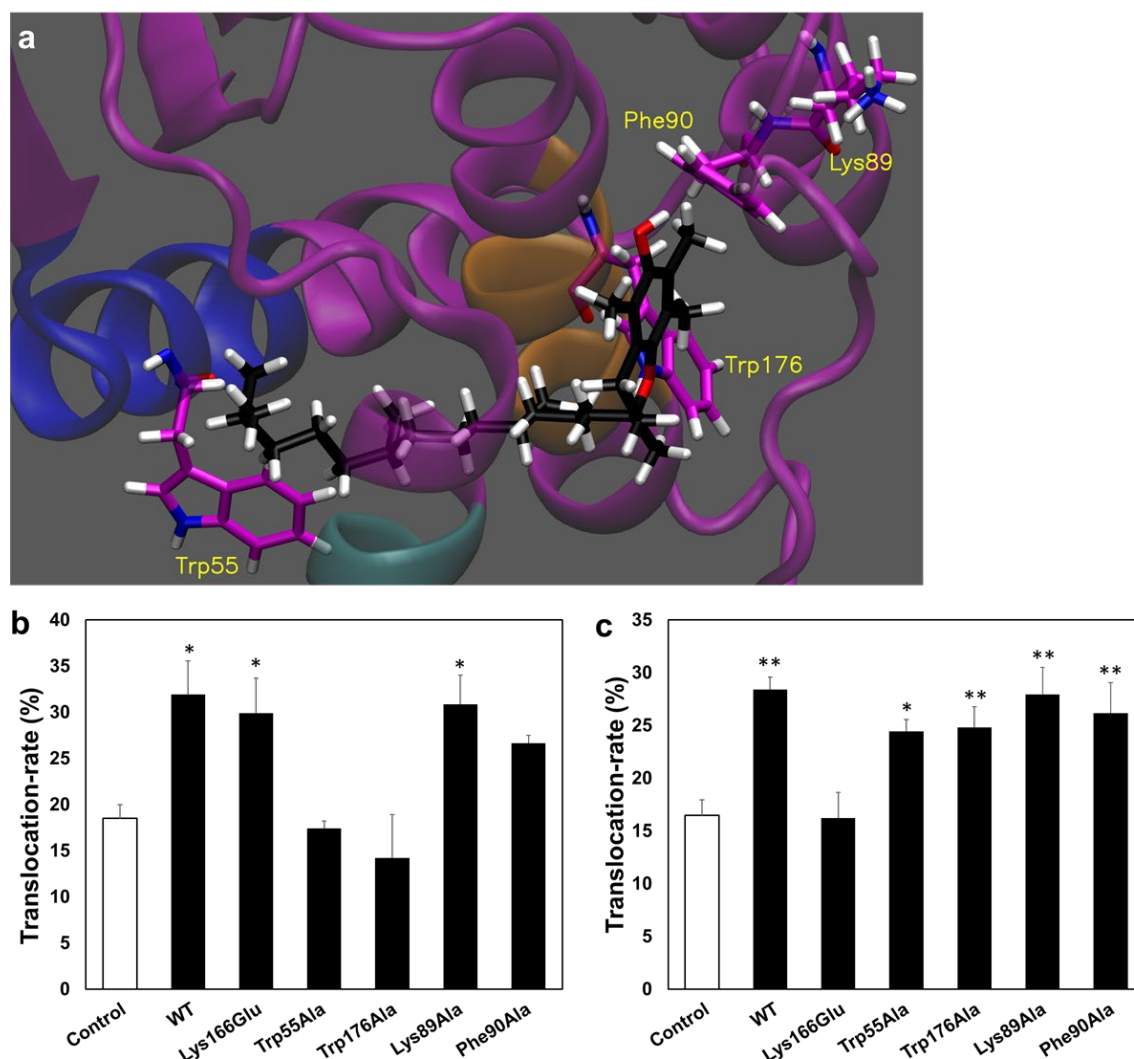


Fig. 5. The effect of mutants in the α Toc binding site on the translocation of DGK α . (a) The position of mutated amino acids in the α Toc binding region is shown. The effect of overexpression of the mutants on the 200 μ M (b) α Toc ($N=3$) or (c) EGCG ($N=3$)-induced translocation of GFP-DGK α in DDT1-MF-2 cells. The Control is the overexpressed FLAG vector alone in the cells. The error bars show SE. Asterisks indicate significance comparing to Control (* p < 0.05, ** p < 0.01).

