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Introduction

The life of eukaryotic cells depends on membranes that define multiple boundaries in cells. The plasma membrane separates cytosol from the extracellular environment, while organelle membranes maintain the characteristic differences between the contents of each organelle and the cytosol. Ion gradients across the plasma and organelle membranes are precisely controlled in living cells to regulate all aspects of cellular biology. Thus, perturbation of such ion concentration differences is often detrimental to cellular homeostasis.

Ionophore natural products are a family of hydrophobic molecules that disrupt ion gradients to exert various biological effects.¹ They are classified as mobile carriers or channels based on the mechanism of ion transport. Mobile carriers bind with ions and the resultant complex diffuses across membranes, whereas channels constitute transmembrane nanopores that permit ion flow through the membrane. Because of the continuous pore, ion transport by channels is generally 10³ to 10⁴-fold faster than that by carriers. Over the past few decades, mobile carriers have attracted intense attention as lead

compounds for developing new anticancer therapeutics. For example, salinomycin selectively targets cancer stem cells to reduce *in vivo* cancer growth and metastasis, and recently entered phase I/II trials.^{2,3} Compared with mobile carriers, the anticancer properties of channels have remained largely unexplored despite their greater ion transport efficiency.⁴⁻⁶

Gramicidin A (**1**, molecular weight = 1882 Da) and polytheonamide B (**5**, 5030 Da) are ion-channel forming natural products (Fig. 1a). Compound **1** is a 15-residue peptide produced by the soil bacterium *Bacillus brevis*,^{7,8} and **5** is a 48-residue peptide isolated from the marine sponge *Theonella swinhonis*.⁹ Although their side chains and molecular sizes are disparate and their natural sources are unrelated, they share a prominent structural feature that only appears in a few natural products. The linear sequences of **1** and **5** comprise alternating D- and L-amino acids except for the achiral glycines. Due to the switching C_α-stereochemistry, all the side chains are on one side of the strand, which forces the sequence into a helix by inter-residue hydrogen bonding. Namely, both peptides fold into a β^{6,3}-helix with a 4 Å diameter pore in a hydrophobic environment (Fig. 1b).¹⁰⁻¹⁶ As a result, **1** and **5** form transmembrane ion channels as a head-to-head dimer and as a monomer, respectively, and allow for the facile diffusion of monovalent cations. The cation selectivities of **1** and **5** are consistent (H⁺ > K⁺ > Na⁺), indicating their similar channel properties.^{17,18} The channel formation of **1** and **5** is attributed to their potent activities against cancer cells: the 50% inhibitory

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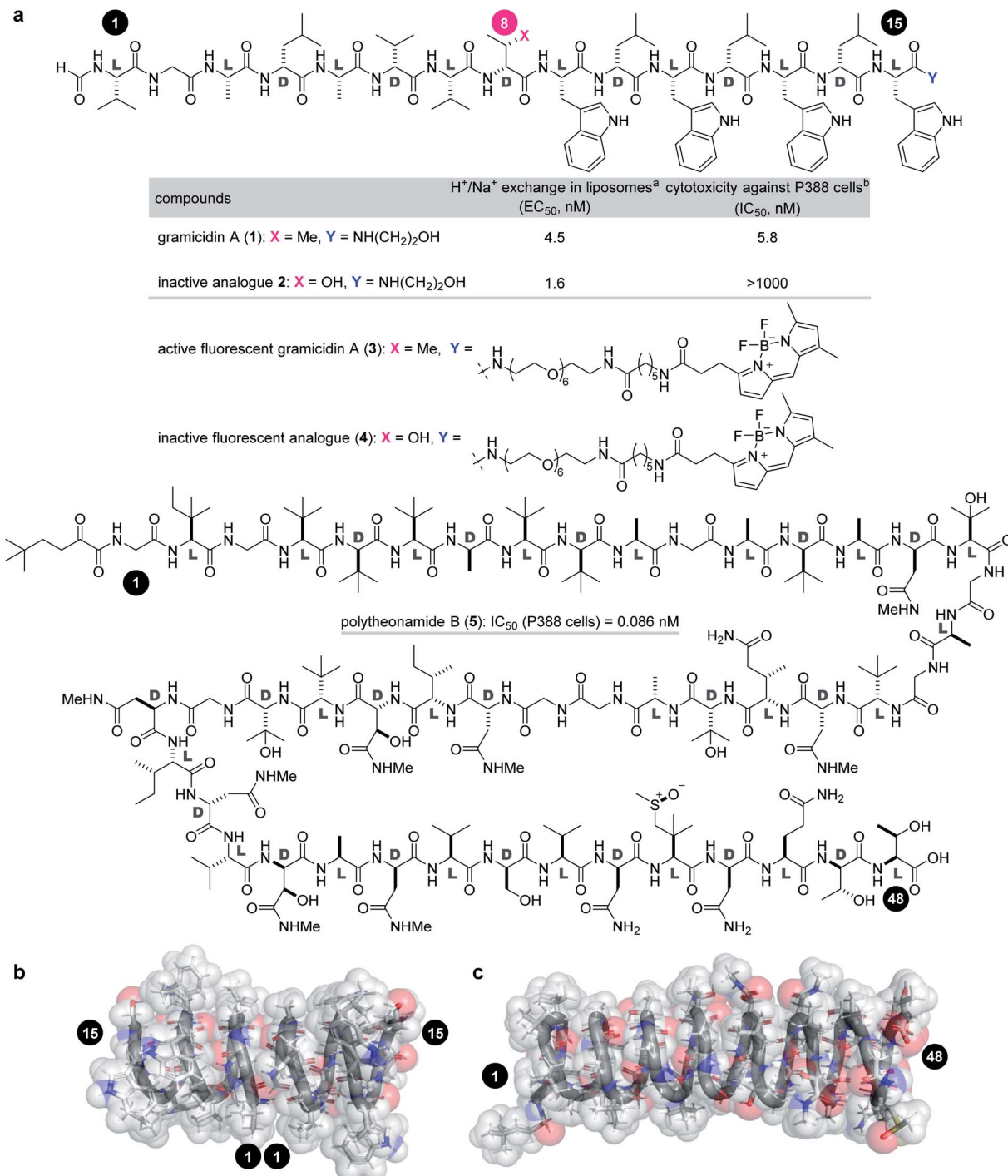


