



# mRNA-LNP Transfection of Platelets is Compatible with Current and Alternative Storage Practices

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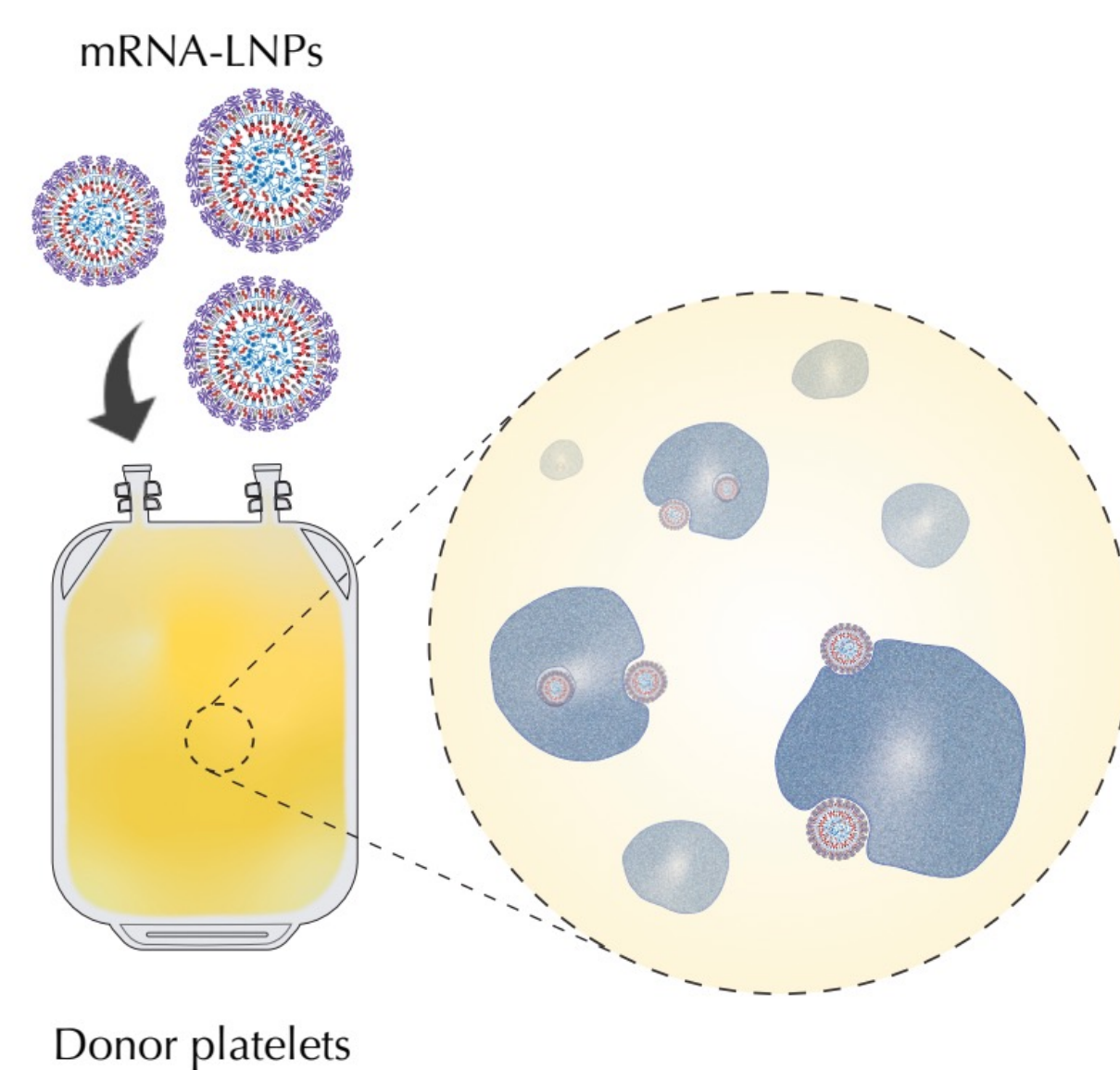
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## INTRODUCTION

Given platelets' integral role in physiology, they have the potential to be utilized as a natural delivery system if enhanced with exogenous cargo. Recently we have demonstrated that optimized lipid nanoparticles (LNPs) can be used to load cargo into platelets. However, this platform was developed using Tyrode's buffer, a crystalloid solution that is not used clinically. We present preliminary data that LNPs can be used to transfect platelets in a setting closer to a clinical environment and is a promising first step for use of enhanced platelets clinically.

