## Lingpeng Meng

List of Publications by Year in descending order

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93 papers

1,141 citations

20 h-index 28 g-index

93 all docs 93
docs citations

93 times ranked 726 citing authors

#	Article	IF	CITATIONS
1	Photocatalytic Reduction of CO <sub>2</sub> with H <sub>2</sub> O Mediated by Ce-Tailored Bismuth Oxybromide Surface Frustrated Lewis Pairs. ACS Catalysis, 2022, 12, 4016-4025.	11.2	95
2	Comparative studies on group III $ f $ -hole and $ f $ -hole interactions. Journal of Computational Chemistry, 2016, 37, 1321-1327.	3.3	49
3	Metal–Metal and Metal–Ligand Bonds in (Î- <sup>5</sup> -C <sub>5</sub> H <sub>5</sub> ) <sub>2</sub> M <sub>2</sub> (M = Be, Mg, Ca, Ni, Cu, Zn). Organometallics, 2013, 32, 1060-1066.	2.3	43
4	A computational study on the nature of the halogen bond between sulfides and dihalogen molecules. Structural Chemistry, 2011, 22, 567-576.	2.0	40
5	Insight into the pseudo Ï€-hole interactions in the M <sub>3</sub> H <sub>6</sub> â<¯(NCF) <sub>n</sub> (M) Tj ET	[Qq1 1 0.7	(84314 rg <mark>BT</mark>
6	The enhancing effects of group V $l$ f-hole interactions on the Fâ $l$ O halogen bond. Physical Chemistry Chemical Physics, 2014, 16, 19282-19289.	2.8	29
7	The cooperativity between the $\ddot{l}f$ -hole and $\ddot{l}\in$ -hole interactions in the ClO···XONO2/XONO···NH3 (XÂ=ÂCl, complexes. Structural Chemistry, 2015, 26, 213-221.	Br. 1) 2.0	29
8	Topological Characteristics of Electron Density Distribution in SSXY â†' XSSY (X or Y = F, Cl, Br, I) Isomerization Reactions. Journal of Physical Chemistry A, 2007, 111, 9093-9101.	2.5	27
9	N···I Halogen Bonding Interactions: Influence of Lewis Bases on Their Strength and Characters. Journal of Physical Chemistry A, 2014, 118, 7058-7065.	2.5	27
10	The competition of Y⋯o and X⋬n halogen bonds to enhance the group V σ-hole interaction in the NCY⋬oPH <sub>3</sub> ⋬NCX and OPH <sub>3</sub> ⋬NCX⋬NCY (X, YF, Cl, and Br) complexes. J Computational Chemistry, 2015, 36, 1349-1358.	on of	27
11	Studies on Reactions INCX → IXCN (X = O, S, and Se). Inorganic Chemistry, 2004, 43, 5311-5320.	4.0	26
12	Insight into the lithium/hydrogen bonding in (CH2)2XLiY/HY (X: C=CH2, O, S; Y=F, Cl, Br) complexes. Journal of Molecular Modeling, 2011, 17, 757-767.	1.8	26
13	Insight into the nature of the interactions of furan and thiophene with hydrogen halides and lithium halides: ab initio and QTAIM studies. Journal of Molecular Modeling, 2011, 17, 2907-2918.	1.8	24
14	The Role of Molecular Electrostatic Potentials in the Formation of a Halogen Bond in Furanâ‹â‹â‹XY and Thiopheneâ‹â‹â‹XY Complexes. ChemPhysChem, 2011, 12, 1080-1087.	2.1	24
15	A comprehensive analysis of P···π pnicogen bonds: substitution effects and comparison with Br···π halogen bonds. Journal of Molecular Modeling, 2015, 21, 143.	1.8	24
16	Theoretical Investigation on Stability and Isomerizations of CH3SO Isomers. Journal of Physical Chemistry A, 2007, 111, 2343-2350.	2.5	23
17	First-principles studies of the structures and properties of Al- and Ag-substituted Mg2Ni alloys and their hydrides. International Journal of Hydrogen Energy, 2010, 35, 10349-10358.	7.1	23
18	Organocatalysis by Halogen, Chalcogen, and Pnictogen Bond Donors in Halide Abstraction Reactions: An Alternative to Hydrogen Bond-Based Catalysis. Journal of Physical Chemistry A, 2020, 124, 3815-3824.	2.5	23

