Bis-Thioether-Containing Lipid Chains in Cationic Amphiphiles: Physicochemical Properties and Applications in Gene Delivery
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To cite this version:

HAL Id: hal-02278044
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Submitted on 18 Nov 2019

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Bis-thioether containing lipid chains in cationic amphiphiles: physicochemical properties and application for gene delivery


Abstract: Cationic amphiphiles featuring two thioether functions in each lipid chain of bicatenar cationic amphiphiles are herein reported for the first time. The physico-chemical properties and transfection abilities of these new amphiphiles were compared with those of already reported analogues featuring either (i) saturated, (ii) unsaturated or (iii) mono-thioether containing lipid chains. The homogeneity of the series of new compounds allowed to clearly underscore the effect of bis-thioether containing lipid chains. This study shows that besides previous strategies based on unsaturation or ramification, the incorporation of two thioether functions per lipid chain constitutes an original complementary alternative to tune the supramolecular properties of amphiphilic compounds. The potential of this strategy was evaluated in the context of gene delivery and report that two cationic amphiphiles (i.e. 4a and 4b) can be proposed as new efficient transfection reagents.

Introduction

Non-polymeric amphiphilic compounds are molecules constituted by a polar head group covalently attached to one (monocatenar) or several (poly-catenar) hydrophobic chains. There exist many classes of amphiphiles including natural derivatives (e.g. phospholipids, sphingolipids, fatty acids)[1] or synthetic compounds (e.g. bio-inspired phospholipids,[2] Bola lipids,[3] gemini[4]) in which amphiphiles were used for a multitude of applications including the extraction and stabilization of proteins,[5] the stabilization of interfaces (e.g. coaservates[6]), the design of nano-reactors,[7] the production of vesicles,[8] the production of carriers of nucleic acids delivery,[9] others were designed for their own pharmaceutical properties (e.g. bactericidal agents,[10] modulation of ion channels activities,[11] anticancer properties[12]). Whatever the application, the physico-chemical properties of the amphiphiles must be finely tune to optimize their potential. To this regards, it must be noticing that the structure of the hydrophobic domain has a dramatic effect on the supramolecular properties of the amphiphilic compounds. As an illustration, the number of lipid chains and their length deeply impact the Critical Micellar Concentration (CMC) and the shape of the amphiphiles can impact the structure of the assemblies (e.g. lamellar versus hexagonal). For nucleic acids delivery, cationic amphiphiles[2,13,14] were evaluated for different purposes (lung transfection,[15,16] tendon healing,[17] cancer therapy[18] anticancer vaccination[19-21]) and the current developments offer news therapeutic perspectives. It must be emphasized that one of the interest of synthetic vectors, when compared to viral vectors, is that they can be produced on large scale and following simple purification processes. To improve the efficacies of cationic amphiphiles for nucleic acids delivery, both the modification of the structure of the polar head group (e.g. nature of the cationic charge[22] polycationic moieties[23,24]) and the structure of the hydrophobic domain were explored.[25] In the last case, it was shown that cationic amphiphiles containing saturated lipid chains produced ineffective pDNA carriers.[26,27] This is likely explained by the fact that the lipoplexes (association of cationic amphiphiles with nucleic acids), generally enter into cells by endocytosis pathway.[28] Subsequently, nucleic acid must escape from endosomes to avoid degradation. This process is likely facilitated by lipid mixing between the cationic amphiphiles and the lipids of the endosomal membrane[29]. Therefore, the introduction of some upset within the lipid chains should produce less tight supramolecular packing favoring better lipid mixing. A first strategy to disturb the packing of the lipid chains consists to use unsaturated lipid chains. For that purpose, oleyl chains (C18:1) that possess one unsaturation with a Z configuration are widely employed.[30] As a second strategy, naturally available branched (e.g. phytanyl chains)[31,32] or synthetically prepared ramified (e.g. through thiol-ene click chemistry) lipid chains can be used.[33] The presence of lateral moieties decreases the strength of the packing[34-36] but can also affect the supramolecular organization (e.g. production of inverted hexagonal phase). Beside these two strategies (unsaturation and ramification) the incorporation of poly-unsaturated lipid chains in the structure of cationic amphiphiles was also investigated. These various derivatives were used for pDNA[16,37,38] or siRNA delivery (the first formulation used for siRNA delivery was recently approved by the FDA for hereditary transthyretin-mediated hATTR amyloidosis). [39,40] It must be however noticed that polyunsaturated lipid chains can be unstable due to their sensitivity to oxidation. Very recently, we investigated an alternative to the use of unsaturation or ramification to modulate the supramolecular packing of cationic amphiphiles. For that purpose, we constructed new lipid chains featuring one thioether function within a linear and saturated hydrophobic chain[41]. We demonstrated that the presence of one thioether function did not modify the amphiphilic nature of the molecules but affected their physico-chemical properties (e.g. fluidity and molecular surface area); efficient gene carriers could thus be obtained depending on the position of the thioether function within the lipid chain. These results echo other studies showing that the incorporation of a thioether functional group affected the temperature of fusion of ionic liquids[42] or the...
temperature of transition ($T_m$) of bolaamphiphiles.\textsuperscript{[43]} In the current work, we further investigate the use of thioether function to prepare cationic amphiphiles featuring two such functions in each hydrophobic chain of bicatenar amphiphiles. The first goal of this study was to propose a simple synthesis procedure to prepare lipid chains incorporating two thioether functions and then to incorporate these new lipid chains in the structure of cationic amphiphiles. The second goal consisted to evaluate the different physico-chemical behavior of the new cationic amphiphiles when compared to similar amphiphiles (identical polar head groups) but possessing either saturated, mono-unsaturated lipid chains or lipid chain containing a single thioether function. The third goal consisted to evaluate the transfection efficacies and the cytotoxicity of the new cationic amphiphiles. We also compared their transfection efficacies to the other benchmark (commercial or from our laboratory i.e. “Brest Synthetic Vectors”) compounds.