The involvement of 67LR in the anticancer effect of tocopherols

To seek another physiological role of α Toc-67LR binding, we focused on the anticancer effect of tocopherols. It has been reported that tocopherols, especially γ Toc, δ Toc, and tocopherol derivatives, exhibit the anticancer effect [52–56]. However, the mechanisms are not completely understood. On the one hand, it is well known that EGCG exhibits anticancer effects, and 67LR is a target of the effect [21,22]. Since all tocopherols and derivatives induce the DGK α translocation, these compounds can bind to 67LR. Therefore, we ask if the anticancer effect of tocopherols is mediated through 67LR. It is known that the expression of 67LR is upregulated with the progression of human colon cancer [23]. We treated HCT116 human colon cancer cells with various concentrations of EGCG and α Toc and measured cell viability. EGCG significantly suppressed the cell viability at 50 μ M (Fig. 6a), and α Toc showed a significant effect on the viability at greater than 100 μ M, although the effect was weak (Fig. 6b). Then, we tested the effect of 50 μ M EGCG and 100 μ M α Toc on the viability of HCT116 cells with or without siRNA knockdown of 67LR. EGCG and α Toc were treated 24 h after siRNA transfection for 48 h. We confirmed that the expression level of 67LR was downregulated by 48h after siRNA-transfection, although the expression was recovered by 72h (Fig. 6c). The effect of EGCG on HCT116 cells was eliminated by the knockdown of 67LR (Fig. 6d). Similarly, the effect of α Toc on HCT116 was also eliminated by the 67LR knockdown (Fig. 6e). These results revealed that 67LR is involved at least in the observed cell growth inhibition effects of 100 μ M α Toc in addition to the DGK α activation effect. However, the role of 67LR in the anticancer effects of α Toc as well as other tocopherol species requires further investigation.

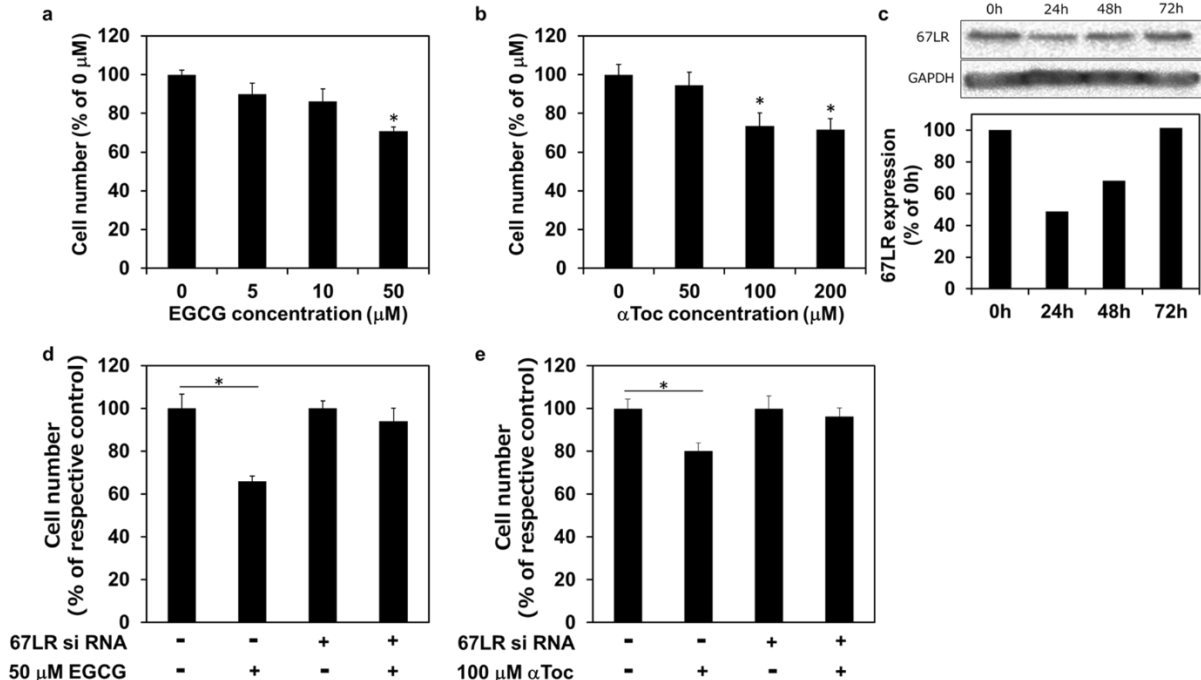


Fig. 6. Effect of the knockdown of 67LR on the anticancer effect of EGCG and α Toc.