Fig. 1 Structures and functions of channel-forming peptidic natural products. (a) Structures of gramicidin A (1), inactive analogue 2, active fluorescent probe 3, inactive fluorescent probe 4, and polytheonamide B (5). (b) Head-to-head $\beta^{6,3}$ -helical dimer structure of 1 (PDB ID 1MAG). (c) $\beta^{6,3}$ -Helical structure of 5 (PDB ID 2RQO). Numbers highlighted in filled circles are residue numbers. ^a EC_{50} value of H^+/Na^+ transport activity toward pH-gradient liposomes comprising egg yolk phosphatidylcholine and egg yolk phosphatidylglycerol (19 : 1). ^b IC_{50} value against the P388 mouse leukemia cell line.

concentrations (IC_{50}) against P388 mouse leukemia cells are 5.8 nM for 1 and 0.086 nM for 5.

We have engaged in the synthesis, biological evaluation, and functional modulation of channel-forming natural products over the past decade. In 2018, we reported the full solid-phase

total synthesis of polytheonamide B (5).^{19–21} The efficient synthesis allowed us to uncover a unique dual mode of action of 5 in MCF-7 human breast cancer cells. Compound 5 elicits free cation transport in the plasma membrane, and then localizes in the lysosomes to diminish the H^+ gradient across the lysosomal



membranes. Depolarization across the plasma membrane and neutralization of the lysosomes are considered to trigger the apoptotic cell death pathway.

We became interested in clarifying whether gramicidin A (**1**) would emulate the dual mode of action of structurally- and functionally-related **5**. Although potent antitumor effects of **1** have been demonstrated by *in vitro* assays using cancer cells and by *in vivo* experiments using a xenograft mouse model,^{22–27} the precise time-dependent mechanisms of the activity of **1** remained unclear. Herein, we report our investigation of the spatiotemporal cellular behavior of **1** using MCF-7 cells. Compound **1** depolarized not only the plasma membrane, but also the inner mitochondrial membrane after accumulating in the mitochondria. The resultant loss of the mitochondrial transmembrane potential attenuated the production of adenosine triphosphate (ATP), which led to eventual cell cycle arrest, ultimately resulting in the inhibition of cancer cell growth. These results revealed a new intricate mode of action of **1** and the functional differences between **1** and **5** for the first time.

Results and discussion

Preparation of **1** and chemical probes **2–4**

Investigation of the biological behavior of **1** required suitable chemical probes. We first selected an inactive analogue of **1** to differentiate **1**-specific and **1**-nonspecific functions in the assays. During our previous high-throughput screening of thousands of synthetic gramicidin A analogues, we discovered various compounds with altered activities.^{28,29} Among them, **2** was found to be inactive against P388 cells (>1000 nM for **2**, Fig. 1a). Despite its negligible toxicity, **2** differs from **1** only in one substituent at residue-8 (C_{β} -Me for **1**, C_{β} -OH for **2**) and retains the ion-channel activity against liposomes ($EC_{50} = 4.5$ nM for **1**, 1.6 nM for **2**). Because of its similar physico-chemical properties with **1**, analogue **2** was expected to serve as a suitable probe for negative control experiments. Hence, pure **1** and **2** were prepared by Fmoc-based solid-phase peptide synthesis (SPPS).^{28,30}

To analyze the cellular uptake and distribution by live cell imaging, fluorescent probes **3** and **4** were designed based on active **1** and inactive **2**, respectively (Fig. 1a). The C-terminus was chosen as the modification site rather than the N-terminus so as not to disturb the head-to-head dimer formation. Moreover, a hydrophobic fluorophore and linker were selected to retain the intrinsic lipophilic properties of **1** and **2**. Therefore, 4,4-difluoro-4-bora-3a,4a-diaza-s-indacene (BODIPY)³¹ was envisioned to be conjugated with the C-termini of **1** and **2** through a tether consisting of polyethylene glycol (PEG)-6 and C6-alkyl chains. Compounds **3** and **4** were synthesized by SPPS, followed by C-terminal modification.

Scheme 1 illustrates the routes to **3** and **4**. The 15-mer peptides were elongated from Fmoc-L-Trp(Boc)-trityl-ChemMatrix resin (**6**)³² via cycles of piperidine-mediated Fmoc removal and microwave-assisted amidation^{33–35} at 60 °C or 40 °C using N_{α} -Fmoc-protected amino acids in the presence of *O*-(7-aza-1*H*-benzotriazol-1-yl)-*N,N,N',N'*-tetramethyluronium

hexafluorophosphate (HATU) and 1-hydroxy-7-azabenzotriazole (HOAt).³⁶ The N-terminus was then deprotected and formylated with *p*-nitrophenyl formate to afford **7a** and **7b**. A mixture of $(CF_3)_2CHOH$ and CH_2Cl_2 cleaved **7a** and **7b** from the resin, thereby releasing **8a** and **8b**.

