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Structure and interaction with lipid membrane models of Semliki Forest virus fusion peptide

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A R T I C L E   I N F O

Article history:
Received 22 March 2016
Received in revised form 21 June 2016
Accepted 12 July 2016
Available online 15 July 2016

Keywords:
SFV fusion peptide
Lipid-peptide interaction
Secondary structure
Viral fusion mechanism
IR spectroscopy
Ellipsometry

A B S T R A C T

Semliki Forest virus (SFV) is a well-characterized alphavirus that infects cells via endocytosis and an acid-triggered fusion step using class II fusion proteins. Membrane fusion is mediated by the viral spike protein, a heterotrimer of two transmembrane subunits, E1 and E2, and a peripheral protein, E3. Sequence analysis of the E1 ectodomain of a number of alphaviruses demonstrated the presence of a highly conserved hydrophobic domain on the E1 ectodomain. This sequence was proposed to be the fusion peptide of SFV and is believed to be the domain of E1 that interacts with the target membrane and triggers fusion. Here, we investigate the structure and the interaction with lipid membrane models of the \textit{SFV21} sequence from SFV, using optical method (ellipsometry) and vibrational spectroscopy approaches (Polarization Modulation Infra-Red Reflection Absorption Spectroscopy, PMIRRAS, and polarized ATR-FTIR). We demonstrate a structural flexibility of \textit{SFV21} sequence whether the lateral pressure and the lipid environment. In a lipid environment that mimics eukaryotic cell membranes, a conformational transition from an \(\alpha\)-helix to a \(\beta\)-sheet is induced in the presence of lipid by increasing the peptide to lipid ratio, which leads to important perturbations in the membrane organisation.

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1. Introduction

Enveloped viruses must carry out a membrane fusion reaction between the virus and cellular membranes in order to infect the host cell. This fusion event is driven by viral fusion proteins present at the extracellular surface of the virions. Viral membrane fusion proteins have been grouped into three classes based on shared structural features. The class I fusion proteins are characterized by a post-translational cleavage which produces an amino-terminal fusion peptide as observed in orthomyxoviruses, paramyxoviruses, filoviruses, coronaviruses and retroviruses. The fusogenic structural transition leads to a trimeric assembly of \(\alpha\)-helical coiled coil hairpins in the post fusion state. The class II contains fusion proteins of alphaviruses and flaviviruses. Conversely to class I proteins, they are not cleaved after synthesis and the fusion peptide is internal. They are found to be trimers of hairpins composed of \(\beta\)-sheets in the post fusion state. The class III contains fusion proteins of vesicular stomatitis virus (VSV), herpes simplex virus, and baculovirus, and shares structural features of both class I and class II (for review see references [1–5]).

These fusion proteins play a key role in the fusion process undergoing a large conformational change [1,2,4,6–8]. These conformational changes are triggered either by the interaction of the proteins with specific cell receptors or after cellular internalization, through the receptor mediated uptake by the endocytic pathway, when the acidic condition of the endosome activates the fusion protein [9,10]. However, while their structural features differ, all virus fusion proteins whose post-fusion structures have been defined mediate fusion through a common mechanism. Conformational changes allow the exposure of a region of the virus fusion protein that inserts directly into the viral and/or cell membranes to lead to their destabilization. These regions are named “fusion peptides” (FP) and are generally \(N\)-terminal or internal regions of 16 to 36 residues that are relatively hydrophobic in nature and are highly conserved within virus families [6].

Due to their important role in fusion process, numerous studies were carried out on FP to investigate membrane-associated structures and processes functioning in viral fusion [11]. If class I FPs were intensively studied (HIV [12–18], Influenza [19–22], SARS [23,24], Ebola [25–27], Paramyxovirus [28,29]), it is noteworthy that similar studies of the FPs from class II fusion proteins are much more limited. One can mention a recent structural study on the FP of the Dengue virus which belongs to the flavivirus family [30] and one about the putative FP of Chikungunya virus which belongs to the alphavirus family [31], both performed by NMR in DPC micelles.

Abbreviations: DMPC, dimirystoyl-phosphatidylcholine; DMPG, dimirystoyl-phosphatidylglycerol; DMPE, dimirystoyl-phosphatidylethanolamine; PS, phosphatidylserine; PMIRRAS, polarization modulation infra-red reflection absorption spectroscopy; ATR, attenuated total reflexion spectroscopy; BAM, Brewster angle microscopy; RI, lipid to peptide ratio.

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http://dx.doi.org/10.1016/j.bbamem.2016.07.003
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In the present study, we will focus on FP of the Semliki Forest virus. Semliki Forest virus (SFV) is a well-characterized alphavirus that infects cells via endocytosis and an acid-triggered fusion step using class II fusion proteins and whose fusion is characterized by a striking requirement of cholesterol in the target membrane (for review, see reference [3]). Membrane fusion is then mediated by the viral spike protein, a heterotrimer of two transmembrane subunits, E1 and E2, and a peripheral protein, E3. The pre- and post-fusion structures of the SFV E1 protein have been determined [32–34]. Sequence analysis of the E1 ectodomain of a number of alphaviruses demonstrated the presence of a highly conserved hydrophobic domain on the E1 ectodomain that lies ~80 residues from the N terminus, between amino acids 79 and 97. This sequence was originally proposed to be the fusion peptide of SFV and is believed to be the domain of E1 that interacts with the target membrane and triggers fusion [35–38]. Here, we propose to characterize the structure and the interaction with lipid membrane models of 76YQCKVYGVPFMWGAYCFC sequence from SFV, named SFV21, using an optical method (ellipsometry) and vibrational spectroscopy (PMIRRAS and polarized ATR-FTIR).

