

Supplemental Figures for: A Two-Step Screen Identifies a Small Molecule that Disrupts Membrane Voltage and is Effective Against Growing and Persister Gram-Negative Bacteria

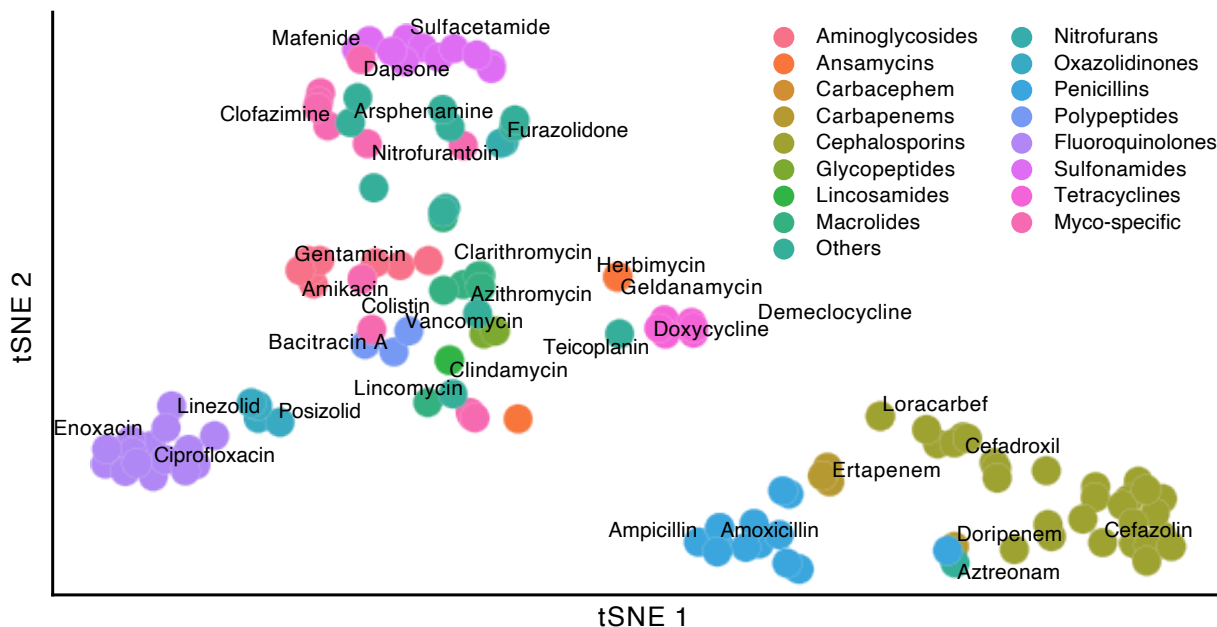


Figure S1. tSNE clustering of antibiotics independent of the ChemBridge library. Antibiotic classes are colored as shown in the key and two representatives of each family are annotated. Clustering is based on MACCS fingerprints and distances were calculated using Tanimoto similarity.

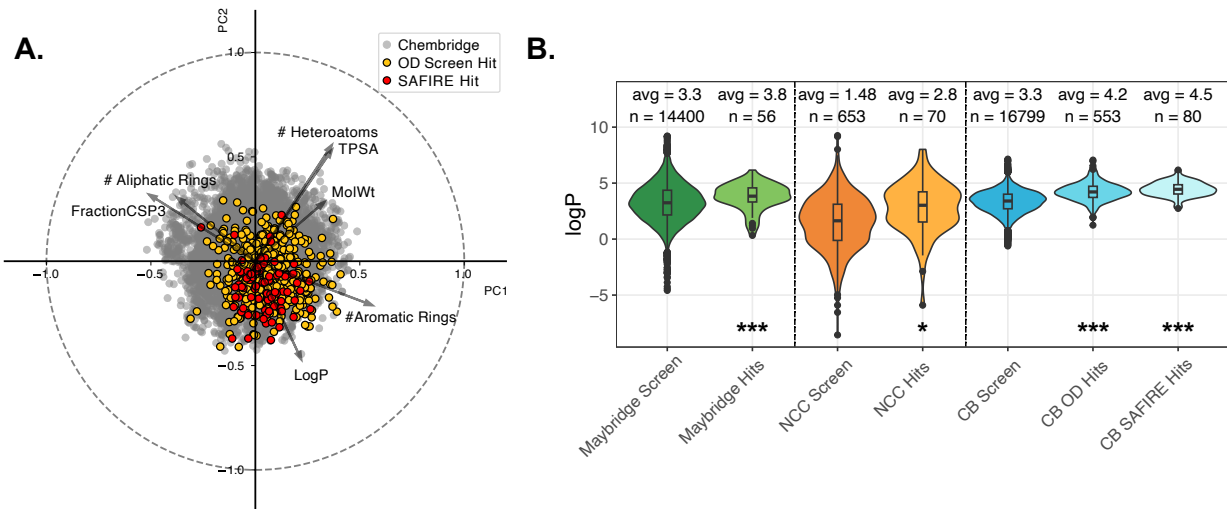


Figure S2. Slight enrichment of logP values found in this and previous SAFIRE screens.

