Rowan University Rowan Digital Works

Stratford Campus Research Day

25th Annual Research Day

May 6th, 12:00 AM

Lipoxin A 4 (LxA 4) Promotes Reduction and Antibiotic Efficacy Against Pseudomonas aeruginosa Biofilm

Julianne M. Thornton Rowan University

Jean Walker *Rowan University*

Prem Y.K. Sundarasivarao Rowan University

Bernd Spur Rowan University

Ana Rodriguez *Rowan University*

Follow this and additional works at: https://rdw.rowan.edu/stratford_research_day See next page for additional authors Part of the Bacterial Infections and Mycoses Commons, Bacteriology Commons, Biological Phenomena, Cell Phenomena, and Immunity Commons, Medical Cell Biology Commons, and the Medicinal and Pharmaceutical Chemistry Commons Let us know how access to this document benefits you - share your thoughts on our feedback form.

Thornton, Julianne M.; Walker, Jean; Sundarasivarao, Prem Y.K.; Spur, Bernd; Rodriguez, Ana; and Yin, Kingsley, "Lipoxin A 4 (LxA 4) Promotes Reduction and Antibiotic Efficacy Against Pseudomonas aeruginosa Biofilm" (2021). *Stratford Campus Research Day*. 52. https://rdw.rowan.edu/stratford_research_day/2021/may6/52

This Poster is brought to you for free and open access by the Conferences, Events, and Symposia at Rowan Digital Works. It has been accepted for inclusion in Stratford Campus Research Day by an authorized administrator of Rowan Digital Works.

Author(s)

Julianne M. Thornton, Jean Walker, Prem Y.K. Sundarasivarao, Bernd Spur, Ana Rodriguez, and Kingsley Yin

Lipoxin A₄ (LxA₄) Promotes Reduction and Antibiotic **ROWAN UNIVERSITY** School of National Institutes of Health Efficacy Against Pseudomonas aeruginosa Biofilm Osteopathic Medicine Turning Discovery Into Health Julianne M. Thornton, Jean Walker, Prem Y. K. Sundarasivarao, Bernd Spur, Ana Rodriguez, Kingsley Yin

Abstract

Pseudomonas aeruginosa (P. aeruginosa) is an opportunistic bacterium commonly found in wound infections and airways of cystic fibrosis patients. P. aeruginosa readily forms biofilms which can reduce the efficacy of antibiotics used to eradicate the pathogen. We have previously shown that a Specialized Pro-resolving Mediator (SPM), Lipoxin A4 (LxA₄) is a quorum sensing inhibitor which can reduce P. aeruginosa virulence. In this study, we examined the direct actions of LxA₄ and RvD₂ on *P. aeruginosa* biofilm formation and virulence gene expression. The influence of LxA₄ on antibiotic efficacy and the combined effects on biofilm formation were also investigated. LxA₄ and RvD₂ reduced P. aeruginosa biofilm formation and virulence gene expression. LxA₄ increased ciprofloxacin inhibition on biofilm formation but did not affect ciprofloxacin's action on non-adherent bacteria. On the other hand, LxA₄ increased bacterial killing action of imipenem but did not affect imipenem's action on biofilm. We also found that LxA₄ can increase ciprofloxacin's bacterial killing ability in established biofilm. Together these results suggest that LxA₄ has direct effects on *P. aeruginosa* biofilm formation and can increase antibiotic efficacy directly.