2. Materials and methods

2.1. Lipids and peptide

Lipids (dimyristoyl-phosphatidylcholine (DMPC), dimyristoyl-phosphatidylglycerol (DMPG), dimyristoyl-phosphatidylethanolamine (DMPE) and cholesterol) were purchased from Sigma. SFV21 peptide: YQCKVYGVPFMWGAYCFC was purchased from GenScript (Piscataway, USA). SFV21 was solubilized in hexafluoroisopropanol (HFIP) from Sigma at micromolar concentrations. Peptide concentrations were calculated from the absorbance at 280 nm using a molar extinction coefficient \( \varepsilon \) of 11,200 M\(^{-1}\) cm\(^{-1}\). The molar extinction coefficient of the peptide has been determined using the formula: 

\[
\varepsilon_{280nm} = 5600 \times n_W + 1400 \times n_Y
\]

which takes into account the individual extinction coefficient of the amino acids W and Y. A Raman spectrum of a dried film of SFV21 after incubation in buffer (Supplementary data S1) was performed to insure that the three cysteines in the sequence remain non oxidized in the experimental conditions used (no band characteristic of S-S bond around 520–540 cm\(^{-1}\), only a band characteristic of S-H bond in the 2500–2600 cm\(^{-1}\) region).

2.2. Film formation and surface pressure measurements

Monolayer experiments were performed on a computer-controlled Langmuir film balance (Nima Technology, Coventry, England). The rectangular trough and the barrier were made of Teflon. The surface pressure (\( \pi \)) was measured by the Wilhelmy method using a filter paper plate. The trough was filled with a saline solution (150 mM NaCl, pH 5.6) using ultra pure water (MilliQ, Millipore). The experiments were carried out at 22 ± 2 °C. Pure peptide films were obtained by spreading of few μl of HFIP peptide stock solutions at the air/water interface. The mixed lipids/peptides at the desired lipid to peptide ratio, Ri were obtained by co-spreading of the lipid/peptide water interface. The mixed lipids/peptides at the de-spread (no band characteristic of S-S bond around 520 cm\(^{-1}\)) only, and the sequence remain non oxydized in the experimental conditions used for pure lipid SUVs, lipids were first dissolved in chloroform and mixed at the desired ratio. The solvent was then evaporated giving a lipid film. SUV were then prepared by tip sonication after direct hydration of the lipid film with a solution of D\(_2\)O containing 150 mM NaCl. For the preparation of the SUVs containing the mixed fusion peptide/lipids, the fusion peptide powder was first dissolved in HFIP then mixed with the lipids in chloroform at the desired ratio. Solvents were evaporated before hydration by D\(_2\)O and tip sonication for SUV formation. SUV were then burst on a germanium ATR crystal to form a single bilayer whose deposition is controlled by the measurement of the absolute IR intensity [12]. The experiments were carried out at 22 ± 2 °C.

2.3. PMIRRAS spectroscopy

PMIRRAS spectra were recorded on a Nicolet Nexus 870 spectrometer equipped with a photovoltaic HgCdTe detector cooled at 77 K. Spectra were obtained by performing 800 scans at a resolution of 8 cm\(^{-1}\). Briefly, PMIRRAS combines FT-IR reflection spectroscopy with fast polarization modulation of the incident beam between parallel (p) and perpendicular (s) polarizations. The differential reflectivity spectrum is obtained using a two-channel processing signal. To remove the contribution of liquid water absorption and the dependence on Bessel functions, the monolayer spectra are divided by that of the subphase. With an incidence angle of 75°, transition moments preferentially oriented in the plane of the interface give intense and upward oriented bands, while perpendicular ones give weaker and downward oriented bands [39].

2.4. ATR spectroscopy of SFV21 in bulk solid state and inserted in a lipid bilayer

For bulk solid spectra, a film of SFV21 was obtained by evaporation of the peptide solution in HFIP on a germanium ATR crystal. For experiments in lipid bilayer, small unilamellar vesicles (SUV) composed of DMPC:DMPG:Cholesterol:DMPE (6:2:5:3) were prepared alone or mixed with the fusion peptide at a defined molar ratio, Ri. For pure lipid SUVs, lipids were first dissolved in chloroform and mixed at the desired ratio. The solvent was then evaporated giving a lipid film. SUV were then prepared by tip sonication after direct hydration of the lipid film with a solution of D\(_2\)O containing 150 mM NaCl. For the preparation of the SUVs containing the mixed fusion peptide/lipids, the fusion peptide powder was first dissolved in HFIP then mixed with the lipids in chloroform at the desired ratio. Solvents were evaporated before hydration by D\(_2\)O and tip sonication for SUV formation. SUV were then burst on a germanium ATR crystal to form a single bilayer whose deposition is controlled by the measurement of the absolute IR intensity [12]. The experiments were carried out at 22 ± 2 °C.

