

# Christoph Junghans

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Deputy Group Leader

Applied Computer Science Group

Computer, Computational, & Statistical Sciences Division

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Web: [www.lanl.gov/junghans](http://www.lanl.gov/junghans)

## Research Interests

- **Currently:** Scientific software development
- Software design and management – core developer of the coarse-graining package [VOTCA](#)
- Porting and optimization of scientific codes ([VOTCA](#), [Cabana](#), [FleCSI](#))
- Novel software development workflow for Exascale computing
- Systematic coarse-graining of liquids and implementation of new methodologies (e.g. thermostats)
- Development of adaptive resolution simulation approaches (changing the degrees of freedom on the fly)
- Generalized ensemble methods to study phase transitions (multicanonical techniques)

## Education

- Oct 2010 **Dr. rer. nat. (Ph.D.) in Physics**, *magna cum laude*,  
Johannes Gutenberg University of Mainz, Germany  
[Between the Scales: Water from different Perspectives](#)  
An adaptive resolution molecular dynamics study using various coarse-graining methods
- Nov 2006 **Diploma (M.Sc.) in Physics**, *summa cum laude*, University of Leipzig, Germany  
[Aggregation of Mesoscopic Protein-like Heteropolymers](#)  
A generalized ensemble Monte Carlo study using multicanonical techniques

## Work Experience

- July 2018 - Present **Deputy Group Leader**, Applied Computer Science Group, Computer, Computational, and Statistical Sciences Division, Los Alamos National Lab
- Feb. 2016 - Present **CNLS Affiliate**, Center for Nonlinear Studies, Theoretical Division, Los Alamos National Lab
- Jan. 2017 - July 2018 **Deputy Team Leader**, Co-Design Team, Computer, Computational, and Statistical Sciences Division, Los Alamos National Lab
- May 2014 - July 2018 **Staff Scientist**, Computer, Computational, and Statistical Sciences Division, Los Alamos National Lab
- Mar 2013 – Apr 2014 **Director’s Postdoctoral Fellow**, Theoretical Division, Los Alamos National Lab
- Nov 2011 – Feb 2013 **Postdoctoral research assistant**, Theoretical Division, Los Alamos National Lab
- May 2012 – June 2012 **Member**, The Kavli Institute for Theoretical Physics, University of California, Santa Barbara, CA
- Nov 2010 – Oct 2011 **Postdoctoral research assistant**, Polymer Theory Group, Max Planck Institute for Polymer Research, Mainz, Germany
- Jan 2007 – Oct 2010 **Ph.D. Student**, Polymer Theory Group, Max Planck Institute for Polymer Research, Mainz, Germany
- Jan 2009 – July 2009 **Specialist for Application Performance & Deep Computing**, Internship, IBM Systems & Technology Group Europe
- Oct 2003 – Sept 2006 **Student assistant**, Institute for Theoretical Physics, University of Leipzig
- Aug 2005 – Oct 2005 **Research student** in the “Scientific Computing” program, Jülich Supercomputing Centre, Germany
- Sep 2004 **Student assistant**, Institute for Meteorology, University of Leipzig
- Feb 2003 – April 2003 **Research student**, Chair for Fluid Mechanics, Martin Luther University of Halle-Wittenberg, Germany

## Open Source Projects

|                           |   |                      |
|---------------------------|---|----------------------|
| <a href="#">VOTCA</a>     | Versatile Toolkit for coarse-graining applications        | Core Developer       |
| <a href="#">FleSCI</a>    | Flexible Computer Science Infrastructure                  | Developer            |
| <a href="#">Gentoo</a>    | Advanced Linux distribution                               | Developer            |
| <a href="#">Fedora</a>    | Linux distribution  | Contributor          |
| <a href="#">OpenSuse</a>  | Linux distribution  | Contributor          |
| <a href="#">ESPReso++</a> | Successor of the ESPReso simulation package               | Developer            |
| <a href="#">GroMaCS</a>   | Versatile package to perform molecular dynamics           | Developer (inactive) |
| <a href="#">ESPReso</a>   | Extensible Simulation Package for Research on Soft matter | Developer (inactive) |

More information can be found on [my GitHub profile](#) and [my OpenHUB profile](#).