The effects of various concentrations of (a) EGCG ($N=3$) or (b) α Toc ($N=3$) on the viability of HCT116 cells. The value is shown as a percentage of that seen with no addition (0 μ M). (c) Changes in the expression of 67LR after siRNA transfection. The effects of the

knockdown of 67LR on the **(d)** 50 μM EGCG ($N=3$) or **(e)** 100 μM αToc ($N=3$) -induced decrease of HCT116 cell viability. The error bars show SE. Asterisks indicate significance comparing to respective control ($*p < 0.05$).

DISCUSSION

It has been of longstanding scientific interest to understand exactly how vitamin E and, in particular, αToc expresses its non-antioxidant functions, such as activation of specific signal transduction. Several studies have shown that αToc interacts with plasma membranes and expresses its function by affecting membrane formation and stability [57,58]. However, the mechanism of just how αToc is involved in various signal transduction processes, especially at a structural level, was not understood. In this study, we have demonstrated that αToc directly binds to 67LR, and we have identified a putative αToc binding site on 67LR. HDX/MS and MD simulations were used to reveal the binding mode of αToc . In addition, we have discovered that 67LR functions as an αToc receptor as well as an EGCG receptor. These results clearly indicate for the first time that 67LR is one of the pathways involved in the function of αToc and other tocopherols. Moreover, we have demonstrated in detail the binding mode of EGCG to its previously proposed binding site on 67LR by employing HDX/MS and MD simulations.

Several downstream signaling pathways of 67LR have been reported [21,59]. However, it is still unclear how 67LR activates its downstream pathways. Our previous study revealed that the Src family tyrosine kinase is involved in the αToc -induced DGK α activation [10]. Recently, we have reported that Src family tyrosine kinase interacts with 67LR and mediates the EGCG-induced DGK α activation as with αToc [25]. Further, Kumazoe *et al.* also reported the implementation of Src in the signaling pathway of 67LR [25]. Therefore, the Src family tyrosine kinase should be one of the downstream pathways of αToc -67LR interaction. However, the binding sites of 67LR for EGCG and αToc are different from one another. Therefore, it is still an open question how two different binding sites trigger the same signaling pathway.

It has been recognized that αToc initiates signal transduction by inhibiting PKC in cells, and the effect was not attributed to its antioxidant effect. However, the detailed mechanisms were enigmatic [5,60]. We have demonstrated herein that αToc -67LR interaction activates DGK α , which can effectively inhibit PKC activity by converting diacylglycerol (DAG), an activator of PKC into phosphatidic acid (PA) [61]. Therefore, activation of the αToc -67LR-DGK α pathway can indirectly lead to PKC inhibition, thereby resulting in a downstream physiological role for αToc .

By employing the ITC approach, we demonstrated the K_d value of αToc to 67LR as 788 ± 174 nM. The K_d value of EGCG has been shown by the surface plasmon resonance (SPR) as 39.9 nM [18]. The affinity of αToc seemed to be relatively lower than that of EGCG. However, since αToc is a vitamin, the basal concentration of αToc in the human plasma is kept at 10 to 30 μM [62]. In contrast, the maximum plasma concentration of EGCG does not exceed 1 μM even after several cups of tea consumption [63]. Regarding the difference in the plasma concentration, the gap in the affinity of these two ligands to 67LR seemed to be reasonable. The affinity of αToc to α -TTP has been reported as 36.1 nM [64]. As described above, α -TTP plays a central role in the absorption of αToc , and the deficiency or impaired

function of α -TTP cause AVED [6–8]. Therefore, it is proper that α -TTP possesses a higher affinity for α Toc than that of 67LR, which is one of the pathways expressing the function of α Toc. Also, it is well known that α Toc is mainly associating with lipoproteins in human plasma and transported into the cell through lipoprotein receptors [65]. Therefore, how monomeric α Toc accesses the cell membrane where 67LR is expressed requires future investigation.

To evaluate the translocation of GFP-DGK α in this study, we used a higher concentration of α Toc to observe a clear translocation. Note that GFP-DGK α translocation was observed at a physiological concentration (10 μ M) of α Toc (data not shown). Indeed, we reported that α Toc ameliorates diabetic nephropathy through DGK α *in vivo* using DGK α knockout mice [11], and that oral consumption of α Toc ameliorated diabetic nephropathy in mice [66]. Also, the inhibition of diabetes-induced PKC activation is observed in mice with the oral administration of α Toc [66]. Therefore, a physiological concentration of α Toc should activate DGK α through 67LR *in vivo*.

