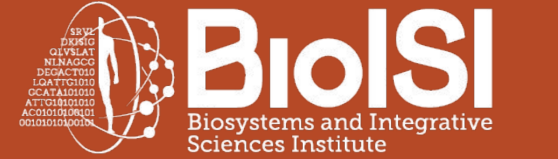
Refining an umbrella-sampling protocol to describe membrane PAINS

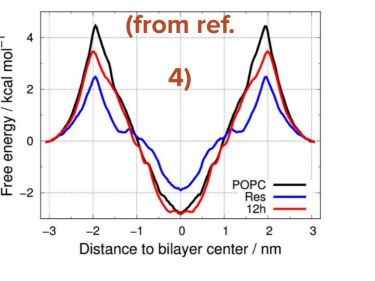




Pedro R. Magalhães, Pedro Reis, Diogo Vila-Viçosa, Miguel Machuqueiro, Bruno L. Victor 🖂 prmagalhaes@fc.ul.pt BioISI: Biosystems and Integrative Sciences Institute; Faculdade de Ciências, Universidade de Lisboa, 1749-016 Lisboa, Portugal

PAINS • Pan assay interference compounds are molecules that show activity in multiple types of assays by interfering with the assay readout rather than through specific compound/target interactions.^{1,2} Membrane PAINS are compounds that interact directly and nonspecifically with lipid membranes, promoting changes in their biophysical properties and affecting the function of mechanosensitive membrane proteins.³

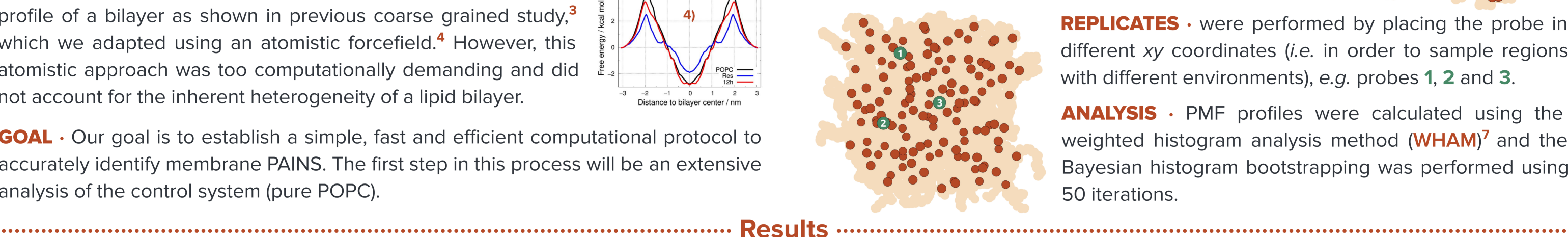
Membrane PAINS will shift the potential of mean force (PMF) profile of a bilayer as shown in previous coarse grained study,³ which we adapted using an atomistic forcefield.⁴ However, this atomistic approach was too computationally demanding and did not account for the inherent heterogeneity of a lipid bilayer.



GOAL • Our goal is to establish a simple, fast and efficient computational protocol to accurately identify membrane PAINS. The first step in this process will be an extensive analysis of the control system (pure POPC).