## Awards

|           |  |
|-----------|--|
| Nov 2019  | Top 10% most cited PLOS ONE papers published in 2015                                     |
| Sep 2017  | LAAP Award in recognition of outstanding contributions to the ISTI Summer School Program |
| Feb 2017  | Postdoc Distinguished Mentor Award, LANL   |
| July 2016 | LAAP Achievement Award in appreciation for active service on the WSST team               |
| Jan 2015  | Classified as outstanding researcher, USCIS  |
| Jan 2014  | Most cited article published in EPJE in the past 5 years                                 |
| Mar 2013  | Los Alamos National Laboratory Director's Post-Doctoral Fellowship                       |
| May 2012  | Member of the Kavli Institute for Theoretical Physics                                    |
| June 2010 | Participant of the 60th Lindau Nobel Laureate Meeting                                    |
| Nov 2006  | Diploma with honor (highest possible grade)  |
| Nov 2005  | Wolfgang Natonek award, University of Leipzig  |
| Oct 2003  | Teubner award, Department for Physics and Earth Science, University of Leipzig           |
| July 2000 | Book award of the German Physical Society (DPG) for high school graduates                |

## Professional Skills

### *Managerial*

- Line management, since July 2018, Group of 65+ staff scientists and Postdocs, \$40M+ budget
- Team management, 1.5 years, technical management of 12+ team member
- Project management, multiple lead roles in Exascale Computing Project (ECP), Laboratory Directed Research & Development (LDRD) and Advanced Simulation & Computing (ASC) projects

### *Referee for Scientific Journals*

- ACS: J. Chem. Inf. Model. & J. Phys. Chem.
- AIP: J. Chem. Phys.
- Elsevier: Comp. Phys. Comm.
- RSC: Phys. Chem. Chem. Phys.
- and others

### *Code Review*

- [Gromacs](#)
- [OpenSuse Build Service](#)
- [Gentoo Science Project](#)

*Conferences*

- Member of the Organization Committee for the [Salishan Conference on High Speed Computing](#)
- Chairman for APS annual meetings

*Representative*

|                       |  |
|-----------------------|--|
| Nov 2018 - Present    | Member of the Information Science and Technology Institute (ISTI) council, LANL  |
| July 2015 - June 2016 | Member of the Worker Safety & Security Team (WSST) for CCS-7, LANL   |
| July 2007 - Dec 2008  | PhD representative of the theory group, Max Planck Institute for Polymer Research, Mainz   |
| Nov 2007 - Nov 2008   | PhD representative of the MPI for Polymer Research   |
| Oct 2002 - Sep 2005   | Elected member of student government (Fachschaftsrat) of the Department for Physics and Earth Science, University of Leipzig - Extensive committee work in this period incl. spokesperson, treasurer and member of several faculty/university committees |

**Computer Skills**

|                         |  |
|-------------------------|--|
| <b>General</b>          | Linux, AIX, Mac OS, DOS, Windows           |
| <b>Programming</b>      | C, Fortran, MPI, OpenMP, C++               |
| <b>Scripting</b>        | Shell, awk, Perl, expect, PHP, tcl, Python |
| <b>Markup Languages</b> | HTML, latex, mediawiki, txt2tags           |

**Teaching Experience***Teaching Assistant*

|                     |  |
|---------------------|--|
| Apr 2008 - Sep 2008 | Theoretical physics III, Johannes Gutenberg University of Mainz                |
| Oct 2007 - Mar 2008 | Theoretical physics II, Johannes Gutenberg University of Mainz                 |
| Apr 2007 - Sep 2007 | Theoretical physics I, Johannes Gutenberg University of Mainz                  |
| Apr 2006 - Sep 2006 | Introduction to computer simulations I, University of Leipzig                  |
| Sep 2004 - Oct 2004 | Mathematical preparation course for first-year students, University of Leipzig |