Next, the linker and fluorophore were introduced in a stepwise fashion. Carboxylic acids **8a** and **8b** were condensed with mono-Boc-protected diamine **9** by the action of 3-(diethoxyphosphoryloxy)-1,2,3-benzotriazin-4(3*H*)-one (DEPBT)³⁷ and 2,4,6-collidine to generate **10a** and **10b**, respectively. The five Boc groups of **10a**, and the five Boc groups and one *t*-Bu group of **10b** were simultaneously removed by treating with trifluoroacetic acid (TFA), *i*-Pr₃SiH,³⁸ and H_2O to furnish **11a** and **11b**. Finally, BODIPY moiety **12** was condensed with the amine of **11a** and **11b** using (benzotriazol-1-yloxy) tripyrrolidinophosphonium hexafluorophosphate (PyBOP)³⁹ and HOAt, thereby giving rise to fluorescent probes **3** and **4** in 10% and 17% yields, respectively, over 34 steps from **6**.

Growth inhibition activities of **1** and chemical probes **2–4** against cancer cells

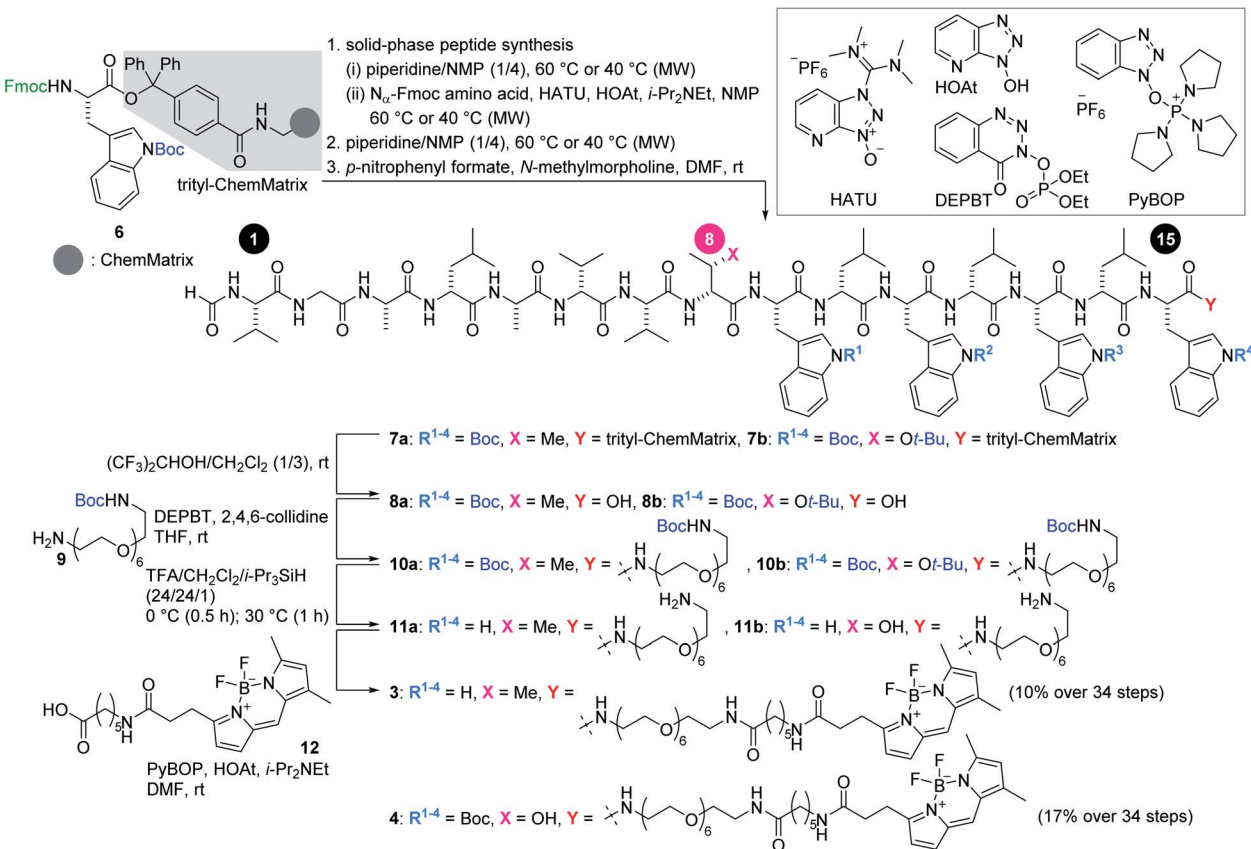
We applied **1** to JFCR39 cancer panel screening to gain preliminary insight into the mode of action (Fig. S1†).^{40,41} Among the 39 cancer cell lines of the panel, BSY-1 and MCF-7 (breast), U-251 (central nervous system), and DMS114 (lung) were the most susceptible strains. To our surprise, a COMPARE analysis of the growth inhibition fingerprints of **1** and **5** (Fig. S2†) revealed no correlation between them ($R = -0.175$).⁴² These results indicated that the mechanism of action of **1** completely differs from that of **5**.

Because of its high sensitivity to **1**, MCF-7 cells were chosen for further study. As an initial step, the activities of **1–4** against MCF-7 cells were assessed by quantifying the 50% growth inhibition concentration (GI_{50}) and 50% lethal concentration (LC_{50}) values (Table 1).⁴³ Interestingly, parent **1** exhibited potent growth inhibition against MCF-7 cells ($GI_{50} = 42$ nM), but was not lethal up to 5000 nM ($LC_{50} > 5000$ nM). Thus, **1** is strong as a cytostatic agent, yet weak as a cytotoxin.⁴⁴ In sharp contrast, the GI_{50} and LD_{50} values of **5** were different only by 19-fold ($GI_{50} = 0.69$ nM vs. $LC_{50} = 13$ nM), demonstrating the high cytotoxic activity of **5** and the distinct effects between **1** and **5**. The growth inhibition activity ($GI_{50} = 990$ nM) of **2** was attenuated 24-fold compared with **1**, and no cytotoxicity ($LC_{50} > 5000$ nM) was observed. These data validated that **2** functioned as a structurally similar inactive analogue. Fluorescent probes **3** ($GI_{50} = 170$ nM, $LC_{50} > 3000$ nM) and **4** ($GI_{50} = 2800$ nM, $LC_{50} > 5000$ nM) maintained the activity profiles of **1** and **2**, respectively, suggesting that the cellular behavior of **3/4** reflect those of **1/2**.