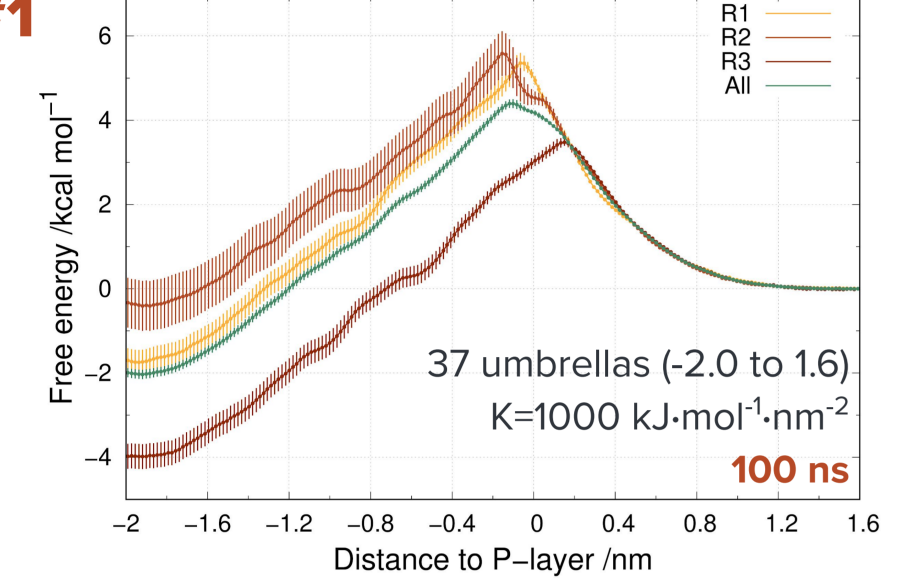
SOFTWARE · GROMACS 2018.6⁵ / GROMOS 54A7⁶

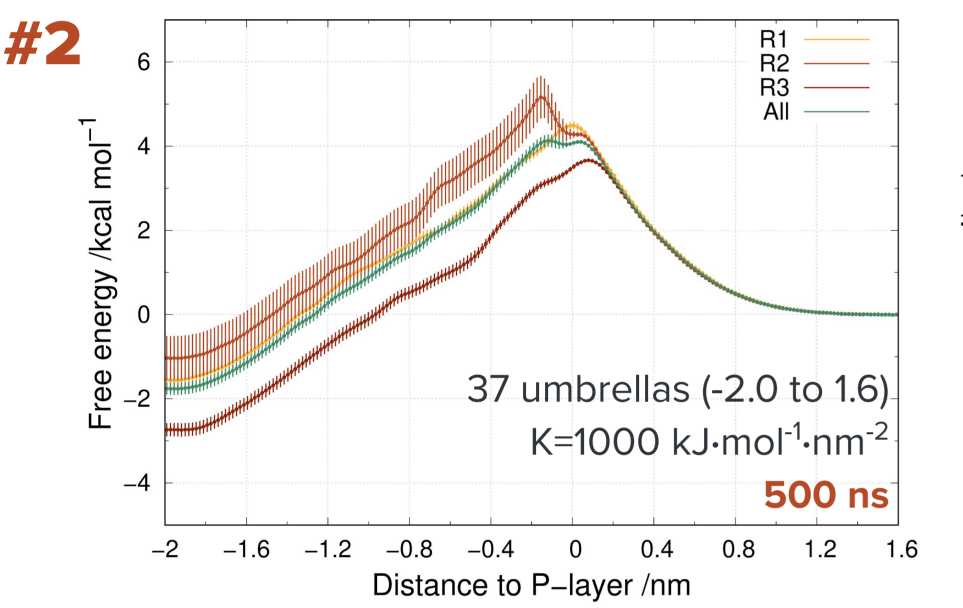
SETUP • We built a 128-POPC bilayer with a probe analogous to a benzene molecule.⁴ The probe was placed in different positions relative to a P-layer (0.0 nm), from the solvent (1.6 nm) to the center of the membrane (-2.0 nm), spaced 0.1 nm.

