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*[[insert\_date]]*

Dear Editor of *Proceedings of the National Academy of Sciences*,

Please find our enclosed manuscript, which we hope you will consider for publication in your journal. There is a growing need to transcend conventional annotations on protein structure (such as residues involved in protein-protein interactions, post-translational modifications, stability, etc.) in order to better understand the conservation patterns that are increasingly coming to light through next-generation sequencing initiatives: deep sequencing has unearthed a class of protein elements that are strongly conserved, despite the fact that available structural annotations generally fail to explain such conservation. Knowledge relevant to dynamic behavior (and in particular, allostery) may often provide the missing conceptual link. In the first study of its kind, we develop an integrative and highly efficient framework that directly uses knowledge of protein structures and conformational changes in order to identify potential allosteric residues on both the surface and the within interior, and followed by the identification of hundreds of alternative conformations throughout the PDB, we apply this framework to study the conservation of the potential allosteric residues identified on a large scale, with conservation being evaluated using multiple metrics and datasets to investigate inter-species and human-specific patterns of constraint. We not only find that our identified residues tend to be conserved using multiple measures and sources of data, but they may also sometimes help to explain otherwise poorly understood human disease-associated variants. Finally, we make our framework available to the scientific community with a newly introduced and publically accessible software tool (STRESS) that is easy to use.

We list a number of suitable reviewers for this work.

 Yours sincerely,

 Mark Gerstein

 Albert L. Williams Professor

 of Biomedical Informatics