Depolarization of the plasma membrane by gramicidin A (**1**) and inactive analogue **2**

To uncover the origin of the strong cytostatic activity of gramicidin A (**1**), we first analyzed the depolarizing effect of active **1** and inactive **2** on the plasma membrane of MCF-7 cells.⁴⁵ DiBAC₄(3)⁴⁶ was adopted as a fluorescent reporter in this assay to evaluate the time-dependent depolarization (Fig. 2a). As





Scheme 1 Solid-phase synthesis of fluorescent probes **3** and **4**. Boc = *tert*-butoxycarbonyl, DEPBT = 3-(diethoxyphosphoryloxy)-1,2,3-benzotriazin-4(3*H*)-one, HATU = *O*-(7-aza-1*H*-benzotriazol-1-yl)-*N,N,N',N'*-tetramethyluronium hexafluorophosphate, HOAt = 1-hydroxy-7-azabenzotriazole, MW = microwave, PyBOP = (benzotriazol-1-yloxy)tritypyrrolidinophosphonium hexafluorophosphate, *t*-Bu = *tert*-butyl, TFA = trifluoroacetic acid.

Table 1 Cytostatic and cytotoxic activities of **1–5** against MCF-7 cells

Compounds	GI ₅₀ (nM) ^a	LC ₅₀ (nM) ^a
1	42 ± 2	>5000
2	990 ± 130	>5000
3	170 ± 20	>3000
4	2800 ± 600	>5000
5	0.69 ± 0.09	13 ± 2

^a The 50% growth inhibitory concentration (GI₅₀, nM) and 50% lethal concentration (LC₅₀, nM) were determined by sulforhodamine B assay. Values are displayed as mean ± SD of three independent experiments.

expected, natural product **1** rapidly enhanced fluorescence over 1 h at concentrations between 12.3 nM to 333 nM (Fig. 2b). The 50% effective concentration (EC₅₀) was calculated to be 82 nM (Table 2). The comparable EC₅₀ and GI₅₀ (42 nM) values supported the well-accepted mode of action of the channel forming **1**: disruption of the ion concentration gradient across the plasma membrane impaired the cellular functions. Alternatively, when inactive probe **2** was added to MCF-7 cells, no significant fluorescence change was observed up to 3000 nM (EC₅₀ > 3000 nM) (Fig. 2b and Table 2). Considering the similar

ion transport activities of **1** and **2** in the liposome assay (see Fig. 1a), inhibition of the ion-conducting channels of **2** in the MCF-7 cell assay is attributable to the different membrane environment between liposomes and MCF-7 cells. The data also disclosed that the subtle structural switch of the side chain of **1** and **2** resulted in the drastic functional change.

Mitochondrial and lysosomal localization of fluorescent probes **3** and **4** and mitophagy induction by **1** and **3**

The cellular distribution was visualized by imaging studies of fluorescent probes **3** and **4** by confocal fluorescence microscopy (Fig. 3). When 50 nM of active probe **3** and inactive probe **4** were individually incubated with MCF-7 cells, **3** and **4** differentially localized in the intracellular structures.⁴⁷ To determine the selective subcellular localization of **3** and **4**, time-dependent localization analyses were conducted for **3** and **4** using two fluorescent organelle markers; MitoTracker Red CMXROS (Fig. 3a), which covalently labels mitochondria,⁴⁸ and Lysotracker Red DND-99, which accumulates in acidic lysosomes (Fig. 3b). Pearson's correlation coefficient (*R*) was calculated to evaluate the degree of colocalization of **3** and **4** with these markers.^{49,50}

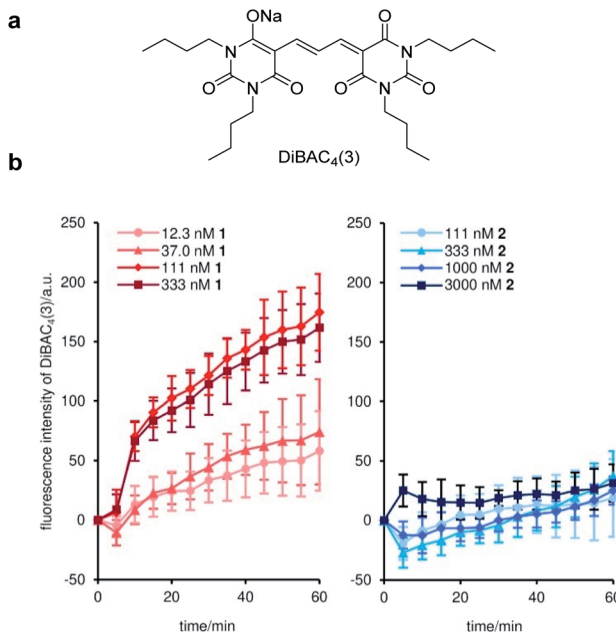


Fig. 2 Time-dependent depolarization of the plasma membrane of MCF-7 cells by 1 and 2. (a) Structure of DiBAC₃(4). (b) Time courses of the changes in the fluorescence intensity of DiBAC₃(4) upon the addition of 1 (left) and 2 (right). Each plot is displayed as mean \pm SD of three replicates. a.u. = arbitrary unit.

Table 2 Plasma membrane depolarizing activity against MCF-7 cells

Compounds	EC ₅₀ (nM) ^a
1	82 \pm 22 ^b
2	>3000

^a The EC₅₀ value was determined from the fluorescence emission of DiBAC₄(3) for 1 h. ^b Values are displayed as mean \pm SD of three independent experiments.