To reach these goals, the new cationic amphiphiles (compounds 4a-c; figure 1) were systematically compared to compound BSV-S12 (which is an efficient vector for gene delivery and that features one thioether function per lipid chain\textsuperscript{[41]} and the references compounds BSV36 (incorporating oleyl lipid chains) and BSV101 (featuring stearyl lipid chains).\textsuperscript{2} It is worth noticing that all these compounds possess a closely related chemical structures; they all possess the same polar headgroup (trimethylammonium) and spacer (phosphoramidate) varying only according to the number and position of thioether functions within lipid chains of exactly the same length (incorporating 18 atoms in its backbone). These structural similarities are helpful to draw conclusions on the impact of the molecular structural variations (presence of two thioether functions and their location in the hydrophobic chains).

Figure 1. Chemical structure of the new cationic amphiphiles possessing two thioether functions per lipid chain (4a-c) and the reference structure BSV-S12 (one thioether function per lipid chain), BSV36 (oleyl chains) and BSV101 (stearyl chains).\textsuperscript{2}

Results and Discussion

The reaction of alkylthiol with a mono-substituted alkene (terminal alkene) in presence of UV light and with a radical initiator (photo-click reaction) is a regio-selective reaction that produce the linear compound possessing one thioether function.\textsuperscript{[44,45]} This reaction is very efficient and was applied for the synthesis of polymers\textsuperscript{[46]} or dendrimers.\textsuperscript{[47]} With the aim to incorporate two thioether functions within the hydrophobic chains of cationic amphiphiles we investigated their construction by applying a double thiol-ene reaction that involved one $\omega$-alkyl-di-thiol and two compounds possessing each a terminal alkene. One of these alkenes possesses one alcohol function in terminal position (Scheme 1) whereas the second alkene is non-functionalized. The thiol-ene reaction was achieved in presence of DMPA (2,2-Dimethoxy-2-phenylacetophenone) in presence of UV light (365 nm) for 18 hours without solvent. We anticipated that this reaction produced three products (statistical reaction): one functionalized with two alcohol functions (side products $A_a-c$), the desired products (1a-c) possessing one alcohol function and the second side product $B_a-c$ that is devoid of any alcohol function. Thanks to their different polarity, these compounds were separated by chromatography to produce compounds 1a-c (Scheme 1) in correct yield (45 to 53 %) and reasonable amount (1.2 to 1.5 g per synthesis). It must be noted that the stoichiometry of the reaction was optimized. We observed that the use 1/1/1 stoichiometry (alkene/dithiol/alkenol) did not produced the best yields (10 %) likely because the alkenes (pentene and hexene) are volatile. In consequence, we increased the quantity of alkene up to 4 equivalents (4/1/2 stoichiometry respectively for alkene/dithiol/alkenol) to obtain better yields (43-53%).