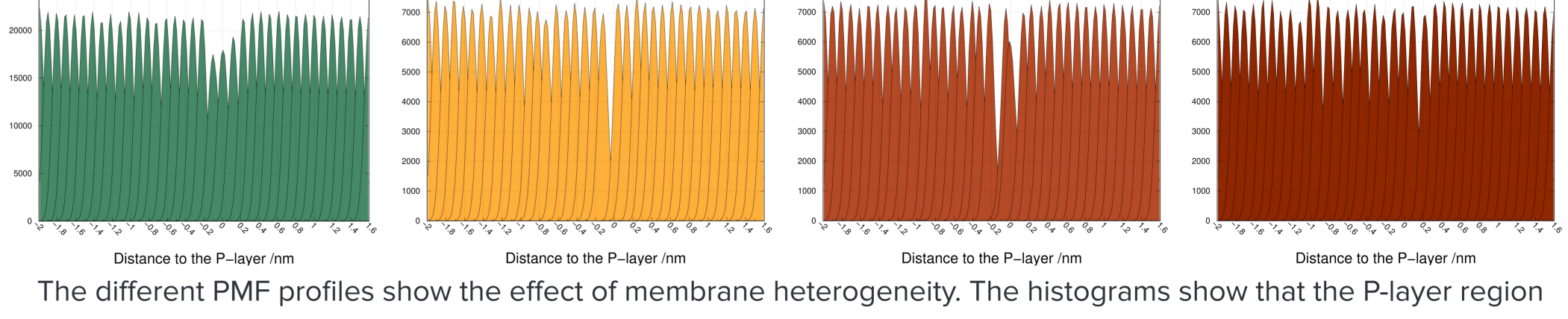


REPLICATES • were performed by placing the probe in different xy coordinates (*i.e.* in order to sample regions with different environments), e.g. probes 1, 2 and 3.

ANALYSIS • PMF profiles were calculated using the weighted histogram analysis method (WHAM)⁷ and the Bayesian histogram bootstrapping was performed using 50 iterations.

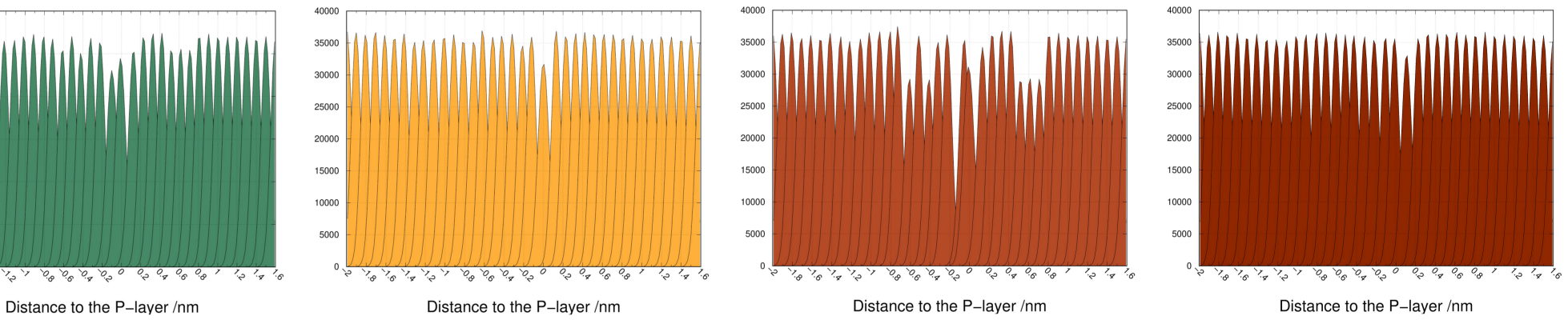






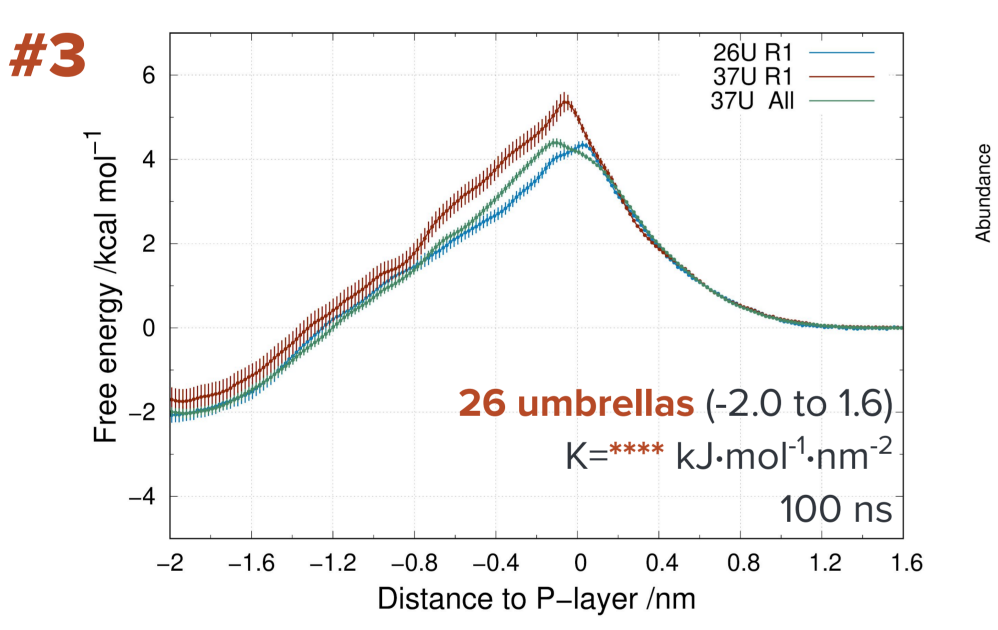
may not be sampled properly with this combination of umbrellas/K.