The images of active **3** and inactive **4** in MCF-7 cells were obtained at 4 h, 8 h, and 24 h after incubation in the presence of MitoTracker and Lysotracker. After 4 h, the fluorescence signal of **3** (shown in green) mainly overlapped with that of MitoTracker (Fig. 3c, $R = 0.62$) (shown in red) and partly with that of Lysotracker (Fig. 3d, $R = 0.42$) (shown in red), establishing mitochondria as the major localization site of **3**.⁵¹ Interestingly, the relative distribution of **3** between the mitochondria and lysosomes changed over time. During the 24 h incubation, fluorescence from the mitochondria time-dependently decreased ($R = 0.54$ for 8 h, $R = 0.41$ for 24 h), while fluorescence from the lysosomes increased ($R = 0.40$ for 8 h, $R = 0.65$ for 24 h). On the other hand, the negative control probe **4** selectively localized to the lysosomes (Fig. 3f) rather than the mitochondria (Fig. 3e) in the same experimental time frame ($R = 0.17$ –0.24 with MitoTracker vs. $R = 0.64$ –0.86 with Lysotracker). These imaging data together suggest the importance of mitochondria as an intracellular target site for gramicidin A (**1**) to exert its potent growth inhibition activity. Thus, the

intracellular behavior of **1** should be distinct from that of lysosome-accumulating polytheonamide B (**5**).

Next, we tested if the internalization of **3** and **4** involves an active pathway (Fig. S3 and S4†). Along with glycolysis (2-deoxy-D-glucose) and oxidative phosphorylation (NaN₃) inhibitors, 5-(N-ethyl-N-isopropyl)-amiloride markedly reduced the uptake of **3** and **4** within the cells.⁵² The results confirmed that **3** and **4** were not passively transported but were internalized by energy-dependent macropinocytosis.⁵³

During the imaging experiments (Fig. 3c and d), the green fluorescence from **3** appeared to move from the mitochondria to the lysosomes. We hypothesized that the phenomenon corresponded to the selective autophagy of mitochondria, known as mitophagy.⁵⁴ Mitophagy is an important mitochondrial quality control mechanism that eliminates damaged mitochondria by lysosome-mediated degradation. To examine this hypothesis, we utilized a mitochondria-selective pH-sensitive dye (Mtphagy dye, Fig. 4a).⁵⁵ Mtphagy dye conjugates with mitochondria and shows weak fluorescence under neutral conditions. Upon mitophagy, it becomes highly fluorescent by the acidification of mitochondrial fragments in the lysosomes. Indeed, the red signals from Mtphagy dye were markedly enhanced after 24 h incubation of **3** (50 nM) and well-merged with the green signal from **3** (Fig. 4b, $R = 0.62$). When inactive **4** (50 nM) was applied, the fluorescence from the Mtphagy dye was not changed, and did not overlap with the fluorescence from **4** ($R = 0.34$). The imaging data analyses for active **3** and inactive **4** in Fig. 3 and 4 show the selective time-dependent behavior of **3**: **3** first accumulated in the mitochondria and then localized in acidic lysosomes *via* mitophagy.

The observed mitophagy was attributable to the H⁺ transport activity of **3**, as disruption of the electrochemical H⁺ gradient of the inner mitochondrial membrane (IMM) causes mitochondrial malfunction and subsequent mitophagy. To further investigate the correlation between mitochondrial depolarization and mitophagy, the function of the parent natural product **1** was compared with that of carbonyl cyanide *m*-chlorophenylhydrazone (CCCP, Fig. 4a), which is known to disrupt the H⁺ gradient of the IMM.⁵⁶ In this imaging study, green Lysotracker dye and red Mtphagy dye were co-incubated with MCF-7 cells for 24 h in the absence and presence of CCCP or **1**. Whereas the fluorescence signals from Lysotracker and Mtphagy dye barely overlapped ($R = 0.17$) in the negative control experiment (DMSO), treatment with CCCP (10 μ M) caused a significant overlap ($R = 0.57$) after 24 h (Fig. 4c). When gramicidin A (**1**, 100 nM) was used (Fig. 4d), the green and red colors were merged in a time-dependent fashion. Upon 24 h incubation, a high correlation value was obtained ($R = 0.52$), indicating that **1** had a potent mitophagy-inducing effect similar to CCCP. Applying the same concentration of inactive **2** (100 nM, $R = 0.19$, Fig. 4d) did not promote mitophagy after 24 h, further demonstrating that the mitophagy was selectively elicited by active **1**. The results in Fig. 4 strongly indicated that the localization of ion-channel forming **1** to mitochondria and disruption of the H⁺ gradient across the IMM prompted the cellular stress response, resulting in fusion of mitochondrial fragments with lysosomes *via* the mitophagy process.