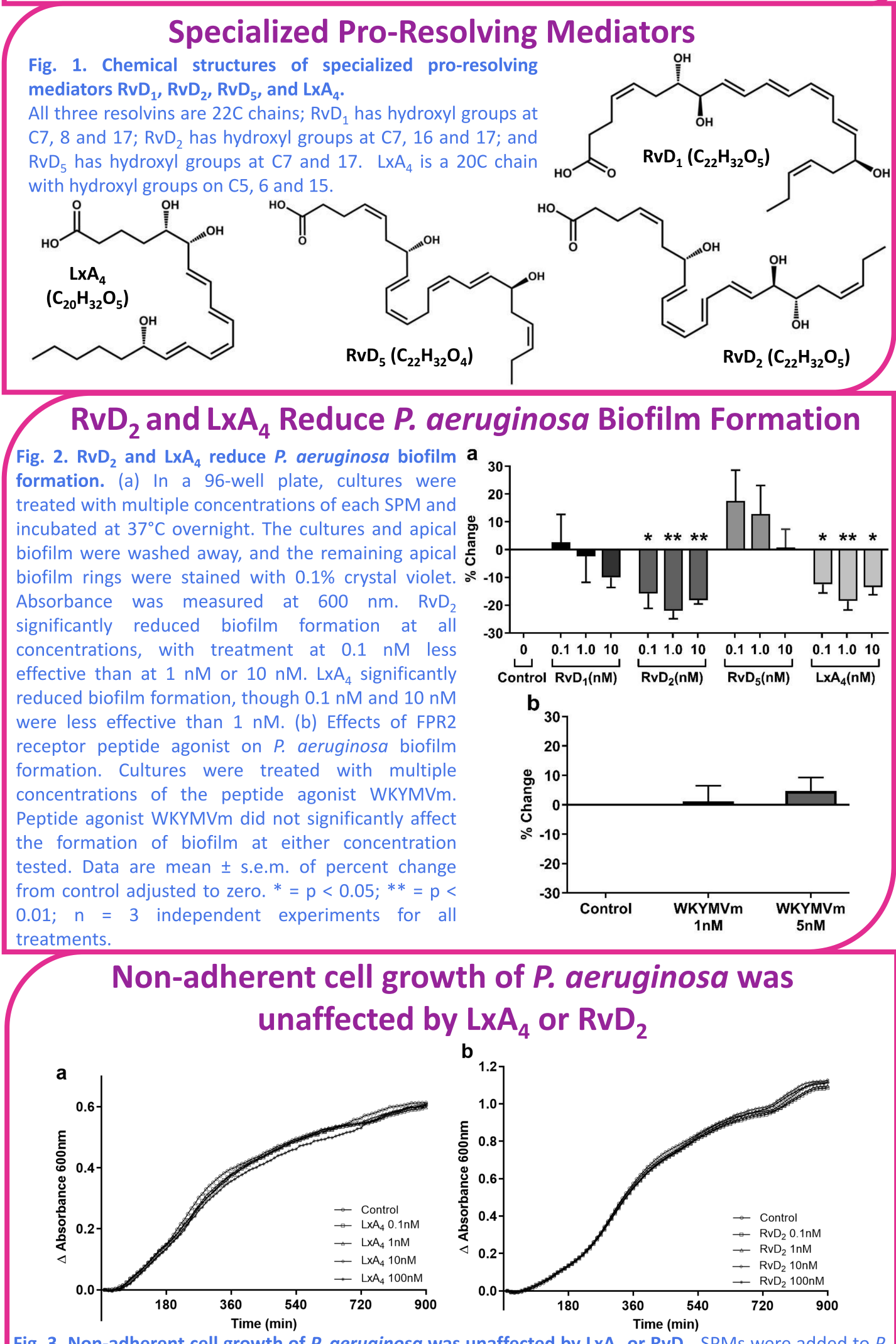
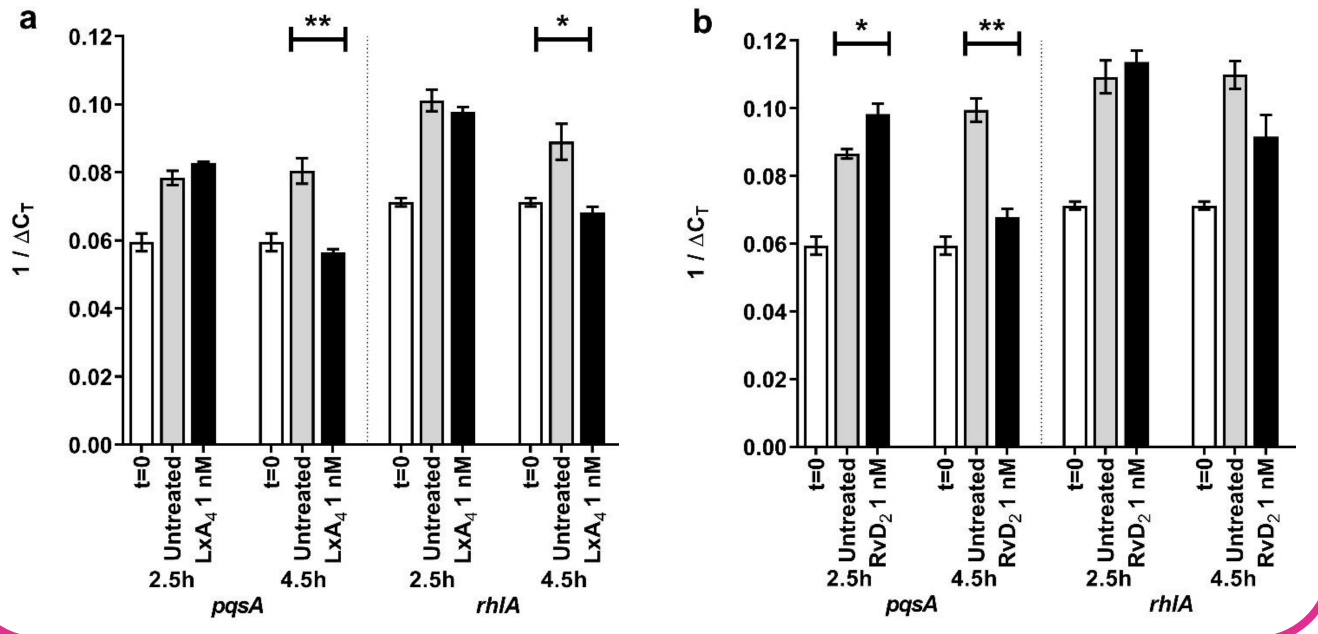


