# Deciphering the distinct biocontrol activities of lipopeptides fengycin and surfactin through their differential impact on lipid membranes

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### **Statistical summary:**

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# **Graphical Abstract**



# **Highlights**

- SRF and FGC's biological effects correlate with distinct impacts on membrane lipids
- Antifungal activity was observed for FGC but not for SRF
- Stimulation of plant immunity was observed for SRF but not for FGC
- FGC permeabilizes membranes with little impact on lipid packing
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# **Abstract**

Lipopeptides produced by beneficial bacilli present promising alternatives to chemical pesticides for plant biocontrol purposes. Our research explores the distinct plant biocontrol activities of lipopeptides surfactin (SRF) and fengycin (FGC) by examining their interactions with lipid membranes.

Our study shows that FGC exhibits a direct antagonistic activity against *Botrytis cinerea* and no marked immune-eliciting activity in *Arabidopsis thaliana* while SRF only demonstrates an ability to stimulate plant immunity. It also reveals that SRF and FGC exhibit diverse effects on membrane integrity and lipid packing. SRF primarily influences membrane physical state without significant membrane permeabilization, while FGC permeabilizes membranes without significantly affecting lipid packing. From our results, we can suggest that the direct antagonistic activity of lipopeptides is linked to their capacity to permeabilize lipid membrane while the stimulation of plant immunity is more likely the result of their ability to alter the mechanical properties of the membrane.

Our work also explores how membrane lipid composition modulates the activities of SRF and FGC. Sterols negatively impact both lipopeptides' activities while sphingolipids mitigate the effects on membrane lipid packing but enhance membrane leakage.

In conclusion, our findings emphasize the importance of considering both membrane lipid packing and leakage mechanisms in predicting the biological effects of lipopeptides. It also sheds light on the intricate interplay between the membrane composition and the effectiveness of the lipopeptides, providing insights for targeted biocontrol agent design.

**Keywords (max 6):** Lipopeptides, membrane activity, antifungal activity, plant immunity

# **1. Introduction**

Biocontrol is a promising option to reduce the use of chemical pesticides that rises concerns regarding the impacts on human health and ecosystem degradation [1–4]. This approach consists in using natural mechanisms with the application of living organisms or derived molecules to reduce the effect of detrimental organisms and/or enhance growth of useful organisms such as crops [5]. Biocontrol agents can act either by direct antagonism against pests or by stimulating plant immunity to improve their responses against pathogen invasion [5]. The latter approach relies on a process called induced systemic resistance (ISR), where plant defense are potentiated by exogenous molecules, called elicitors, that can be synthetic or arising from living organisms such as microbes or plants [6–9].

In this context, cyclic lipopeptides (CLPs) produced by plant beneficial bacilli showed to be promising candidates as biocontrol agents as they present both direct antimicrobial activities and ability to stimulate plant defense [9–11]. Among the CLPs produced by *Bacillus*, the two most widely conserved families are surfactin (SRF) and fengycin (FGC) [12]. Each family presents specific structural traits **(Figure 1)** that can influence their conformation, their interaction with membranes, but also their biological activities [13,14].



**Figure 1** Representative structure of surfactin and fengycin families. The canonical surfactin family displays a heptapeptide with two negative charges closed by a fatty acid chain (length ranging from C12 to C17) with a lactone bond. The fengycin family has a decapeptide with three charges (one positive and two negative) closed by an internal lactone ring and linked to a fatty acid chain with a length ranging from C14 to C18. Amino acid in red are charged residues and the ones in blue are neutral residues. [13]

While FGC presents antifungal and antimicrobial activities, SRF is efficient against enveloped viruses [15,16] but does not show consistent antifungal activities [17–24] and requires much higher concentrations for antibacterial properties [13], with an effect mainly observed in synergy with other antimicrobial compounds [25]. The reported ability to stimulate plant immunity also varies between CLPs. In pathosystems involving dicotyledonous plants, ISRactivity of *Bacillus* CLPs showed to be driven by the presence of SRF and, to a lower extent, of FGC [17,26,27]. In rice, a monocotyledonous plant, CLPs-mediated ISR showed a strong dependence to FGC with a minor contribution of SRF [18].

The biological activities of CLPs are supposed to originate from their ability to interact with lipids of the plasma membrane (PM) [9,14,28]. Due to their amphiphilic properties, CLPs readily insert into cell PM which can cause pore formation and membrane disruption contributing to their antimicrobial activities [14]. For this mechanism, the lipid composition of the targeted organism is important as it can affect the interaction of CLPs with PM and is

supposed to drive the specificity of CLPs antimicrobial activities [14,23,29–32]. For plant immune stimulation, the mode of action is far to be understood but one hypothesis is that the insertion of CLPs would alter PM lipid organization triggering defense induction [9,10,33].

Lipids of PM present a huge molecular diversity that varies across kingdom and species and can be separated in three main classes: glycerolipids, sterols and sphingolipids [34–36]. These lipids also present a complex organization inside PM with an heterogenous lipid distribution between the two leaflets of PM (referred as membrane lipid asymmetry) but also lateral heterogeneities with the presence of membrane lipid domains, or raft, enriched in their sphingolipid and/or sterol content [35,37–39]. Accumulating evidence have shown the importance of this PM lipid segregation in biological processes [38–40]. However, the huge lipid molecular diversity of biological membranes makes complex the understanding of molecular determinants governing the role of PM lipids in cellular functions. Hence, artificial biomimetic membrane models, such as liposomes or lipid vesicles, supported lipid bilayers and lipid monolayers, have been developed as complementary tools allowing a fine-tune control on their lipid composition [34,40]. These models allow to disentangle more easily the relationship between a specific lipid molecule or lipid organization and the membrane activity of exogenous molecules. Nevertheless, as a simplified representation of biological systems, the relevance of membrane model studies has to be confirmed by biological studies [29,41]. In this study, we assessed how the effect of FGC and SRF on lipid membranes can be related

to their biocontrol activities. In a first step, we compared their biocontrol activities by studying both their direct antagonism against the fungal pathogen *Botrytis cinerea* and their ability to stimulate *Arabidopsis thaliana* (hereafter *Arabidopsis*) immune response. In a second step, the membrane activity of the two CLPs was evaluated using liposomes as membrane models, with an easily tunable lipid composition, to detect the specificities in the interaction mechanism between CLPs and membrane lipids. Altogether, biological and biophysical results allowed us to delve into the significance of specific lipids or lipid properties in the eliciting and antifungal activities of CLPs.