formulated as liposomal solutions by hydration of a lipid film. After a hydration period of 18h at 4°C, the solutions were sonicated (30 min.) thus producing homogeneous solutions that were characterized by Dynamic Light Scattering (DLS) and zeta potential measurements (Table 1 and SI4). We have previously observed that the cationic lipophosphoramidates possessing one thioether function per hydrophobic lipid chain produced liposomes featuring a sized ranging from 75 to 94 nm[41] whereas the cationic amphiphiles possessing the same polar head group but oleyl or stearyl lipid chains produced liposomes with size around 200 nm according to DLS measurements. This trend was confirmed herein, since the cationic amphiphiles 4a-c possess a size ranging from 82 to 89 nm (Table 1). We can therefore conclude that the phosphoramidate featuring hydrophobic chains functionalized with one or two thioether functions per lipid chain produced, according to our protocol, liposomes featuring a sized below 100 nm. These liposomes, prepared from 4a-c, were kept at 4°C for 3 months and only a few differences were observed by DLS and zeta measurements attesting of their stabilities (supporting information SI5). These results suggest that the incorporation of this type of hydrophobic chain could be a strategy to produce liposomes with a small size (<100 nm). It also suggest that this type of lipid chain accept high curvature when compared to cationic amphiphiles featuring oleyl or stearyl chains. This property can likely be related to the flexibility of the lipid chain.[50]

Regarding the zeta potential, all the liposomal solutions prepared have a strong positive zeta potential values as usually observed for cationic liposomes (Table 1).

Then, we recorded the compression isotherms at the air-water interface for the new cationic amphiphiles 4a-c and the three benchmark compounds (supporting information SI6). The more important data extracted from these isotherms are shown in table 2. First we observed that the minimal surface area at which the surface pressure started to increase \( A_0 \) of the cationic amphiphiles 4a-c is sensible (lift-off value; Table 2) is ranging from 154 to 227 Å²/molecule for the compounds 4a-c (a similar value is observed for BSV S12: 180 Å²/molecule ) whereas this value is 80 Å²/molecule for compound BSV101 (stearyl chains). The presence of one or two thioether functions per lipid chain (compounds 4a-c and BSV S12) produces amphiphiles that cover a more important surface than BSV101. Weak hydrogen bonds involving the thioether functions and the water surface could explain this behavior. The molecular surface area at the collapse is another

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Table 1: Size and zeta potential of the new cationic amphiphiles 4a-c and the benchmark compounds BSV-S12, BSV36 and BSV101.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Size (nm)</th>
<th>Pdi</th>
<th>Zeta (mV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>4a (BSV-S4,13)</td>
<td>89.3 ± 2.1</td>
<td>0.34</td>
<td>+ 69 ± 3</td>
</tr>
<tr>
<td>4b (BSV-S5,12)</td>
<td>83.5 ± 1.7</td>
<td>0.28</td>
<td>+ 44 ± 2</td>
</tr>
<tr>
<td>4c (BSV-S7,12)</td>
<td>82.1 ± 3.8</td>
<td>0.38</td>
<td>+ 71 ± 4</td>
</tr>
<tr>
<td>BSV-S12</td>
<td>94.0 ± 1.6</td>
<td>0.34</td>
<td>+ 69 ± 1</td>
</tr>
<tr>
<td>BSV36</td>
<td>200.3 ± 4.5</td>
<td>0.35</td>
<td>+ 48 ± 2</td>
</tr>
<tr>
<td>BSV101</td>
<td>198.3 ± 3.3</td>
<td>0.31</td>
<td>+ 59 ± 1</td>
</tr>
</tbody>
</table>