ATR spectra were recorded on a Nicolet 6700 spectrometer Thermo Scientific equipped with a MCT detector cooled at 77 K. Since ATR spectroscopy is sensitive to the orientation of the structures [40–42], spectra were recorded with parallel (p) and perpendicular (s) polarizations of the incident light with respect to the ATR plate. 400 scans were recorded at a resolution of 8 cm\(^{-1}\). All the orientation information is then contained in the dichroic ratio \( R_{\pi} = \frac{A_p}{A_s} \), where \( A_p \) respectively represents the absorbance of the considered band at p or s polarization of the incident light.

The determination of the position of the band maximum was obtained by second-derivative method and the decomposition of the amide I band in single components characteristic of each secondary structure has been calculated using Omnic or GRAMS V.5 software (Galactic Software).

2.5. Ellipsometric measurements

The morphology of pure peptides or mixed lipids with the peptides were observed at the air/water interface using an ellipsometer iElli2000 microscope (NFT, Göttingen, Germany) mounted on a Langmuir trough. The microscope was equipped with doubled frequency Nd:Yag laser (532 nm, 50 mW), a polarizer, a compensator, an analyzer and a CCD camera. The spatial resolution of the pictures with the 10x magnification lens was about 2 μm and the image size 600 × 450 μm. For ellipsometric measurements, the imaging ellipsometer works at an incidence angle close to the Brewster angle (54.6°). It operates using the principle of a classical null ellipsometer [43].

The ellipsometric angles (\( \Delta, \Psi \)) are determined from polarizer and analyzer azimuths at extinction; \( \Delta \) and \( \Psi \) are related to the optical properties of the sample. In ultra-thin film conditions, \( \Delta \) is proportional to the film thickness. The comparison of the experimental data with a computerized optical model included in the ellipsometer software allowed to provide an estimation of the film thickness when a value of refractive index can be obtained. Since it is difficult to determine an experimental refraction index value, we choose 1.50 for the SFV21 peptides as it was used in a previous work concerning the fusion peptide FP23 of HIV and EBO17 and EBO24 of Ebola [12,27]. Indeed, in the literature, variations between 1.45 [44] and 1.53 [45,46] can be encountered.
for protein and peptide depending on the amount of water taken into account. Concerning lipids, anisotropic refractive indices between 1.44 [47] and 1.50 [48] were reported for DMPC in the literature. Therefore, we use an intermediate value of 1.46 to perform thickness estimations [12,27].

3. Results

3.1.1. Structure of SFV21 in the bulk solid state by ATR spectroscopy

ATR FT-IR absorption spectrum of SFV21 in the bulk solid state displays an intense asymmetric amide I band centered around 1625 cm⁻¹ (Fig. 1). The amide I region was analyzed and the secondary structure content was evaluated from the analysis of the amide I band shape and curve fitting [40]. The decomposition of the spectrum shows that the main band around 1625 cm⁻¹ is well fitted by two components at 1628 and 1615 cm⁻¹. Associated with another weak contribution around 1694 cm⁻¹, these three bands are characteristic of antiparallel β-sheets which contribute for 49.0% of the secondary structure. Another contribution at 1656 cm⁻¹ can be attributed to α-helix and represents 31.5% of the structure. Two other minor contributions are observed at 1644 cm⁻¹ and 1677 cm⁻¹, they are assigned respectively to random structure and β-turn and contribute for 6.3% and 13.2% respectively [40,41,49,50]. The amide II mode absorbs around 1520 cm⁻¹.

3.2. Morphological and structural study of the SFV21 at the air/water interface

The Brewster angle microscope (BAM) images of the pure SFV21 layer at increasing surface pressures from 2 to 20 mN/m (Fig. 2) displays a progressive increase of the averaged normalized grey level until 20 mN/m. Every picture shows a stable homogeneous surface of the peptide film appearing brighter during compression. The thickness of the layer was estimated by ellipsometry using a constant refractive index of 1.50 for the peptide (cf Materials and methods). As the peptide film is compressed, estimated thickness increases from 21 ± 1 Å (Δ = 347.40°, Ψ = 2.39°) at 2 mN/m to 24 ± 1 Å (Δ = 345.23°, Ψ = 2.40°) at 12 mN/m and 26 ± 1 Å (Δ = 343.29°, Ψ = 2.40°) at 20 mN/m.

The PMIRRAS spectra at the air/water interface of the SFV21 peptide film during compression from 1 to 20 mN/m displayed well resolved amide I bands between 1600 and 1700 cm⁻¹ and amide II bands centered around 1530 cm⁻¹ (Fig. 3). Whatever the surface pressure from 2 to 20 mN/m, the main amide I band is centered around 1626 cm⁻¹ with another contribution around 1695 cm⁻¹ reflecting a main content of antiparallel β-sheets. Shoulders and secondary derivative (not shown) display also contributions around 1645 cm⁻¹, 1655 cm⁻¹ and 1670 cm⁻¹ respectively characteristic of random, α-helical and β-turn structures [41,49,50]. During compression of the peptide film (up to 20 mN/m), since their contributions remain positive, one can assume that antiparallel β-sheets are mainly oriented flat on the air/water interface plane [51,52] whatever the lateral pressure. Simultaneously, the Amide I/Amide II ratio decreases from 1.9 to 1.4 during the compression from 2 to 12 mN/m, and remains constant around 1.4 at 20 mN/m. Due to the specificity of the PMIRRAS selection rules, this change in the amide I/amide II ratio could be attributed both to a change in the orientation of one or more of the minor structural components or to a concomitant change in their proportion.