# **2. Materials and methods**

#### *Lipopeptide purification*

Surfactin (> 99% purity of a mix of homologues C12/C13/C14/C15 in relative proportions 7/17/45/31%) and fengycin (> 99% purity of a mix of isoform A and B in relative proportion 52/48 %) were purified from spent supernatant of *B. velezensis* liquid culture as previously described [42–44].

Purity of the CLPs was checked by Ultra-Performance Liquid Chromatography (UPLC - Acquity H-class, Waters s.a., Zellik, Belgium) coupled to a single quadrupole mass spectrometer (Waters SQD mass analyzer) on an ACQUITY UPLC® BEH  $C_{18}$  1.7 µm column. Elution was performed at 40°C with a constant flow rate of 0.6 mL/min using a gradient of acetonitrile and water both acidified with 0.1% formic acid as follows: 1 min at 30%, from 30% to 95% in 3.4 min and maintained at 95 % for 2 min. Compounds were detected in electrospray positive ion mode by setting SQD parameters as follows: cone voltage 120 V, source temperature 130°C; desolvation temperature 400°C, and nitrogen flow: 1000 L.h<sup>-1</sup>.

#### *Plant growth conditions*

Seeds of *Arabidopsis* Columbia ecotype Col-0 were disinfected for 2 minutes in ethanol 75 %, 6 minutes in bleach 5° and rinsed three times in sterilized water. After the disinfection steps, seeds were grown on half strength Murashige and Skoog medium (M0222, Duchefa Biochimie) with addition of 1% (w/v) sucrose and 1.4 % (w/v) agar, with a photoperiod of 12 hours (100  $\mu$ mol s<sup>-1</sup> m<sup>-2</sup>) and a temperature of 21°C.

For systemic immune activation assay, germinated seedlings were then transferred in Araponics systems containing nutritive solution (0.25% (v/v) FLORAMICRO®, 0.25% (v/v) FLORABLOOM®, 0.25% (v/v) FLORAGRO®; General Hydroponics®) and were grown during 3 weeks at 21°C under a photoperiod of 12h (100  $\mu$ mol s<sup>-1</sup> m<sup>-2</sup>).

### *Mycelial growth inhibition*

*B. cinerea* isolate R16 [45] was taken out of the glycerol stock at -80°C and plated on Potato Dextrose Agar (PDA, Merck). The pathogen was then grown for 10 days in a chamber (23°C, 16D:8N photoperiod) before use in the experiment. Round Petri dishes (Ø 90mm) were filled with 35mL of sterilized PDA containing 2.5, 10 or 20µM of either SRF or FGC diluted in ethanol, or ethanol alone for the negative control. A 0.8mm wide mycelial plug of *B. cinerea* grown for 10 days was placed in the center of each dish with the mycelium facing down. The dishes were sealed and placed for 5 days in the same growing chamber. The radius of each fungus was measured every 24h. The Growth Inhibition factor (GI) was calculated at each time point as:  $GI = \frac{Rc - R}{Rc}$  $\frac{c-n}{Rc}$  where Rc is the radius of the control and R is the radius of the treatment.

#### *Systemic immune activation assays*

Four weeks-old plants grown in araponics were transferred in 50ml falcons covered with aluminum foil and containing 45ml of nutritive solution. SRF and FGC were then added in the falcons at a working concentration of 10µM to perform treatment at the root level. The day after, leaf disks of 5mm were cut form the fourth to the sixth leaves of the rosette and were transferred into a white 96-well microplate containing 150µl of deionized water. The plate was incubated at room temperature in the dark overnight.

Measurement of apoplastic UPLC production was performed via a luminol-based chemiluminescence assay. Water was carefully removed and replaced by 90µl of fresh deionized water before adding 10  $\mu$  of a 200  $\mu$ M luminol L-012 + 10  $\mu$ g/ml horseradish peroxydase solution. Luminescence signals were measured using a Spark Tecan multiplate reader. First, the background luminescence level was measured every minute during 5min before adding 2µl of a 5mg/ml chitin suspension. After the addition chitin suspension, luminescence signals were measured every minute for 60min (integration time = 1000 ms).

#### *Root Protoplast isolation*

Roots from 12 to 15 days-old seedlings were cut into 1-2 mm segments and transferred to protoplasting solution (20 mM MES pH 5.7, 0.4 M mannitol, 20 mM KCl, 10 mM CaCl<sub>2</sub>, 0.1 %) (w/v) BSA, 1.5 % (w/v) cellulase R10, 0.4 % (w/v) macerozyme R10) for 5 hours at room temperature and in the dark. The suspension was then filtrated on gauze to eliminate root debris and the filtrate was centrifuged 6 min at 800 rcf. The supernatant was discarded and the pelleted protoplasts were rinsed once with W5 solution (4mM MES pH 5.7, 154 mM NaCl, 125 mM CaCl<sub>2</sub>, 5mM KCI) before being resuspended in WI<sub>Ca</sub> solution (2mM MES pH 5.7, 0.5) M mannitol, 20 mM KCl,  $2mM$  CaCl<sub>2</sub>) at a suitable concentration [46].

