Hitoshi Goto

List of Publications by Year in descending order

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68 papers 2,533 citations

304743

22

h-index

189892 50 g-index

71 all docs

71 docs citations

71 times ranked

2458 citing authors

#	Article	IF	CITATIONS
1	Preparation of photonic molecular trains via soft-crystal polymerization of lanthanide complexes. Nature Communications, 2022, 13, .	12.8	7
2	Crystal Structure Optimization Method for Evaluation of Mechanical Properties of Soft Crystals. Nihon Kessho Gakkaishi, 2021, 63, 63-68.	0.0	1
3	Improving the Accuracy of Crystal Structure Prediction Using FMO Crystal Energy: An Example of Target XXIII. Journal of Computer Chemistry Japan, 2021, 20, 92-93.	0.1	O
4	Validation Study of QSAR/DNN Models Using the Competition Datasets. Molecular Informatics, 2020, 39, 1900154.	2.5	10
5	Selective Formation and SHG Intensity of Noncentrosymmetric and Centrosymmetric $1,1,2,2$ -Tetramethyl- $1-(4-(N,N-dimethylamino)phenyl)-2-(2\hat{a}\in \mathbb{Z}^2-cyanophenyl)disilane Crystals under External Stimuli. Journal of Physical Chemistry C, 2020, 124, 17450-17458.$	3.1	13
6	Thermosalience in Macrocycle-Based Soft Crystals via Anisotropic Deformation of Disilanyl Architecture. Journal of the American Chemical Society, 2020, 142, 12651-12657.	13.7	44
7	Charge mobility calculation of organic semiconductors without use of experimental single-crystal data. Scientific Reports, 2020, 10, 2524.	3.3	13
8	Crystal Structure Prediction; From the Present to the Future. Nihon Kessho Gakkaishi, 2020, 62, 260-268.	0.0	1
9	Computational chemical analysis of Ru(II)â€Pheox–catalyzed highly enantioselective intramolecular cyclopropanation reactions. Chirality, 2019, 31, 52-61.	2.6	14
10	Soft Crystal Force Field for Reproducing the Crystal Structures of Aryl Gold Isocyanide Complexes. Journal of Computer Chemistry Japan, 2018, 17, 155-157.	0.1	2
11	Highly stereoselective spirocyclopropanation of various diazooxindoles with olefins catalyzed using Ru(<scp>ii</scp>)-complex. RSC Advances, 2018, 8, 39865-39869.	3.6	19
12	Ligand Exchange Reaction on a Ru(II)–Pheox Complex as a Mechanistic Study of Catalytic Reactions. ACS Omega, 2018, 3, 11286-11289.	3.5	10
13	Numerical calculation on a two-step subdiffusion behavior of lateral protein movement in plasma membranes. Physical Review E, 2017, 96, 042410.	2.1	3
14	Chiral Olefin and Molecular Motor. Journal of Computer Chemistry Japan, 2017, 16, A21-A22.	0.1	0
15	Investigation of a Virtual Nested Two-dimensional Lattice Model for Representing the Diffusive Motion of a Transmembrane Protein in Cell Membrane. Journal of Computer Chemistry Japan, 2016, 15, 229-231.	0.1	1
16	Molecular activity prediction using deep learning software library. , 2016, , .		8
17	Report on the sixth blind test of organic crystal structure prediction methods. Acta Crystallographica Section B: Structural Science, Crystal Engineering and Materials, 2016, 72, 439-459.	1.1	445
18	High-speed prediction of crystal structures for organic molecules. AIP Conference Proceedings, 2015,	0.4	11