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19	Interplay between halogen bonds and hydrogen bonds in OH/SH···HOX···HY (X = Cl, Br; Y = F, Cl, Br) complexes. Journal of Molecular Modeling, 2013, 19, 1069-1077.	1.8	21
20	Ab initio and AIM studies on typical Ï€â€type and pseudoâ€Ï€â€type halogen bonds: Comparison with hydrogen bonds. International Journal of Quantum Chemistry, 2011, 111, 3725-3740.	2.0	20
21	The Role of Ï€ Electrons in the Formation of Benzeneâ€Containing Lithiumâ€Bonded Complexes. ChemPhysChem, 2011, 12, 3584-3590.	2.1	19
22	Effect of Sr substitution on the property and stability of CH <sub>3</sub> NH <sub>3</sub> Snl <sub>perovskite: A firstâ€principles investigation. International Journal of Energy Research, 2020, 44, 5765-5778.</sub>	4.5	19
23	Influence of the Liâ‹â‹â‹ï∈ Interaction on the H/Xâ‹â‹â‹ï∈ Interactions in HOLiâ‹â‹â‹C‹sub>6⟨/sub>H⟨sub>â‹â‹â‹HOX/XOH (X=F, Cl, Br, I) Complexes. ChemPhysChem, 1591-1600.	<b>2</b> 013, 14,	18
24	The mutual influence between Ï€-hole pnicogen bonds and σ-hole halogen bonds in complexes of PO2Cl and XCN/C6H6 (XÂ=ÂF, Cl, Br). Structural Chemistry, 2016, 27, 1427-1437.	2.0	18
25	Topological Characterization of HXO2 (X = Cl, Br, I) Isomerization. Journal of Physical Chemistry A, 2007, 111, 1530-1535.	2.5	17
26	Assessment of intermolecular interactions at three sites of the arylalkyne in phenylacetyleneâ€containing lithiumâ€bonded complexes: Ab initio and QTAIM studies. Journal of Computational Chemistry, 2012, 33, 1321-1327.	3.3	15
27	Improvement in dehydrogenation performance ofÂMg(BH4)2·2NH3 doped with transition metal: First-principles investigation. International Journal of Hydrogen Energy, 2015, 40, 8721-8731.	7.1	15
28	Understanding the Properties of Inorganic Benzenes Based on π-Electron Densities. Journal of Physical Chemistry A, 2015, 119, 2091-2097.	2.5	15
29	Influences of the substituents on the M–M bonding in Cp <sub>4</sub> Al <sub>4</sub> and Cp <sub>2</sub> M <sub>2</sub> X <sub>2</sub> (M = B, Al, Ga; Cp = C <sub>5</sub> H <sub>5</sub> , X =) Tj ETC	2 <b>գ</b> 131 0.78	4 <b>3</b> 14 rgB
30	Comparison of Anionâ€Anion Halogen Bonds with Neutralâ€Anion Halogen Bonds in the Gas Phase and Polar Solvents. ChemPlusChem, 2021, 86, 232-240.	2.8	15
31	AIM Studies on Reactions FNCX → FXCN (X = O, S, and Se). Journal of Physical Chemistry A, 2004, 108, 10527-10534.	2.5	14
32	Topological studies on IRC paths of X+H2?XH+H reactions. Journal of Computational Chemistry, 1997, 18, 1167-1174.	3.3	13
33	Substituent effects on the intramolecular hydrogen bond in 1-hydroxyanthraquinone: AIM and NBO analyses. Structural Chemistry, 2012, 23, 1233-1240.	2.0	13
34	Insight into the Ï€â€hole···πâ€electrons tetrel bonds between F <sub>2</sub> ZO (Z = C, Si, Ge) and unsaturated hydrocarbons. International Journal of Quantum Chemistry, 2018, 118, e25521.	2.0	13
35	Predicting the halogenâ€ <i>n</i> ( <i>n</i> = 3–6) synthons to form the "windmill―pattern bonding based on the halogenâ€bonded interactions. Journal of Computational Chemistry, 2019, 40, 1219-1226.	3.3	13
36	Influences of Al, Ti and Nb doping on the structure and hydrogen storage property of Mg(BH 4 ) 2 (001) surface – A theoretical study. International Journal of Hydrogen Energy, 2015, 40, 10516-10526.	7.1	12

