

Aalborg Universitet

Interatomic potential parameterization using particle swarm optimization: Case study of glassy silica

Christensen, Rasmus; Sørensen, Søren Strandskov; Liu, Han; Li, Kevin; Bauchy, Mathieu; Smedskjær, Morten Mattrup

Publication date: 2022

Document Version Other version

Link to publication from Aalborg University

Citation for published version (APA): Christensen, R., Sørensen, S. S., Liu, H., Li, K., Bauchy, M., & Smedskjær, M. M. (2022). *Interatomic potential* parameterization using particle swarm optimization: Case study of glassy silica. Poster presented at 26th International Congress on Glass, Berlin, Germany.

Copyright and moral rights for the publications made accessible in the public portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

- Users may download and print one copy of any publication from the public portal for the purpose of private study or research.
- You may not further distribute the material or use it for any profit-making activity or commercial gain
 You may freely distribute the URL identifying the publication in the public portal -

Take down policy
If you believe that this document breaches copyright please contact us at vbn@aub.aau.dk providing details, and we will remove access to the work immediately and investigate your claim.

Interatomic potential parameterization using particle swarm optimization: Case study of glassy silica

R. Christensen^{1*}, S. S. Sørensen¹, H. Liu², K. Li², M. Bauchy², M. M. Smedskjaer¹

Classical molecular dynamics simulations of glassy materials rely on the availability of accurate yet computationally efficient interatomic force fields. The parameterization of new potentials remains challenging due to the non-convex nature of the accompanying optimization problem, which renders the traditional optimization methods inefficient or subject to bias. In this study, we present a new parameterization method based on particle swarm optimization (PSO), which is a stochastic population-based optimization method. Using glassy silica as a case study, we introduce two interatomic potentials using PSO, which are parameterized so as to match structural features obtained from ab initio simulations and experimental neutron diffraction data. We find that the PSO algorithm is highly efficient at searching for and identifying viable potential parameters that reproduce the structural features used as the target in the parameterization. The presented approach is very general and can be easily applied to other interatomic potential parameterization schemes.

¹ Department of Chemistry and Bioscience, Aalborg University, DK-9220 Aalborg, Denmark.,

² Department of Civil and Environmental Engineering, University of California, Los Angeles, California 90095, USA.

^{*}rasmusc@bio.aau.dk