

# Józef Adam Liwo

## List of publications and invited presentations

### Original papers in ISI journals

283. A. G. Lipska, A. K. Sieradzan, S. Atmaca, C. Czaplewski, **A. Liwo**. Toward Consistent Physics-Based Modeling of Local Backbone Structures and Chirality Change of Proteins in Coarse-Grained Approaches. *J. Phys. Chem. Lett.*, **2023**, 14, 9824-9833.
282. L. Borges-Araújo, I. Patmanidis, A. P. Singh, L. H. S. Santos, A. K. Sieradzan, S. Vanni, C. Czaplewski, S. Pantano, W. Shinoda, L. Monticelli, Pragmatic Coarse-Graining of Proteins: Models and Applications. **A. Liwo**, S. J. Marrink, P. C. T. Souza. *J. Chem. Theory Comput.* **2023**, 19, 7112-7135.
281. A Danielsson, S. A. Samsonov, **A. Liwo**, A. K. Sieradzan. Extension of the SUGRES-1P Coarse-Grained Model of Polysaccharides to Heparin. *J. Chem. Theory Comput.*, **2023**, 19, 6023-6036.
280. M. F. Lensink, G. Brysbaert, N. Raouraoua, P. A. Bates, M. Giulini, R. V. Honorato, C. van Noort, J. M. C. Teixeira, A. M. J. J. Bonvin, R. Kong, H. Shi, X. Lu, S. Chang, J. Liu, Z. Guo, X. Chen, A. Morehead, R. S. Roy, T. Wu, N. Giri, F. Quadir, C. Chen, J. Cheng, C. A. Del Carpio, E. Ichiishi, L. A. Rodriguez-Lumbrieras, J. Fernandez-Recio, A. Harmalkar, L.-S. Chu, S. Canner, R. Smanta, J. J. Gray, H. Li, P. Lin, J. He, H. Tao, S.-Y. Huang, J. Roel-Touris, B. Jimenez-Garcia, C. W. Christoffer, A. J. Jain, Y. Kagaya, H. Kannan, T. Nakamura, G. Terashi, J. C. Verburgt, Y. Zhang, Z. Zhang, H. Fujuta, M. Sekijima, D. Kihara, O. Khan, S. Kotelnikov, U. Ghani, D. Padhorny, D. Beglov, S. Vajda, D. Kozakov, S. S. Negi, T. Ricciardelli, D. Barradas-Bautista, Z. Cao, M. Chawla, L. Cavallo, R. Oliva, R. Yin, M. Cheung, J. D Guest, J. Lee, B. G. Pierce, B. Shor, T. Cohen, M. Halfon, D. Schneidman-Duhovny, S. Zhu, R. Yin, Y. Sun, Y. Shen, M. Maszota-Zieleniak, K. K. Bojarski, E. A. Lubecka, M. Marcisz, A. Danielsson, L. Dziadek, M. Gaardlos, A. Gieldon, **A. Liwo**, S. A. Samsonov, R. Slusarz, K. Zieba, A. K. Sieradzan, C. Czaplewski, S. Kobayashi, Y. Miyakawa, Y. Kiyota, M. Takeda-Shitaka, K. Olechnovic, L. Valancauskas, J. Dapkus, C. Venclovas, B. Wallner, L. Yang, C. Hou, X. He, S. Guo, S. Jiang, X. Ma, R. Duan, L. Qui, X. Xu, X. Zou, S. Velankar, S. J. Wodak. Impact of AlphaFold on structure prediction of protein complexes: The CASP15-CAPRI experiment. *Proteins: Struct. Funct. Bioinf.*, **2023**, 91, 1658-1683.
279. M. Maszota-Zieleniak, **A. Liwo**, S. Ricard-Blum, S. A. Samsonov. Interplay of heparan sulfate chains with the core proteins of syndecans 2 and 4. *Proteoglycan Res.*, **2023**, 1, e10.
278. K. M. Ocetkiewicz, C. Czaplewski, H. Krawczyk, A. G. Lipska, **A. Liwo**, J. Proficz, A. K. Sieradzan, P. Czarnul. UNRES-GPU for Physics-Based Coarse-Grained Simulations of Protein Systems at Biological Time-and Size-Scales. *Bioinformatics*, **2023**, 39, btad391.
277. strains. A. G. Lipska, A. K. Sieradzan, C. Czaplewski, A. D. Lipińska, K. M. Ocetkiewicz, J. Proficz, P. Czarnul, H. Krawczyk, **A. Liwo**. Long-time scale simulations of virus-like particles from three human-norovirus *J. Comput. Chem.*, **2023**, 44, 1470-1483.