*Mentoring*

|                     |   |
|---------------------|---|
| June 2019 - Present | Postdoctoral research assistant, Los Alamos National Laboratory                       |
| Feb 2018 - Present  | Postdoctoral Research Assistant, Los Alamos National Laboratory                       |
| May 2018 - Feb 2019 | Postdoctoral research assistant (now Staff Scientist), Los Alamos National Laboratory |
| Feb 2017 - Jan 2019 | Postdoc. Research Assistant (now Staff Scientist), Los Alamos National Laboratory     |
| Sep 2018            | Member of PhD committee, University of Reims Champagne-Ardenne                        |
| Jun 2018 - Aug 2018 | ECP Summer Student, Los Alamos National Laboratory                                    |
| May 2018 - Aug 2018 | Co-Design Summer School (6 Students), Los Alamos National Laboratory                  |
| May 2018 - Aug 2018 | ASC Summer Student, Los Alamos National Laboratory                                    |
| Feb 2016 - May 2018 | Postdoc. Research Assistant (now Staff Scientist), Los Alamos National Laboratory     |
| Dec 2016 - Apr 2018 | Postdoc. Research Assistant (now Staff Scientist), Los Alamos National Laboratory     |
| Jun 2017 - Aug 2017 | High School Internship Student, Los Alamos National Laboratory                        |
| Apr 2017 - Jun 2017 | ASC Summer Student, Los Alamos National Laboratory                                    |
| May 2017 - Jul 2017 | Data-Science at Scale Summer Student, Los Alamos National Laboratory                  |
| May 2017 - Jul 2017 | Co-Design Summer School (6 Students), Los Alamos National Laboratory                  |
| May 2015 - May 2017 | Metropolis Postdoc Fellow (now Staff Scientist), Los Alamos National Laboratory       |
| Aug 2016 - Apr 2017 | Postdoc research assistant (now Staff Scientist), Los Alamos National Laboratory      |

|                     |   |
|---------------------|---|
| Jul 2016 - Sep 2016 | CNLS Summer Student, Los Alamos National Laboratory                             |
| Jun 2016 - Aug 2016 | Co-Design Summer School (7 Students), Los Alamos National Laboratory            |
| Jul 2015 - Sep 2015 | IC Summer Student, Los Alamos National Laboratory                               |
| Jun 2015 - Aug 2015 | Co-Design Summer School (6 Students), Los Alamos National Laboratory            |
| Jul 2014 - Sep 2014 | CNLS Summer Student, Los Alamos National Laboratory                             |
| Jun 2014 - Aug 2014 | Co-Design Summer School (6 Students), Los Alamos National Laboratory            |
| Jul 2013 - Aug 2013 | Co-Design Summer School (6 Students), Los Alamos National Laboratory            |
| Jun 2010 - Aug 2010 | Summer Student, University of Mainz   |
| Oct 2005 - Mar 2006 | Simulation methods in generalized ensembles (2 Students), University of Leipzig |

### *Tutorials (invited)*

|           |  |
|-----------|--|
| Oct. 2016 | “Coarse-Graining with VOTCA” (Tutorial), <a href="#">CECAM Workshop “Multiscale Simulations of Soft Matter with Hands-On Tutorials on ESPResSo++ and VOTCA”</a> , Mainz, Germany |
| May 2016  | Gitlab Hands-On course, ASC/NGC Program, LANL  |
| Oct. 2015 | “Coarse-Graining with VOTCA” (Tutorial), <a href="#">CECAM Summer School “Simulating Soft and Active Matter with ESPResSo, ESPResSo++ and VOTCA”</a> , ICP Stuttgart             |
| Sep. 2015 | Git Hands-On course, ASC/NGC Program, LANL   |
| Nov. 2014 | Git IC Tutorial, Institutional Computing, LANL   |
| Oct. 2014 | Hands-on: VOTCA, <a href="#">CECAM “School on Multiscale Modeling and Use of Espresso++ and VOTCA”</a> , Mainz, Germany  |
| Aug. 2014 | Gromacs Tutorial, <a href="#">The Eighth q-bio Summer School</a> , Albuquerque, NM (LA-UR 14-26188)  |
| June 2014 | Gromacs IC Tutorial, Institutional Computing, LANL (LA-UR 14-24814)  |
| Oct. 2013 | “Systematic Coarse-Graining with VOTCA” (Tutorial), <a href="#">CECAM Summer School “Simulating Soft Matter with ESPResSo, ESPResSo++ and VOTCA”</a> , ICP Stuttgart             |
| July 2013 | Gromacs Tutorial, <a href="#">The Seventh q-bio Summer School</a> , Santa Fe, NM   |
| Oct. 2012 | “Systematic Coarse-Graining with VOTCA” (Tutorial), <a href="#">CECAM Workshop “Simulating Soft Matter with ESPResSo, ESPResSo++ and VOTCA”</a> , ICP Stuttgart                  |
| July 2012 | Gromacs Tutorial, <a href="#">The Sixth q-bio Summer School</a> , Santa Fe, NM   |
| Oct. 2011 | Hands-on: VOTCA, <a href="#">CECAM Workshop “Coarse-grained Simulation of Biological Soft Matter Systems using ESPResSo”</a> , ICP Stuttgart                                     |
| May 2010  | Votca Workshop, <a href="#">CSI Darmstadt</a>  |