It has been reported that the anticancer effect of α Toc is weak or not observed *in vivo* [53,54,67,68]. Indeed, in this study, less than 100 μ M α Toc which is above the physiological concentration, did not significantly affect the HCT116 cell viability. On the one hand, it has been reported that γ Toc and δ Toc not α Toc showed an anticancer effect *in vivo* and suggested that the ability to trap reactive oxygen and nitrogen species of them make a difference in the effect [54]. Since all tocopherols showed a similar effect on the DGK α translocation (Fig. 1f), they might have a similar affinity to 67LR. Accordingly, although there is some contribution, the contribution of 67LR to the anticancer effect of tocopherols seemed not to be dominant. For the physiological role of α Toc-67LR binding other than DGK α activation will be needed further investigations.

This study successfully employed HDX/MS to experimentally determine how EGCG and α Toc bind to separate and distinct binding sites on 67LR, one hydrophobic and one hydrophilic. In the HDX/MS experiments, the degree of inhibition of deuterium exchange by α Toc was less than was observed by EGCG (Fig. 3f, g), though this was on different peptides at different sites. Since the source of deuterium is D₂O in the HDX/MS experiments, water accessibility contributes to the rate and amount of deuterium exchange. Therefore, more water-accessible regions can be more sensitive to the inhibition in the deuterium exchange in the presence of ligands or morphological changes. Indeed, the EGCG binding region is highly water accessible whereas, the α Toc binding region exists in a hydrophobic part of 67LR, which is closed until α Toc accesses the binding site. These differences in physical characteristics might contribute to the difference in the degree of inhibition during deuterium exchange.

It has been reported that 67LR is formed by homodimers of 37LRP, and dimerization is important for membrane localization [16,47,51,69]. However, the actual formation of a dimer and the membrane binding conformation are not entirely understood. We attempted to use the putative formation of a 67LR homodimer suggested by the crystal structure for the MD simulations. However, the dimer structure did not properly fit in the lipid bilayer *in silico*. Therefore, we had to take advantage of the monomer structure of 67LR (37LRP). However, we successfully observed direct binding between monomer 67LR and α Toc and the binding mode between monomer 67LR and EGCG and α Toc in the MD simulations. These facts

suggest that dimer formation is important for activating 67LR downstream or localization at the membrane, but not for binding to α Toc and EGCG.

At the beginning of the binding of α Toc to 67LR, the chroman ring structure faced to Phe203 and Trp176 which are at the entrance of the α Toc binding site, and this step appeared to be critical to guide α Toc to the hydrophobic site. Indeed, overexpressing the Trp176Ala mutant failed to increase the α Toc-induced translocation of DGK α (Fig. 5b), and the importance of the chroman ring structure was consistent with a previous report [10]. Thereafter, the phytyl moiety entered into the binding site. In other words, both the chroman ring structure and the phytyl moiety are essential for α Toc binding to 67LR, and the position of the methyl moiety on the chroman ring structure seems to not contribute to the binding. Indeed, docking simulation using the other tocopherols revealed that the binding poses and binding scores of all tocopherols are similar to each other (Fig. S11). In other words, all tocopherols could bind to 67LR with a similar affinity regardless of the position and number of the methyl groups on the chroman ring. This result was consistent with the result that the other tocopherols induced translocation of DGK α as well as α Toc (Fig. 1e, f) and suggesting that 67LR could well be the receptor for all tocopherols.

Amino acid residues 161 to 180 are well known as a laminin-binding domain named “peptide G,” and the reported EGCG binding motif was contained within peptide G [50,70]. The binding mode revealed that Lys166 and Asn165 in the previously predicted EGCG binding motif and Asp151 that was not known as a binding residue, are critical for binding with galloyl moiety of EGCG. It is well known that 67LR binds only to catechins with a galloyl moiety [18], and these residues may contribute to the selectivity of 67LR for catechins. In addition to these interactions, the MD simulation showed that the hydroxy group on the B ring of EGCG firmly and frequently interacted with Asp126 which has not been reported as a binding region. Also, Asp151 formed a hydrogen bond with the B ring in the second simulation (Fig. S10d). It is reported that EGCG has the best affinity to 67LR among catechins, and catechins which have three hydroxy groups on the B ring have a higher affinity than catechins which have two hydroxy groups on the B ring [71]. Thus, the variation in the affinity may occur because of the differences in the number of hydroxy groups on the B ring, which can interact with these residues. Indeed, all three hydroxy groups on the B ring were occupied with hydrogen bonds with Asp126 and Asp151 in the binding mode from the second simulation (Fig S10d). In other words, Asp126 and Asp151 contribute to the strength of the 67LR affinity to the catechins based on the number of hydroxy groups on the B ring.