#### *Calcium measurements with Fluo-4 AM on protoplasts*

Protoplasts suspension (1-3 x10<sup>5</sup> protoplasts / mL) isolated from roots of *Arabidopsis* col-0 was incubated for 1 hour with 5 µM of fluo-4 AM (ThermoFischer) (from a 5 mM stock solution in DMSO). The suspension was then centrifuge at 700 rcf and the supernatant was discarded to eliminate the remaining free fluo-4 AM. The protoplasts were resuspended in fresh  $Wl<sub>Ca</sub>$ solution and were incubated for 1 hour more. Then, wells of black 96-well microplates (Greiner Bio-One™ CellStar™, Fischer Scientific) were loaded with 100 µL of protoplasts solution per well. After the addition of 25 µL of 5 times concentrated treatment (i.e. a concentration of 50 or 100 µM of CLPs to reach a final concentration of 10 or 20 µM, respectively), the fluorescence was recorded every 15 seconds using Spark® microplate reader (Tecan) with an excitation wavelength at 465 ( $\pm$ 35) nm and an emission wavelength at 535 ( $\pm$ 25) nm.

The values obtained were then converted as normalized fluorescence increase ( $F/F_0$ ) by dividing the fluorescence measured at each time points (F) by the fluorescence measured at the first time point  $(F_0)$ .

#### *Composition of biomimetic liposomes*

To mimic fungal PM, liposomes composed of 53 mol% 1-palmitoyl-2-oleoyl-glycero-3 phosphocholine (POPC), 22 mol% of 1-palmitoyl-2-oleoyl-sn-glycero-3-phospho-(10-racglycerol) (POPG) and 25 mol% of ergosterol were used. This model was based on the one developed by Monnier and co-workers to represent *B. cinerea* PM [47] with the presence of high amounts of ergosterol and a high ratio of anionic/zwitterionic phospholipids.

For liposomes mimicking plant PM, we used a lipid composition of 60 mol% 1-palmitoyl-2 linoleoyl-sn-glycero-3-phosphocholine (PLPC) to be representative of plant phospholipid acyl chains composed mainly of palmitic and linoleic acids [35,48], 20 mol% sitosterol (Sito) as it is the most abundant sterol in plants [35] and 20 mol% D-glucosyl-ß-1,1'-N-palmitoyl-D-erythrosphingosine (GluCer) as a commercially available plant sphingolipid [34] and an important lipid class for plant PM in terms of composition but also with a key role in its organisation [35,49].

### *Liposome preparation*

1-palmitoyl-2-oleoyl-glycero-3-phosphocholine (POPC), 1-palmitoyl-2-oleoyl-sn-glycero-3 phospho-(10-rac-glycerol) (POPG), ergosterol, 1-palmitoyl-2-linoleoyl-sn-glycero-3 phosphocholine (PLPC), sitosterol (Sito) and D-glucosyl-ß-1,1'-N-palmitoyl-D-erythrosphingosine (GluCer) were purchased from Avanti Polar Lipids and used without further purification. The different lipid mixtures of POPC-POPG (70:30 molar ratio), POPC-POPGergosterol (53:22:25 molar ratio), PLPC, PLPC-Sito (80:20 molar ratio) and PLPC-Sito-GluCer (60:20:20 molar ratio) were dispersed in a chloroform/methanol (Scharlau Lab Co.) (2/1; v/v) solution and dried under reduced pressure in a rotary evaporator and then kept under vacuum overnight. For laurdan and dynamic light scattering (DLS) analysis, the dried lipid films were then hydrated to 1 mM of lipids in MES 10 mM - NaCl 150 mM buffer at pH 5.8 during 1 h at 45°C with vortex mixing applied every 15 min and then subjected to five freeze/thaw cycles. For HPTS-DPX measurement, the lipid film was hydrated similarly but the lipid concentration was 3 mM and the hydration buffer also contained 30 mM of 8-hydroxypyrene-1,3,6 trisulfonic acid (HPTS) and 50 mM of p-xylene-bis-pyridinium bromide (DPX). The dispersions were finally extruded fifteen times through two stacked Nuclepore 100 nm polycarbonate filters using a Lipex Biomembranes (Vancouver, BC) extruder to obtain LUVs.

#### *HPTS-DPX leakage assays*

Membrane leakage assays were based on measurements of the release of a fluorescent dye (in this case HPTS) previously co-encapsulated inside liposomes with a quencher (in this case, DPX). Upon leakage, the dye and its quencher become highly diluted in extra-vesicular media which dissociates the dye and the quencher leading to an increase in fluorescence [50].

Following extrusion, liposome suspension was flowed through a Sephadex-G75 (Cytiva) gel purification column to remove the unencapsulated dye. The Sephadex gel was obtained by hydrating 1 g of Sephadex G75 with 30 mL of distilled water at 150 °C for 2 h under shaking at 200 rpm with a magnetic stirrer. The Sephadex gel was conditioned with 20 mL of MES 10 mM - NaCl 150 mM buffer at pH 5.8 before liposome purification.

After purification, liposome suspension was diluted to a concentration of 100-200 µM of lipid and this solution was loaded in wells of a black 96-well microplate (Greiner Bio-One™ CellStar™, Fischer Scientific) with a volume of 100µL per well. The fluorescence was recorded once before treatment and every 2 min until 44 min after the addition of 25 µL of 5 times concentrated treatments (i.e. 12.5, 25, 50, 75, 100, 125 and 250µM of CLPs to reach 2.5, 5, 10, 15, 20, 25 and 50 µM of final concentration, respectively) using Spark® microplate reader (Tecan) with an excitation wavelength at 465(±35) nm and an emission wavelength at 535(±25) nm.