## Talks

### *Colloquium Talks*

|           |   |
|-----------|---|
| July 2011 | “Locality Analysis via Adaptive Resolution Simulations”, Theoriekolloquium, <a href="#">Faculty of Natural Sciences II, Martin Luther University Halle-Wittenberg</a> |
|-----------|---|

### *Invited Talks*

|           |  |
|-----------|--|
| Oct. 2016 | “Recent developments in VOTCA”, <a href="#">CECAM Workshop “Multiscale Simulations of Soft Matter with Hands-On Tutorials on ESPResSo++ and VOTCA”</a> , Mainz, Germany          |
| Oct. 2015 | “Recent and Future Developments of VOTCA”, <a href="#">CECAM Summer School “Simulating Soft and Active Matter with ESPResSo, ESPResSo++ and VOTCA”</a> , ICP Stuttgart           |
| Oct. 2014 | “Introduction to coarse-graining of liquids and soft matter using VOTCA”, <a href="#">CECAM “School on Multiscale Modeling and Use of Espresso++ and VOTCA”</a> , Mainz, Germany |
| Oct. 2013 | “Recent and Future Developments of VOTCA”, <a href="#">CECAM Summer School “Simulating Soft Matter with ESPResSo, ESPResSo++ and VOTCA”</a> , ICP Stuttgart                      |
| Oct. 2012 | “Multi-scale modeling using AdResS”, <a href="#">CECAM Workshop “Simulating Soft Matter with ESPResSo, ESPResSo++ and VOTCA”</a> , ICP Stuttgart                                 |

- June 2012 “Molecular Dynamics using Gromacs”, Intel EPOCH, Workshop, Portland, OR
- June 2012 “Towards a Unified Framework for Coarse-graining Particle-based Simulations”, [KITP Conference: “Modeling Soft Matter: Linking Multiple Length and Time Scales”](#), UCSB
- Oct. 2011 “Multi-scale modeling using AdResS”, [CECAM Workshop “Coarse-grained Simulation of Biological Soft Matter Systems using ESPResSo”](#), ICP Stuttgart
- Oct. 2010 “Multi-scale modeling using AdResS”, [CECAM Workshop “Simulating Soft Matter with ESPResSo”](#), ICP Stuttgart
- Nov. 2007 “Controlling material properties using a thermostat”, [CompPhys07](#), ITP Leipzig

