

# Mikael E Trellet

## List of Publications by Year in descending order

Source: <https://exaly.com/author-pdf/639930/publications.pdf>

Version: 2024-02-01

23

papers

3,367

citations

516710

16

h-index

677142

22

g-index

25

all docs

25

docs citations

25

times ranked

6259

citing authors

| #  | ARTICLE  | IF   | CITATIONS |
|----|--|------|-----------|
| 1  | <scp>PDB</scp> tools</scp> web: A user-friendly interface for the manipulation of <scp>PDB</scp> files. <i>Proteins: Structure, Function and Bioinformatics</i> , 2021, 89, 330-335.               | 2.6  | 15        |
| 2  | An overview of dataâ€driven HADDOCK strategies in CAPRI rounds 38â€45. <i>Proteins: Structure, Function and Bioinformatics</i> , 2020, 88, 1029-1036.  | 2.6  | 11        |
| 3  | Inhibition of the integrated stress response by viral proteins that block p-eIF2â€“eIF2B association. <i>Nature Microbiology</i> , 2020, 5, 1361-1373.   | 13.3 | 39        |
| 4  | Proteinâ€“Protein Modeling Using Cryo-EM Restraints. <i>Methods in Molecular Biology</i> , 2020, 2112, 145-162.  | 0.9  | 3         |
| 5  | Blind prediction of homoâ€and heteroâ€protein complexes: The CASP13â€CAPRI experiment. <i>Proteins: Structure, Function and Bioinformatics</i> , 2019, 87, 1200-1221.                              | 2.6  | 99        |
| 6  | Sharing Data from Molecular Simulations. <i>Journal of Chemical Information and Modeling</i> , 2019, 59, 4093-4099.  | 5.4  | 26        |
| 7  | Folding Then Binding vs Folding Through Binding in Macrocyclic Peptide Inhibitors of Human Pancreatic Î±-Amylase. <i>ACS Chemical Biology</i> , 2019, 14, 1751-1759.                               | 3.4  | 16        |
| 8  | West-Life: A Virtual Research Environment for structural biology. <i>Journal of Structural Biology: X</i> , 2019, 1, 100006.   | 1.3  | 2         |
| 9  | Large-scale prediction of binding affinity in proteinâ€“small ligand complexes: the PRODIGY-LIG web server. <i>Bioinformatics</i> , 2019, 35, 1585-1587.   | 4.1  | 130       |
| 10 | Performance of HADDOCK and a simple contact-based proteinâ€“ligand binding affinity predictor in the D3R Grand Challenge 2. <i>Journal of Computer-Aided Molecular Design</i> , 2018, 32, 175-185. | 2.9  | 97        |
| 11 | Semantics for an Integrative and Immersive Pipeline Combining Visualization and Analysis of Molecular Data. <i>Journal of Integrative Bioinformatics</i> , 2018, 15, .                             | 1.5  | 20        |
| 12 | pdb-tools: a swiss army knife for molecular structures. <i>F1000Research</i> , 2018, 7, 1961.  | 1.6  | 99        |
| 13 | SpotOn: High Accuracy Identification of Protein-Protein Interface Hot-Spots. <i>Scientific Reports</i> , 2017, 7, 8007.  | 3.3  | 77        |
| 14 | Sense and simplicity in <scp>HADDOCK</scp> scoring: Lessons from <scp>CASPâ€CAPRI</scp> round 1. <i>Proteins: Structure, Function and Bioinformatics</i> , 2017, 85, 417-423.                      | 2.6  | 44        |
| 15 | The HADDOCK2.2 Web Server: User-Friendly Integrative Modeling of Biomolecular Complexes. <i>Journal of Molecular Biology</i> , 2016, 428, 720-725.   | 4.2  | 2,071     |
| 16 | Content and task based navigation for structural biology in 3D environments. , 2015, , .   |      | 4         |
| 17 | Information-Driven Modeling of Protein-Peptide Complexes. <i>Methods in Molecular Biology</i> , 2015, 1268, 221-239.   | 0.9  | 24        |
| 18 | ExaViz: a flexible framework to analyse, steer and interact with molecular dynamics simulations. <i>Faraday Discussions</i> , 2014, 169, 119-142.  | 3.2  | 11        |

| #  | ARTICLE   | IF   | CITATIONS |
|----|---|------|-----------|
| 19 | Defining the limits of homology modeling in informationâ€“driven protein docking. Proteins: Structure, Function and Bioinformatics, 2013, 81, 2119-2128.                            | 2.6  | 63        |
| 20 | A Unified Conformational Selection and Induced Fit Approach to Protein-Peptide Docking. PLoS ONE, 2013, 8, e58769.  | 2.5  | 163       |
| 21 | Community-wide evaluation of methods for predicting the effect of mutations on protein-protein interactions. Proteins: Structure, Function and Bioinformatics, 2013, 81, 1980-1987. | 2.6  | 87        |
| 22 | Clustering biomolecular complexes by residue contacts similarity. Proteins: Structure, Function and Bioinformatics, 2012, 80, 1810-1817.  | 2.6  | 103       |
| 23 | Posttranslational Modification of Pili upon Cell Contact Triggers <i>&lt; i&gt;N. meningitidis&lt;/i&gt;</i> Dissemination. Science, 2011, 331, 778-782.                            | 12.6 | 162       |