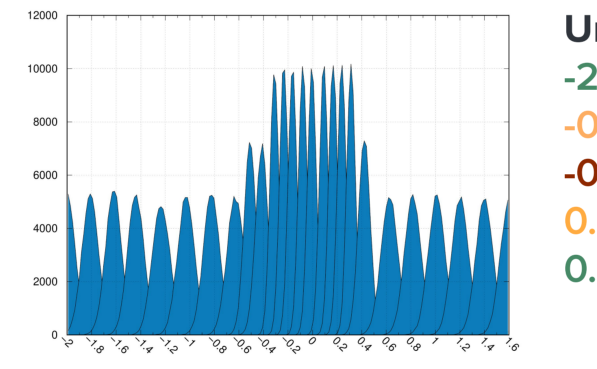
What if we increase the sampling?



Increasing the simulation time helps the PMFs converge, but the lack of sampling in the P-layer region remains. The computational cost becomes prohibitive.

What if we adjust the number of umbrellas?



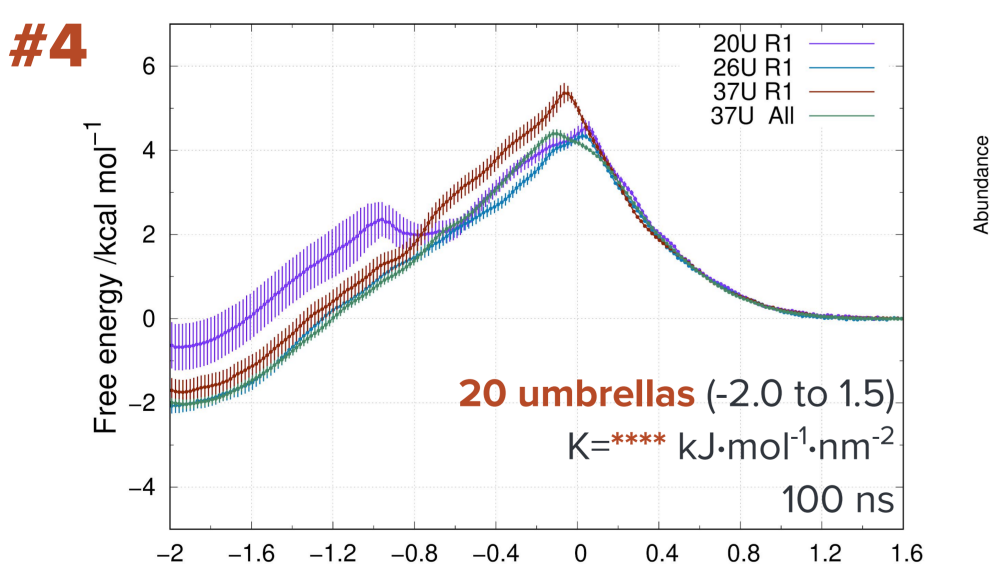


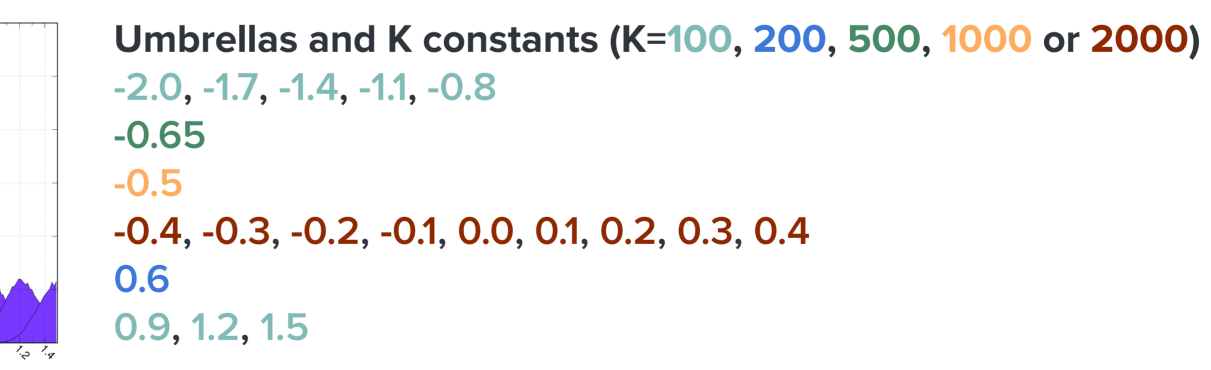
Distance to the P-layer /nm

Umbrellas and K constants (K=500, 1000 or 2000) -2.0, -1.8, -1.6, -1.4, -1.2, -1.0, -0.8, -0.6 -0.5, -0.4 -0.3, -0.225, -0.15, -0.075, 0.0, 0.075, 0.15, 0.225, 0.30.4 0.6, 0.8, 1.0, 1.2, 1.4, 1.6

Some issues persist in the region of interest, but the PMF profile looks very similar to the one with 37 umbrellas, at a reduced computational cost.