#### *Seminar Talks (invited)*

- Sep. 2018 “Co-Design at Los Alamos National Laboratory”, [University of Reims Champagne-Ardenne](#)
- June 2017 “Parallel Runtime Environments with Cloud Database: Performance Study for HMM with Adaptive Sampling”, Nambe Meeting, Los Alamos, NM
- Oct. 2016 “Co-Design at Los Alamos National Laboratory”, [MPCDF](#)
- June 2016 ASC/NGC Gitlab Tutorial, Parallel Computing Summer Research Internship Program, LANL
- Sep. 2012 “Adaptive Resolution Simulations of C60 in Toluene”, [Biomolecular Modeling Group, UMD](#)
- Aug. 2012 “Multi-scale Modeling using the Adaptive Resolution Scheme”, [Sandia National Laboratory](#), Albuquerque, NM
- May 2012 “Particle-based multi-scale simulations using the Adaptive Resolution Scheme”, [IBM Research, Almaden](#)
- May 2012 “Coarse-graining Using the VOTCA Package”, [KITP Program: “Physical Principles of Multiscale Modeling, Analysis and Simulation in Soft Condensed Matter”](#), UCSB
- July 2011 “Locality Analysis via Adaptive Resolution Simulations”, [DFH-UFA & TKM Seminar, Institut für Theoretische Physik, University of Leipzig](#)
- June 2011 “Versatile object-oriented toolkit for coarse-graining applications”, [Department of Materials and Environmental Chemistry, Stockholm University](#)
- Apr. 2011 “Adaptive resolution simulations of aqueous solutions”, Séminaire de Physique, [Biophysique et Physique Statistique, UPV Metz](#)
- Mar. 2011 “Adaptive resolution simulations of water”, [Theoretical and Computational Biophysics Group, UIUC](#)
- Mar. 2011 “Adaptive resolution simulations of aqueous solutions”, [Laufer Center for Physical and Quantitative Biology, Stony Brook University](#)
- Jan. 2011 “Versatile object-oriented toolkit for coarse-graining applications”, [Theoretical and Computational Biophysics Department, MPI-BPC](#)
- Nov. 2009 “Versatile object-oriented toolkit for coarse-graining applications”, Computational Biology Cluster Seminar, IFF-2, [FZ Jülich](#)
- July 2009 “New HPC libraries for performance improvements”, HPC Seminar, IBM Systems & Technology Group
- June 2007 “Extensions to Dissipative Particle Dynamics”, [Faculty of Science, Lund University](#)

#### *Contributed Talks*

- Aug. 2017 “LAMMPS on CMake”, [August 2017 LAMMPS Users’ Workshop and Symposium](#), Albuquerque, NM
- Feb. 2017 “Using Asynchronous Runtimes to Support Multiscale Multiphysics On the Trinity Supercomputer” (with R. Pavel), Programming Models and Co-Design Meeting, Livermore, CA

- Feb. 2016 “Tabasco: A realistic multi-scale proxy app for Material science” (with S. Mniszewski), Programming Models and Co-Design Meeting, Los Alamos, NM
- Feb. 2016 “Actually Using Runtimes for Meaningful Computation: Multiphysics, not Fibonacci” (with R. Pavel), Programming Models and Co-Design Meeting, Los Alamos, NM
- Aug. 2015 “Coarse-graining using VOTCA and LAMMPS”, [August 2015 LAMMPS Users’ Workshop and Symposium](#), Albuquerque, NM
- May 2015 “Parallel Runtime Environments with Cloud Database: Performance Study for HMM with Adaptive Sampling”, [13th Annual Workshop on Charm++ and its Applications](#), Champaign-Urbana
- Mar. 2014 “Discrete Event-based Performance Prediction for Temperature Accelerated Dynamics”, [APS March Meeting 2014](#), Denver
- Sep. 2013 “Adaptive Resolution Simulations of C60 in Toluene”, [Gromacs USA Workshop and Conference](#), UVA
- Aug. 2013 “Introduction to coarse-graining of liquids and soft matter”, [August 2013 LAMMPS Users’ Workshop and Symposium](#), Albuquerque, NM
- Mar. 2013 “Recent developments in the VOTCA package for coarse-graining”, [APS March Meeting 2013](#), Baltimore
- Sep. 2012 “Multi-scale simulations of fullerenes in aqueous solution employing the Adaptive Resolution Scheme”, [IWCMM XXII](#)
- Mar. 2012 “Structure formation of toluene around C60: Application of the Adaptive Resolution Scheme”, [APS March Meeting 2012](#), Boston
- Feb. 2012 “Local structure formation of toluene around a C60 fullerene: An adaptive resolution study”, [25th Annual CSP Workshop](#), [Center for Simulational Physics](#), [University of Georgia](#)
- Nov. 2009 “Versatile object-oriented toolkit for coarse-graining applications”, [CompPhys09](#), [ITP Leipzig](#)
- Nov. 2008 “Comparative atomistic and coarse-grained study of water: simulation details vs. simulation feasibility”, [CompPhys08](#), [ITP Leipzig](#)
- June 2008 “Adjusting dynamic material properties by a thermostat”, [ICMMES 2008](#), [University of Amsterdam](#)
- Feb. 2008 “Controlling material properties using a thermostat”, [DPG Spring Meeting 2008](#) Berlin