276. **A. Liwo**, M. Pyrka, C. Czaplewski, X. Peng, A. J. Niemi. Long-Time Dynamics of Selected Molecular-Motor Components Using a Physics-Based Coarse-Grained Approach. *Biomolecules*, **2023**, 13, 941.
275. A. K. Sieradzan, J. Sans-Duñó, E. A. Lubecka, C. Czaplewski, A. G. Lipska, H. Leszczyński, K. M. Ocetkiewicz, J. Proficz, P. Czarnul, H. Krawczyk, **A. Liwo**. Optimization of parallel implementation of UNRES package for coarse-grained simulations to treat large proteins. *J. Comput. Chem.*, **2023**, 44, 602-625.
274. S.-J. Chen, M. Hassan, R. L. Jernigan, K. Jia, D. Kihara, A. Kloczkowski, S. Kotelnikov, D. Kozakov, J. Liang, **A. Liwo**, S. Matysiak, J. Meller, C. Micheletti, J. C. Mitchell, S. Mondal, R. Nussinov, K.-i. Okazaki, D. Padhorny, J. Skolnick, T. R. Sosnick, G. Stan, I. Vakser, X. Zou, G. D. Rose. Protein folds vs. protein folding: Differing questions, different challenges. *Proc. Natl. Acad. Sci.*, **2023**, 120, e2214423119.
273. peptides. C. Schulze, A. Danielsson, **A. Liwo**, D. Huster, S. A. Samsonov, A. Penk. Ligand binding of interleukin-8: a comparison of glycosaminoglycans and acidic *Phys. Chem. Chem. Phys.*, **2023**, 25, 24930-24947.
272. R. Ślusarz, E. A. Lubecka, C. Czaplewski, **A. Liwo**. Improvements and new functionalities of UNRES server for coarse-grained modeling of protein structure, dynamics, and interactions. *Front. Mol. Biosci.*, **2022**, 9, 1071428.
271. E. A. Lubecka, **A. Liwo**. A coarse-grained approach to NMR-data-assisted modeling of protein structures. *J. Comput. Chem.*, **2022**, 43, 2047-2059.
270. C. Sikorska, **A. Liwo**. Origin of Correlations between Local Conformational States of Consecutive Amino Acid Residues and Their Role in Shaping Protein Structures and in Allostery *J. Phys. Chem. B*, **2022**, 126, 9493-9505.
269. I. Biskupek, C. Czaplewski, J. Sawicka, E. Ilowska, M. Dzierżyńska, S. Rodziewicz-Motowidło, **A. Liwo**. Prediction of Aggregation of Biologically-Active Peptides with the UNRES Coarse-Grained Model. *Biomolecules*, **2022**, 12, 1140.
268. I. Biskupek, A. Sieradzan, C. Czaplewski, **A. Liwo**, A. Lesner, A. Giełdoń. Theoretical investigation of the coronavirus SARS-CoV-2 (COVID-19) infection mechanism and selectivity. *Molecules*, **2022**, 2080.
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266. C. Czaplewski, Z. Gong, E.A. Lubecka, K. Xue, C. Tang, **A. Liwo**. Recent Developments in Data-Assisted Modeling of Flexible Proteins. *Front. Mol. Biosci.*, **2021**, 8, 765562.
265. **A. Liwo**, C. Czaplewski, A.K. Sieradzan, A.G. Lipska, S.A. Samsonov, R.K. Murarka. Theory and Practice of Coarse-Grained Molecular Dynamics of Biologically Important Systems. *Biomolecules*, **2021**, 11, 1347.