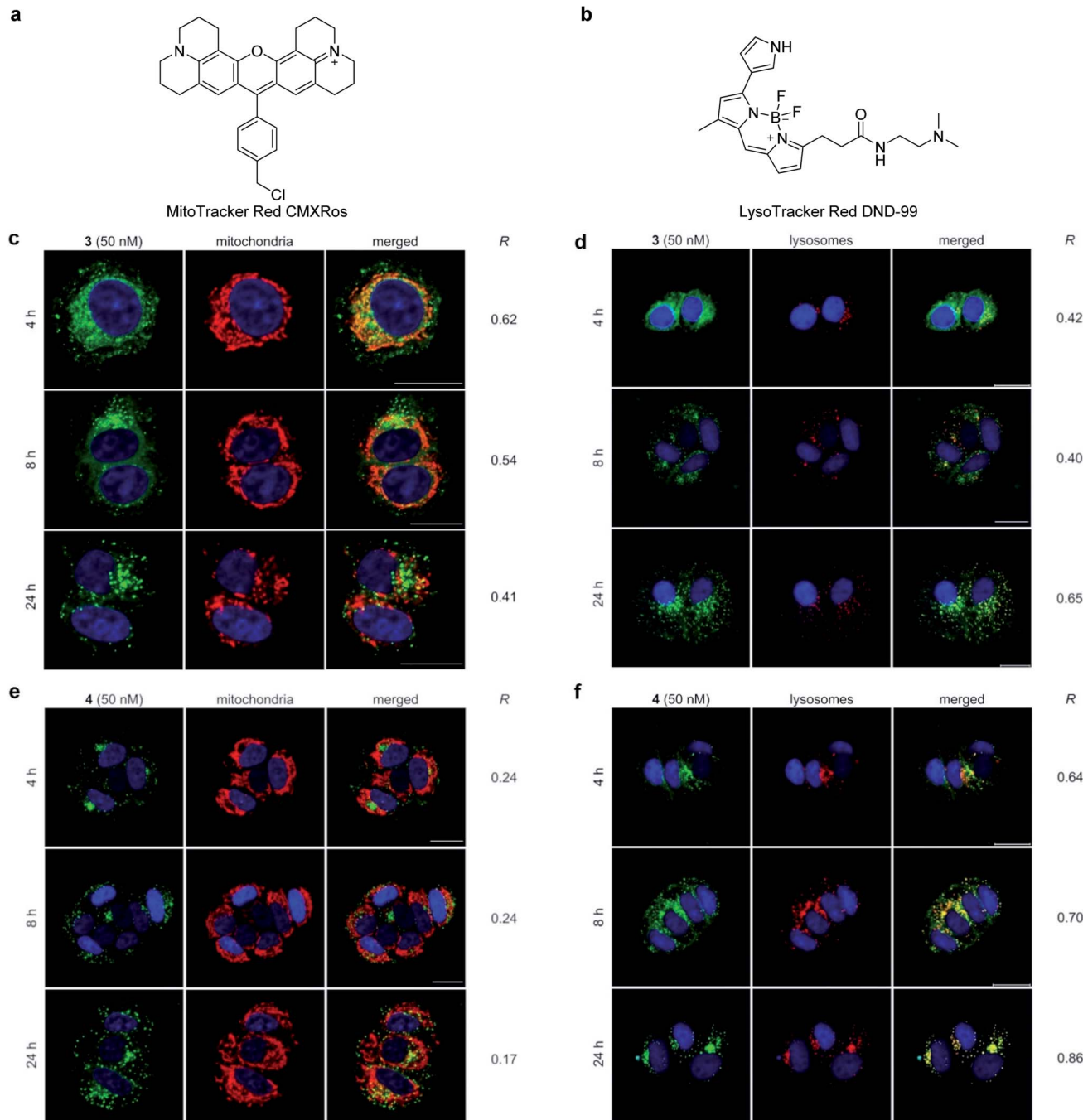


Fig. 3 Time-dependent intracellular localization of fluorescent probes **3** and **4** in the presence of organelle markers in MCF-7 cells. (a) Structure of a mitochondrial marker, MitoTracker Red CMXRos. (b) Structure of a lysosomal marker, Lysotracker Red DND-99. (c) **3** with MitoTracker. (d) **3** with Lysotracker. (e) **4** with MitoTracker. (f) **4** with Lysotracker. The signals from the probes and organelle markers are green and red, respectively. Colocalization of the probes and organelle markers is shown as the merged images. The nuclei were stained with Hoechst 33342. The probes (50 nM) were incubated for 4 h, 8 h, and 24 h. Scale bar represents 20 μ m. *R* value represents Pearson's correlation coefficients calculated from the green and red fluorescence in each merged image.

Mitochondrial depolarization effects of gramicidin A

Disruption of the H^+ gradient across IMM of MCF-7 by **1** was quantified in separate experiments using tetramethylrhodamine methyl ester (TMRM, Fig. 5a). Cationic TMRM accumulates in the negatively polarized IMM and disperses from the

mitochondria upon depolarization, thereby acting as a fluorescent reporter of the H^+ gradient. Specifically, the TMRM-stained area per cell, which is calculated by image analysis, is proportional to the degree of the retained H^+ gradient. When active **1** (100 nM) was employed, the H^+ gradient gradually decreased