## Publications

h-index: [18 \(Google Scholar\)](#), [15 \(ResearcherID\)](#)

### *Reviewed Papers*

- 34. A. Diaw, K. Barros, J. Haack, **C. Junghans**, B. Keenan, Y. W. Li, D. Livescu, N. Lubbers, M. McKerns, R. S. Pavel, D. Rosenberger, I. Sagert, and T. C. Germann, *Multiscale simulation of plasma flows using active learning*, [Phys. Rev. E](#) **102**, 023310 (2020).
- 33. N. Lubbers, A. Agarwal, Y. Chen, S. Son, M. Mehana, Q. Kang, S. Karra, **C. Junghans**, T. C. Germann, and H. S. Viswanathan, *Modeling and scale-bridging using machine learning: nanoconfinement effects in porous media*, [Sci. Rep.](#) **10**, 13312 (2020).
- 32. G. Tirimbo, V. Sundaram, O. Caylak, W. Scharpach, J. Sijen, **C. Junghans**, J. Brown, F. Zapata Ruiz, N. Renaud, J. Wehner, and B. Baumeier, *Excited-State Electronic Structure of Molecules Using Many-Body Green’s Functions: Quasiparticles and Electron-Hole Excitations with VOTCA-XTP*, [J. Chem. Phys.](#) **152**, 114103 (2020).