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1. F. Kasprzykowski, P. Kania, S. Ołdziej, **A. Liwo**, M. Tarnowska, Z. Grzonka, M. Abrahamsson, I. Olafsson, A. Grubb and J. Trojnar. Inhibition of cysteine proteinases by a series of peptidyl-diazomethanes structurally related to the binding center of cystatins. In *Peptides 1990*, eds. E. Ginolt, D. Andrew, ESCOM, Leiden 1991, pp. 799-800.

**Józef Adam Liwo**  
**Invited presentations**

50. Long-time dynamics of selected molecular-motor components using a physics-based coarse-grained approach. 21th KIAS Conference on Protein Structure and Function, Seoul, South Korea, October 11 – 14, 2023.
49. Long-time dynamics of selected molecular-motor components using a physics-based coarse-grained approach. 7th Telluride Workshop on Coarse-Grained Modeling of Structure and Dynamics of Biomacromolecules, July 24 – 28, 2023, Telluride, USA.
48. Recent developments and applications of the UNICORN coarse-grained model of biological macromolecules. Workshop on Topology, Physics, and Chemistry of Soft Matter: Eutopia IV, 5 – 9 September 2022, Trento, Italy.
47. Origin of through-sequence correlations between local conformational states of amino-acid residues and their role in shaping protein structures and in allostery. 20th KIAS Conference on Protein Structure and Function, Seoul, South Korea, September 14 – 17, 2022.
46. Origin of through-sequence correlations between local conformational states of amino-acid residues and their role in shaping protein structures and in allostery. 6th Telluride Workshop on Coarse-Grained Modeling of Structure and Dynamics of Biomacromolecules, July 25 – 28, 2022, Telluride, USA.
45. Theory and practice of coarse graining, Mathematics of Life, MoL2021, 13 – 16 September 2021, Hisarya, Bulgaria & online.
44. Data-assisted modeling of protein structures with the coarse-grained UNRES model . 14th Community Wide Experiment on the Critical Assessment of Techniques for Protein Structure Prediction CASP14, Virtual Meeting, November 30 – December 4, 2020.
43. The Unified Coarse Grained model of biological macromolecules: principles and applications. XLVIII Annual Meeting of the Argentinian Biophysical Society, San Luis, Argentina, Nov. 27 – 29, 2019.
42. The scale-consistent UNICORN package for large-scale simulations of biological macromolecules. E-CAM Extended Software Development Workshop: Inverse Molecular Design & Inference: building a Molecular Foundry I, Clifden, Galway, Ireland, November 1 – 9, 2019.
41. A scale-consistent approach to coarse graining biomolecular systems. The 19th KIAS Conference on Protein Structure and Function, Korea Institute for Advanced Study, Seoul, Korea, September 26-September 28, 2019.
40. New scale-consistent UNRES force field for protein simulatios. CECAM Protein Simulations – Current State of Art, Tel Aviv, Israel, October 22-24, 2019.
39. Simulations of protein structure, dynamics, and thermodynamics with the coarse-grained UNRES force field and massively parallel computers. HPC 2019 Bulgaria, September 2-6, 2019, Borovets, Bulgaria.