This experiment suggests that SFV21 remains mainly folded into antiparallel β-sheets laying flat at the air/water interface during compression and decomposition, and that the contribution of other minor structural components can slightly change both in proportion and orientation versus lateral pressure changes.

3.3. Morphological and structural study of SFV21 inserted into a lipid monolayer

To investigate the changes in morphology induced by SFV21 upon interaction with the DMPC:DMPC:cholesterol:DMPE monolayer at the molar ratio 6:2:5:3, we first studied the morphology of the lipid alone in the surface pressure range from 5 to 30 mN/m. Such a complex lipid composition was used to reflect the variety of lipids present in eukaryotic cell membranes [53]. PC and PE polar heads and cholesterol are main components in this type of membrane. We replaced the negatively charged PS polar headgroup generally encountered in natural membranes by PG in our model since the asymmetric stretching band of the PS carboxylate group overlaps with the amide I band of proteins and thus interferes with IR measurements. We choose myristoyl fatty acid chains for their stability at the air water interface and to avoid oxidation processes like encountered with C18:1 like DOPC at the interface. This composition was also chosen to allow comparison with the FP23 fusion peptide of HIV and the Ebola fusion peptide previously studied in these conditions with the same techniques [12,27].

BAM pictures (Fig. 4, first row micrographs) show that the surface is homogeneous whatever the surface pressure during compression and decomposition. Such an observation is surprising since the monolayer contains DMPE and cholesterol which display high phase transition temperatures (>-49 °C) compared to DMPC and DMPC (23 °C), then one can expect to observe segregation of lipids in the monolayer. Taking into account the large content of cholesterol in the monolayer, the observation of such a homogeneous film can probably be explained by attractive interactions between the different phospholipids and cholesterol as previously reported [54]. Ellipsometric measurements and thickness estimations using a refractive index of 1.46 for the lipids (cf Materials and methods) leads to 16 ± 1 Å (Δ = 350.93°, Ψ = 2.39°) for the thickness of the lipid monolayer at 5 mN/m, 17 ± 1 Å (Δ = 350.75°, Ψ = 2.40°) at 15 mN/m and 18 ± 1 Å (Δ = 349.88°, Ψ = 2.39°) at 30 mN/m.

BAM pictures of SFV21 inserted into a lipid monolayer are recorded after co-deposition at the air/water interface of a peptide/lipid (DMPC:DMPC:Cholesterol:DMPE 6:2:5:3) mixture at a defined lipid to peptide ratio, Ri = 75 or Ri = 20. When SFV21 is inserted into the
lipid monolayer at low surface pressure (5 mN/m), whatever the Ri, an homogeneous surface is observed with gray levels and ellipsometric measurements comparable to what is observed for lipids alone. For higher lateral pressures, important changes in the morphology of the layer are observed compared to the lipid alone (Fig. 4). Whatever the pressure and the lipid/peptide ratio Ri (75 or 20) heterogeneous surfaces are observed which are characterized by a brighter network/phase in a surrounding less dense phase (dark background). The brightness of the denser phase increases with lateral pressure from 15 to 30 mN/m but is comparable whatever the SFV21 content (Ri from 75 to 20). Estimations performed using ellipsometric measurements lead to thicknesses around 19 ± 1 Å at Π = 15 mN/m and 30 ± 2 Å at Π = 30 mN/m for both Ri (75 and 20). For the less dense phase, whatever the Ri (75 or 20) and the surface pressure (15 to 30 mN/m), thicknesses are highly comparable to what was obtained for the lipid alone (17 ± 1 Å at 15 mN/m and 19 ± 1 Å at 30 mN/m). The decomposition down to 15 mN/m shows a reversible behaviour both in morphology and thicknesses and further decomposition down to 5 mN/m (not shown) leads back to a homogenization of the surface.