31. H. V. Guzman, N. Tretyakov, H. Kobayashi, A. C. Fogarty, K. Kreis, J. Krajniak, **C. Junghans**, K. Kremer, and T. Stuehn, *ESPREsSo++ 2.0: Advanced methods for multiscale molecular simulation*, *Comp. Phys. Comm.* **238**, 66 (2019).
30. J. Wehner, L. Brombacher, J. Brown, **C. Junghans**, O. Caylak, Y. Khalak, P. Madhikar, G. Tirimbo, and B. Baumeier, *Electronic Excitations in Complex Molecular Environments: Many-Body Green's Functions Theory in VOTCA-XTP*, *J. Chem. Theo. Comp.* **14**, 6253 (2018).
29. C. Krekeler, A. Agarwal, **C. Junghans**, M. Praprotnik, and L. Delle Site, *Adaptive Resolution Molecular Dynamics Technique: Down to the Essential*, *J. Chem. Phys.* **149**, 024104 (2018).
28. H. V. Guzman, **C. Junghans**, K. Kremer, and T. Stuehn, *Scalable and fast heterogeneous molecular simulation with predictive parallelization schemes*, *Phys. Rev. E* **96**, 053311 (2017).
27. S. K. Gutierrez, K. Davis, D. Arnold, R. S. Baker, R. W. Robey, P. McCormick, D. Holladay, J. A. Dahl, J. Zerr, F. Weik, and **C. Junghans**, *Accommodating Thread-Level Heterogeneity in Coupled Parallel Applications*, in: *Proceedings of 2017 IEEE International Parallel and Distributed Processing Symposium (IPDPS)*, Orlando, FL, USA, 469 (2017).
26. **C. Junghans**, A. Agarwal, and L. Delle Site, *Computational efficiency and Amdahl's law for the adaptive resolution simulation technique*, *Comp. Phys. Comm.* **215**, 20 (2017).
25. T. E. de Oliveira, P. A. Netz, K. Kremer, **C. Junghans**, and D. Mukherji, *C-IBI: Targeting cumulative coordination within an iterative protocol to derive coarse-grained models of (multi-component) complex fluids*, *J. Chem. Phys.* **144**, 174106 (2016).
24. R. S. Pavel, A. L. McPherson, T. C. Germann, and **C. Junghans**, *Database Assisted Distribution to Improve Fault Tolerance for Multiphysics Applications*, in: *Proceedings of the 2nd International Workshop on Hardware-Software Co-Design for High Performance Computing 2015 (Co-HPC '15)*, ACM, New York, NY, USA, Article 4.
23. S. Y. Mashayak, M. N. Jochum, K. Koschke, N. R. Aluru, V. Rühle, and **C. Junghans**, *Relative entropy and optimization-driven coarse-graining methods in VOTCA*, *PLoS one* **10**, e131754 (2015).
22. S. M. Mniszewski, **C. Junghans**, A. F. Voter, D. Perez, and S. J. Eidenbenz, *TADSim: Discrete Event-based Performance Prediction for Temperature Accelerated Dynamics*, *Trans. Mod. Comp. Sim.* **25**, 15 (2015).
21. D. G. Roehm, R. S. Pavel, K. Barros, B. Rouet-Leduc, A. L. McPherson, T. C. Germann, and **C. Junghans**, *Distributed Database Kriging for Adaptive Sampling*, *Comp. Phys. Comm.* **192**, 138 (2015).
20. S. Bevc, **C. Junghans**, and M. Praprotnik, *STOCK: Structure Mapper and Online Coarse-Graining Kit for Molecular Simulations*, *J. Comp. Chem.* **36**, 467 (2015).
19. **C. Junghans**, D. Perez and T. Vogel, *Molecular Dynamics in the Multicanonical Ensemble: Equivalence of Wang-Landau Sampling, Statistical Temperature Molecular Dynamics, and Metadynamics*, *J. Chem. Theo. Comp.* **10**, 1843 (2014).
18. B. Rouet-Leduc, K. Barros, E. Cieren, V. Elango, **C. Junghans**, T. Lookman, J. Mohd-Yusof, R. S. Pavel, A. Y. Rivera, D. Roehm, A. L. McPherson, T. C. Germann, *Spatial adaptive sampling in multiscale simulation*, *Comp. Phys. Comm.* **185**, 1857 (2014).
17. A. Nagarajan, **C. Junghans** and S. Matysiak, *Multiscale simulation of liquid water using a four-to-one mapping for coarse-graining*, *J. Chem. Theo. Comp.* **9**, 5168 (2013).

