

原著論文/著書リスト

原著論文/全て査読有 (A200 以下が採録決定済/A206 以下が arXiv 済)

- * A206; " Topological data analysis for revealing structural origin of density anomalies in silica glass",
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A. Tirelli, [K. Nakano](#),
arXiv: 2208.06378 (2022)

- * A205; " High- T_c superconductivity of clathrate Y_3EuH_{24} ",
(クラスレート構造イットリウム・ユーロピウム水素化物の高温超伝導)

[A. Ghaffar](#), [P. Song](#), [K. Nakano](#), [K. Hongo](#), [R. Maezono](#),
arXiv:2205.05906 (2022)

- * A204; " Quantum phase diagram of high-pressure hydrogen",
(第一原理量子モンテカルロ法を利用した高圧水素の量子相図研究)

Lorenzo Monacelli, Michele Casula, [Kosuke Nakano](#), Sandro Sorella, Francesco Mauri,
arXiv:2202.05740 (2022)

- * A203; "High- T_c ternary metal hydrides, YKH_{12} and $LaKH_{12}$, discovered by machine learning",
(機械学習的探索で発見された高温超伝導 3 元水素化物)

[P. Song](#), H. Zhufeng, P. Baptista de Castro, [K. Nakano](#), [K. Hongo](#), Y. Takano, [R. Maezono](#),
arXiv:2103.00193 (2021)

- * A202; "First Principles Calculations of Superconducting Critical Temperature of $ThCr_2Si_2$ -Publisher Structure",
($ThCr_2Si_2$ 型化合物の超伝導転移温度に関するハイスループットスクリーニング)

[G.S. Sinaga](#), [K. Utimula](#), [K. Nakano](#), [K. Hongo](#), [R. Maezono](#),
(in preparation for J. Phys. Chem. C (2022/IF= 3.7))
arxiv.org/abs/1911.10716

*A201; "Importance of vdW and long-range exchange interactions to DFT-predicted docking energies between plumbagin and cyclodextrins"

(シクロデキストリンのホストゲスト結合予見における電子相関)

[T. Ichibha](#), [O. Srihakulung](#), [G. Chao](#), [A. T. Hanindriyo](#), [L. Lawtrakul](#), [K. Hongo](#), [R. Maezono](#),
arXiv:1904.02503(2019)

* A200; "Kinetic effects of ball-milling precursors on the synthesis pathway of KSbF₄",

(酸素空孔を有する Ti ドープ ZnO 単層の NO ガスセンシングに関する第一原理解析)

[R. Hisasue](#), [T. Saquai](#), [T. Ichiba](#), [Y. Fujii](#), [A. Yamashita](#), [K. Hongo](#), [R. Maezono](#), [K. Tadanaga](#), [A. Miura*](#),
Ceram. Int., 51, 33653-33660 (2025) [Q1-journal]
DOI : 10.1016/j.ceramint.2025.05.096

* A199; "Density functional theory insights into NO gas sensing of Ti-doped ZnO monolayer with oxygen vacancy",

(酸素空孔を有する Ti ドープ ZnO 単層の NO ガスセンシングに関する第一原理解析)

[Z. Mahmoudi](#), [T. Ichibha](#), [R. Maezono](#), [M. Abbasnejad*](#),
J. Appl. Phys., 137, 184303 (2025) [Q2-journal]
DOI : 10.1063/5.0256255

* A198; "Hand Milling Induced Phase Transition for Marcasite-type Carbodiimide",

(マルカサイト型カルボジイミドにおける手粉碎誘起の相転移)

[Y. Yamamoto](#), [K. Kume](#), [S. Miyazaki](#), [A. Shinozaki](#), [P. Song](#), [S. S. Hasan](#), [K. Hongo](#), [R. Maezono](#), [H. Ubukata](#), [H. Kageyama*](#), [M. Higuchi](#), [Y. Masubuchi*](#),
J. Am. Chem. Soc., 147, 11390-11398 (2025) [Q1-journal]
DOI : 10.1021/jacs.5c00962

* A197; "Superconductivity in o-MAX phases",

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[M. Keivanloo](#), [M. Sandoghchi](#), [M. R. Mohammadizadeh](#), [M. Kawamura](#), [H. Raebiger](#), [K. Hongo](#), [R. Maezono](#), [M. Khazaei*](#),
Nanoscale, 17, 5341-5349 (2025) [Q1-journal]
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* A196; "Quantum machine learning for ABO₃ perovskite structure prediction",