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19	Theoretical Electronic Circular Dichroism Study of 1,3â€Diene Derivatives for the Elucidation of ECD Spectra of 1,3â€Cyclohexadiene and Its Derivatives. Chirality, 2015, 27, 476-478.	2.6	2
20	Development of Visualization Technique of Biological Information for Mobile Terminal Devices. Journal of Computer Chemistry Japan, 2014, 13, 175-176.	0.1	0
21	Development of Education Curriculum for First Level Simulation Engineering in Chemistry - Questionnaire Research and Core-Curriculum Proposal Journal of Computer Chemistry Japan, 2014, 13, 179-180.	0.1	0
22	Nomenclature of Aldohexopyranose Conformations IncludingRotational Isomers of Hydroxyl Groups. Journal of Computer Chemistry Japan, 2014, 13, 263-267.	0.1	0
23	Vacancy profile in reverse osmosis membranes studied by positron annihilation lifetime measurements and molecular dynamics simulations. Journal of Physics: Conference Series, 2013, 443, 012050.	0.4	7
24	Mode-selective internal conversion of perylene. Molecular Physics, 2011, 109, 1831-1840.	1.7	14
25	Geometrical structure of benzene and naphthalene: Ultrahigh-resolution laser spectroscopy and <i>ab</i> 倉 <i>initio</i> calculation. Journal of Chemical Physics, 2011, 135, 054305.	3.0	51
26	Journal of Computer Chemistry, Japan:Conversion of Japanese Word Manuscript into the XML Format and Web Publication Management System. Journal of Computer Chemistry Japan, 2011, 10, 141-146.	0.1	3
27	Ultrahigh-resolution laser spectroscopy of the S1 1B2↕S0 1A transition of perylene. Journal of Molecular Spectroscopy, 2010, 260, 72-76.	1.2	14
28	Structure and excited-state dynamics of anthracene: Ultrahigh-resolution spectroscopy and theoretical calculation. Journal of Chemical Physics, 2009, 130, 134315.	3.0	47
29	Conformational Polymorphism Analysis of Aspirin Crystal with a Crystal Calculation Method. Journal of Computer Chemistry Japan, 2008, 7, 151-164.	0.1	13
30	Parallelization of Crystal Calculation for Large-Scale Molecular Crystal Structure Analysis. Journal of Computer Aided Chemistry, 2008, 9, 8-16.	0.3	15
31	Drug discovery using grid technology. , 2006, , 227-248.		2
32	Molecular mechanics (CONFLEX/MM3) search/minimization study of the conformations of ornoside and escuside. Journal of Molecular Structure, 2005, 748, 9-16.	3.6	3
33	Proposed mechanism for diterpene synthases in the formation of phomactatriene and taxadiene. Organic and Biomolecular Chemistry, 2005, 3, 2713.	2.8	30
34	Implementation and performance evaluation of CONFLEX-G. , 2004, , .		3
35	C100 IPR fullerenes: temperature-dependent relative stabilities based on the Gibbs function. Chemical Physics, 2004, 306, 93-104.	1.9	40
36	Theoretical Studies on the Relative Stabilities of C96IPR Fullerenes. Journal of Physical Chemistry A, 2004, 108, 4479-4484.	2.5	40

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37	Synthesis, Resolution, and Absolute Stereochemistry of (â^')-Blestriarene C. Journal of Organic Chemistry, 2003, 68, 2099-2108.	3.2	62
38	Theoretical investigations on relative stabilities of fullerene C94. Journal of Chemical Physics, 2003, 118, 10534-10540.	3.0	26
39	Continuously Growing Spiral Carbon Nanoparticles as the Intermediates in the Formation of Fullerenes and Nanoonions. Journal of Physical Chemistry B, 2002, 106, 7135-7138.	2.6	51
40	Synthesis, Absolute Configuration, and Enantiomeric Enrichment of a Cruciferous Oxindole Phytoalexin, (S)-(â^')-Spirobrassinin, and Its Oxazoline Analog. Journal of Organic Chemistry, 2001, 66, 3940-3947.	3.2	170
41	Cooperative Enhancement of Water Binding to Crownophane by Multiple Hydrogen Bonds:  Analysis by High Level ab Initio Calculations. Journal of the American Chemical Society, 2001, 123, 4255-4258.	13.7	48
42	Spiral carbon nanoparticles. AIP Conference Proceedings, 2001, , .	0.4	0
43	Computing the relative gas-phase populations of C60 and C70: beyond the traditional ÎHf,298o scale. Journal of Molecular Graphics and Modelling, 2001, 19, 216-221.	2.4	29
44	Chemistry of Unique Chiral Olefins. 4. Theoretical Studies of the Racemization Mechanism oftrans- andcis-1,1 ,2,2 ,3,3 ,4,4 -Octahydro-4,4 -biphenanthrylidenes. Journal of Organic Chemistry, 1999, 64 1667-1674.	,3.2	36
45	A frontier mode-following method for mapping saddle points of conformational interconversion in flexible molecules starting from the energy minimum. Chemical Physics Letters, 1998, 292, 254-258.	2.6	13
46	Absolute configuration of chiral fullerenes and covalent derivatives from their calculated circular dichroism spectra. Journal of the Chemical Society Perkin Transactions II, 1998, , 1719-1724.	0.9	40
47	Is the CD Exciton Chirality Method Applicable to Chiral $1,1\hat{a}\in \tilde{\ }$ -Biphenanthryl Compounds?. Journal of the American Chemical Society, 1998, 120, 9086-9087.	13.7	31
48	Cluster conformational analysis of a seco acid used in Woodward's total synthesis of erythromycin A. Computational and Theoretical Chemistry, 1997, 398-399, 229-236.	1.5	9
49	Electronic structures and redox properties of silylmethylated C60. Tetrahedron, 1996, 52, 5053-5064.	1.9	13
50	The phaseon line: a new criterion for the stability of fullerenes., 1996, 1, 151-162.		3
51	Prediction of favorable isomeric structures for the C $<$ sub $>$ 100 $<$ /sub $>$ to C $<$ sub $>$ 120 $<$ /sub $>$ giant fullerenes. An application of the phason line criteria. , 1996, 1, 163-171.		13
52	Conformational Analysis of Long Chain Seco-Acids Used in Woodward's Total Synthesis of Erythromycin A — Conformational Space Search as the Basis of Molecular Modeling. , 1996, , 189-197.		1
53	Photoinduced molecular transformations. Part 155. General synthesis of macrocyclic Retones based on a ring expansion involving a selective β-scission of alkoxyl radicals, its application to a new synthesis of (±)-isocaryophyllene and (±)-caryophyllene, and a conformational analysis of the two sesquiterpenes and the radical intermediate in the synthesis by MM3 calculations. Journal of the	0.9	24
54	Chemical Society Perkin Transactions 1, 1995, , 69-81. How many conformers are there for small n-alkanes? Consequences of asymmetric deformation in GC′ segment. Tetrahedron, 1993, 49, 387-396.	1.9	57