## BIG QUESTION

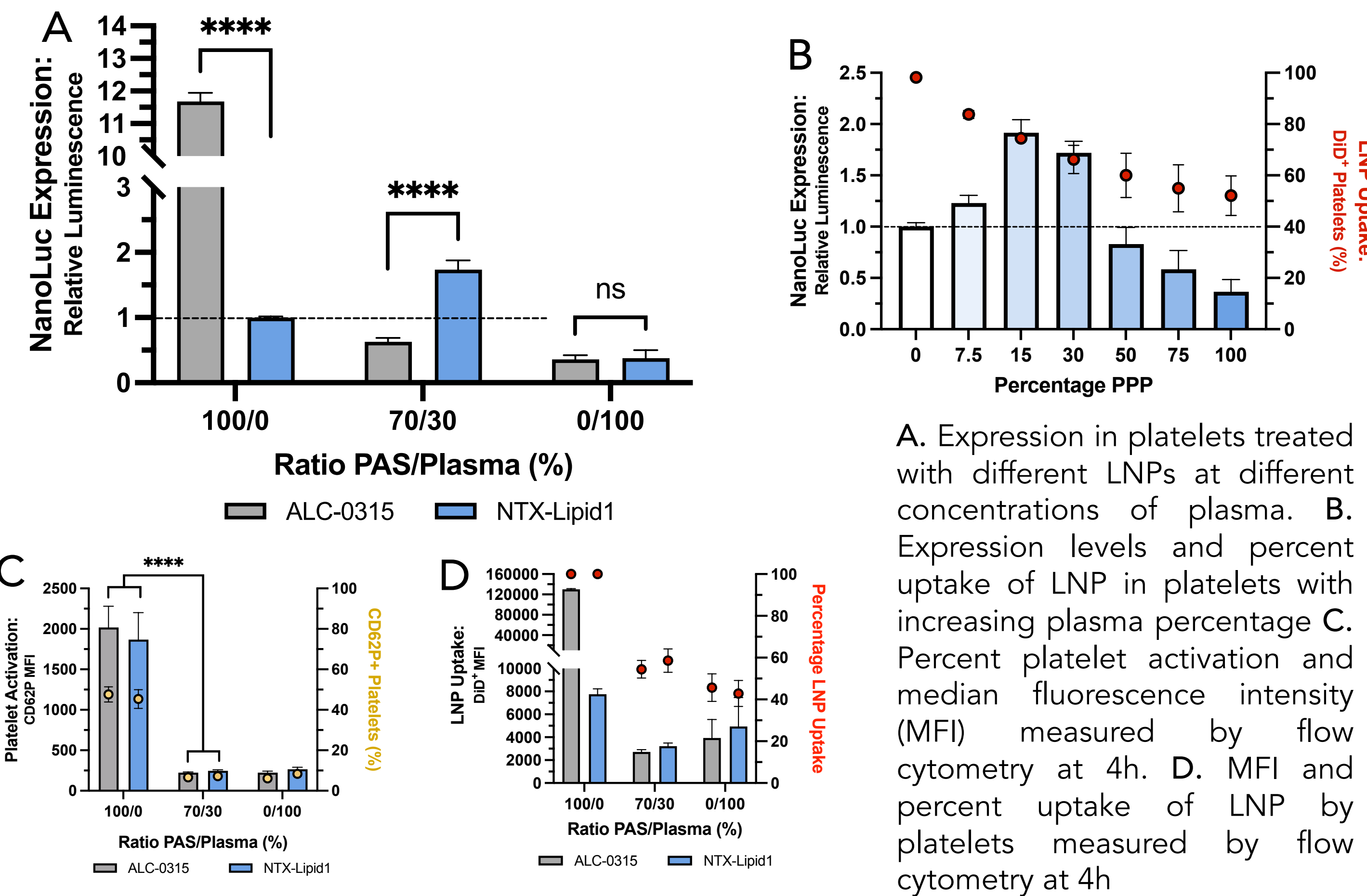
Can we engineer platelets in clinically relevant systems?