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[M. Akrom*](#), [S. Rustad*](#), [H. K. Dipojono*](#), [R. Maezono](#), [H. Kasai](#),
Comput. Mater. Sci., 250, 113694 (2025) [Q1-journal]
DOI : 10.1016/j.commatsci.2025.113694

* A195; "Nickel Vanadate Cathode Induced In Situ Phase Transition for Improved Zinc Storage by Low Migration Barrier and Zn²⁺/H⁺

Co-Insertion Mechanism",

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H. Bandi, A. K. Kakarla, [R. Dahule](#), [R. Maezono](#), D. Narsimulu, R. Shanthappa, J. S. Yu*,
Small, 21, (2025) [Q1-journal]
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* A194; "W/Mo/Cr Doping Modulates the Negative–Positive Inversion Gas Sensing Behavior of VO₂(M1)",

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L. Miao, Y. Xue, [P. Song](#), T. Hasegawa, A. Okawa, [R. Maezono](#), T. Sekino, S. Yin*,
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DOI : 10.1021/acssensors.4c03006

* A193; "Physics-informed Data-driven Discovery of Polymer Crystals with High Thermal Conductivity",

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[R. Dahule*](#), [K. Oqmhula](#), [R. Maezono](#), [K. Hongo*](#),
ACS Appl. Polym. Mater., 7, 1431-1439, (2025) [Q1-journal]
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* A192; "Stability and Electronic Properties of SnS/ZnS Interfaces: A First-Principles Investigation",

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[R. Dahule*](#), B. Gao, [K. Hongo](#), Emila Panda, Yanming Ma, [R. Maezono*](#),
J. Phys. Chem. C., 129, 3158-3167, (2025) [Q1-journal]
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* A191; "Novel Selectivity: Target of Gas Sensing Defined by Behavior",

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L. Miao, [P. Song](#), Y. Xue, Z. Hou, T. Hasegawa, A. Okawa, T. Goto, Y. Seo, [R. Maezono](#), T. Sekino, S. Yin*,
Adv. Mater., 37, 2413023, (2024) [Q1-journal]
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* A190; "A machine learning approach for forecasting the efficacy of pyridazine corrosion inhibitors",

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G. A. Trisnapradika, M. Akrom, S. Rustad, H. K. Dipojono, [R. Maezono](#), H. Kasai*,
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* A189; "Theoretical Insights into High-*T_c* Superconductivity of Structurally Ordered YThH₁₈: A First-Principles Study",

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A. Ghaffar*, P. Song, R. Maezono, K. Hongo*,
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- * A188; "Computational Study of the Structural, Mechanical, Electronic, Optical and Thermal properties of BaLiX (X =P, As, Sb) perovskites",
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M. Z. Rahman, [S. S. Hasan](#), M. S. Akter, N. M. Mukhtar, N. Absar, M. A. Hasan, [T. Ichibha](#), [R. Maezono](#), [K. Hongo](#), M. A. Islam*,
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- * A187; "Using reinforcement learning to autonomously identify the source of errors for agents in group missions",
(原因系を効率よく切り分ける検証行動計画を自動的に策定する強化学習)

[K. Utimula](#)*, K-T. Hayaschi, T. J. Bihl, [K. Nakano](#), [K. Hongo](#), [R. Maezono](#),
Front. Control Eng., 5, (2024) [Q2-journal]
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- * A186; "Solid-liquid phase boundary of oxide solid solutions using neural network potentials",
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K. Hyodo, [K. Hongo](#), [T. Ichibha](#), [R. Maezono](#)*,
J. Alloys Compd., 1006, 176227, (2024) [Q1-journal]
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- * A185; "Unveiling the NIR modulation performance enhancement of VO₂ endowed by oxygen vacancy elimination",
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Y. Xue, L. Miao, [P. Song](#), T. Hasegawa, A. Okawa, [R. Maezono](#), T. Sekino, S. Yin*,
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- * A184; "Structural predictions and phonon-mediated superconductivity in platinum hydride under low pressure: Insight from first-principles calculations",
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P. Tsuppayakorn-aek, [P. Song](#), W. Sukmas, [R. Maezono](#), T. Bovornratanaraks,
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- * A183; "Ca substitution effects on structure transformation and physical properties of (Eu,Ca)FeAs₂ co-doped with La and Co",

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S. P. K. Naik*, S. Alberto, K. Kataoka, Y. Gotoh, [T. Ichibha](#), [K. Hongo](#), [R. Maezono](#), T. Nishio, H. Ogino*,
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* A182; "Enhancing MnBi₂Te₄ Stability by Doping",