Fig. 3. Non-adherent cell growth of *P. aeruginosa* was unaffected by LxA₄ or RvD₂. SPMs were added to *P.* aeruginosa cultures in 96-well plates and incubated with orbital shaking at 37°C in a microplate reader for 15 hr. Absorbance was measured every 10 min. Neither LxA₄ (a) nor RvD₂ (b) showed any significant effect on non-adherent cell growth in minimal media. n = 3 (LxA₄) and n = 4 (RvD₂) independent experiments.

Rowan University School of Osteopathic Medicine, Stratford, NJ, USA

LxA₄ and RvD₂ Reduce Two **Quorum-sensing Genes' Expression**

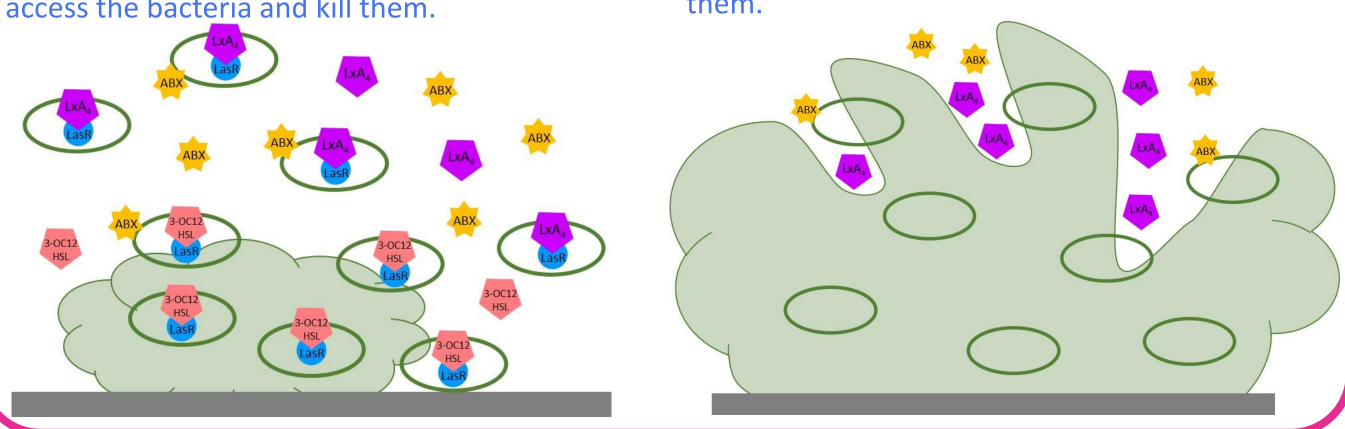
Fig. 4. Effects of SPMs on *Pseudomonas aeruginosa* quorum-sensing virulence gene expression. (a) $1/\Delta CT$ of two virulence genes' expression when treated with 1 nM LxA₄. LxA₄ significantly reduced both genes' expression at 4.5 hr. (b) $1/\Delta CT$ of two virulence genes' expression when treated with 1 nM RvD₂. RvD₂ significantly reduced *pqsA* gene expression at 4.5 hr. Data are mean ± s.e.m., with p-values determined by unpaired t-test comparing untreated control to treatment group of same timepoint. * p < 0.05, ** p < 0.01. n = 3 independent experiments.



