Anandamide-ceramide interactions in a membrane environment: Molecular dynamic simulations data

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Anandamide is a lipid neurotransmitter that interacts with various plasma membrane lipids. The data here consists of molecular dynamics simulations of anandamide, C18-ceramide and cholesterol performed in vacuo and within a hydrated palmitoyl-oleoyl-phosphatidylcholine (POPC)/cholesterol membrane. Several models of anandamide/cholesterol and anandamide/ceramide complexes are presented. The energy of interaction and the nature of the intermolecular forces involved in each of these complexes are detailed. The impact of water molecules hydrating the POPC/cholesterol membrane for the stability of the anandamide/cholesterol and anandamide/ceramide complexes is also analyzed. From a total number of 1920 water molecules stochastically merged with the lipid matrix, 48 were eventually redistributed around the polar head groups of the anandamide/ceramide complex, whereas only 15 reached with the anandamide/cholesterol complex. The interpretation of this dataset is presented in the accompanying article “Ceramide binding to anandamide increases its half-life and potentiates its cytotoxicity in human neuroblastoma cells” [1].

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Anandamide (AEA) is a lipid neurotransmitter derived from arachidonic acid that inserts in the plasma membrane of target cells prior to exerting its biological effects [2]. Therefore, the way AEA interacts with membrane lipids is of critical importance to understand its mechanism of action at the molecular level. Previous data showed that AEA interacts with cholesterol and uses this lipid as an efficient transport shuttle to cross the plasma membrane [3]. The dataset presented here contains a series of snapshots of molecular dynamics studies of AEA interacting with either cholesterol or ceramide in a palmitoyl-oleoyl-phosphatidylcholine membrane containing dispersed cholesterol molecules (typical Ld phase [4,5]) and hydrated by several layers of water molecules.

2. Experimental design, materials and methods

2.1. Molecular modeling simulations

Geometry optimization of each cholesterol/AEA and C18-ceramide/AEA complexes was first achieved with Polak-Ribiere algorithm [6]. The complexes were then merged within a phosphatidylcholine (POPC) matrix containing cholesterol molecules (molecular ratio POPC:cholesterol 10:1) as previously described [7]. Molecular dynamics simulations with the CHARMM force field were performed in vacuo or in presence of several layers of water molecules with the Hyperchem program (ChemCAD, Obernay, France) [8,9]. A step-by-step description of these molecular modeling
approaches has been published by Di Scala and Fantini [9]. Further interesting insight in the modeling approach for membrane lipids can be found in the literature [10–12].

2.2. Molecular modeling of the AEA/C18-ceramide and AEA/cholesterol complexes in vacuo

Molecular modeling simulations have been first carried out in vacuo to assess if a stable AEA/ceramide complex could be formed within a POPC/cholesterol membrane (starting conditions consisting of 120 POPC molecules stochastically merged with 12 cholesterol, for a molar ratio of 10 POPC per cholesterol). The respective conformations of AEA and ceramide in the membrane environment are consistent with the formation of two types of binary AEA/ceramide complexes (models I and II) which are not mutually exclusive and can even be associated in a ternary complex (model III) constituted by one AEA molecule interacting with two ceramide molecules (Fig. 1). The AEA/ceramide complex (model I) is stabilized by numerous van der Waals interactions and a hydrogen bond involving the carbonyl group of ceramide and the NH group of AEA at a distance of 2.3 Å. The
alternative AEA/ceramide complex (model II) is also stabilized by numerous van der Waals interactions and a hydrogen bond involving the carbonyl group of AEA and the NH group of ceramide at a distance of 2.9 Å. Overall, the energy of interaction of these AEA/ceramide complexes was estimated to -43.0 kJ mol\(^{-1}\) (model I) and -47.7 kJ mol\(^{-1}\) (model II). In the ternary complex (model III), the two hydrogen bounds described in models I and II coexist, yet with interatomic distances slightly modified (2.0 Å and 2.8 Å). The energy of interaction of the ternary complex was -84.6 kJ mol\(^{-1}\), a value close to the sum of the energy of interaction of each binary complex (-90.7 kJ mol\(^{-1}\)). For comparison, the energy of interaction of the AEA/cholesterol complex (model IV) was significantly lower (-30.3 kJ mol\(^{-1}\)) [3]. The conformation of cholesterol in this complex is consistent with previously characterized cholesterol conformers in membrane environment [9].

2.3. Molecular modeling of the AEA/C18-ceramide and AEA/cholesterol complexes within a hydrated membrane

In a second series of experiments, we have performed similar rounds of molecular dynamics simulations in presence of several layers of water molecules (initial conditions with 1920 molecules of water randomly distributed above the polar head groups of the lipid matrix). Two snapshots of these simulations are presented in Fig. 2A (after 2 ns) and 2B (after 10 ns). The presence of water molecules hydrating the lipid membrane reinforced the association of AEA with both cholesterol and ceramide.
Indeed, the AEA/ceramide complex was hydrated with 48 molecules of water which appeared to be homogeneously distributed around the polar head groups of both lipids (Fig. 2B). In contrast, the hydration shell of the AEA/cholesterol complex involved only 15 water molecules, all interacting with the terminal OH group of AEA. The difference of the hydration level of both complexes is especially visible in Fig. 2B with the concerned water molecules in tube rendering. Apart from these selective hydration properties, the AEA/ceramide and AEA/cholesterol complexes that were simulated within a fully hydrated POPC/cholesterol membrane displayed energy interaction values similar to those obtained in vacuo.

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