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K. Saritas*, [A. Ghaffar](#), J. T. Krogel, [T. Ichibha](#), [K. Hongo](#), [R. Maezono](#), F. A. Reboredo,
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DOI : 10.1021/acs.jpcc.4c00848

* A181; "Thermal Decomposition of Oxygen-containing Ta₃N₅",

(タンタル窒化物への酸素ドーピングに関する状態図解析)

N. Moharana, C. Ghosh, A. Dasgupta, [R. Maezono](#), S. Sarma, Ravi Kumar, K.C.Hari Kumar,
J. Am. Ceram. Soc., 107, 6342-6352(2024) [Q1-journal]
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* A180; "Substitutional Doping Strategies for Fermi Level Depinning and Enhanced Interface Quality in WS₂-Metal Contacts",

(ドーピングした遷移金属ダイカルゴゲナイトの界面に関する第一原理解析)

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* A179; "Key Role of Metal-to-Metal Charge Transfer Transition between Mo⁶⁺ and Bi³⁺ for Enhancement in NIR Luminescence of Gd₂MoO₆:Bi,Yb Nanophosphor",

(ナノ蛍光体の発光を増強させる金属イオン間の電荷移動)

T. Hangai, *T. Hasegawa, J. Xu, T. Nakanishi, T. Takeda, [K. Nakano](#), [K. Hongo](#), [R. Maezono](#), T. Goto, Y. Sato, A. Okawa, S. Yin,
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* A178; "Single crystal growth and physical properties of La, Co doped (Eu,Ca)FeAs₂",

(新規鉄系化合物単結晶の基礎物性解析)

*S. P. K. Naik, S. Alberto, K. Kataoka, Y. Gotoh, [T. Ichibha](#), [K. Hongo](#), [R. Maezono](#), T. Nishio, *H. Ogino,
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* A177; "Multi-emission of Ce³⁺ from Single Crystallographic Site Induced by Disorder of Ions",

(秩序の乱れに起因した Ce^{3+} からの発光バリエーション)

T. Yasunaga, *M. Kobayashi, [K. Oqmhula](#), [H. Qi](#), [T. Ichibha](#), [K. Hongo](#), S. Yamamoto, [R. Maezono](#), M. Mitsuishi, M. Osada, *H. Kato, M. Kakihana,
Inorg. Chem. 63, 1288 – 1295 (2024) [Q1-journal]
DOI : 10.1021/acs.inorgchem.3c03789

- * A176; "Locality Error Free Effective Core Potentials for 3d Transition Metal Elements Developed for the Diffusion Monte Carlo Method",
(量子拡散モンテカルロ法用途の有効各電荷の開発--3d 遷移金属における局所近似誤差問題の解決)

*[T. Ichibha](#), [Y. Nikaido](#), C. M Bennett, J. T Krogel, [K. Hongo](#), [R. Maezono](#), *F. A Reboredo,
J. Chem. Phys. 159 164114 (2023) [Q1-journal]
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- * A175; "(La,Th)H₁₀: Potential high- T_c (242 K) superconductors Stabilized Thermodynamically below 200 GPa",
(高圧下におけるランタン・トリウム水素化物の高温超伝導)

*[P. Song](#), A. P. Durajski, Z. Hou, [A. Ghaffar](#), [R. Dahule](#), R. Szcześniak, [K. Hongo](#), [R. Maezono](#),
J. Phys. Chem. C . 128 2656–2665 (2024) [Q1-journal]
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- * A174; "First-Principles Investigation of Stability and Superconductivity in Ternary Yttrium – Praseodymium Hydrides under High Pressure",
(遺伝的アルゴリズムを用いた高圧下の Y-Pr 水素化物における超伝導と合成安定性の予見)

*[K. S. Qin](#), *[P. Song](#), *[K. Hongo](#), *[R. Maezono](#),
J. Phys. Chem. C 127 21242-21249 (2023) [Q1-journal]
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- * A173; "Structure, optical, and electrical properties of layered oxychalcogenide $\text{Sr}_2\text{ZnCu}_2(\text{S}_{1-x}\text{Se}_x)_2\text{O}_2$ ($0 \leq x \leq 1$) compounds",
(層状カルコゲナイト酸化物の構造と光学物性、電子物性)

T. Kato, Y. Iwasa, S. Pavan K. Naik, S. Ishida, Y. Higashi, I. Hase, T. Nishio, [K. Hongo](#), [R. Maezono](#), *H. Ogino,
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- * A172; "Recognition of spatial finiteness in meniscus splitting based on evaporative interface fluctuations",
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L. Wu, I. Saitoh, [K. Hongo](#), *K. Okeyoshi,
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- * A171; "Stiffer Bonding of Armchair Edge in Single-Layer Molybdenum Disulfide Nanoribbons",