Hypotheses: How LxA Can Assist **Antibiotics Against Biofilms**

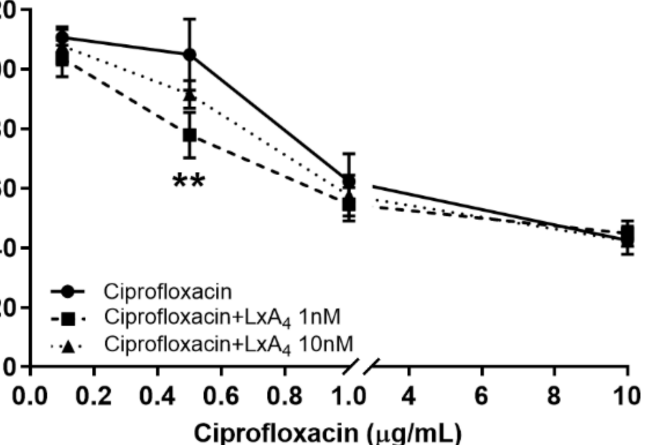
Biofilm Formation: LxA₄ antagonizes the LasR receptor, blocking the quorum-sensing auto-3-OC12 HSL, reducing biofilm inducer formation. Antibiotics (ABX) are then able to access the bacteria and kill them.

Established Biofilm: LxA₄, as a lipid, can integrate with the biofilm matrix, weakening its integrity. This provides an opportunity for antibiotics (ABX) to access the bacteria within the biofilm and kill them.



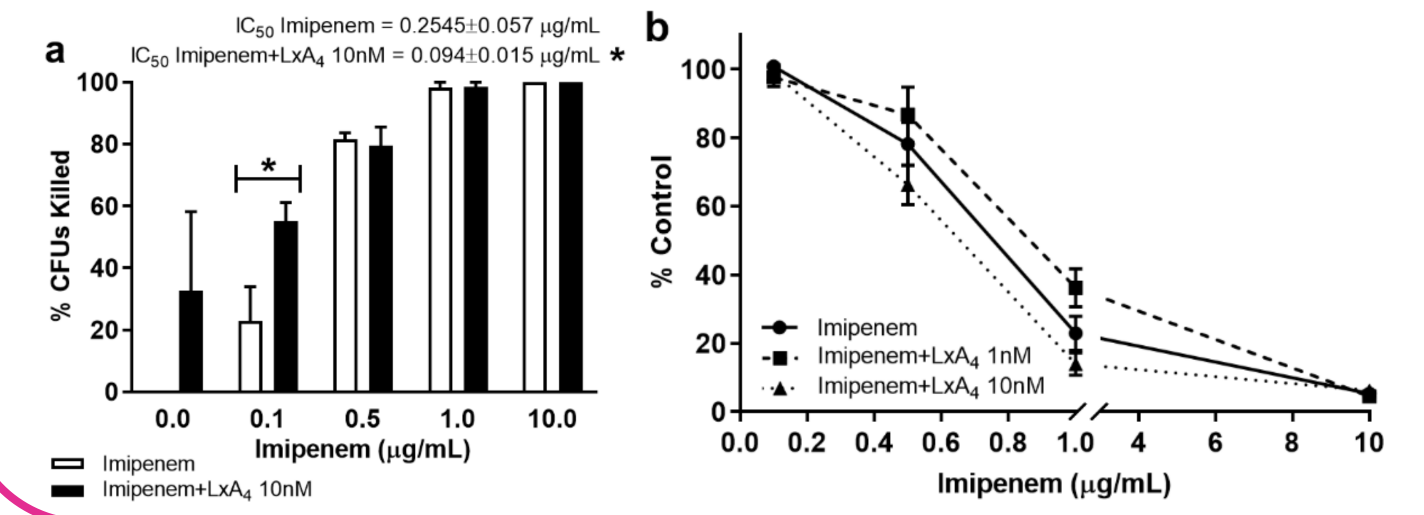
LxA₄ Enhances Ciprofloxacin Effects on Biofilm Formation But Has No Affect on Planktonic Cell Growth IC_{50} Cipro = 0.025 \pm 0.004 µg/mL **b** 6 IC₅₀ Cipro+LxA₄ 1nM = 0.040±0.019 μg/mL 120 г 100-100 80· Killed 60-**4**0 60. FUs **20**· ΰ % 20-Ciprofloxacin -20 Ciprofloxacin+LxA₄ 1nM 0.0 1.0 0.5 Ciprofloxacin (µg/mL)