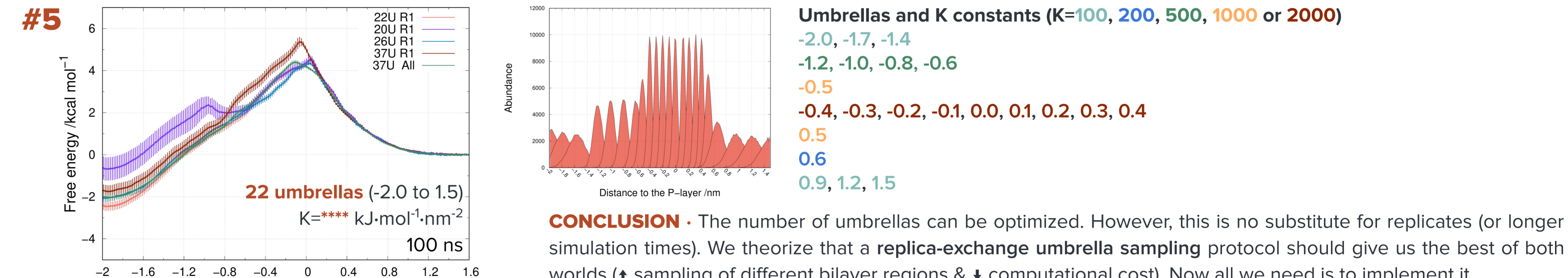
Can we further optimize this?





Several regions (not just in the center of interest) now lack sampling, which greatly impacts the PMF profile. Can we fix this by using umbrellas from previous runs to fill in the gaps?

Distance to P-layer /nm



Distance to the P-layer /nm

simulation times). We theorize that a replica-exchange umbrella sampling protocol should give us the best of both worlds (+ sampling of different bilayer regions & + computational cost). Now all we need is to implement it.



PTDC/MED-QUI/29036/2017

Distance to P-layer /nm

Refs. 1 Nelson, K.M. Baell, J.B. et al. (2010) et al. (2017) JMC 53:2719 JMC 60:1620



Jesus, A.R. et al. (2017) JMC 60, 568 🔲 🖬





調整