PMIRRAS spectra of SFV21 inserted into a DMPC:DMPG:cholesterol:DMPE (6:2:5:3) lipid monolayer at Ri = 20 and 75 were recorded as previously described. Since the results obtained for both Ri are very comparable, only the spectra at Ri = 20 are presented in Fig. 5. The mixed monolayer was compressed from 5 to 40 mN/m then decompressed down to 5 mN/m. PMIRRAS spectra were recorded at different surface pressures (Fig. 5A). Every spectrum shows a band centered around 1732 cm⁻¹ characteristics of the ν(C=O) ester stretching vibration [55]. The ν(ν(CH₂)) bending vibration of acyl chains around 1465 cm⁻¹ can also be observed. Whatever the lateral pressure, in the amide I region, the main contribution around 1627 cm⁻¹ is characteristic of β-sheets. A shoulder around 1650 cm⁻¹, characteristic of α-helix, is present whatever the lateral pressure and increases with compression. The weak amide II contribution whatever the spectrum shows that the structures are mainly laying flat at the interface and that the orientation does not significantly changes with lateral pressure increase [56]. To have a better insight on the secondary structure, the difference spectrum between lipids and absence and presence of SFV21 at comparable lateral pressure was calculated (Fig. 5B). For this, separate PMIRRAS measurements with monolayers of the pure lipids were registered at the same surface pressure and we assume that these spectra can be used for the subtraction. The data clearly show that the two main secondary structure contributions are due to the 1650 cm⁻¹ and 1627 cm⁻¹ bands, and that their respective proportions vary depending on the applied lateral pressure. One can observe that the position of the alpha helical and beta sheet components of the amide I band in Fig. 5A remains unchanged in all spectra whereas some differences in position and shape of some of the amide I bands appear in Fig. 5B. This can mainly be explained by the fact that this region of the spectrum (broad negative dip around 1650 cm⁻¹ associated with the deformation vibrational mode of liquid water [57]) is sensitive to the interface coverage, then small differences in the pure lipid film organisation compared to the mixed SFV21/lipid film can lead to small distortions. To minimize these effects and assuming that the main contributions to the secondary structures are due to these bands and that the orientation does not change significantly, the area of the band was estimated and the respective proportion of both secondary structures calculated using three independent experiments. The results are summarized in Table 1 for Ri = 20 and Ri = 75. The values show that whatever the Ri, at low lateral pressure, the β-sheet structure is largely predominant. The β-sheet proportion decreases with lateral pressure increase while the α-helical proportion increases. The α-helical proportion becomes predominant from 15 mN/m (55%) at Ri = 75, while it slowly increases up to 48% for 35mN/m at Ri = 20. These results show a progressive secondary structure transition from β-sheet to α-helix induced by compression of the monolayer, the behaviour is more pronounced for the higher Ri (75), then the lower SFV21 content. This behaviour demonstrates a structural flexibility of the SFV21 sequence and shows that the conformation is both sensitive to lateral pressure and to the presence of lipids. During decompression, the transition is not reversible since the α-helix predominates compared to the β-sheet whatever the Ri.

To estimate the effect of the SFV21 incorporation on the lipid chain organization, the position of the band corresponding to asymmetric stretching modes of the methylene groups, νas(CH₂), was measured.
we observe a slight decrease in the wavenumber of the ν\textsubscript{as}(CH\textsubscript{2}) when the lateral pressure increases from 5 to 35 mM/m, due to the progressive organization of the lipid chains during compression. It is then interesting to notice that at low lateral pressures (< 15 mM/m), there is no noticeable difference in wavenumber of the ν\textsubscript{as}(CH\textsubscript{2}) in absence or presence of SFV21, while at higher pressures (> 25 mM/m), the wavenumber of ν\textsubscript{as}(CH\textsubscript{2}) is higher in presence of SFV21 than for the lipid monolayer alone. This shows a more important disorder in the lipid chain packing in the monolayer due to the SFV21 incorporation. The higher the SFV21 concentration (the lower the Ri), the more important the disorder.

3.4. Study of SFV21 inserted into lipid bilayers

We further investigated the behaviour of SFV21 inserted into a single supported lipid bilayer made of the same lipid mixture DMPC:DMPG:cholesterol:DMPE (6:2:5:3). We hence performed polarized ATR experiments in order to analyze the structure of the fusion peptide in such membrane models as well as perturbations induced by the peptide on the organization of the lipids.