Percentage of membrane leakage was defined as  $(F_f-F_0)/(F_{max}-F_0)_x$ 100, where  $F_t$  is the fluorescence measured at each time point,  $F_0$  is the fluorescence measured before the addition of treatments and  $F_{\text{max}}$  is the maximum fluorescence signal obtained after complete membrane disruption with 0.2% of Triton X-100.

#### *Laurdan polarization on root protoplasts and lipid vesicles*

Protoplast suspension (1-3  $_{x}10^{5}$  protoplasts / mL) or 100 µM lipid vesicle preparation (from a stock solution at 1 mM) was incubated with 1 µM of laurdan (Sigma-Aldrich) for 1h30. Then, wells of a black 96-well microplate (Greiner Bio-One™ CellStar™, Fischer Scientific) were loaded with 100 µL of protoplasts or lipid vesicle solution per well.

The fluorescence was then recorded twice before treatment and every 2 min until 44 min (for liposomes) or every 2.5 min for 20 min (for protoplasts) following the addition of 25 µL of 5 times concentrated treatment (i.e. 12.5, 25, 50, 100 and 250µM of CLPs to reach 2.5, 5, 10, 20 and 50 µM of final concentration, respectively) using Spark® microplate reader (Tecan) with an excitation wavelength at  $360(\pm 35)$  nm and emission wavelengths at  $430(\pm 20)$  nm and 485(±20) nm.

The generalized polarization (GP) was defined as GP =  $\frac{(I_{430nm}-I_{485nm})}{(I_{430nm}-I_{485nm})}$  $\frac{(1430nm^{-1}485nm)}{(I_{430nm}+I_{485nm})}$ , were  $I_{430nm}$  and  $I_{485nm}$ represents the blank-subtracted fluorescence intensities at emission wavelengths of 430 nm and 485 nm respectively. Variation of GP (ΔGP) is defined as the subtraction of GP measured at each time point following treatment and GP measured before treatment.

Blank measurements were carried out on samples containing the different concentrations of CLPs but which were not stained with laurdan. In all blanks tested, no specific fluorescence signal was observed for FGC or SRF.

#### **Dynamic light scattering measurements on liposomes**

Dynamic light scattering (DLS) measurements were performed on liposome solution at different concentrations of SRF or FGC (0, 10, 20, 40, 75 and 187.5 µM). One mL of vesicle solution (lipid concentration of 300 µM) was placed in a cuvette for DLS measurement. The scattered intensity was measured at 25.0 °C using a Zetasizer Nano DLS instrument (Malvern Panalytical) to obtain the liposome size and the derived count rate, in kilo counts per second (kcps). After a first measurements to get liposome size and derived count rate without CLPs, SRF or FGC from a stock solution of 0.5mM were added successively in the liposome solution to obtain the evolution of liposome size with increasing CLP concentration. Measurements were performed three times on each sample to ensure reproducibility (eight acquisitions per measurement with an acquisition time of 10 to 15 seconds).

#### **3. Results**

#### *3.1. Antifungal property and stimulation of plant cell immune responses*

The direct antagonistic activity of SRF and FGC was evaluated against *B.cinerea,* a fungal pathogen of high scientific and economic importance responsible for both pre- and postharvest crop losses [51]. The effect of SRF and FGC was evaluated by comparing the growth of the pathogen on PDA amended with several CLPs concentrations to its growth on PDA without CLPs **(Figure 2a, b)**. In presence of FGC, we observed a significant reduction of *B. cinerea* growth in the same order of magnitude for all the concentrations used, with an inhibition of growth (GI) after 72 hours of about 50% in presence of FGC. The fungal growth in presence of SRF did not show any difference with growth in absence of CLPs. Hence, in accordance with previous studies [17–24], biocontrol activity of SRF does not rely on a direct antagonism while FGC can inhibit the fungal growth through a direct toxicity against fungi.

In our experimental conditions, the effect of FGC on fungal growth did not show a typical doseresponse effect, in accordance with the work by Wise and coworkers [23] on different fungal strains including *B. cinerea*. Nevertheless, our range of concentrations tested, all higher than the critical micellar concentration of fengycin [52], remains limited compared to other studies [21,22] which may explain the absence of a visible dose-response effect.



**Figure 2** Impact of surfactin (SRF) and fengycin (FGC) on the mycelial growth of *B. cinerea* (a) Time-course evolution of *B. cinerea* mycelial growth on PDA in presence of 0.1% ethanol (control), 2.5µM, 10µM and 20µM SRF and 2.5µM, 10µM and 20µM FGC. Images show representative pictures of *B. cinerea* mycelial growth after 72 hours in presence of 0.1% ethanol (negative control), 10µM SRF and 10µM FGC. Scale bar in upper right represents 15 mm. (b) Growth inhibition factor (GI) of *B. cinerea growth* with surfactin and fengycin treatments after 72 hours. Letters indicate statistically different groups at  $α = 0.05$  (one-way ANOVA and Tukey's multiple-comparison post-test). Data in (a) and (b) represents mean  $\pm$  SD (n=6) from two independent experiments.

The stimulation of plant immunity by CLPs was evaluated by analyzing the systemic activation of immune response of *Arabidopsis* pre-treated with CLPs and by analyzing the variation of cytosolic calcium ([Ca<sup>2+</sup><sub>cyto</sub>]) in protoplasts from *Arabidopsis* roots.

To assess the systemic immune activation (SIA), we studied the potentiation of immune responses in leaves of plant pre-treated with CLPs at the root level. After pre-treatments, we measured the production of apoplastic reactive oxygen species (ROSapo) in leaf disks of plant in response to a stimulation with chitin, a well-known plant elicitor [53]. The SIA assays revealed that *Arabidopsis* plants pre-treated at the root level with 10µM SRF, a concentration previously determined as the minimal active concentration [17,54], show an increased  $\text{ROS}_{\text{apo}}$ production in response to chitin treatment compared with non-pretreated plants **(Figure 3a, c)**. In contrast, plants pre-treated with 10  $\mu$ M FGC show a similar ROS<sub>apo</sub> production in response to chitin than plants pre-treated with control solution **(Figure 3b, c)**.