38. Physics-based scale- and geometry-consistent coarse-grained potentials. Workshop on Multi-scale Modelling, Leiden, The Netherlands, June 24-June 28, 2019.
37. Using the UNRES server and the standalone UNRES package in SAXS-data-assisted modeling of protein structure. simSAS 2019, Grenoble France, April 8-11/12, 2019
36. Data-assisted prediction of protein structures with the coarse-grained UNRES force field. CASP13 Meeting, Iberostar Paraiso Maya, Mexico, December 1-4, 2018.
35. Physics-based scale- and geometry-consistent coarse-grained potentials. 26th Conference on Current Trends in Computational Chemistry (2018 CCTCC) Jackson, MS (USA), 9-10 November 2018.
34. Systematic design of physics-based scale-consistent coarse-grained potentials for the simulations of biomolecules and nanostructures, Bio-, chem-, and nanoinformatics approaches to study bionano interface Dublin, Ireland, May 23-25, 2018.
33. Implementation of geometry-consistent local and correlation potentials in the UNRES force field. Biomolecules and Nanostructures 6, Podlesice, Poland, May 10-14, 2017
32. Geometry-consistent expressions for energy terms in coarse-grained force fields for (bio)polymers with UNRES as an example. Workshop on Physics and Biology of Proteins, Natal, Brazil, June 12-30, 2017
31. A general method for the derivation of effective energy expressions in coarse-grained force fields. 3rd Korean-Polish Conference on Protein Folding: Theoretical and Experimental Approaches, Seoul, Republic of Korea, February 4 – 9, 2017
30. The Unified Coarse-Grained Model for large-scale simulations of biological macromolecules. The 16th KIAS Conference on Protein Structure and Function, Seoul, Republic of Korea, September 22 – September 24, 2016.
29. How do the local and long-range interactions encode the three-dimensional structures of biological macromolecules: a coarse-grained perspective. 6th Visegrad Symposium on Structural Systems Biology, Warsaw, Poland, 19-21 June, 2016.
28. A rigorous approach to derive analytical expressions for the effective energy terms in coarse-grained force fields. 3rd International Conference on Protein and RNA Structure Prediction Punta Cana, Dominican Republic, Dec 14-18, 2015.
27. The UNRES coarse-grained model of polypeptide chains: theory and applications. Invited lecture presented at the Department of Physics, Beijing Institute of Technology, Beijing, P.R. China, September 17, 2015.
26. Maximum-likelihood calibration of force fields. 1st Korean-Polish Conference on Protein Folding: Theoretical and Experimental Approaches, Seoul, Rep. of Korea, May 24-28, 2015.
25. A unified coarse-grained model for biomolecular simulations. 2015 Workshop of Wuhan Center of Physical Biology, Wuhan, P.R. China, September 7, 2015.

24. A rigorous approach to the derivation of analytical potentials in physics-based coarse-grained force fields. 3rd International Workshop on Theoretical and Computational Physics, Complex Systems and Interdisciplinary Physics, Da Lat, Vietnam, July 27-30, 2015.
23. The first Nobel Prize in computational chemistry. An invited lecture presented during the quarterly Colloquium at the Institute of Physics, Polish Academy of Sciences, Warsaw, Poland, February 25, 2014.
22. A maximum-likelihood approach to force-field calibration. Protein Folding Conference 2014, Grand Palladium Resort & Spa, Bavaro Hotel Punta Cana, Dominican Republic, July 16-19, 2014.
21. Essential role of mean-field electrostatic interactions in free modeling of protein and nucleic acid structures at coarse-grained level. Protein & RNA Structure Prediction Conference 2013, Occidental Grand Xcaret, Playa del Carmen Mexico, December 1-5, 2013.
20. Mean field dipole-dipole interactions as essential factors in the formation of biomolecular architectures. Biomolecules and Nanostructures 4, Pułtusk, Poland, May 15-19, 2013.
19. A simple coarse-grained model of nucleic acids reveals the essential role of mean-field dipole-dipole interactions between nucleic-acid bases and double-helix formation. Fifth Korea-Japan Seminars on Biomolecular Sciences: Experiments and Simulations. High1 Resort, Republic of Korea, February 24-16, 2013.
18. Coarse grained description of biomolecular systems. 10th Workshop on Bioinformatics and 5th Symposium of the Polish Bioinformatics Society, Gdańsk, Poland, May 25-26, 2012.
17. Coarse-grained models for proteins. Nordita Workshop “Dynamics of Biomolecular Processes: From Atomistic Representations to Coarse-Grained Models”, Stockholm, Sweden, February 27 - March 23, 2012; a series of 4 invited lectures.
16. Construction and application of coarse-grained force fields for biomacromolecules, Multipole Approaches to Structural Biology, Warsaw, Poland, November 16-19, 2011.
15. From atomistic simulations to network description of biological systems, Workshop “From Computational Biophysics to Systems Biology” 2011 (CBSB11), Juelich, Germany, July 20-22, 2011.
14. The nature of the conformational ensemble at the transition temperature: insights from simulations and experiment. Third Korea-Japan Seminars on Biomolecular Sciences: Experiments and Simulations, Hotel Lotte, Jeju, Korea, February 26, March 1, 2011.
13. Use of the UNRES force field and massively parallel computers in millisecond scale simulations of protein dynamics. Workshop on Bioinformatics (BIT09), Toruń, Poland May 23-25, 2009.
12. Towards simulations of structure and dynamics of large proteins with the UNRES force field. The 8th KIAS - Yonsei Conference on Protein Structure and Function, Seoul, Korea, October 9-11, 2008.