## GOALS

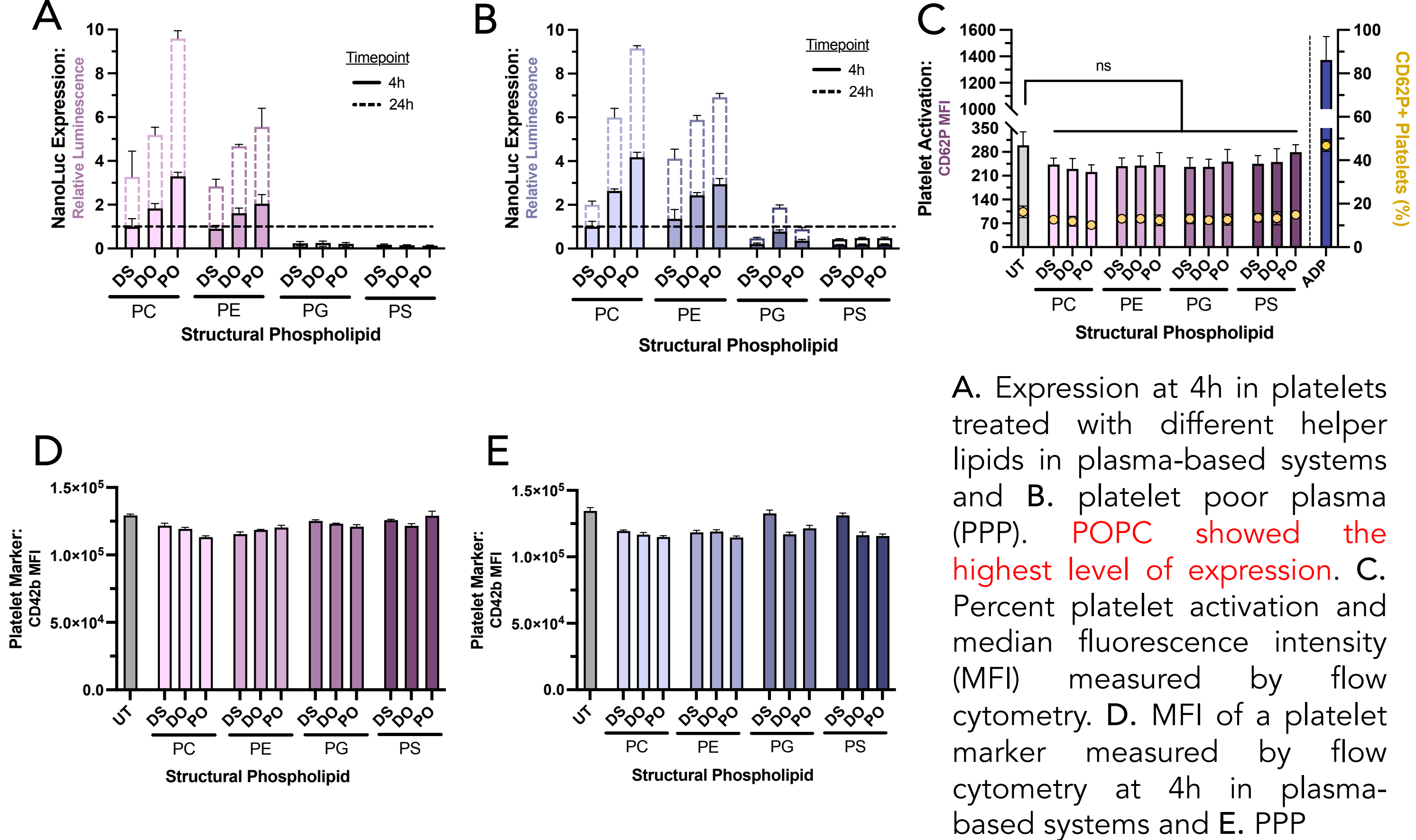
- 1 Screen different lipid nanoparticles formulations for optimal delivery in plasma and plasma-based systems
- 2 Show mRNA-LNP treatments are scalable to high and physiological unit concentrations of platelets
- 3 Characterize mRNA-LNP platelets to show they are responsive to different agonists