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Scheme 1. Synthesis of the new cationic amphiphiles 4a-c.
parameter that indicates how the amphiphiles can reduced their surface area before the monolayer was disrupted. To this respect, the compression isotherms clearly show that the amphiphiles 4a-c occupied a lower surface area at the collapse (from 25 to 41 Å²/molecule) when compared to BSV36 (75 Å²/molecule) or to a less extent BSV101 (45 Å²/molecule) or BSV-S12 (45 Å²/molecule). The differences observed between 4a-c and BSV-S12 indicate that the presence of a second thioether function per lipid chain can provide additional effects. This feature is also consistent with the formation of small liposomes because such supramolecular assemblies require, in the inner leaflet, amphiphilic compounds that occupy a reduced surface area whereas in the outer leaflet the lipid chain have more space due to the curvature. Finally, the surface pressure at the collapse indicates how the supramolecular assembly supports the lateral compression and also indicates on its stability. The more is the pressure at the collapse the more is the film stability. To this respect, the highest surface pressure at the collapse is logically observed for BSV101 that possess C18:0 lipid chains. The cationic amphiphiles 4a-c feature the lowest value (42 to 43 ±0.5 mN/m). These values are even lower than for BSV36 (oleyl chains) and BSV-S12 (one thioether function per hydrophobic chain).

Altogether, the compression isotherms indicate that the hydrophobic chains of compounds 4a-c are more adaptable to compressive stress than the other lipid chains included in this study (stearyl, oleyl, and mono-thioether lipid chains). The position of the two thioether functions within the hydrophobic chain has in this series of compound a limited impact since compounds 4a-c feature very close behavior at the air-water interface except for the lift-off with values ranging from 154 to 227 Å²/molecule.

The reflectivity of the surface (reflective index and thickness) is the property of the film which is detected by measuring the ellipsometric angle (Δmax). These measurements were achieved all along the compression isotherms (simultaneously with the surface pressure) but two important values (at the lowest compression A0 (Δmin), and at the highest compression (Δmax) just before the collapse) are reported in table 3 for each sample (supporting information SI7). The ellipsometric values are correlated to the thickness of the lipid surface at the air-water interface and only the comparison of these values worth to be discussed. It must be emphasis that we hypothesize that all the lipid chains have a comparable reflective index despite the presence of sulfur atom in compound 4a-c and no sulfur atom in compounds BSV36 and BSV101. First, the delta max values revealed that the presence of one unsaturation (BSV36) deeply limit the thickness of the lipid layer at the highest compression (5.8° for BSV36 when compared to almost 10° for 4a-c). The delta max values also indicates that the presence of two thioether functions per lipid chain produced thicker lipid layers. These layers are even thicker than with lipid chain containing only one thioether function (BSV-S12). These results indicate that under lateral pressure the lipid chains possessing two thioether functions can adopt a straight conformation likely orthogonal to the surface thus producing a lipid layer having a comparable thickness than with saturated lipid chain (BSV101). Second, at the lift-off pressure (A0), as expected all delta values are less than those recorded at higher pressure because the lipid chains have more space and therefore the lipid monolayer is less thick. However, we can emphasis some marked differences depending on the structure of the lipid chain. With the saturated lipid chains (BSV101), the ellipsometric angle is up to 3 times higher than for the compounds 4a-c. This suggests that compounds 4a-c at low compression pressure can cover more surface than with the saturated lipid chains (BSV101). This could be explained by the presence of weak hydrogen bonds between the thioether function and the water surface. The presence of one unsaturation (BSV36) produces reflectivity comparable to those recorded for compounds

![Figure 2. Compression isotherms for the new compounds 4a-c](image)

| Table 2. Principal data extracted from the compression isotherm experiments |
|-------------------|-------------------|-------------------|
| Lift off value (n=2 or 3) Å²/molecule | Collapse molecular area (n=2 or 3) Å²/molecule | Collapse surface pressure (n=2 or 3) mN/m |
| 4a (BSV-S4,13) | 165 | 41 | 42 |
| 4b (BSV-S5,12) | 154 | 25 | 45 |
| 4c (BSV-S7,12) | 227 | 30 | 42 |
| BSV-S12 | 180 | 45 | 47 |
| BSV36 | 160 | 75 | 47 |
| BSV101 | 80 | 45 | 58 |