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C. Liu, [K. Hongo](#), [R. Maezono](#), *J. Zhang, *Y. Oshima,
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* A170; "Biophysical properties of fibril structure of the toxic conformer of amyloid- β 42: characterization by atomic force microscopy and molecular docking",

(毒性型アミロイドベータのフィブリル構造に関するドッキングシミュレーションと原子力顕微鏡観測)

R. Biyani, K. Hirata, [K. Ogumhula](#), A. Yurtsever, [K. Hongo](#), [R. Maezono](#), M. Takagi, *T. Fukuma, *M. Biyani,
ACS Appl. Mater. Interfaces, 15, 27789–27800 (2023) [Q1-journal]
DOI : 10.1021/acsami.3c06460

* A169; "Existence of La-site antisite defects in LaMO_3 (M = Mn, Fe, and Co) predicted with many-body diffusion quantum Monte Carlo",

(LaMO_3 型ペロブスカイト中アンチサイト欠陥 -- 量子拡散モンテカルロ法による存在予見)

[T. Ichibha](#), S. Yoon, J. M. Ok, M. Yoon, H. N. Lee, F. A. Reboredo,
Sci. Rep., 13 6703 (2023) [Q1-journal]
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* A168; "High-pressure BaCN_2 phases explored by genetic algorithm",

(遺伝アルゴリズムを用いた金属カルボジイミド結晶の構造探索)

[P. Song](#), [M. Khawaguch](#), Y. Masubuchi, [K. Ogumhula](#), [K. Nakano](#), [R. Maezono](#), [K. Hongo](#),
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* A167; "Evolutionary Algorithm Directed Synthesis of Mixed Anion Compounds LaF_2X (X = Br, I) and LaFI_2 .",

(遺伝的アルゴリズムを援用したフッ化物複合アニオン物質の合成)

D. Kato, [P. Song](#), H. Taguro, C. Tassel, H. Ubukata, K. Miyazaki, T. Abe, K. Nakano, [T. Ichibha](#), [K. Hongo](#), *[R. Maezono](#), *H. Kageyama,,
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DOI : 10.1002/anie.202301416

* A166; "Towards chemical accuracy using the Jastrow correlated antisymmetrized geminal power ansatz",

(ジェミナル型多体波動関数で化学的精度を実現する電子状態計算)

[A. Raghav](#), [R. Maezono](#), [K. Hongo](#), S. Sorella, [K. Nakano](#),
J. Chem. Theory Comput. 19, 2222-2229 (2023) [Q1-journal]
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* A165; "Thermodynamic Understanding of Impurity Phase Segregation in a $\text{PdCrO}_2/\text{CuCrO}_2$ Heterostructure",

(LaMO₃ 型ペロブスカイト中アンチサイト欠陥 -- 量子拡散モンテカルロ法による存在予見)

[T. Ichibha](#), [S. Yoon](#), [J. M. Ok](#), [M. Yoon](#), [H. N. Lee](#), [F. A. Reboredo](#),
Advanced Physics Research (2023) ,
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- * A164; "First-Principles-Based Insight into Electrochemical Reactivity in A Cobalt-Carbonate-Hydrate Pseudocapacitor",
(電気化学キャパシタ材料の蓄電特性をスーパーコンピュータを用いたシミュレーションで解明)

[K. Ogmhula](#), [T. Toma](#), [R. Maezono](#), [K. Hongo](#),
ACS Omega 8, 6743-6752 (2023) [Q1-journal]
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- * A163; "Mechanistic insights and importance of hydrophobicity in cationic polymers for cancer therapy",
(抗ガン高分子の分子設計指針に新たな光～カチオン性と疎水性の相乗効果で高い細胞障害性が発現～)

[N. Kumar](#), [K. Ogmhula](#), [K. Hongo](#), [K. Takagi](#), [S. Yusa](#), [R. Rajan](#), [K. Matsumura](#),
J. Mater. Chem. B. 11, 1456-1468 (2023) [Q1-journal]
DOI : 10.1039/D2TB02059A

- * A162; "Order–disorder competition in equiatomic 3d–transition–metal quaternary alloys: Phase stability and electronic structure",
(3d 遷移金属等組成 4 元系合金における秩序-無秩序競合：相安定性と電子状態)