A) PCA of the 16,799 Chembridge Combiset library compounds was performed on the physiochemical properties depicted in the plot. Molecular features were computed with RDKit and Z-score normalized with sklearn python package prior to PCA. Implementation of PCA was performed with scikit-learn.

B) Mean and range of calculated logP values of compounds from the present study and two previous SAFIRE screens. Maybridge Hitfinder v11 Library (Maybridge; Reens et al. 2018), National Institutes of Health National Clinical Collection (NCC; Nagy et al. 2020), and Chembridge Combiset Library (CB; this study). LogP values were calculated with RDkit. The number of compounds in each library and set of hit compounds is annotated (n). The violin plot is scaled by width. The boxplots span the interquartile range with the horizontal line indicating the median logP value and outlier data plotted as points. A two-sample t-test was performed on the logP values from the hit compounds versus the parent library for each screen. $P < 0.05$, 0.005, 0.0005 denoted by *, **, and ***, respectively.

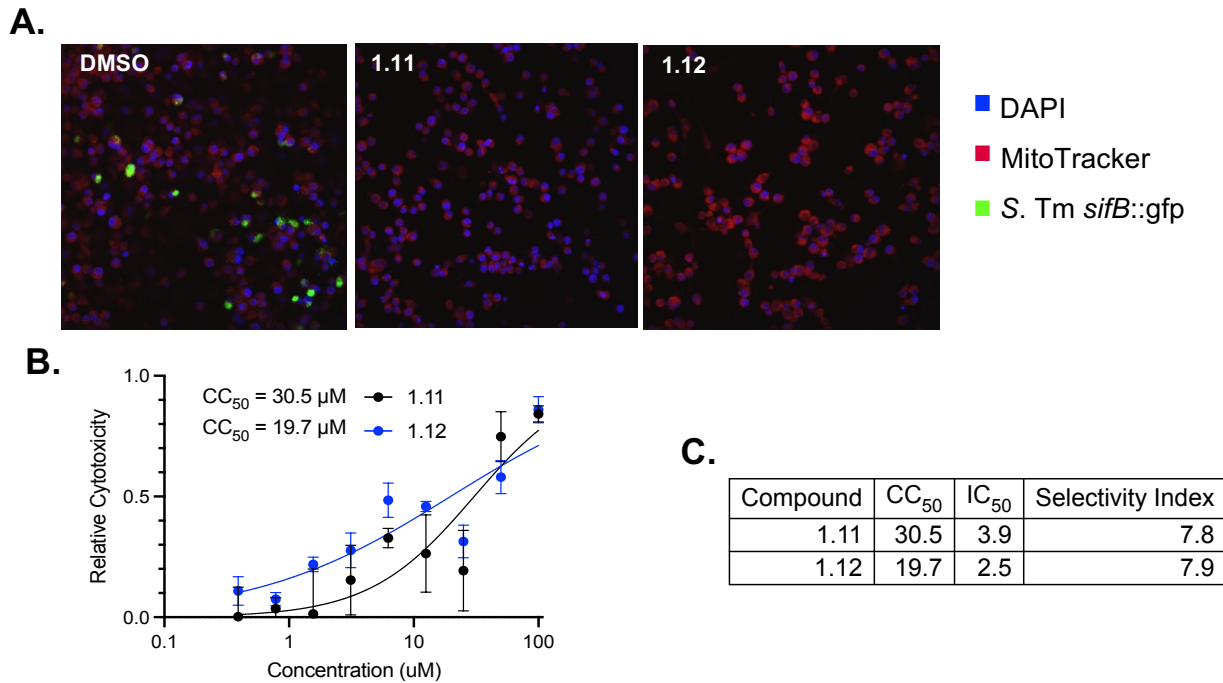


Figure S3. Measures of host cell toxicity for compounds CB1.11 and CB1.12.

A) Representative images from the SAFIRE assay, in which RAW264.7 macrophage-like cells were treated with 25 μM of compound 2 hours after infection, incubated with MitoTracker Red after 17.5 hours, stained with DAPI, fixed, and imaged after 18 hours of infection.

B) The lactate dehydrogenase (LDH) membrane integrity assay performed with RAW 264.7 cells at 16 hours after compound treatment. The key denotes half maximal cell cytotoxicity (CC_{50}) toxicity. Data are normalized to DMSO controls. Mean +/- SEM of three biological replicates.