Fig. 5. LxA₄ treatment can aid the efficacy of the antibiotic ciprofloxacin in reducing biofilm formation. Cultures were treated in a 96-well plate overnight with multiple concentrations of ciprofloxacin and LxA₄. The cultures and apical biofilm were washed away, and the remaining apical biofilm rings were stained with 0.1% crystal violet. Absorbance was measured at 600 nm. Some cultures were recovered from the 96-well plate, diluted and spread on tryptic soy agar plates to incubate overnight at 37°C. Colonies were counted the next day. (a) LxA₄ does not have an effect when combined with ciprofloxacin on non-adherent cell growth, determined by colony forming units (CFUs). (b) When combined with LxA₄, biofilm formation is significantly reduced in ciprofloxacin treatments at and above bactericidal doses (0.5 µg/mL). CFU data are mean ± s.e.m. of percent change from control adjusted to zero. Biofilm data are mean ± s.e.m. percent of control. ** = p < 0.01; CFUs n = 4 independent experiments; biofilm n = 6 (1 nM) and n = 3 (10 nM) independent experiments.



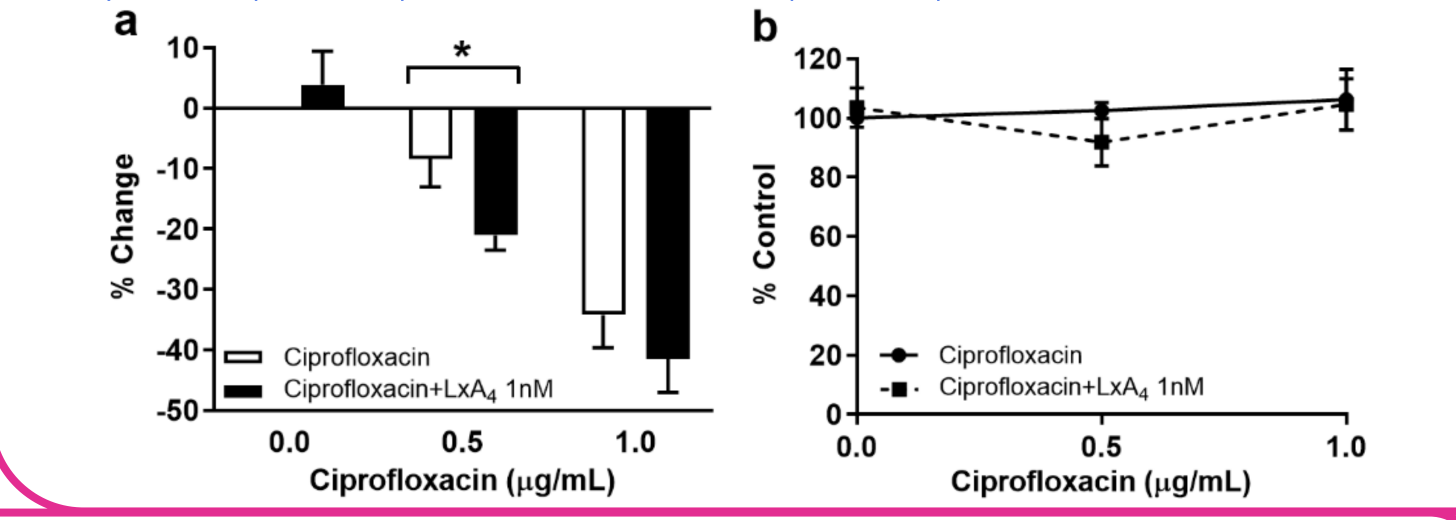
LxA₄ Does Not Affect Imipenem Efficacy **Against Biofilm Formation But Enhances Suppression of Planktonic Growth**