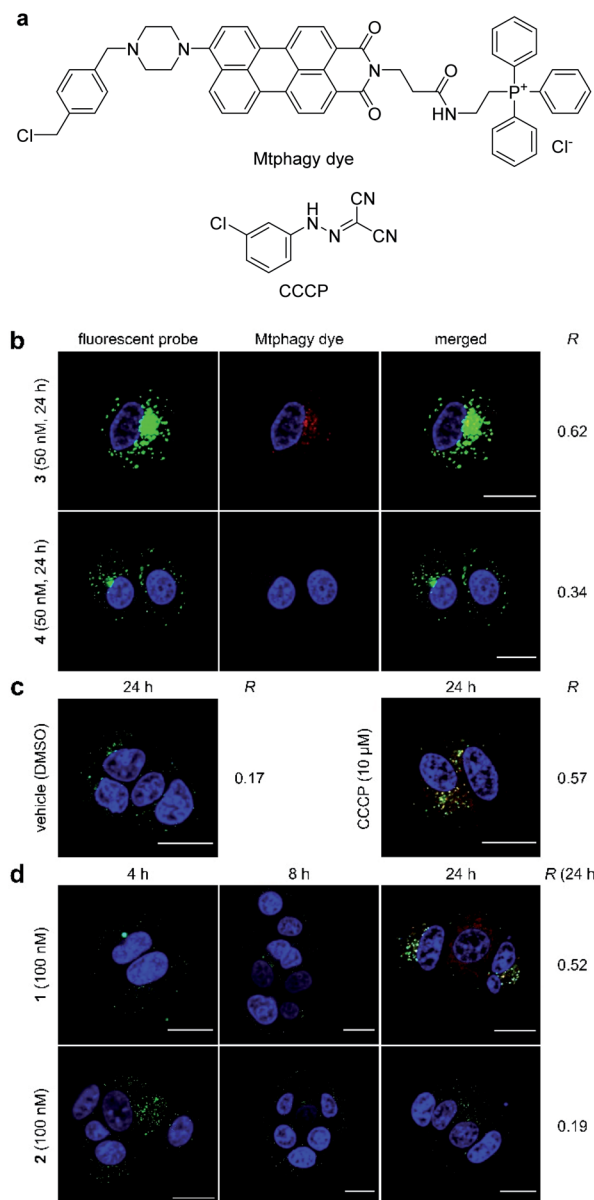


Fig. 4 Mitophagy induced by **1–4**. (a) Structures of Mtphagy dye and carbonyl cyanide *m*-chlorophenylhydrazone (CCCP). (b) Confocal fluorescent images obtained using fluorescent probe **3** or **4** (50 nM) in the presence of Mtphagy dye (red) after 24 h incubation with MCF-7 cells. (c) Confocal fluorescent images obtained using Mtphagy dye (red) and lysosome-selective dye (Lyso dye, green). Vehicle (DMSO) and CCCP (10 μ M) were added and incubated for 24 h. (d) Confocal fluorescent images obtained using Mtphagy dye and Lyso dye after incubation with **1** or **2** (100 nM) for 4, 8, and 24 h. Subcellular regions colored in yellow in the images indicate colocalization of the Mtphagy dye and synthetic probes (b) or Lyso dye (c, d). The nuclei were stained with Hoechst 33342. Scale bar represents 20 μ m. *R* value represents Pearson's correlation coefficients calculated from the green and red fluorescence in each merged image.

during the 24 h incubation (Fig. 5b, red), strongly suggesting the H^+ channel activity of **1** in the IMM. On the other hand, inactive **2** did not depolarize the IMM (Fig. 5b, cyan), even though the same concentration was added. Since **1** decreased

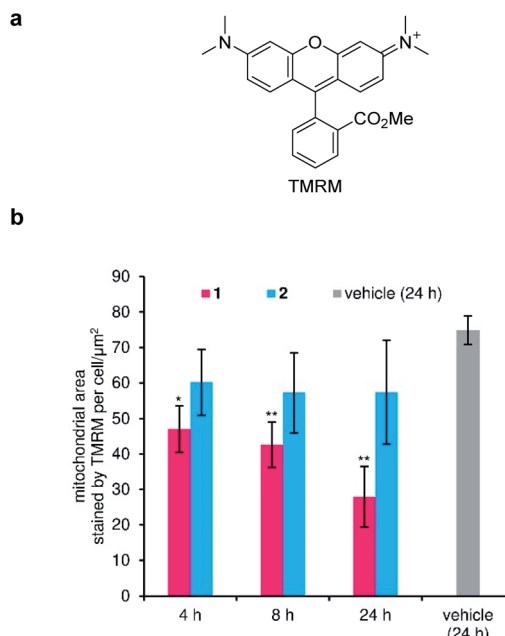


Fig. 5 Depolarization activity of the inner mitochondrial membrane (IMM) by **1**. (a) Structure of tetramethylrhodamine methyl ester (TMRM). (b) Time-dependent change in the fluorescence from TMRM. Compounds (100 nM) were added and incubated for 4, 8, and 24 h. Each plot was made from three images as mean \pm SD. Asterisks indicate significant differences compared with vehicle control (DMSO, 24 h) by Dunnett's test. * P < 0.05, ** P < 0.01.

the H^+ gradient around its GI_{50} value (82 nM), the mitochondrial depolarization effect of **1** must associate with its potent cytostatic activity. Here, we revealed that ion-channel forming **1** depolarized not only the plasma membrane, but also the mitochondrial membrane.^{57–61}

Inhibition of ATP production in mitochondria by gramicidin A

Mitochondria fuel cellular function through the synthesis of ATP by oxidative phosphorylation. The H^+ gradient across the IMM creates potential energy, which is utilized by F_0F_1 -ATP synthase to produce ATP.⁶² As depolarization of the IMM by **1** was expected to decrease ATP synthesis, we next analyzed perturbation of the mitochondrial ATP production rate by **1**. In this assay, MCF-7 cells were permeabilized with digitonin to give a solution of intact mitochondria, which was applied to an enzymatic ATP-detection system.^{63,64} As a positive control, we adopted oligomycin A, an inhibitor of F_0F_1 -ATP synthase (Fig. 6a).⁶⁵ Under these conditions, oligomycin potently decreased the ATP concentration with an EC_{50} value of 1.3 ± 0.5 nM (black line, Fig. 6b). Alternatively, compound **1** inhibited ATP production in a dose-dependent manner (pink line), and the EC_{50} was calculated to be 67 ± 5 nM.⁶⁶ This value was comparable to the required concentrations for the IMM depolarization (100 nM), mitophagy-induction (100 nM), and cytostatic activity (42 nM), corroborating the interconnection of these series of events.