16. S. Bevc, **C. Junghans**, K. Kremer and M. Praprotnik, *Adaptive resolution simulation of salt solutions*, *New J. Phys.*, **15**, 105007 (2013).
15. P. Ganguly, D. Mukherji, **C. Junghans** and N. F. A. van der Vegt, *Kirkwood-Buff coarse-grained force fields for aqueous solutions*, *J. Chem. Theo. Comp.*, **8**, 1802 (2012).
14. S. Fritsch, S. Poblete, **C. Junghans**, G. Ciccotti, L. Delle Site and K. Kremer, *Adaptive resolution molecular dynamics simulation through coupling to an internal particle reservoir*, *Phys. Rev. Lett.* **108**, 170602 (2012).
13. S. Fritsch, **C. Junghans** and K. Kremer, *Structure formation of toluene around C60: Implementation of the Adaptive Resolution Scheme (AdResS) into GROMACS*, *J. Chem. Theo. Comp.* **8**, 398 (2012).
12. V. Rühle and **C. Junghans**, *Hybrid approaches to coarse-graining using the VOTCA package: liquid hexane*, *Macromol. Theory Simul.*, **20**, 472 (2011).
11. **C. Junghans**, W. Janke and M. Bachmann, *Hierarchies in Nucleation Transitions*, *Comp. Phys. Comm.*, **182**, 1937 (2011).
10. B. P. Lambeth, Jr., **C. Junghans**, K. Kremer, C. Clementi, and L. Delle Site, *Communication: On the Locality of Hydrogen Bond Networks at Hydrophobic Interface*, *J. Chem. Phys.* **133**, 221101 (2010).
9. **C. Junghans** and S. Poblete, *A reference implementation of the adaptive resolution scheme in ESPResSo*, *Comp. Phys. Comm.* **181**, 1449 (2010).
8. V. Rühle, **C. Junghans**, A. Lukyanov, K. Kremer and D. Andrienko, *Versatile Object-oriented Toolkit for Coarse-graining Applications*, *J. Chem. Theo. Comp.* **5**, 3211 (2009).
7. **C. Junghans**, M. Bachmann and W. Janke, *Statistical Mechanics of Aggregation and Crystallization for Semiflexible Polymers*, *Europhys. Lett.* **87**, 40002 (2009).
6. H. Wang, **C. Junghans** and K. Kremer, *Comparative atomistic and coarse-grained study of water: What do we lose by coarse-graining?*, *Euro. Phys. J. E* **28**, 221 (2009). “Most cited article published in the journal in the past 5 years” (2014)
5. M. Praprotnik, **C. Junghans**, L. Delle Site and K. Kremer, *Simulation approaches to soft matter: Generic statistical properties vs. chemical details*, *Comp. Phys. Comm.* **179**, 51 (2008).
4. **C. Junghans**, M. Bachmann and W. Janke, *Thermodynamics of Peptide Aggregation Processes: An Analysis from Perspectives of Three Statistical Ensembles*, *J. Chem. Phys.* **128**, 085103 (2008) .
3. **C. Junghans**, M. Praprotnik and K. Kremer, *Transport properties controlled by a thermostat: An extended dissipative particle dynamics thermostat*, *Soft Matter* **4**, 156 (2008).
2. **C. Junghans**, M. Bachmann and W. Janke, *Microcanonical Analyses of Peptide Aggregation Processes*, *Phys. Rev. Lett.* **97**, 218103 (2006).
1. **C. Junghans** and U. H. E. Hansmann, *Numerical Comparison of Wang Landau Sampling and Parallel Tempering for Met-enkephalin*, *Int. J. Mod. Phys. C* **17**, 817 (2006).

More information can be found on [my ResearcherID profile](#), [my Google Scholar profile](#), [my ORCID profile](#), [my ResearchGate profile](#) or [my Impactstory profile](#).



*Book Chapters*

3. L. Delle Site, A. Agarwal, **C. Junghans**, and H. Wang, *Adaptive Resolution Simulation as a Grand Canonical Molecular Dynamics Scheme: Principles, Applications, Perspectives*, in: C. Clementi (Ed.), *Multiscale Methods in Molecular Biophysics*, Series in Computational Biophysics, CRC Press (2016), ISBN 1482225700.
2. E. Apol, R. Apostolov, H.J.C. Berendsen, A. van Buuren, P. Bjelkmar, R. van Drunen, A. Feenstra, S. Fritsch, G. Groenhof, **C. Junghans**, P. Kasson, P. Larsson, P. Meulenhoff, T. Murtola, S. Pall, S. Pronk, R. Schulz, M. Shirts, A. Sijbers, P. Tieleman, B. Hess, D. van der Spoel, and E. Lindahl, *Gromacs User Manual Version 4.6*, [gromacs.org](http://gromacs.org) (2013).
1. **C. Junghans**, M. Praprotnik and L. Delle Site, *Adaptive Resolution Schemes*, in: J. Grotendorst, N. Attig, S. Blügel and D. Marx (Eds.), *Multiscale Simulation Methods in Molecular Sciences*, **NIC Series Vol. 42**, Jülich (2009), 359.

*Other Publications*

6. *Balancing the load: Los Alamos researchers develop code to distribute computation more efficiently and across increasing numbers of supercomputer processors*. [ASCR Discovery Story](#), [May 2018](#), [HPCwire](#), [insideHPC](#) and [LANL Science Brief](#)
5. **C. Junghans**, A. K. Hüttel and U. Müller, *Gentoo Linux: Quelltexte und Rolling Releases*, [c't Magazin](#) **16**, 162 (2012).
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