The pure or mixed SF21/lipid bilayers adsorbed on a germanium ATR crystal were obtained by spontaneous fusion of small unilamellar vesicles (SUV) of pure lipid or mixed SF21/lipid at the defined Ri as described in the Materials and methods section. Fig. 6 displays the ATR spectra obtained in the 1800–1500 cm\textsuperscript{−1} region. The band around 1735 cm\textsuperscript{−1} corresponds to the ν(C = O) ester of the lipids while the 1700–1600 cm\textsuperscript{−1} domain corresponds to the amide I. The amide I region was analyzed and the secondary structure content was evaluated from the analysis of the amide I band shape and curve fitting (Supplementary data S2) [39]. The 1685 ± 1 cm\textsuperscript{−1}, 1632 ± 1 cm\textsuperscript{−1} and 1624 ± 1 cm\textsuperscript{−1} bands were assigned to β-sheets, the 1664 cm\textsuperscript{−1} and 1653 ± 1 cm\textsuperscript{−1} bands to α-helices, the 1642 ± 2 cm\textsuperscript{−1} band to random structures and the 1675 ± 2 cm\textsuperscript{−1} band to β-turns. The results, presented in Table 3, show that the predominant structures are the α-helices and the β-sheets. There is a progressive interconversion from the predominant α-helices (41.0% at Ri = 150) to the β-sheets (37.2% at Ri = 20) with peptide concentration. The other structures, β-turns and random, counts respectively for around 10% and 20% and their proportion remains constant whatever the Ri. Using the p- and s-polarized spectra, we then estimated the dichroic ratio of the amide I band of the α-helix (1664/1653 cm\textsuperscript{−1}) and of the β-sheet (1632/1624 cm\textsuperscript{−1}) presented in Table 3. The evolution of R\textsubscript{ATR} (amide I) for the α-helix from 1.44 (Ri = 150) up to values between 1.8 and 2.0 demonstrates a progressive reorientation of the α-helix tilt angle from around 70° to 0°–20° compared to the interface normal [12]. This means that when diluted in the lipids, the helix is mainly laying flat on the membrane while its becomes perpendicular to the plane of the bilayer at higher concentration. Concerning the β-sheet, the weak variation of R\textsubscript{ATR} (amide I) allows to estimate a tilt angle around 35° relative to the membrane plane, this orientation does not significantly depends on the peptide concentration [12].
To investigate the consequence of SFV21 bilayer incorporation on the organization of the lipid acyl chains in the bilayer, the shift of the bands corresponding to asymmetric and symmetric stretching modes of the methylene groups, νs(CH2) and νas(CH2), respectively, were measured (Table 4). Indeed, the wavenumbers of these bands are known to be sensitive to changes in the configuration of the acyl chains, in chain mobility and packing [53–56]. The νs(CH2) and νas(CH2) of 2851.7 and 2920.7 cm⁻¹ respectively for the bilayer alone are characteristic of quite well ordered acyl chains, which is consistent with a bilayer composed of phospholipids with phase transition temperature higher than 23 °C. When SFV21 is added to the bilayer, there is an increase up to 1.8 ± 0.3 cm⁻¹ at Ri = 20 in the wavenumbers of the νs(CH2) and νas(CH2) which indicates a decrease in chain order due to SFV21 peptide incorporation (Table 4). This is confirmed by the evolution of the dichroic ratio (RATR) of the νs(CH2) band of lipid chains which was measured on the polarized ATR spectra (Table 4). Indeed, since the transition moment for the νs(CH2) lies perpendicular to the chain axis, the introduction of order or disorder alters the direction of the main chain with respect to the normal of the ATR crystal. The dichroic ratio is then expected to provide a sensitive and quantitative indicator of acyl chain order by estimating the average C—C—C angle of the lipid carbon chains with the normal to the interface [12,27]. Here, the RATR increases when SFV21 interacts with the lipids, compared to the lipid alone (Table 4). This corresponds to a reorientation of the lipid chains from approximately 45° defined to the membrane normal to 60–65° in presence of SFV21 at Ri = 150 and 20. A RATR value of 1.93 obtained at Ri = 75 is characteristic of an important disorder of the bilayer, close to the isotropy [12]. Conversely, the dichroic ratio of the ν(C = O) ester band of the lipids increases from 1.67 for the lipid alone to around 2.00 in the presence of SFV21 at Ri = 75 and 20 which is characteristic of an important increase of disorder. Altogether, this experiment is showing SFV21 insertion leads to disorder increase in both the acyl chains and the glycerol region of the lipids, but the glycerol region is sensitive at higher Ri compared to the acyl chains.

4. Discussion

4.1. Influence of environmental conditions on SFV21 conformation

The structural studies of SFV21 show a sensitivity of its conformation whether the conditions used. Indeed, for SFV21 in bulk state or alone at the air/water interface whatever the compression state, the main conformation is an antiparallel β-sheet. In presence of lipids (monolayer at air/water interface and bilayer), SFV21 undergoes a conformational change from α-helix to antiparallel β-sheet.

Two parameters play a role in the SFV21 structuration: the lateral pressure and the presence of lipids. At low lateral pressure at the air/water interface (with or without lipid), the main conformation of SFV21 is the antiparallel β-sheet. This structure may be favoured by the large surface available since β-sheets occupy a larger area at the

### Table 1

| Ti (mN/m) | Ri = 20 | Ri = 75 |
|-----------|---------|---------|
|           | 1650 cm⁻¹ (α-helix) | 1627 cm⁻¹ (β-sheet) | 1650 cm⁻¹ (α-helix) | 1627 cm⁻¹ (β-sheet) |
| 5         | 29.5 ± 0.4 | 70.5 ± 0.4 | 29.5 ± 0.4 | 70.5 ± 0.4 |
| 15        | 40.5 ± 0.4 | 59.5 ± 0.4 | 55.5 ± 0.4 | 44.5 ± 0.4 |
| 25        | 41.0 ± 0.4 | 59.0 ± 0.4 | – – | – – |
| 30        | – – | – – | 62.0 ± 0.4 | 38.0 ± 0.4 |
| 35        | 48.0 ± 0.4 | 52.0 ± 0.4 | – – | – – |
| 15 D      | – – | – – | 74.5 ± 0.4 | 25.5 ± 0.4 |
| 10 D      | 45.0 ± 0.4 | 55.0 ± 0.4 | – – | – – |

