## Igor Yu Gotlib

List of Publications by Year in descending order

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ICOP YU COTUR

| #  | Article                                                                                                                                                                                  | IF  | CITATIONS |
|----|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----|-----------|
| 1  | Properties of coexisting fluid phases of a binary system methanolî—,ethane by computer simulation. Fluid<br>Phase Equilibria, 1997, 129, 1-13.                                           | 2.5 | 12        |
| 2  | Computer simulation of ionic transport in silver iodide within carbon nanotubes. Solid State Ionics, 2011, 188, 6-14.                                                                    | 2.7 | 10        |
| 3  | Properties of Coexisting Phases for the Ethanolâ^'Ethane Binary System by Computer Simulation.<br>Journal of Physical Chemistry B, 1999, 103, 7681-7686.                                 | 2.6 | 7         |
| 4  | Molecular Dynamics Simulation of Poly(p-xylylene):  Bulk Phase and a Single Molecule. Journal of<br>Physical Chemistry C, 2007, 111, 6613-6620.                                          | 3.1 | 6         |
| 5  | Association Equilibrium for Cross-Associating Chains in a Good Solvent: Crowding and Other<br>Nonideality Effects. Journal of Physical Chemistry B, 2016, 120, 7234-7243.                | 2.6 | 6         |
| 6  | Structure and Ionic Transport Properties of Agl1–xBrxwithin Single-Wall Carbon Nanotubes from<br>Molecular Dynamics Simulation. Journal of Physical Chemistry C, 2012, 116, 19554-19570. | 3.1 | 5         |
| 7  | Molecular Thermodynamic Modeling of Self-Assembly into Branches and Spatial Networks in Solution. Journal of Chemical & Engineering Data, 2016, 61, 4013-4022.                           | 1.9 | 5         |
| 8  | Title is missing!. Inorganic Materials, 2001, 37, 975-979.                                                                                                                               | 0.8 | 4         |
| 9  | Molecular Dynamics Simulations of Li x Mn2O4 Spinel Solid Solutions with Simple Potential Models.<br>Inorganic Materials, 2003, 39, 404-408.                                             | 0.8 | 4         |
| 10 | Steric asymmetry vs charge asymmetry in dilute solution containing large weakly charged ions. Fluid<br>Phase Equilibria, 2016, 428, 203-211.                                             | 2.5 | 4         |
| 11 | Molecular dynamics simulation of the Ba1â^'xGdxF2+x system in a wide temperature range. Solid State<br>Ionics, 2003, 159, 49-62.                                                         | 2.7 | 3         |
| 12 | Molecular dynamics simulation of fluorite- and tysonite-type solid electrolytes. Computational<br>Materials Science, 2006, 36, 73-78.                                                    | 3.0 | 3         |
| 13 | Title is missing!. Inorganic Materials, 2003, 39, 291-298.                                                                                                                               | 0.8 | 2         |
| 14 | Molecular dynamics simulation of silver bromide nanostructures in single-walled carbon nanotubes.<br>Physics of the Solid State, 2011, 53, 2375-2384.                                    | 0.6 | 2         |
| 15 | Effects of the linear chain structure on the cross-association equilibrium between chainlike molecules in a good solvent. Fluid Phase Equilibria, 2017, 454, 116-121.                    | 2.5 | 2         |
| 16 | Computer simulation of Agl nanostructures in single-wall carbon nanotubes. Inorganic Materials, 2010, 46, 1375-1383.                                                                     | 0.8 | 1         |
| 17 | Molecular dynamics simulation of SnF2 nanostructures in the internal channels of single-walled carbon nanotubes. Physics of the Solid State, 2014, 56, 1472-1482.                        | 0.6 | 1         |
| 18 | Association Equilibrium and Gelation in Solutions of Cross-Associating Chains Containing Inactive Fragments. Polymer Science - Series C, 2018, 60, 95-105.                               | 1.7 | 1         |

| #  | Article                                                                                                                                  | IF  | CITATIONS |
|----|------------------------------------------------------------------------------------------------------------------------------------------|-----|-----------|
| 19 | Association kinetics and equilibrium in solutions of cross-associating chains that contain inactive spacers. Polymer, 2020, 187, 122085. | 3.8 | 0         |