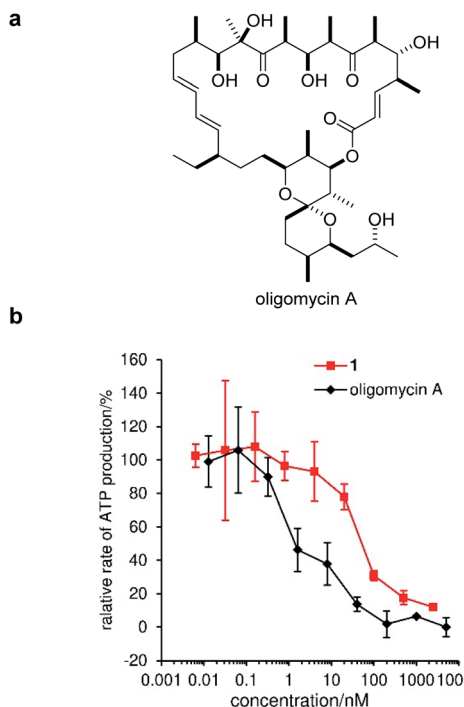


Fig. 6 Inhibition of ATP production by **1** using membrane-permeabilized MCF-7 cells. (a) Structure of oligomycin A. (b) Concentration-dependent inhibition of ATP production. The ATP production rate was normalized against the rates obtained by adding vehicle (DMSO) as 100% and 5 μ M oligomycin A as 0%. Each plot is displayed as mean \pm SD of three replicates. The EC₅₀ values of **1** and oligomycin A were calculated as 67 ± 5 nM (pink) and 1.3 ± 0.5 nM, respectively, by sigmoidal curve fittings.

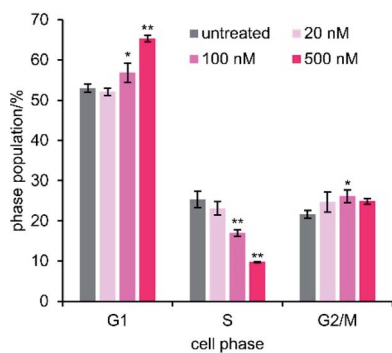


Fig. 7 Cell cycle analysis. MCF-7 cells were treated with **1** for 48 h. The values are displayed as mean \pm SD of three independent experiments. Asterisks indicate significant differences compared with an untreated control by Dunnett's test. * $P < 0.05$, ** $P < 0.01$.

Cell cycle arrest by gramicidin A

Since ATP production in MCF-7 cells consists of 80% oxidative phosphorylation,⁶⁷ mitochondrial malfunction would suppress cell growth due to the cellular energy depletion.⁶⁸ To decipher the details of the cytostatic activity of **1**, we examined the arrest of cell cycle progression by **1** (Fig. 7).^{69,70} Flow cytometric analysis of the cell population was performed after 48 h treatment of

MCF-7 cells with varied concentrations of **1** (20–500 nM). The proportion of cells in G1 was dose-dependently increased in proportion to the decrease in S. On the other hand, the change in the G2/M phase was smaller than that in G1 and S phases. Because it is known that an energy-sensitive check exists at the G1/S border, these data indicated that the depleted ATP levels induced by **1** did not satisfy the energy requirement for passage through G1 into S phase. The cell cycle arrest at the energetic checkpoint again supported the significance of the depolarization of mitochondria and the ensuing ATP reduction as the mode of action of **1**.

Conclusions

The present study on gramicidin A (**1**) integrated total synthesis, imaging analysis, and functional investigation to decipher the unprecedented spatiotemporal behavior of **1** in MCF-7 cancer cells. We first synthetically prepared 15-mer peptide **1** and its BODIPY-conjugated fluorescent probe **3** as well as inactive analogue **2** and its fluorescent probe **4**. Application of the synthetic **1–4** revealed the multiple cellular functions of **1** that ultimately result in the growth inhibition of MCF-7 cells at a nanomolar concentration. Compound **1** quickly depolarized the plasma membrane within 1 h, and then selectively localized to the mitochondria and reduced the H⁺ gradient of the IMM within 24 h. Consequently, the ion balance across both the plasma and mitochondrial membranes was disrupted, presumably by the facile ion transport through the pore of the head-to-head $\beta^{6,3}$ -helical dimer of **1**. The ion-channel function of **1** in the IMM leads to additional cellular responses: *e.g.*, eradication of the H⁺ gradient decreases the ATP concentration, and the resultant ATP depletion causes cell cycle arrest at the G1 energetic checkpoint. Moreover, the mitochondrial malfunction triggered elimination of the entire organelle by fusion with the lysosomes through mitophagy. Although how **1** achieves selective localization to the mitochondria remains to be elucidated at the molecular level,⁷¹ we uncovered the intricate time-dependent cellular behavior of potent cytostatic agent **1** for the first time. Mitochondria are energy-producing organelles with essential functions in cell biology, and are promising therapeutic targets for treating cancer.^{72,73} Because **1** has unique mitochondrial function and shows no structural similarity with typical mitochondria-targeting compounds,^{74,75} **1** and its derivatives should serve as new platform structures for the development of novel mitochondrial modulators and anti-cancer agents.

While **1**, inactive analogue **2**, and polytheonamide B (**5**) all share the D,L-alternating sequences and function as ion channels, their cellular behaviors are distinct from each other. Despite a difference in only one substituent (Me for **1**, OH for **2**), **2** does not accumulate to the mitochondria and possesses 24-fold weaker cytostatic activity against MCF-7 cells. The 48-mer peptide **5** rapidly diminishes the potential across the plasma membrane, localizes to acidic lysosomes, and neutralizes the lysosomal H⁺ gradient, causing apoptotic cell death of MCF-7 cells. These contrasting behaviors among

the three related peptides are likely programmed by the side chain structures and the overall sequence of their amino acid components. Therefore, the structural and functional differences of **1**, **2**, and **5** offer exciting opportunities for potentially controlling their ion channel function, organelle selectivity, anticancer activity, and even antibacterial activity by structural alterations. Such investigation should lead to the development of tailor-made $\beta^{6,3}$ -helical ion channels for selectively modulating the ion balance of various cells and organelles.

Data availability

The experimental procedures and additional data can be found in the ESI.[†]

Author contributions

H. I. and M. I. conceived and designed the study. Y.-W. X. performed the synthesis of probes and the assays. S. D. performed the cancer panel assay. H. I. and M. I. co-wrote the paper.

Conflicts of interest

There are no conflicts to declare.

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