### Table 2

| Ti (mN/m) | Lipid monolayer | + SFV21 Ri = 75 | + SFV21 Ri = 20 |
|-----------|----------------|----------------|----------------|
| 5         | 2922.9 ± 0.4 | 2921.9 ± 0.4 | 2921.8 ± 0.4 |
| 15        | 2921.2 ± 0.4 | 2921.4 ± 0.4 | 2921.9 ± 0.4 |
| 25        | 2919.5 ± 0.4 | 2921.1 ± 0.4 | 2921.4 ± 0.4 |
| 35        | 2919.5 ± 0.4 | 2920.1 ± 0.4 | 2921.4 ± 0.4 |
interface than α-helices and by the fact that the hydrophobicity of the air/water interface may favour changes in the hydrogen bonding interactions [61,62]. Increasing lateral pressure for pure SFV21 monolayer at the air water interface only leads to weak changes in the proportions and/or orientation of the minor structural components as shown by the ATR spectra of the amide I region. Lipid composition: DMPC:DMPG:cholesterol:DMPE and/or orientation of the minor structural components as shown by the air water interface only leads to weak changes in the proportions Wavenumbers, secondary structure contents and dichroïc ratio for Amide I band for SFV21 Table 3 actions [61,62] air/water interface may favour changes in the hydrogen bonding inter-

Fig. 6. ATR isotropic spectra of the amide I region of SFV21 inserted into a lipid bilayer DMPC:DMPC:cholesterol:DMPE (6:2:5:3) hydrated with D_2O and 150 mM NaCl, pH 5.6.

Table 3

| Secondary structure | Wavenumber (cm\(^{-1}\)) | % (±1.0) | R\_ATR (Amide I) (±0.05) |
|---------------------|------------------------|---------|------------------------|
| \(\text{Ri} = 150\) | \(\beta\)-Sheet | 1624–1633–1685 | 28.7 | 1.55 |
| | Random | 1642 | 19.7 | – |
| | \(\alpha\)-Helix | 1654–1663 | 43.0 | 1.44 |
| | B-Turn | 1677 | 8.6 | – |
| \(\text{Ri} = 75\) | \(\beta\)-Sheet | 1624–1633–1681 | 31.5 | 1.45 |
| | Random | 1643 | 21.2 | – |
| | \(\alpha\)-Helix | 1653–1664 | 37.8 | 1.80 |
| | \(\beta\)-Turn | 1673 | 9.5 | – |
| \(\text{Ri} = 20\) | \(\beta\)-Sheet | 1617–1631–1685 | 37.2 | 1.50 |
| | Random | 1642 | 20.6 | – |
| | \(\alpha\)-Helix | 1653–1664 | 33.1 | 2.05 |
| | \(\beta\)-Turn | 1675 | 9.1 | – |

Table 4

| \(v_\text{as} (\text{CH}_2)\) (cm\(^{-1}\)) | \(v_\text{s} (\text{CH}_2)\) (cm\(^{-1}\)) | \(R_{\text{ATR}}(\nu_\text{as}(\text{C=O}))\) | \(R_{\text{ATR}}(\nu(\text{C=O}))\) |
|------------------------|------------------------|------------------------|------------------------|
| SUV alone | 2920.7 ± 0.4 | 2851.7 ± 0.3 | 1.34 ± 0.05 | 1.67 ± 0.05 |
| \(\text{Ri} = 150\) | 2921.4 ± 0.4 | 2851.5 ± 0.3 | 1.73 ± 0.05 | 1.72 ± 0.05 |
| \(\text{Ri} = 75\) | 2922.2 ± 0.4 | 2852.3 ± 0.3 | 1.85 ± 0.05 | 2.02 ± 0.05 |
| \(\text{Ri} = 20\) | 2922.5 ± 0.4 | 2853.5 ± 0.3 | 1.93 ± 0.05 | 1.97 ± 0.05 |

membrane fusion has been extensively studied, and cholesterol demonstrated to be necessary for very different fusion processes as neuronal exocytosis [63] and viral entry. Indeed several recent studies on viruses as different as Influenza [64], Human or Feline Immunodeficiency Virus [65–68], Herpes Simplex virus [69] demonstrate the potential role of cholesterol or cholesterol-enriched domains (rafts) as possible platforms for the entry of viruses into cells.

It is also interesting to compare our results on SFV21 to these obtained for the putative fusion peptide VT18 of Chikungunya alphavirus using NMR spectroscopy in zwitterionic lipid environments [31]. VT18 is an 18 residue-long peptide which shares 13 residues with SFV21 (VYPFMWGGAYCFC). In the conditions used (DPC micelles at diluted lipid to peptide ratio around 250), authors do not observe significant contribution of α-helices as we observe in lipid bilayers in diluted conditions but rather extended or β-type conformations for most of the residues. Authors compare the NMR derived DPC bound structure of VT18 with that of the conformation deduced in the context of protein complexes (prefusion state) obtained by X-ray crystallography [70]. They show that the mica-bound structure of the fusion peptide, VT18, markedly differs in terms of packing interactions of some aromatic residues compared to the the X-ray structure and conclude that the lipidic environment of DPC micelles led to a significant conformational reorganization of the aromatic residues in the fusion peptide. This result is in agreement with the observation that the conformation of fusion peptides is very sensitive to the lipid environment as well as the experimental conditions used. Concerning VT18, authors also observe that residues MWGG adopt a type I β-turn conformation with the N- and C-termini pointing in an antiparallel orientation. In our case, we show that in lipid bilayer environment, the proportions of random and β-turn structure remain rather constant whatever the Ri (±20%) and ≈9% respectively). It cannot be excluded that this relatively high but conserved proportion of random and β-turn could be due to this part of the sequence. One can also speculate that the presence of such a β-turn will result in a partial folding of SFV21 into intramolecular antiparallel β-sheet for the residues close to the turn, the structural plasticity resulting from the N- and C- terminal part of the sequence and from the 8 residues which are not present in the VT18 sequence.