## 1 LNPs CAN TRANSFECT PLATELETS IN PLASMA AND PLASMA-BASED SYSTEMS



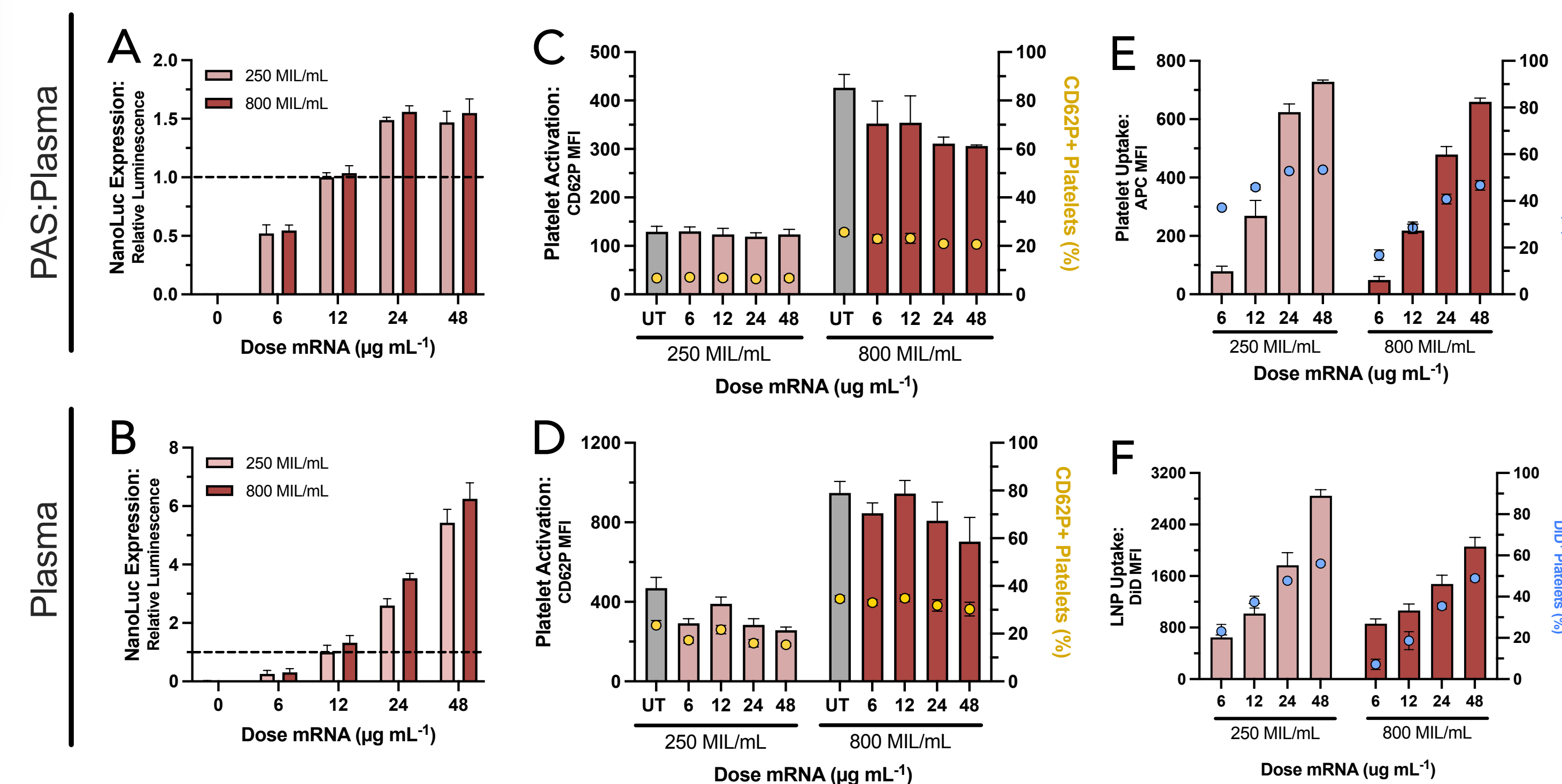
**Key Takeaway:** LNPs can be optimized for transfection of platelets in plasma