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37	Enhancing Ïf∫Ï€-type copper( <scp>i</scp> )â <rthiophene (metal="Li," 1283-1291.<="" 2015,="" 44,="" by="" ca,="" dalton="" doping="" interactions="" k,="" metal="" na,="" sc).="" td="" transactions,=""><td>3.3</td><td>12</td></rthiophene>	3.3	12
38	Coinage metal dimers as the noncovalent interaction acceptors: study of the $lf$ -lump interactions. Physical Chemistry Chemical Physics, 2019, 21, 21152-21161.	2.8	11
39	Cooperativity between S···π and Rg···π in the OCS···C <sub>6</sub> H <sub>6</sub> ····Rg (Rg = He	, Ne, Ar, a 2.5	nd) Tj ETQq
40	Electron Structure of Arx′ZnZnArx′ (Arx′ = C <sub>6</sub> H <sub>3</sub> -2,) Tj ETQq0 0 0 rgBT /Overloo Hydrogen/Sodium Atoms on Zn–Zn Interactions. Organometallics, 2012, 31, 6582-6588.	ck 10 Tf 50 2.3	0 627 Td (6 9
41	First-principles study of hydrogen vacancies in lithium amide doped with titanium and niobium. International Journal of Hydrogen Energy, 2013, 38, 11303-11312.	7.1	9
42	Dinuclear first-row transition metal–(C8Me6)2complexes: metal–metal and metal–ligand bonds determined by the d electron configuration of the metal atom. New Journal of Chemistry, 2016, 40, 1988-1996.	2.8	9
43	Comparison of pnicogen and tetrel bonds in complexes containing CX2 carbenes (X = F, Cl, Br, OH,) Tj ETQq1 1 0.	784314 rg 2.8	gBT /Overlo
44	Discovery of Ïf-hole interactions involving ylides. Journal of Molecular Modeling, 2013, 19, 4887-4895.	1.8	8
45	Influence of titanium and nickel dopants on the dehydrogenation properties of Mg(AlH4)2: Electronic structure mechanisms. International Journal of Hydrogen Energy, 2014, 39, 9276-9287.	7.1	8
46	Effects of Mg and Al doping on the electronic structure and dehydrogenation of LiBH4·NH3. International Journal of Hydrogen Energy, 2014, 39, 17144-17152.	7.1	8
47	Nature of the M–M bonding (MÂ=ÂCr, Mo, and W) in [CpM(CO)3]2: Covalent single bond or noncovalent interaction?. Journal of Organometallic Chemistry, 2014, 769, 106-111.	1.8	7
48	Influence of Cr/Zr doping on the electronic structure and hydrogen storage properties of the Mg2Ni (010) surface: A first principles study. Journal of Alloys and Compounds, 2014, 601, 280-288.	<b>5.</b> 5	7
49	Aromaticity analysis of fused heterometallacycles containing M M(M = Cr, Mo and W) quintuple bond. Chemical Physics Letters, 2019, 731, 136600.	2.6	7
50	The mechanism of ringâ€opening polymerization of Lâ€lactide by ICl <sub>3</sub> catalysts: Halogen bond catalysis or participating in reactions?. Journal of Computational Chemistry, 2019, 40, 2827-2833.	3.3	7
51	Mechanism and Stereoselectivity of the Elementometalation Process of Activated Alkyne RCCR(RCO 2) Tj	<u></u> ξТQq1 1	0 <sub>7</sub> 784314 r
52	Improving the stability of perovskite by covering graphene on <scp> FAPbI <sub>3</sub> </scp> surface. International Journal of Energy Research, 2021, 45, 10808-10820.	4.5	7
53	Enhancing the stability of perovskites by constructing heterojunctions of graphene/MASnI <sub>3</sub> . Physical Chemistry Chemical Physics, 2020, 22, 3724-3733.	2.8	6
54	Control of the Regioselectivity of Alkyne Hydrostannylation by Tuning the Metal Pair of Heterobimetallic Catalysts: A Theoretical Study. Organometallics, 2021, 40, 654-662.	2.3	6