It is then interesting to see that the conformational flexibility observed for SFV21 fusion peptide of class II is also shared by viral fusion peptide of class I (HIV1, paramyxovirus, Ebola [12–27]) and SNARE proteins involved in intracellular fusion process [71]. These similarities on structural properties shared by peptides or proteins involved in totally different fusion processes may play an important role in the fusion mechanism.

4.2. SFV21 interaction with model membranes

The present study shows that, whatever the lipid membrane model used, monolayer at lateral pressure close to "biological equivalent pressure" of a bilayer around 30–35 mN/m or lipid bilayer, the structural behaviour of SFV21 fusion peptide is comparable (Table 5). A conformational transition from an α-helix to a β-sheet is induced by a change in the peptide to lipid ratio in the membrane. At low concentration in lipids, SFV21 interacts with lipids mainly as α-helices. As the
local peptide concentration increases in the membrane, the proportion of $\alpha$-helix drops off in favor of a mainly antiparallel $\beta$-sheet structure. The formation of antiparallel $\beta$-sheet structure is probably due to peptide self-assembling triggered by close peptide contacts induced by increasing their concentration (Fig. 7). This is confirmed by the aggregates which are observed in presence of SFV21 by BAM and ellipsometry. This concentration dependent effect on peptide conformation might be of biological relevance during the formation of the trimers of hairpins composed of $\beta$-sheets as seen in the post-fusion state of class II fusion proteins. Formation of fusion oligomers was already suggested for class I viruses like HIV to be linked to fusion activity for envelope protein-mediated fusion and for liposome fusion mediated by fusion peptides [72,73]. For instance, in the case of the fusion peptide of the Sendai virus [74], the peptide could self-assemble in its membrane-bound state, thus suggesting its role in assisting the assembly of the envelope protein of the virus.

This structural plasticity with formation of $\beta$-sheets as fusogenic conformation, may then promote membrane topological changes during fusion. Indeed, we observe by ATR spectroscopy that while the proportion of the $\beta$-sheets increases with the SFV21 concentration, the $\beta$-sheets remain tilted compared to the interface plane of the bilayer (35°), which lead to important disorder in the lipid chain organisation as observed by the increase of the $R_{\text{ATR}}$ and of the wavenumber of bands characteristic of the stretching of the chain acyl groups. Conversely, while the proportion of $\alpha$-helix decreases with SFV21 concentration increase, their orientation becomes perpendicular to the plane of the lipid bilayer, which may result in a weaker perturbation of the lipid chain organization but a destabilization of the lipid head groups [12]. It is also interesting to notice that we do not observe this orientational change with the monolayer lipid model. Such a difference may be due to the fact that a lipid monolayer does not match correctly the hydrophobic thickness of a membrane and does not allow a proper orientation of SFV21 $\alpha$-helix which is then too long to match the hydrophobic part of the lipids. Therefore, even if the monolayer system allows the detection of the structural plasticity of SFV21, the lipid bilayer model seems to be more relevant in mimicking the biological membrane and the orientation of SFV21 in such a system will be more representative of the reality.

Such a process of lipid bilayer disorganization should be essential during membrane fusion processes by generating new lipid phases that are thought to be associated with initial steps of membrane fusion by increasing the negative curvature strain, favoring the formation of inverted phases [75,76].

To conclude, the present study shows the structural plasticity of SFV21 fusion peptide and demonstrates its ability to transit from a mainly $\alpha$-helical structure to antiparallel $\beta$-sheets leading to a destabilization at large scale of the lipid membrane. Such a process is probably combined with concerted effects due to various domains of the viral fusion proteins, besides the fusion peptide, which could cooperate in destabilizing the apposing viral and plasma membranes. Our results show then the interest of this type of fundamental study to highlight some specific structural properties (flexibility, plasticity) shared by fusion peptides belonging to different fusion processes (viral fusion, intracellular fusion) which could play a key role in the fusion mechanism.

### Transparency document

The Transparency document associated with this article can be found, in online version.

### Acknowledgements

We acknowledge ANR-09-BLAN-0385-02 for its financial support. We acknowledge Etienne Harté for registration of the Raman spectrum and Dr. Isabel Alves for its careful reading of the manuscript.

### Appendix A. Supplementary data

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.bbamem.2016.07.003.

| $\alpha/(\alpha + \beta)$ % | Lipid monolayer | Lipid bilayer |
|--------------------------|----------------|--------------|
| $R_i = 150$              | –              | 60           |
| $R_i = 75$               | 62             | 58           |
| $R_i = 20$               | 48             | 47           |

Fig. 7. Molecular cartoon of the SFV21 structure and orientation in the lipid mono/bi-layer as a function of the peptide concentration.
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