As described above, it is well recognized that EGCG functions as an anticancer agent through 67LR, and EGCG-67LR binding is expected as an attractive target for treating various cancer [19–23]. However, the low stability and absorption of EGCG in the human plasma has been a critical issue for the usage of EGCG as an anticancer agent. In the present study, we showed the detailed binding mode of EGCG for the first time, and the binding mode was reasonable and consistent with previous reports. Therefore, now it should be possible to produce optimal agonists of 67LR based on our binding model using structure-based drug design approaches.

CONCLUSION

This study clearly shows for the first time that α Toc directly binds to the hydrophobic binding site of 67LR, which is different from the hydrophilic binding site for EGCG, and that 67LR mediates DGK α activation (Fig. 7). Also, it is suggested that 67LR contributes to the

anticancer effect of α Toc at least in vitro at a high concentration of α Toc (Fig. 7). This explains one of the pathways for how α Toc expresses non-oxidative functions through the cell surface receptor, and provides important insights that should help establish the complete picture of the functioning of vitamin E in metabolism.

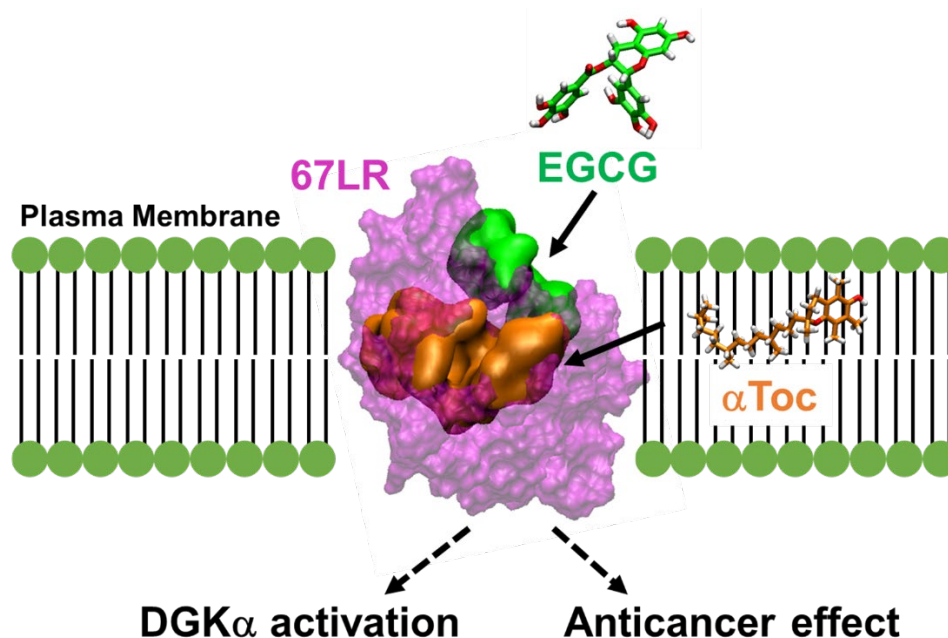


Fig. 7 Model for the EGCG and α Toc binding sites on 67LR and its functions. The transparent purple surface shows the entire 67LR. The green surface shows a hydrophilic EGCG binding site facing the outer environment. The orange surface shows a hydrophobic α Toc binding site embedded in the membrane.

SUPPORTING INFORMATION

The Supporting Tables, Figures, and Movies are available online.

AUTHOR CONTRIBUTIONS

D. H. performed most of the experiments and analyzed most of the data. V. D. M. helped with HDX/MS and performed MD simulations. S. O. performed a mutant experiment. L. W. analyzed α Toc by HPLC. S. L. analyzed the HDX/MS samples. H. T. and H. A. provided information about 67LR and α -TTP. S. U., M. Y. and H. A. advised on the experiments. E. A. D. advised on the HDX/MS and MD simulations. D. H., V. D. M., E. A. D., and Y. S. wrote the manuscript. D. H., V. D. M., and Y. S. conceived the project. Y. S. supervised the research.

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