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55	Viewpoint $11~\hat{a}$ e" approaches to the global minimum problem. Computational and Theoretical Chemistry, 1993, 285, 157-168.	1.5	42
56	Combined molecular mechanics (MM2) and molecular orbital (AM1) study of reriplanone-a and analogues. Evaluation of biological activity from electronic properties and geometries. Part 2. Journal of the Chemical Society Perkin Transactions II, 1993, , 1167-1173.	0.9	3
57	An efficient algorithm for searching low-energy conformers of cyclic and acyclic molecules. Journal of the Chemical Society Perkin Transactions II, 1993, , 187-198.	0.9	351
58	Origin of regioselectivity in the O-methylation of erythromycin as elucidated with the aid of computational conformational space search. Journal of the Chemical Society Perkin Transactions II, 1993, , 1647-1654.	0.9	13
59	Combined molecular mechanics (MM2) and molecular orbital (AM1) study of periplanone-B and analogues. Evaluation of biological activity from electronic properties and geometries. Journal of the Chemical Society Perkin Transactions II, 1992, , 811-818.	0.9	7
60	A revised nomenclature for the ring conformation and a note on the conformational distance in cyclododecane. Tetrahedron, 1992, 48, 7131-7144.	1.9	14
61	Investigation of intramolecular interactions in n-alkanes. Cooperative energy increments associated with GG and GTG' [G = gauche, T = trans] sequences. Journal of the American Chemical Society, 1991, 113, 4665-4671.	13.7	105
62	Chiral Synthesis of Polyketide-Derived Natural Products. 33. Stereoselective Total Synthesis of 16-Membered Macrolide Aglycons, Leuconolides and Maridonolides. Macrocyclic Stereocontrol Based on Conformational Analysis of the 16-Membered Macrolide Ring Chemical and Pharmaceutical Bulletin, 1991, 39, 2819-2829.	1.3	4
63	Conformational analyses of periplanone analogs by molecular mechanics calculations. Journal of Chemical Ecology, 1991, 17, 779-788.	1.8	5
64	FAST ALGORITHM FOR COVERING CONFORMATIONAL SPACE BY MOLECULAR MECHANICS., 1991, , 411-416.		2
65	Conformational analysis of serricornin: Application of molecular mechanics calculations to stereochemical assignment of serricornin, sex pheromone of cigarette beetle (Lasioderma serricorne) Tj ETQq1 1	0. 7.8 4314	· rgBT /Overl
66	Chiral synthesis of polyketide-derived natural products. 28. Synthesis of 16-membered macrolide aglycons, carbonolide A, leuconolides, and maridonolides, via carbonolide B type compounds by virtue of completely stereoselective epoxidation and reduction based on the conformational control of macrolide rings with protecting groups. Journal of Organic Chemistry, 1990, 55, 1129-1132.	3.2	12
67	Corner flapping: a simple and fast algorithm for exhaustive generation of ring conformations. Journal of the American Chemical Society, 1989, 111, 8950-8951.	13.7	445
68	Application of molecular mechanics to natural product chemistry. Pure and Applied Chemistry, 1989, 61, 597-600.	1.9	12