



FIGURE 3: Dynamical Cross Correlational Map of the protein residue backbone of 1B7Y\_B\_408.

simulated a total of about 1,500 nanoseconds (1.5 milliseconds) worth of simulations involving 87 protein loop classifications associated with aaRS domains delimiting the specificities of the genetic code. Loop molecular dynamic simulations of these multi-domain proteins have yielded impressive results which have laid the **groundwork** for analyses involving single-domain metabolic metaconsensus enzymes selected using comparative bioinformatics techniques grounded in sequence, structure, and metabolic reactions.

### NEXT GENERATION WORK

Our preliminary experiments indicate an intricate set of patterns among structure, function, and dynamics of proteins. These patterns have the potential to **uncover** evolutionary drivers that may shed light on a basic yet confounding phenomenon in nature: does structure dictate function or vice versa? We aim to expand our analysis to bigger datasets concerning proteins in signaling networks. Also, we are interested in performing machine learning analyses of dynamic simulation datasets to dissect distinct molecular dynamic patterns along an evolutionary timeline of protein domains.

### PUBLICATIONS AND DATA SETS

Mughal, F., G. Caetano-Anollés and F. Gräter. Mining the evolutionary dynamics of protein loop structure and its role in biological functions. *Blue Waters Annual Report* (Urbana, Illinois, 2015), pp.130-131.

Caetano-Anollés, G., F. Gräter, C. Debès, D. Mercadante, and F. Mughal. The dynamics of protein disorder and its evolution: Understanding single molecule FRET experiments of disordered proteins. *Blue Waters Annual Report* (Urbana, Illinois, 2014), pp. 100-101.

## IMPROVING THE ACCURACY OF DRUG PERMEABILITY CALCULATIONS

**Allocation:** Illinois/25.0 Knh  
**PI:** Christopher Chipot<sup>1</sup>  
**Co-PI:** Jeffrey Comer<sup>2</sup>

<sup>1</sup>University Illinois at Urbana-Champaign

<sup>2</sup>Kansas State University

### EXECUTIVE SUMMARY

The inhomogeneous solubility-diffusion model has provided a convenient framework for understanding membrane permeation by drug molecules. This

model shows the relationship between the resistance to permeation in the direction normal to the membrane to the position-dependent diffusivity of the drug and the one-dimensional free-energy profile

underlying its translocation from the bulk aqueous phase to the interior of the lipid environment. For the **first time**, we provide a model for membrane permeation of a drug that, in stark contrast with the solubility-diffusion model, does not assume a lack of long-range correlations in time and space. Our model allows for better understanding of permeation dynamics for molecules exhibiting subdiffusive behavior on the characteristic timescales of their permeation. Our simulations suggest that this subdiffusive behavior is a result of permeation being governed by the spontaneous formation of voids within the membrane, which leads to intermittent large displacements of a permeant that is otherwise nearly immobile.

### INTRODUCTION

In the search of novel therapeutic agents, many chemical compounds able to bind a given target with very high affinity are eventually discarded on account of their cytotoxicity, propensity to associate with potassium channel human Ether-à-go-go-Related Gene (hERG), or poor bioavailability. Predicting these properties at an early stage of drug discovery, upstream from costly organic syntheses and clinical trials, is desirable. One possible avenue to address high drug-attribution rates [1] consists in quantifying the ability of the substrate to traverse lipid membranes spontaneously, for instance, in the gastrointestinal tract, and reach the targeted protein in an adequate amount. A consistent theoretical model of the lipid membrane permeation process is essential for linking the physicochemical properties of drug candidates to their adsorption and distribution.

### METHODS & RESULTS

The goal of this research is to understand a question central to drug discovery, namely how a drug spontaneously translocates across the biological membrane to reach its designated target. A model that has pervaded the field over the past twenty years is the so-called solubility-diffusion model of passive membrane permeation of small molecules [2]. In this model, the diffusion of the permeant is ordinarily assumed to obey the conventional Smoluchowski diffusion equation, which describes classical diffusion of particles on an inhomogeneous free-energy and diffusivity landscape. However, this

equation cannot accommodate subdiffusive behavior [3], which has long been recognized in other aspects of lipid bilayer dynamics, including lateral diffusion of individual lipids. Using large-scale molecular dynamics simulations of permeation events in a fully hydrated lipid bilayer performed on Blue Waters, we show that subdiffusive behavior is present in the transverse diffusion of a series of alcohols through a pure membrane, remaining relevant on timescales approaching the typical permeation time. We find that a model based on a fractional-order differential equation appropriately describes the motion of the permeant on timescales ranging from 1 picosecond to 1 nanosecond, which cannot be replicated by a single conventional Smoluchowski model. Multiple approaches indicate that the mean squared displacement within the bilayer, in the absence of a net force, depends on time as a power law, namely  $t^{0.7}$ , in contrast with the conventional model where this dependence is strictly linear. Our molecular dynamics simulations bring to light an unexpected phenomenon, linking subdiffusion to the formation of transient voids that spontaneously appear within the hydrophobic region of the bilayer and allow rare, but large displacements of the permeant, which is otherwise virtually immobile. The results of this investigation, which reweaves the fabric of the physical principles underlying membrane permeation by drug molecules, have been reported in a research article recently submitted for publication [4].

### WHY BLUE WATERS

Blue Waters was essential to perform a very large series of independent molecular-dynamics simulations of a membrane assembly in a time-bound fashion.