**Figure 3** Effect of fengycin (FGC) and surfactin (SRF) on plant cell immune stimulation and plasma membrane mechanics (a-b) Kinetics of ROS<sub>apo</sub> production measured with luminescence assay following treatment with 250 µg/mL chitin in leaf disk of *Arabidopsis* pretreated at the root level with 10  $\mu$ M SRF (a) or 10  $\mu$ M FGC (b) compared to control pretreatment (0.1% ethanol). Results show the mean luminescence  $\pm$  SD (n≥6). (c) Total ROS<sub>apo</sub> production measured with luminescence assay following treatment with 250 µg/mL chitin in leaf disk of *Arabidopsis* pre-treated at the root level with 10 µM SRF, 10 µM FGC or control pre-treated (0.1% ethanol). Results show the mean total luminescence  $\pm$  SD (n≥6), obtained by the sum of all detected luminescence values. Letters represent statistically different groups at  $\alpha$  = 0.05 (one-way ANOVA and Neuwman-Keuls multiple comparison post-tests). (d) [Ca2+ cyto] elevation detected with Fluo-4 in root protoplasts of *Arabidopsis* Col-0 after control treatment or treatment with 10 $\mu$ M SRF, 10 $\mu$ M FGC or 20  $\mu$ M FGC. Data shows mean  $\pm$  SD

(n≥10) of normalized fluorescence increase of Fluo-4 five minutes after treatments. Letters represent statistically different groups at  $\alpha$  = 0.05 (one-way ANOVA and Tukey's multiplecomparison post-test). (e) Change of laurdan generalized polarization (ΔGP) measured in *Arabidopsis* root protoplasts after control treatment or treatment with 10µM SRF, 10µM FGC or 20  $\mu$ M FGC. Data are represented as mean GP variation  $\pm$  SD (n=6) compared to GP measured before the addition of treatment.

To complete the analysis of CLP's effect on plant immunity, measurements of  $[Ca^{2+}C_{\text{cyto}}]$ variation in root protoplasts were performed. Treatment with 10µM SRF induced a significant [Ca<sup>2+</sup><sub>cyto</sub>] increase while FGC treatment showed significantly lower [Ca<sup>2+</sup><sub>cyto</sub>] responses, even at concentration twice higher than for SRF (**Figure 3d, Suppl Fig1** for time course measurements).

Since the ability of CLPs to stimulate plant immune responses is believed to originate from their interaction with plant PM, we used the facilitated access to PM allowed by protoplasts to investigate the possible effect of CLPs on PM mechanics. Using the solvatochromic probe laurdan, sensitive to membrane hydration and thus to lipid packing [55], we analyzed the variation of PM fluidity in root protoplasts with the two CLPs. No effect on PM lipid packing was observed for FGC contrary to SRF that increased lipid compaction **(Figure 3e)**.

The correlation between the effect of CLPs on fluidity of protoplast PM and their ability to induce a  $[Ca<sup>2+</sup><sub>cyto</sub>]$  responses and SIA agrees with the hypothesis of plant immune stimulation by CLPs based on their ability to interact with plant PM [33] and suggests that FGC is very poorly susceptible to stimulate immunity in *Arabidopsis* contrary to SRF.

### *3.2. Membrane activity on liposomes mimicking fungal and plant plasma membranes*

The biological activities of CLPs likely rely on their ability to insert into membranes and alter lipid membrane properties [14,29,33]. As SRF and FGC exhibit distinct plant biocontrol activities, we investigated whether distinct characteristics in the membrane activity of CLPs could account for these differences. For this purpose, we used liposomes mimicking fungal or plant PM to assess the effect of CLPs on membrane permeabilization and fluidity. The permeabilization, or leakage, assays were conducted by measuring the efflux of HPTS-DPX initially encapsulated within the liposomes. It indicates the ability of CLPs to disrupt and possibly form pores into membranes. This property has been correlated with induction of cell leakage and cell death leading to antimicrobial properties [29,30,32,41,56]. Membrane fluidity experiments on liposomes were performed with the laurdan probe as on protoplasts. Since analysis of membrane activity with liposomes can depend on the lipid concentration [32,57], we analysed our data using the CLPs-to-lipid molar ratio (ratio between CLPs and lipid concentrations, in molar) rather than CLPs concentration only.

Distinct behaviour between the two CLPs was observed in leakage and fluidity assays **(Figure 4)**. For both models, FGC induced membrane leakage at much lower CLPs-to-lipid ratio than SRF. A leakage was already observed at a ratio of 0.06 for FGC while leakage started progressively at ratios higher than 0.12 for SRF (**Figure 4a, e**). The shape of dye efflux was also different for each CLP on both lipid models. The leakage induced by FGC shows a steep increase at low ratio that stabilizes to a plateau while the leakage induced by SRF increased more progressively with increasing CLPs-to-lipid ratio. In contrast to leakage and in accordance with assays on protoplasts, membrane fluidity was more affected by SRF with a slight effect observable at a ratio of 0.12 that progressively increased while FGC effect on membrane fluidity remained low even at high CLPs-to-lipid ratio **(Figure 4b, f)**.