## 2 LNPs WITH PC HELPER LIPIDS TRANSFECT PLATELETS



**Key Takeaway:** Transfection in plasma and plasma-based systems requires LNPs formulated with PC helper lipids

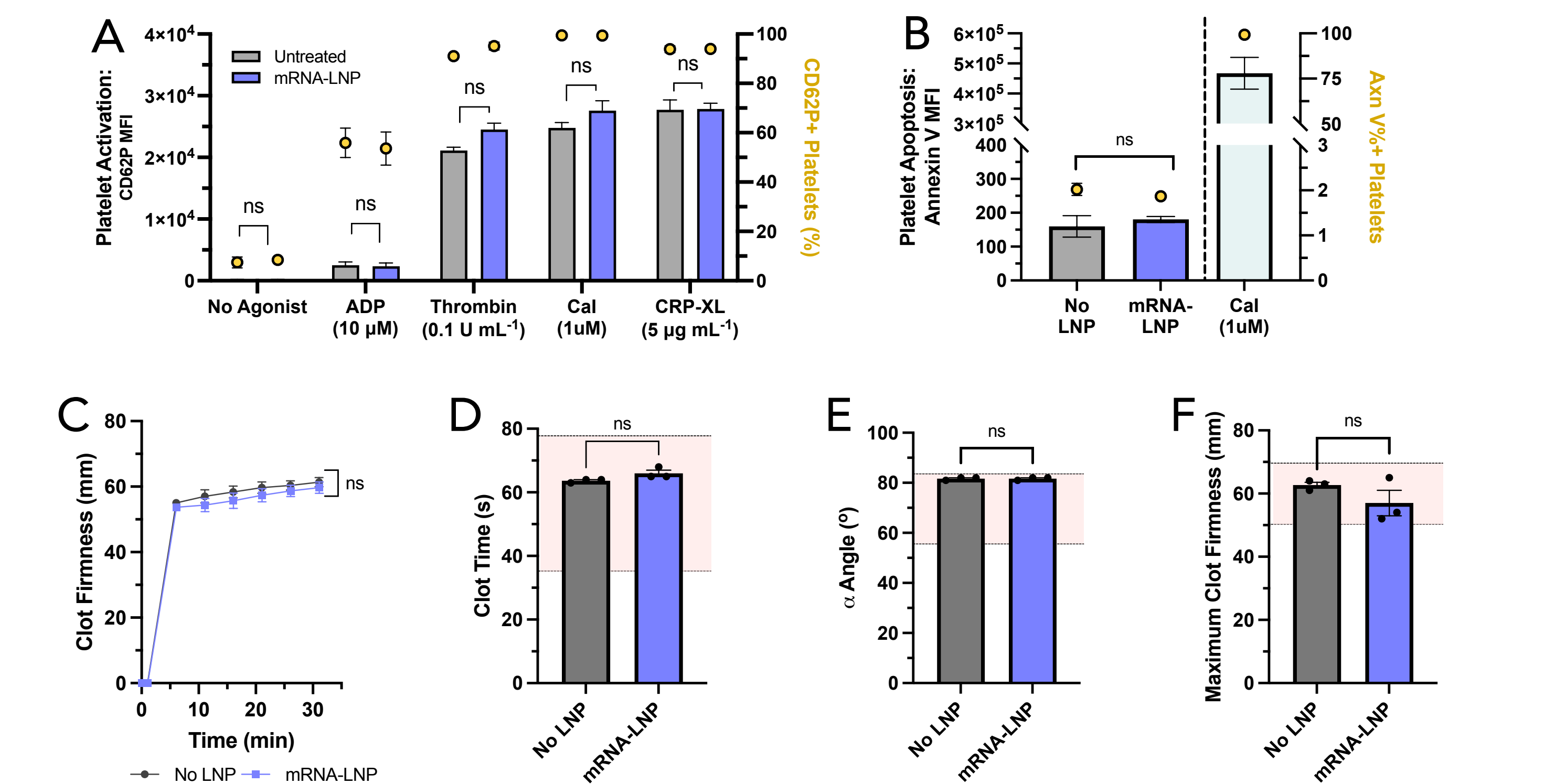
## 3 LNP TREATMENTS ARE SCALABLE



**A.** Expression in platelets treated with different doses of LNP at different concentrations in plasma-based systems and **B.** platelet poor plasma (PPP). **C.** Percent platelet activation and median fluorescence intensity (MFI) measured by flow cytometry in plasma-based systems **D.** in PPP. **E.** Percentage of LNP uptake and MFI measured by flow cytometry in plasma-based systems and **F.** in PPP

**Key Takeaway:** mRNA-LNP transfection is scalable to high and physiological unit concentrations in plasma and plasma-based systems

## 4 LNP TREATED PLATELETS RESPOND TO AGONISTS



**A.** Platelet activation and median fluorescence intensity (MFI) after activation with ADP, thrombin, calcium ionophore (Cal), and CRP-XL measured by flow cytometry. **B.** Platelet apoptosis MFI and percent measured by Annexin V by binding to phosphatidylserine using flow cytometry. **C-F.** ROTEM measurements measuring clot firmness overtime (**C**), clot time (**D**), alpha angle (**E**), and maximum clot firmness (**F**). Red shaded areas represent clinically acceptable ranges

**Key Takeaway:** LNP treated platelets are agonist responsive, and the treatments do not affect their coagulability morphology

## CONCLUSION

LNP formulations can transfect platelets in clinically relevant systems and do not affect their innate responsiveness to agonists

## ACKNOWLEDGMENTS



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