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55	Mechanism and origin of the stereoselectivity of manganese-catalyzed hydrosilylation of alkynes: a DFT study. Catalysis Science and Technology, 2022, 12, 2649-2658.	4.1	6
56	A DFT study of the generation of interstellar species XN (X=Cl, and Br) activated by molecular sieve clusters. Chemical Physics Letters, 2004, 400, 394-400.	2.6	5
57	Regulatory factors and the nature of Cuâ< Cu interaction in copper(i) complexes with NHC and NHCP ligands: a theoretical assessment. New Journal of Chemistry, 2014, 38, 5786-5792.	2.8	5
58	Mechanism and kinetics for the reactions of methacrolein and methyl vinyl ketone with HO <sub>2</sub> radical. New Journal of Chemistry, 2017, 41, 7714-7722.	2.8	5
59	Electronic structure of triangular M <sub>3</sub> (M = B, Al, Ga): nonclassical three-center two electron $\tilde{I} \in \text{bond}$ and $\tilde{I} \in \text{f}$ delocalization. Physical Chemistry Chemical Physics, 2020, 22, 18071-18077.	2.8	5
60	Effects of halogen substitutions on the properties of CH3NH3Sn0.5Pb0.5I3 perovskites. Computational Materials Science, 2020, 177, 109576.	3.0	5
61	A comparative study of some lithium and hydrogen-bonded complexes: <i>Ab initio</i> and QTAIM studies. International Journal of Quantum Chemistry, 2014, 114, 400-408.	2.0	4
62	Enhancing effects of hydrogen/halogen bonds on $\ddot{l}_f$ -hole interactions involving ylide. Journal of Molecular Modeling, 2014, 20, 2282.	1.8	4
63	Chalcogen- and halogen-bonds involving SX2 (X = F, Cl, and Br) with formaldehyde. Journal of Molecular Modeling, 2016, 22, 167.	1.8	4
64	Comparison of the directionality of the halogen, hydrogen, and lithium bonds between HOOOH and XF (XÂ=ÂCl, Br, H, Li). Journal of Molecular Modeling, 2016, 22, 52.	1.8	4
65	Mutual enhancing effects of the Ïf-hole interactions and halogen/hydrogen-bonded interactions in the iodine-ylide containing complexes. Structural Chemistry, 2016, 27, 927-937.	2.0	4
66	The enhancing effects of molecule X (X = PH <sub>2</sub> Cl, SHCl, ClCl) on chalcogen–chalcogen interactions in cyclic trimers Y···A·Y···X (Y = SHCl, SeHCl). International Journal of Quantum Chemis 2017, 117, e25354.	str <b>2</b> y.,0	4
67	Nature of MoH···I bonds in Cp <sub>2</sub> Mo(L)H···I ≡Câ€R Complexes (L=H, CN, PPh <sub>2<td>ub&gt;,) Tj ET 3.5</td><td>Qq1 1 0.784 4</td></sub>	ub>,) Tj ET 3.5	Qq1 1 0.784 4
68	Nb( i PrNPMe 2 ) 3 Fe–PMe 3 : A potential high reactivity heterobimetallic catalyst for acetylene cycloadditions. Applied Organometallic Chemistry, 2020, 34, e5966.	3.5	4
69	Mechanism and regioselectivity of [Cu-Fe] heterobimetallic-catalyzed hydroboration of pyridines: DFT investigation. Molecular Catalysis, 2021, 511, 111722.	2.0	4
70	Halogen Bond Catalysis on Carbonyl–Olefin <scp>Ring losing</scp> Metathesis Reaction: Comparison with Lewis Acid Catalysis. Chinese Journal of Chemistry, 2022, 40, 1275-1284.	4.9	4
71	Charge-regulated regioselective mechanism of bicobalt-catalyzed hydrogermylation of alkynes: DFT investigation. Molecular Catalysis, 2022, 526, 112379.	2.0	4
72	Intriguing E…E' bonding in [Nap(EPh)(E'Ph)] <sup>•+</sup> (E, E'=O, S, Se, Te). International Journal of Quantum Chemistry, 2016, 116, 1090-1096.	2.0	3