[a] Minimal surface area at the surface pressure equal to 0 mN/m.
meaning that at low pressure the thickness of the layer should be comparable. The last column of Table 3 reports the difference between maximal and minimal ellipsometric angle (Δn, °). More this difference is high more the lipid can adapt its conformation to respond to the lateral pressure. To this respect, compounds 4a-c features the highest difference. Considering that the polar head groups are identical in all this series of cationic amphiphiles whereas the volume occupied by the lipid chains varies depending on the lateral pressure we can conclude that these amphiphiles 4a-c could have a positive curvature or a negative curvature depending on the constrain. This adaptability could favor the stabilization of nanoparticles with higher curvatures and could explain the formation of smaller supramolecular assemblies with the compounds 4a-c (Table 1).

Table 3. Ellipsometry results extracted from compression isotherms.

<table>
<thead>
<tr>
<th></th>
<th>Delta min[n]</th>
<th>Delta max[n]</th>
<th>Delta max - Delta min</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(n=2 or 3)</td>
<td>(n=2 or 3)</td>
<td>Δn, °</td>
</tr>
<tr>
<td>4a (BSV-S4,13)</td>
<td>3.6 ±0.1</td>
<td>10.1 ±0.2</td>
<td>6.5 ±0.2</td>
</tr>
<tr>
<td>4b (BSV-S5,12)</td>
<td>2.0 ±0.1</td>
<td>9.8 ±0.1</td>
<td>7.8 ±0.1</td>
</tr>
<tr>
<td>4c (BSV-S7,12)</td>
<td>3.3 ±0.1</td>
<td>10.6 ±0.1</td>
<td>7.3 ±0.1</td>
</tr>
<tr>
<td>BSV-S12</td>
<td>1.5 ±0.1</td>
<td>7.7 ±0.1</td>
<td>6.2 ±0.1</td>
</tr>
<tr>
<td>BSV36</td>
<td>3.5 ±0.1</td>
<td>5.8 ±0.1</td>
<td>2.3 ±0.1</td>
</tr>
<tr>
<td>BSV101</td>
<td>6.1 ±0.1</td>
<td>9.3 ±0.1</td>
<td>3.2 ±0.1</td>
</tr>
</tbody>
</table>

[a] Delta at the Minimal surface area at the surface pressure equal to 0 mN/m. [b] Delta at high pressure (just before the collapse).

Fluorescence anisotropy of a hydrophobic probe is another technic that can be used to assess the rigidity/fluidity of the non-polar domains of supramolecular assembly of amphiphiles. For that purpose the compounds 4a-c were formulated as liposomal solutions and the fluorescent probe diphenylhexatriene (DPH) was added. We recorded the anisotropy of fluorescence of liposomes 4a-c in water from 5 to 65°C (Figure 3 and supporting information S1B). We observed that the compounds 4a-c exhibited a low anisotropy but no transition between a gel to a liquid phase was detected in the range of temperature considered suggesting that the main transition (Tm) was below 5°C. This result contrast with our previous study reporting a transition at 16°C for BSV-S12. We can conclude that the compound 4a-c exist in a fluid phase at temperature upper than 5°C. This result is fully consistent with the compression isotherms that concluded that the amphiphiles 4a-c were very flexible and more fluid than the compounds featuring only one thioether function per lipid chain (BSV-S12) as shown by the pressure at the collapse, which is lower for the compounds 4a-c. We can therefore conclude on a cumulative effect induced by increasing the number of thioether function per lipid chain on the fluidity of the hydrophobic domains.