11. Prediction of structure and simulation of dynamics of protein folding with the mesoscopic UNRES force field, Modelling and Design of Molecular Materials. Piechowice, Poland, June 23-28, 2008.
10. Mesoscopic dynamics with the UNRES force field - a tool for studying the kinetics and thermodynamics of protein folding. NIC Workshop 2007, From Computational Biology to System Biology, Jülich, Germany, May 2-4, 2007.
9. Optimization of a mesoscopic force field for simulation of protein folding pathways. Workshop on Structure and Function of Biomolecules May 13-15, 2006, Będlewo near Poznań, Poland.
8. Ab initio simulations of protein folding pathways by molecular dynamics with the united-residue (UNRES) model of polypeptide chains. 30th FEBS Congress and 9th IUBMB Conference, Budapest, Hungary, 2005; FEBS Journal., **2005**, 272 (Suppl. 1) 359.
7. Design of hierarchical caldera-like potential-energy functions for energy-based prediction of protein structure and simulation of protein folding:application to the UNRES force field. Third KIAS Conference on Protein Structure and Function: Folding Mechanism, Proteomics, and Bioinformatics, Seoul, Korea, 29.09-1.10, 2003, 2003,
6. Energy-based prediction of protein structure with the UNRES force field, First KIAS Conference on Protein Structure and Function: Protein Folding in Post-Genome Era, Korea Institute for Advanced Study, Seoul, Korea, November 28 - November 30, 2001.
5. A knowledge-based united-residue force field for off-lattice calculations of protein structure that recognizes native folds. International Symposium on Theoretical, Experimental Aspects of Protein Folding, San Luis, Argentina, June 17-21, 1996
4. Role of singlet-oxygen binding to anthraquinones in the peroxidating activity of antitumor anthraquinone drugs. 5th International Symposium on Molecular Aspects of Chemotherapy, Gdansk, Poland, August 21-24, 1995
3. Prediction of protein structure using a mean-field united-residue potential determined from protein crystal data. 3-rd Conference Computers in Chemistry '94, Wrocław, Poland, June 23-26, 1994.
2. Application of the Free-Wilson method in structure-activity correlation of bioactive compounds on the example of oxytocin and vasopressin analogs. Theoretical and Experimental Aspects of Molecular Structure, Karpacz, Poland, June 5-10, 1994
1. The role of hydrophobic residue packing in protein folding. 8th Conference of Young Scientists on Organic, Bioorganic Chemistry, Riga, Latvia, 2-9 November 1991.