[H. Mizuseki](#), [R. Sahara](#), [K. Hongo](#),
Sci. Technol. Adv. Mater. 3, 2153632 (2023) [Q1-journal]
DOI : 10.1080/27660400.2022.2153632

- * A161; "Electronic and magnetic properties of pure and Cu doped non-polar ZnO(10 $\bar{1}$ 0) surfaces",
(Cu をドーピングした ZnO 表面の電氣的磁氣的性質に関する第一原理解析)

[E. Irandegani](#), [R. Maezono](#), [M. Abbasnejad](#),
J. Appl. Phys. 132 173903 (2022) [Q2-journal]
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- * A160; "Feature space of XRD patterns constructed by auto-encorder",
(オートエンコーダによって構成された XRD パターンの特徴量空間)

[K. Utimula](#), [M. Yano](#), [H. Kimoto](#), [K. Hongo](#), [K. Nakano](#), [R. Maezono](#),
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- * A159; " Potential high- T_c superconductivity in YCeH₂₀ and LaCeH₂₀ under pressure",

(Ce 系水素化物の高圧化超伝導)

P. Song, Z. Hou, K. Nakano, K. Hongo, R. Maezono,
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- * A158; "Anionic ordering in $\text{Pb}_2\text{Ti}_4\text{O}_9\text{F}_2$ revisited by nuclear magnetic resonance and density functional theory",
(鉛酸フッ化物のローンペアで安定化されるアニオン秩序配列)

K. Oka, T. Ichibha, D. Kato, M. Iwasaki, N. Noma, K. Hongo, R. Maezono, F. A. Reboredo,
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- * A157; "Ab-initio-based Interface Modeling and Statistical Analysis for Estimate of the Water Contact Angle on a Metallic Cu(111) Surface",
(第一原理計算による界面モデリング:金属表面に対する接触角評価)

T. Murono, K. Hongo, K. Nakano, R. Maezono,
Surf. Interfaces 34, 102342 (2022) [Q1-journal]
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- * A156; " Electronic structure and effective mass analysis of doped TiO_2 (anatase) systems using DFT+ U ",
(ドーピングしたチタン酸化物の DFT+U 電子状態計算と有効質量解析)

A. Raghav, K. Hongo, R. Maezono, E. Panda,
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- * A155; "Ab initio molecular dynamics simulation of structural and elastic properties of SiO_2 - P_2O_5 - Al_2O_3 glass",
(ガラス材料の力学特性に関する第一原理分子動力学解析)

Y. Qian, B. Song, J. Jin, G. I. Prayogo, K. Utimula, K. Nakano, R. Maezono, K. Hongo, G. Zhao,
J. Am. Ceram. Soc. 105, 6604-6615(2022) [Q1-journal]
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- * A154; "High pressure hydrogen by machine learning and quantum Monte Carlo",
(厳密な電子状態計算 × 機械学習ポテンシャル:高圧水素における液体-液体相転移の研究)

A. Tirelli, G. Tenti, K. Nakano, S. Sorella,
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DOI : 10.1103/PhysRevB.106.L041105

- * A153; " SHRY: Application of canonical augmentation to the atomic substitution problem",

(SHRY:正準強化法の原子置換問題への応用)

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- * A152; " High pressure behavior of tetragonal barium carbodiimide, BaNCN",
(カルボジイミド無機化合物の結晶構造の X 線回折および密度汎関数法による解析)

Y. Masubuchi, S. Miyazaki, P. Song, T. Yamamoto, K. Nakano, K. Hongo, R. Maezono,
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- * A66; "Selectivity of CO and NO adsorption on ZnO (0002) surfaces: A DFT investigation"

(ZnO 表面における CO 分子/NO 分子の選択的吸収)

Nugraha, A.G. Saputro, M.K. Agusta, B. Yulianto, H.K. Dipojono, F. Rusydi, and R. Maezono,
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K. Nakano, K. Hongo, and R. Maezono,
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(エピタキシャル成長させた酸水素化物薄膜の電氣的性質)

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(CaBe2Ge2 型構造を持つ超伝導体:LaPd2Sb2)

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Y. Nozaki, K. Nakano, T. Yajima, H. Kageyama, B. Frandsen, L. Liu, S. Cheung, T. Goko, Y. J. Uemura, T. S. J. Munsie, T. Medina, G. M. Luke, J. Munevar, D. Nishio-Hamane, and C. M. Brown,
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($d1$ 正方格子を持つ新規ビスマス超伝導体: $\text{BaTi}_2\text{Bi}_2\text{O}$, $(\text{SrF})_2\text{Ti}_2\text{Bi}_2\text{O}$ の合成と物性)

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