**Figure 4** Effect of fengycin (FGC) and surfactin (SRF) on liposomes mimicking fungal (a to d) and plant (e to h) plasma membranes as a function of the CLPs-to-lipid molar ratio. (a, e) Membrane leakage measured with HPTS-DPX in lipid vesicles (final lipid concentration of 80µM) composed of POPC-POPG-ergosterol (53:22:25 molar %) as fungal model (a) and PLPC-sito-GluCer (60:20:20 molar %) as plant model (e) 30 minutes after the addition of increasing concentration of FGC or SRF. Data are represented as % of leakage  $\pm$  SD (n=6 for 0µM, n=4 for the other concentrations) compared to leakage observed with the addition of 0.2% Triton X-100. (b, f) Change of laurdan generalized polarization (GP) in lipid vesicles (final lipid concentration of 80µM) composed of the fungal model (b) and the plant model (f) 10 minutes after the addition of increasing concentration of FGC or SRF. Data are represented as mean GP variation  $\pm$  SD compared to GP measured before the addition of treatment. (n=4 for fungal model, n=10 for plant model – SRF, n=8 for plant model - FGC). (c, d, g, h) Liposomes diameter (c, g) and variation of derived count rates (d, h) (mean  $\pm$  SD, n=6) measured with dynamic light scattering with increasing concentration of FGC or SRF. The initial lipid concentration was 300 µM. All measurements in the figures come from data obtained from at least 2 independent batches of liposomes.

Results from membrane models suggest that FGC effects are more related to its ability to disrupt and/or permeabilize lipid membranes while SRF insertion leads to changes in membrane structure and/or organization rather than membrane permeabilization, especially at low CLPs-to-lipids ratios. To confirm the permeabilizing effect observed for FGC but not SRF, we measured the liposome diameter and the derived count rate by dynamic light scattering (DLS). The derived count rate measures the strength of light scattering signal of the sample and is thus influenced by both the number and the size of liposomes in the medium [58,59] (a higher derived count rate indicates higher number of particles, larger particles, or higher number and larger particles in the medium). In the presence of a permeabilizing effect, a reduction of liposome number and/or size can be expected. In the fungal model, the presence of FGC has low impact on liposome size while SRF shows a slight increase in liposome diameter at high CLPs-to-lipid ratio **(Figure 4d)**. In the plant model, a slight reduction of liposome size was observed at low CLPs-to-lipid ratio for both CLPs. At higher CLPs-to-lipid ratio, an increase of liposome size was observed for SRF while the liposome size remains stable for FGC **(Figure 4h)**. In addition to size measurements, a progressive reduction of the derived count rate was noticed with increasing FGC concentrations in both models **(Figure 4 c, g)**. In the fungal model, the stable liposome size with a decreasing count rate in presence of FGC indicates a reduction in the number of liposomes. This agrees with the hypothesis that FGC acts by permeabilizing liposomes. In the plant model, the effect of FGC at low ratios on the derived count rate can originate from both its impact on liposome size and a permeabilizing effect, explaining a steeper decrease in the plant model compared to the fungal model, while at higher ratios, the decrease in the derived count rate with no change in liposome size also support a permeabilizing mechanism. In contrast, the presence of SRF has only a small effect on the derived count rate in the fungal model and slightly increased the derived count rate at high CLPs-to-lipid ratio in the plant model **(Figure 4c, g).** This increase coincides with an observed increase in liposome size at high SRF-to-lipid ratio. Hence, it is likely that the increase in derived count rate by SRF comes from its impact on liposome size rather than an impact on the liposome number. The absence of effect of SRF on liposome number supports the idea that SRF does not act through a permeabilizing mechanism but modifies the global structure of the liposome bilayer which could lead to liposome fusion at high SRF-to-lipid ratio.

#### *3.3. Lipid specificity of the interaction with biomimetic liposomes*

The lipid composition is a key parameter for CLPs biological activities. In particular, ergosterol was found to be a crucial determinant for the selectivity of the fungicidal activity of CLPs [23,29,60] and sphingolipids of plant PM was suggested as a preferential interactant for SRF [9]. We therefore investigated the lipid specificity of CLPs membrane disturbance through the modulation of liposome lipid composition **(Figure 5)**. For the fungal model, we evaluated the importance of ergosterol by comparing CLPs membrane activity obtained with the model POPC-POPG-ergosterol (53:22:25 molar ratio) with a second model POPC-POPG (70-30 molar ratio) that keeps the same anionic/zwitterionic lipid ratio but does not contain ergosterol. For plant model, we focused on the importance of sphingolipids and sterols and compared CLPs membrane activity on liposomes with the entire lipid composition PLPC-sito-GluCer (60:20:20 molar %), liposomes containing PLPC-sito (80:20 molar %) without sphingolipids and liposomes containing only PLPC.

In all the lipid compositions tested, FGC-induced leakage showed a similar shape with a steep increase in dye efflux at low ratio (between 0.03 and 0.12) that stabilizes around a plateau for ratios higher than 0.12. The presence of a sterol, either ergosterol or sitosterol, reduced the leakage effect of FGC that occurred at higher CLPs-to-lipid ratio than without sterols (0.06 for POPC-POPG-ergosterol and PLPC-sito vs 0.03 for POPC-POPG and PLPC, respectively) and with a lower magnitude (maximum dye efflux of around 40% for POPC-POPG-ergosterol and 75% for PLPC-sito vs 65% for POPC-POPG and 100% for PLPC) **(Figure 5a, b)**. Interestingly, the presence of both sterols and sphingolipids in the plant model suppressed the inhibitory effect of sterols as a similar leakage was observed for liposomes composed of PLPC and PLPC-sito-GluCer. Comparison of FGC-induced leakage between fungal and plant models also revealed a much lower leakage in fungal models compared to plant models.