The physicochemical characterizations of the cationic amphiphiles 4a-c emphasize the specificities induced by the presence of two thioether functions within the lipid chains. We have shown an increase of the fluidity and a capacity to adapt the conformation of the lipid chains when a lateral pressure is applied. All these features prompt us to explore the use of these cationic amphiphiles as nucleic acids’ carriers. Indeed, the adaptability of the conformation of the lipid chains should stabilize the nanoparticles formed by the auto-assembly of a plasmid DNA (pDNA) with cationic amphiphiles (lipoplexes). On the other hand, the fluidity should favor the escape of the lipoplexes from the endosomes after cell internalization. First, we assessed the capacity of the cationic amphiphiles 4a-c to compact pDNA by mixing these amphiphiles at different charge ratio (CR) with a pDNA (pGM144466). DLS measurements of the lipoplexes prepared at different CR (CR = 2, 4, 6) (supporting materials S19) indicated an increase of the size whereas the zeta potential consistently increased from CR=2 to CR=6. As shown with the electrophoresis gel retardation assays (Figure 4A), all three compounds 4a-c were able to compact pDNA. A full compaction was observed at CR=2 for 4b and 4c whereas it was observed at CR=4 for 4a. The addition of the polyanionic polymer dextran sulfate (ds) on preformed lipoplexes (CR=4) allowed to restore the migration of free pDNA as shown in figure 4B. By contrast, only very partial pDNA release was observed after addition of dextran sulfate on BSV101 lipoplexes, suggesting a stronger compaction within such complexes (Supporting Information S10).
The newly reported cationic amphiphiles 4a-c were evaluated as regards their ability to deliver the luciferase-encoding pDNA pGM144 in three different cell lines (A549, C2C12 and 16HBE). These transfections were achieved at different charge ratio (from 0.5 to 8). It was observed that the best CR was 4 in term of transfection efficiencies. Accordingly, we present in figure 5, the transfection results at CR 4 for 4a-c and three reference compounds. All the transfection assays were achieved the same day under identical experimental conditions thus allowing to make a straight comparison of the transfection efficacies. It must be emphasized that BSV36 was previously compared to Lipofectamine [26] and also to the benchmark compound from our laboratory (KLN47) and demonstrated its efficacy [36]. The results obtained indicate that the transfection efficacy was dependent of the cell line (A549 and 16HBE were more easily transfected than C2C12). The most efficient carriers were 4a and 4b, irrespective of the cell line considered.

In parallel, cells viability was measured under identical experimental conditions, i.e. 24 h after transfection with lipoplexes formed at the same CR as before (CR 2, 4 and 6). As for 16HBE and A549 cell lines, the results showed that 4a and 4c were almost safe (with cell viabilities remaining between 80 to 100%). However, 4b used at CR 4 exhibited some toxicity towards A549 (with about 60% of viability). Unexpectedly, for a higher CR, 4b and BSV-S12 were found less toxic towards A549. It is noteworthy that the best transfection efficiencies were measured at CR=4, for which cell viability was in most of the cases the lowest. For instance, compound 4b was both less efficient and less cytotoxic when increasing the CR above 4 (i.e. at CR 6). As for C2C12, it appeared to be more sensitive to all lipoplexes, especially 4b, independently of the CR used.

**Figure 4.** A) DNA binding ability of compounds 4a-c at charge ratio (CR) = 0.5, 1, 2, 4, 6 and 8. B) Gel electrophoresis of lipoplexes prepared at CR=4 in the absence (-ds) and in presence (+ds) of dextran sulfate. The pGM144 pDNA was used in all these experiments.

**Figure 5.** Transfection efficiency of compounds 4a, 4b and 4c in comparison to BSV-S12 and two references BSV101 and BSV36 at CR=4 on three different cell lines (A549, C2C12 and 16HBE). As negative control, free pDNA (pGM144, luciferase-encoding plasmid, 0.25 µg/well) was used. 24 h after deposition, transfection efficiency was determined by luciferase assay. Results are expressed as the Relative Light Units/mg of total proteins, as the mean ± SD of 3 wells.

**Figure 6.** Cell viability of A549, C2C12 and 16HBE 24 h after treatment with lipoplexes formed at different CR (2, 4 and 6). Naked pDNA was used as negative control. Results are expressed as a percentage of the viability determined with un-transfected cells, as the mean ± SD of 3 wells.