Fig. 6. LxA₄ in combination with imipenem significantly decreased non-adherent cell growth but not biofilm formation. Cultures were treated in a 96-well plate overnight with multiple concentrations of imipenem and LxA₄. The cultures and apical biofilm were washed away, and the remaining apical biofilm rings were stained with 0.1% crystal violet. Absorbance was measured at 600 nm. Some cultures were recovered from a 96-well plate, diluted and spread on tryptic soy agar plates to incubate overnight at 37°C. Colonies were counted the next day. (a) LxA₄ (10 nM) in combination with imipenem (0.1 µg/mL) significantly decreased non-adherent cell growth. (b) LxA₄ does not significantly affect the efficacy of imipenem against biofilm formation. These data combined with the ciprofloxacin data (Fig. 4) suggests the synergistic action of LxA₄ with antibiotics is dependent on the class of antibiotics and their mechanism of action. CFU data are mean ± s.e.m. of percent change from control adjusted to zero. Biofilm data are mean ± s.e.m. percent of control. * = p < 0.05; CFUs n = 4 independent experiments; biofilm n = 3 (LxA₄ 1 nM) and n = 4 (LxA₄ 10 nM) independent experiments.



LxA₄ Can Increase the Efficacy of Ciprofloxacin **Against Bacteria of an Established Biofilm**

Fig. 7. Effects of LxA₄ on ciprofloxacin action in pre-formed biofilms. (a) LxA₄ (1 nM) significantly increased the efficacy of ciprofloxacin (0.5 µg/mL) to reduce the amount of metabolically active bacterial cells associated with the biofilm, determined by MTT assay. (b) LxA_{4} does not significantly alter the efficacy of ciprofloxacin to reduce biofilm biomass, determined by crystal violet staining. This suggests that LxA₄ treatment can aid the ciprofloxacin in accessing the bacteria associated with a pre-formed biofilm. Viability data are mean ± s.e.m. of percent change from control adjusted to zero. Biofilm data are mean ± s.e.m. percent of control. * = p < 0.05; viability n = 5 independent experiments; biomass n = 5 independent experiments.



In summary, our results show that LxA₄ and RvD₂ can directly reduce *P. aeruginosa* biofilm formation. LxA₄ and RvD₂ can also downregulate virulence gene expression. LxA₄ enhances the efficacy of antibiotics directly against P. aeruginosa biofilm formation and bacterial cells within existing biofilm. These studies also provide evidence that further investigation into the antimicrobial mechanisms of RvD₂ is warranted. The results suggest that there is relative selectivity in SPM inhibition of biofilm formation.

Future Directions regulation • 2-hit treatments of SPMs ± antibiotics on established biofilms to assess the effects of SPMs on antibiotic ability to kill biofilm-associated persister cells Pre-treatment of THP-1 monocytes with SPMs against established biofilm

immune cell ability to phagocytose biofilm

We thank Rowan University Graduate School of Biomedical Sciences, New Jersey Health Foundation and the NIH (RO1 Al128202)

Conclusions

• SPM treatment of *P. aeruginosa* to evaluate the effects of SPMs on quorum-sensing gene expression and

• Co-incubation of THP-1 monocytes with SPMs against established biofilm to assess the effects of SPMs on

Acknowledgments