For SRF, leakage was more progressive and occurred at higher CLPs-to-lipid ratio than FGC **(Figure 5a, b)**. SRF-induced leakage was also less impacted by the presence of sterols, even though, for plant models, a slight reduction of dye efflux can be noticed for PLPC-sito compared to PLPC. Intriguingly, the presence of sphingolipids increases SRF-induced leakage with a complete leakage reached at a CLPs-to-lipid ratio of 0.25 for PLPC-sito-GluCer liposomes, 0.31 for PLPC and 0.63 for PLPC-sito. Comparison of SRF-induced leakage between fungal and plant models also revealed a slightly lower leakage in the fungal model but the difference was less pronounced than the one observed for FGC.





generalized polarization (GP) in lipid vesicles (final lipid concentration of 80µM) composed of POPC-POPG (70:30 molar %) and POPC-POPG-ergosterol (53:22:25 molar %) in (c) and PLPC, PLPC-sito (60:20 molar %) and PLPC-sito-GluCer (60:20:20 molar %) in (d) 10 minutes after the addition of increasing concentration of FGC (left) or SRF (right). Data are represented as mean GP variation  $\pm$  SD (n $\geq$ 4) compared to GP measured before the addition of treatment. All measurements in the figures comes from data obtained from 2 independent batches of liposomes.

Regarding membrane fluidity, we observed a gradual reduction of FGC and SRF effect with increasing lipid complexity but for all lipid composition, SRF had a higher impact on membrane fluidity than FGC **(Figure 5c, d)**. When looking at the initial lipid packing of the different compositions without CLPs, measured with laurdan generalized polarisation (GP), we observed that models with the highest initial lipid packing (i.e. PLPC-sito-GluCer with an initial GP of -0.11±0.04 and POPC-POPG-ergosterol with an initial GP of -0.16±0.04) are the less sensitive to CLPs-induced rigidification **(Figure 5c, d, Suppl Figure 2)**. At the opposite, the model with only PLPC or POPC-POPG, whose fluidity was the most affected by CLPs, are the most fluid model between the ones tested (with a GP of -0.35±0.06 for PLPC and -0.31±0.03 for POPC-POPG). Hence, a model presenting already a high lipid packing is less prone to undergo further compaction of their lipids.

### **4. Discussion**

### **4.1.** *Disturbance of membrane lipid packing as an additional mechanism to membrane leakage for CLPs biological activities*

SRF and FGC showed to affect lipid membrane through distinct processes. While SRF strongly affects membrane physical state but was less prone to induce membrane permeabilization, FGC showed the opposite with a strong ability to permeabilize membrane without marked effect on its lipid packing.

Differences in membrane activity between SRF and FGC have already been discussed in their leakage mechanism on liposomes, with a graded leakage effect observed for SRF while FGC showed an all-or-none leakage mechanism [29,32,61,62]. A graded leakage is characterized by a progressive and transient release of the dye from all vesicles while the all-or-none leakage shows a more heterogenous dye efflux with a fraction of vesicles releasing all their dyes while the other fraction keeps the dye entirely entrapped. The graded process originates from a phenomenon that dissipates upon leakage such as an asymmetric insertion of molecules in the outer leaflet. This asymmetric insertion causes differential stress between the two leaflets due to imbalance in area density of molecule between leaflets [63]. It leads to transient mechanical failure of the membrane to release the stress, followed by the annealing of membrane [64]. The presence of differential stress into membrane has recently been shown to impact important membrane properties such as tension [65] and could therefore explain the observed impact of SRF on membrane fluidity, which has also been reported previously for DOPC:DPPC vesicles [66]. The all-or-none process arises from the stabilization of defects or pore-like structures in the membrane by covering their edges with a rim enriched in membraneactive molecules [41,64]. This process seems to impact less the membrane lipid packing as we observed lower impact of FGC on membrane fluidity compared to SRF. Such low impact on membrane fluidity of FGC has also been observed on DPPC membranes, where FGC showed to mainly affect the polar headgroup regions but not the hydrophobic core [67]. Nevertheless, the low impact on membrane fluidity does not exclude an impact on the global distribution of membrane domains as it was observed in DOPC:DPPC bilayer [68]. Hence, the effect of CLPs on both membrane leakage and membrane lipid packing could represent two complementary indicators of their specific mechanism of membrane destabilization.

The differences in interaction mechanisms of SRF and FGC with lipid membranes has been suggested as an explanation for their specificities in antimicrobial activities [29,32]. The all-ornone leakage mechanism has indeed shown good correlation with antimicrobial properties which is not the case for the graded leakage. Hence the difference in antifungal activities between SRF and FGC, noted in our study and previous ones [18,21,22,69], can be related to an efficient membrane disruption and/or permeabilization for FGC (in accordance with its allor-none leakage mechanism). In contrast, the effect of SRF on membrane lipid packing and its graded leakage effect may be less effective to disrupt biological membranes. This distinction could also account for antibacterial activities where SRF may require high dose or synergic compounds to be effective [13,25].

However, altering PM physical properties may be responsible for other important biological activities. PM is a flexible and dynamic structure capable of sensing and transducing mechanical signals inside the cell due to its highly dynamic physical properties [70,71]. PM organization has been shown to be crucial for immune responses in plants [72]. Modifying PM physical properties can activate PM signalling proteins [73,74]. This could explain the very low effect of FGC on plant responses as FGC does not alter membrane physical properties. In contrast SRF, able to increase the PM lipid packing, triggers a marked  $[Ca^{2+}C_{\text{cyto}}]$  influx in protoplasts and potentiates ROS<sub>apo</sub> responses in leaves while applied at the root level. Hence, our observations suggest that the distinct effect of SRF and FGC on plant PM properties contributes to their difference in plant eliciting activity.

#### **4.2.** *Modulation of CLPs' membrane activity by membrane lipid composition*

CLPs' activities on both fungal and plant plasma membrane models were negatively impacted by the presence of sterols. Interestingly, the combination sterols-sphingolipids in plant models tended to reduce the impact on membrane lipid packing but it also compensated the sterol effects on leakage and even improved leakage for SRF. Sterols and sphingolipids are known to be key players for membrane organization and lipid packing since changes in their proportion modulate the presence of lipid domains in membrane [35,38,75].