C) Comparison of CC_{50} , IC_{50} , and selectivity index values for compounds CB1.11 and CB1.12.

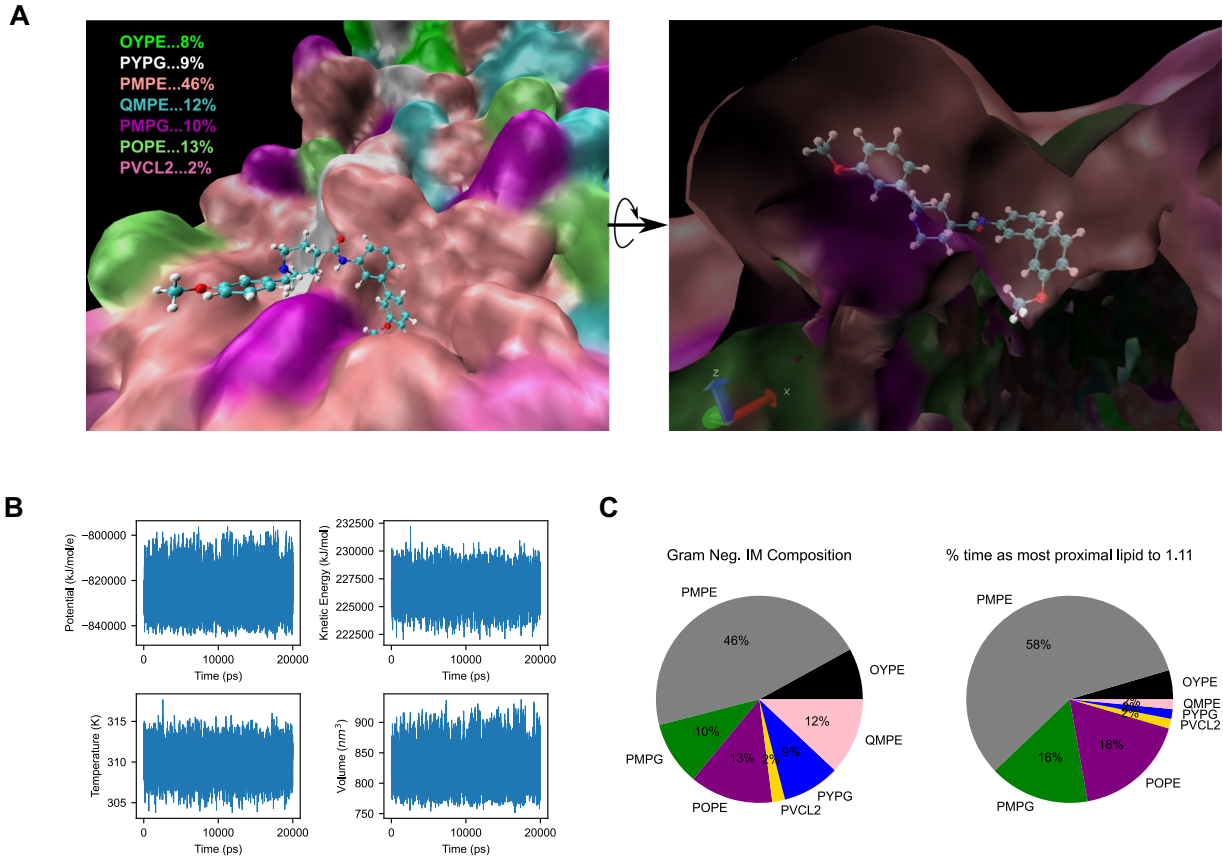


Figure S4. MD simulation, membrane energetics and system properties, and comparison of membrane lipid composition and compound proximity.

A) Model of CB1.11 inserting into the bacterial inner membrane.

B) Stable energetics and system properties for MD simulations. Time trace of potential and kinetic energy, temperature, and volume over 20 ns MD simulation post system minimization and equilibration. Simulations available in the [GitHub repository](#) associated with this paper.

C) (Left) Lipid composition of Gram-negative bacteria. (Right) Percent time lipid group is most proximal to CB1.11 during MD simulation.

Supplemental references:

- Nagy, Toni A., Amy L. Crooks, Joaquin L. J. Quintana, and Corrella S. Detweiler. 2020. “Clofazimine Reduces the Survival of *Salmonella Enterica* in Macrophages and Mice.” *ACS Infectious Diseases* 6 (5): 1238–49. <https://doi.org/10.1021/acsinfecdis.0c00023>.
- Reens, Abigail L., Amy L. Crooks, Chih-Chia Su, et al. 2018. “A Cell-Based Infection Assay Identifies Efflux Pump Modulators That Reduce Bacterial Intracellular Load.” *PLoS Pathogens* 14 (6): e1007115. <https://doi.org/10.1371/journal.ppat.1007115>.