**Conclusions**

The synthesis of hydrophobic chains that incorporate two thioether functions can be readily achieved by using a statistical double thiol-ene photo-click reaction. Accordingly, the structure of the di-thiol used as substrate defines the number of methylene units that separate the two thioether functions in the final hydrophobic chain. Bicatenar cationic amphiphiles possessing these thioether-functionalized hydrophobic chains were synthesized. These cationic amphiphiles can more easily adapt the conformation of their lipid chain to support lateral pressure as illustrated by the lower value of the molecular surface area at the collapse for 4a-c when compared to BSV101, BSV36 and BSV-S12. This adaptability to the lateral pressure is also evidenced by the thickness of the supramolecular assemblies (determined by ellipsometric measurements) at the air-water interface. To this respect, the cationic amphiphiles possessing the new lipid chains (4a-c) feature the highest adaptability when compared to cationic amphiphiles possessing oleyl chains (BSV36) or stearyl chains...
(BSV101). The fluorescence anisotropy also emphasizes the additional effect induced by the presence of two thioether functions per lipid chain as shown by the lower values of the transition between rigid to fluid phase. All these physico-chemical properties suggest that the incorporation of two thioether functions per hydrophobic chain (4a-c), in place of one (BSV-S12), renders the supramolecular assembly more fluid, with a greater adaptability to lateral pressure (lower molecular surface at the collapse, lower value of the pressure at the collapse, higher variation of the ellipsometry angles). The incorporation of two thioether functions in the hydrophobic domain thus represents another alternative for modulating the strength of the resulting supramolecular assemblies. The effect of subsequent properties towards gene transfection activity was evaluated. We report that two of the new compounds described herein (i.e. 4a and 4b) are among the most efficient to transient three animal or human-derived cell lines.

Experimental Section

Synthesis – General protocols

Synthesis of compounds 1a-c: Alkenol (1 equivalent), alkene (4 equivalents) and alkanedithiol (1 equivalent) were placed in a Schlenk tube (pyrex) with 2,2-dimethoxy-2-phenylacetophenone DMPA (0.4 equivalent) and degassed under argon for 10 min. The UV Lamp was a Phillips TL-D 18W/08 BLB (λ = 365 nm) that was placed close to the Schlenk tube (= 1 cm). The mixture was then placed under UV light at 18-20°C for 18h. The crude product was purified using flash chromatography Hexane/AcOEt to give the desired product.

Synthesis of compounds 2a-c: Compound 1 (2.2 equivalent) and diphenylphosphate (1 equivalent) were mixed and heated at 130 °C in a Kugelroch distillation apparatus under reduced pressure (4.2 10⁻² mbar) for 3 h. Phenol is removed by distillation. The undistilled compound correspond to the desired products 2a-c. They were used without further purification.

Synthesis of compounds 3a-c: To a solution of compound 2 (1 equivalent) in CHCl₃ was added N,N-dimethylhexylenediamine (1.1 equivalent), DIFEA (1.1 equivalent) and BrCcI₂ (1 equivalent) at 0°C. The reaction was stirred for 15 min. at 0°C then for 4 h at 20°C. The solvent was evaporated and replaced by diethyl ether. After agitation for 10 min. the solution was filtrated and the filtrated was concentrated. The residue was then redissolved in CH₂Cl₂, washed with water, dried over MgSO₄ filtered and concentrated. The compounds were purified by silica gel chromatography using CH₂Cl₂/Methanol: 90/10 (v/v) as eluent to produce 3a-c as a colourless viscous oil. The purification by chromatography at this step is important because in the next step (see below), the purification consisted to remove all the volatile compounds.

Synthesis of compounds 4a-c: To a solution of compound 3 (1 equivalent) in CH₂Cl₂ was added methyl iodide (4 equivalents). The reaction was stirred overnight at 20°C. The solvent and the excess of methyl iodide were evaporated by leaving under vacuum for 2 to 5 h in order to remove all the volatiles (solvent and excess of methyl iodide) that are the only chemicals present in the crude product. No additional purification was needed. The compound 4a-c were isolated as pale yellow waxes.

Compression isotherms

The Langmuir trough was cleaned with ethanol and water then it was filled with the sub-phase (ultrapure water). A good base line in the (π–A) and (Δ–A) isotherms indicated the cleanliness of the interface. The cationic lipid in CHCl₃/MeOH (2/1) mixture (25 μL) was slowly spread at the air-water interface using a Hamilton micro syringe. After waiting 10 minutes till the solvents evaporate, the isotherm of the lipid monolayer was recorded by compressing the barriers at the rate of 5 cm²/min. The surface pressure was measured using a filter paper held by a Wilhelmy balance connected to a microelectronic feedback system (Nima technology). The experiment is stopped once the collapse is reached.