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73	The ground and excited-state electronic structures of sandwich compounds Cp2(ME)2 contain an (ME)2 four-membered ring (Cp = C5H5; M = Ni, Pd, Pt; E = O, S, Se, Te). New Journal of Chemistry, 2017, 41, 12028-12034.	2.8	3
74	Stability and donor-acceptor bond in dinuclear organometallics CpM1–M2Cl3 (M1, M2 = B, Al, Ga, In;	Cpậ€‰=	- Î∙) Tj ETÇ
75	Insight into the Effects of Electrostatic Potentials on the Conversion Mechanism of the Hydrogen-Bonded Complexes and Carbon-Bonded Complexes: An Ab Initio and Quantum Theory of "Atoms in Molecules―Investigation. ACS Omega, 2019, 4, 231-241.	3.5	3
76	Electronic and Photovoltaic Properties of Superlattices Constructed by Organic–Inorganic Perovskites: a Theoretical Perspective. ACS Applied Energy Materials, 2022, 5, 2430-2441.	5.1	3
77	Mechanism, Stereoselectivity, and Role of O <sub>2</sub> in Aza-Diels–Alder Reactions Catalyzed by Dinuclear Molybdenum Complexes: A Theoretical Study. Inorganic Chemistry, 2022, 61, 4714-4724.	4.0	3
78	Computational studies of $lf$ -type weak interactions between NCO/NCS radicals and XY(X = H, Cl; Y = F, Cl,) Tj ETO	Qq0,00 r	gBT <sub>2</sub> /Overlock
79	Insight into π-hole interactions containing the inorganic heterocyclic compounds S2N2/SN2P2. Journal of Molecular Modeling, 2017, 23, 233.	1.8	2
80	Inorganic benzenes as the noncovalent interaction donor: a study of the π-hole interactions. Journal of Molecular Modeling, 2017, 23, 335.	1.8	2
81	Nature of non-nuclear (3, â^'3) Ï€-attractor and Ï€-bonding: Theoretical analysis on Ï€-electron density. Chemical Physics Letters, 2018, 691, 347-350.	2.6	2
82	Investigation on the intramolecular proton transfer mechanism of [Cp2MHn](Cp = C5H5, M = Mo, W,	Re;) Tj ET 2.4	Qq <u>Q</u> 0 0 rgBT
83	Stereoselectivity and nonmigratory insertion mechanism of dimethylacetylene dicarboxylate into metalloceneâ€hydride of Cp 2 M(L)H [Cp = η 5   5 H 5; M = Nb, V; L = CO, P (OMe) 3]: A DFT study. Applied Organometallic Chemistry, 2020, 34, e5601.	3.5	2
84	Reaction Mechanism and Kinetics study on Addition of CCl 4 to 1â€hexene Catalyzed by Moâ€Mo Quintuplyâ€bond. Applied Organometallic Chemistry, 2020, 34, e5726.	3.5	2
85	Competition and conversion between pnicogen bonds and hydrogen bonds involving prototype organophosphorus compounds. Physical Chemistry Chemical Physics, 2021, 23, 18794-18805.	2.8	2
86	Theoretical investigation on the nature of substituted benzenear AuX interactions: covalent or noncovalent?. New Journal of Chemistry, 0, , .	2.8	2
87	CIˉ as the halogen bond acceptor: studies on strong halogen bonds. Structural Chemistry, 2018, 29, 503-511.	2.0	1
88	The roles of native defects and transition metal additives in the dehydrogenation mechanism of Mg(AlH4)2. International Journal of Hydrogen Energy, 2020, 45, 17625-17636.	7.1	1
89	<scp> BF <sub>3</sub> â€Catalyzed </scp> Mukaiyama aldol reaction of acetaldehyde with 2â€siloxyâ€1â€propene. Journal of Computational Chemistry, 2022, 43, 402-412.	3.3	1
90	Theoretical Studies on the Isomerizations of CH <sub>3</sub> NO <sub>2</sub> . Chinese Journal of Chemistry, 2009, 27, 1025-1030.	4.9	0

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91	AIM and ELF Analyses of Halogen Bonding in NCS:BrCl Complexes. Chinese Journal of Chemistry, 2011, 29, 2416-2420.	4.9	0
92	Substituent-regulated mechanism on reaction Cp2NbH3 (Cp = η5-C5H5) with RCî€,CR (R = COOMe and Me). Dalton Transactions, 2020, 49, 15376-15384.	3.3	0
93	Mechanistic insights into the reaction Cp2Nb(CO)H (Cp = î·5-C5H5) with acetylenedicarboxylic acid (ADCA): DFT studies. Inorganica Chimica Acta, 2021, 514, 119990.	2.4	0