The influence of sterols could come from their ability to increase membrane packing [75]. Previous studies have observed that a higher lipid packing can impede the insertion of SRF [28,66,76] and FGC [44], resulting in a lower membrane disturbance. Sterols were also found to reduce the depth of insertion of other CLPs from the viscosin groups produced by *Pseudomonas* spp. [77]. FGC presents a longer acyl chain than SRF and also possesses two amino acids outside the peptide ring, while SRF has all its amino acids inside the peptide ring (**Figure 1**). This may suggest that acyl chain of FGC will insert more deeply inside the membrane than the acyl chain of SRF. Since sterols are located in the hydrophobic core of the lipid bilayer and hinder a deep insertion of CLPs, they are likely to further affect molecules that insert deeper into the membrane. This could explain why sterols impact FGC-induced leakage but have minimal impact on SRF-induced leakage, as previously observed [29,78]. In the case of FGC, the effect of sterols correlates with its lower antifungal effects on plant pathogen with higher ergosterol content [23].

Regarding the addition of sphingolipids to the sterol-containing model, interaction between sphingolipids and sterols are known to modulate the lateral lipid distribution with the formation of raft domains with more tightly packed lipids [79–81]. The presence of sphingolipids could therefore increase the lateral heterogeneity of the membrane, creating some packing defects at the border of the domains which in turn favors CLPs insertion and membrane disturbance. It has indeed been reported that in presence of phase coexistence, SRF inserts preferentially at the boundary between phases [66] and presents a higher binding affinity for lipid vesicles presenting a phase coexistence [33]. The initial rate of SRF-induced membrane leakage was also higher in DPPC:POPC vesicles [82], which exhibit phase separation [83], compared to POPC vesicles. The importance of phase coexistence has been less studied for FGC but preferential insertion at phases boundary has also been suggested [52]. The lipid phase separation has already been reported as a key parameter for the ability of nystatin, an antifungal drug, to form pore into membrane [84]. In fungal membranes, sphingolipids are important players in domain segregation [38]. Therefore, the inhibition of nystatin activity in *Saccharomyces cerevisiae* mutant with altered sphingolipid composition supports the importance of phase separation for biological activities of nystatin [38,85], and more broadly membrane-active molecules.

Besides the role of sphingolipids and sterols, a lower CLPs-induced leakage was observed in the fungal model compared to the plant model. The differences between these models lie in the presence of the negatively charged lipid POPG in the fungal model and in the difference in the number of unsaturations in the lipid chains, i.e. one unsaturation in the fungal model and two unsaturations in the plant model. The influence of PG presence in the membrane on both SRF and FGC activity has previously been studied and showed to have a negative impact on the FGC-induced leakage and a positive impact on the SRF-induced leakage [29]. This is consistent with the lower FGC-induced leakage we observed in the fungal model compared to the plant model but it contrasts with our slightly lower SRF-induced leakage observed in the fungal model compared to the plant model. A possible explanation for SRF could arise from differences in the number of acyl chain unsaturations that impacts the affinity of SRF for lipid membranes, with a higher affinity for PLPC compared to POPC [33]. Thus, both negatively charged lipids and acyl chain unsaturation may play a role in the activity of CLPs. It nevertheless remains difficult to clearly establish individual contribution of the two components based on our present data.

# **5. Conclusion**

Distinct biocontrol activities of SRF and FGC, whether by stimulating plant immunity or by direct antagonism against plant pathogens, strongly correlates with their specific interaction processes with lipid membranes. CLPs are primarily known for their ability to destabilize membrane, notably through pore formation [14]. Our observations indicate that changes in the membrane physical state also contribute to biological properties of CLPs.

To fully grasp the significance of these membrane-related activities in reducing plant infection by SRF and FGC, further complementary experiments are needed. Other lipopeptides might exert a similar effect, and investigating their capacity to alter membrane physical state could offer additional insights into their potential biological effects. In addition, gaining a deeper understanding of the interplay between membrane activity and biological activity of membraneactive molecules could provide valuable information to the rational design and implementation of CLPs as biocontrol agents.

It is worth noting that this study specifically focused on activities related to plant biocontrol. However, disturbance of membrane lipid organization and structure could unveil other interesting properties, such as antiviral [86–88], antibacterial [89] or antitumoral activities [90]. Therefore, considering the impact on membrane lipid packing alongside the leakage mechanism is essential for understanding and predicting the biological activities of CLPs.

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# **8. Author contribution**

Conceptualization and Methodology: GG, CDC, MD Investigation: GG, TD, FB Resources: MO, MHJ, MD Supervision and Funding acquisition: MD Writing - original draft: GG Writing - review & editing: GG, TD, CDC, MO and MD

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### **Supplementary data**



**Suppl Figure 1** Change of [Ca<sup>2+</sup><sub>cyto</sub>] in *Arabidopsis* root protoplasts in presence of surfactin (SRF) and fengycin (FGC). Detection of [Ca2+ cyto] variation over time with Fluo-4 in *Arabidopsis*  root protoplasts following treatment with 0.1% ethanol (control), 10 µM SRF, 10 µM FGC or 20 µM FGC. Data are represented as mean normalized fluorescence increase  $(± SD, n≥10)$ .



**Suppl Figure 2** Laurdan generalized polarization (GP) of lipid vesicles with the different lipid compositions tested in absence of lipopeptides. Data are presented as mean  $\pm$  SD, n≥5.

Letters represent statistically different groups at  $\alpha$  = 0.05 (Brown-Forsythe and Welch ANOVA and Tamhane's T2 multiple comparison post-test). All measurements in the figures comes from data obtained from at least 2 independent batches of liposomes.