Ellipsometry

The ellipsometric measurements were carried out with a home-made ellipsometer associated to the Langmuir trough and the tensiometer. The software developed recorded simultaneously the value of the surface pressure and the value of the ellipsometric angle during the isotherm compression. The principal of the ellipsometric angle measurement is the following: a polarized He–Ne laser beam (λ = 632.8 nm, Melles Griot, Carlsbad, CA) is reflected by the air/water interface. The incidence angle of the light was 1° away from the Brewster angle (53.12° value obtained for an air/water interface). After reflection, the laser light passed through a λ/4 retardation plate, a Glan-Thompson analyzer, and a photomultiplier. Through a computer-controlled feedback loop, the analyzer automatically rotated toward the extinction light position. In this “null ellipsometer” configuration, the analyzer angle, multiplied by 2, yielded the value of the ellipsometric angle (Δ), i.e., the phase difference between parallel and perpendicular polarization of the reflected light, which reflects the thickness of the molecular film adsorbed at the interface. The laser beam probed a surface of 1 mm² and a depth of the order of 1 μm. Values of Δ were recorded every 4 s with a precision of ±0.5°.

Determination of CAC and Fluorescence anisotropy

Fluorescence was recorded on Agilent Cary Eclipse Fluorescence Spectrophotometer, using automatic polarizers for anisotropy measurements. The fluorescent probe 1,6-diphenylhexatriene (DPH) was used for anisotropy measurements whereas the solvatochromic fluorescent probe Nile red was used for Critical Aggregation Concentration (CAC) determination. Both were purchased from Sigma-Aldrich.

Size and zeta measurements

Size and zeta measurements were achieved with a MALVERN Nano ZS.

Compaction of pDNA

Lipoplexes were prepared by mixing pDNA (pGM144 – 3.7 kb)[54] with each liposomal solution in water, for concentrations corresponding to CR ranging from 0.5 to 8. The resulting mixtures were incubated at room temperature for 30 minutes before being subjected to electrophoresis in a 0.8% agarose gel at 100 V, 90 mA. The gel previously stained with ethidium bromide nuclei acids gel staining (Dominique Deutscher) was visualized using a UV transilluminator (Fisher Bioblock). In order to determine the recover free pDNA from preformed lipoplexes, dextran sulfate was mixed with the latter and incubated for 30 min at room temperature. Electrophoresis was then performed as detailed before.

Transfection

The in vitro reporter gene assay via luciferase measurement was carried out as previously reported.[11] In short, the three cell lines were grown in either EMEM (16HBE) or DMEM (A549 and C2C12) both supplemented
with 10% bovine fetal serum, 1% antibiotic and 1% L-glutamine. All incubations were performed at 37°C in a humidified atmosphere containing 5% CO₂. The day before transfection, the cells were seeded into a 96-well plate at a density of 20,000 cells per well for A549 and C2C12 and 40,000 cells per well for 16HBE. Lipoplexes were prepared as detailed above and then added dropwise into each well; the reference compound BSV36 was used as a positive transfection control whereas naked DNA was used as a negative control. After 24 h at 37°C, the culture medium was removed and the cells were lysed with Passive Lysis Buffer (Promega) prior to running a chemiluminescence assay (Luciferase Assay System, Promega) to determine luciferase expression. The total protein content of each cell lysate was determined using the BC assay kit (Uptima). Finally, data were expressed as relative light units (RLU) per milligram of total proteins (mean ± SD with n=3).

Cell viability
The ViaLight kit (Lonza) was used to determine the ATP content which reflects the number of living cells (transfected or not) in culture. The latter was expressed as percentages relative to the viability of non-transfected cells used as reference (100% cell viability).

Acknowledgements
We thank Association Française contre les Myopathies (AFM-Francaise), Association Gaëtan Saleün (Brest-France), Vaincre La L mucoviscidose (Paris, France), CNRS and INSERM for funding. RG is grateful for a PhD fellowship from Brest Métropole and Association Gaëtan Saleün. We thank the platforms (RMN-RPE and spectrométrie de masse) from the University of Brest.

Keywords: cationic amphiphiles • lipid • thyoether • supramolecular chemistry • transfection

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