Guy Tréglia

List of Publications by Year in descending order

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| 162 | 5,038 | 38 | 64 |
|----------|----------------|--------------|---------------------|
| papers | citations | h-index | g-index |
| 162 | 162 | 162 | 2087 citing authors |
| all docs | docs citations | times ranked | |

| # | Article | IF | CITATIONS |
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| 1 | Surface core level spectroscopy of transition metals: A new tool for the determination of their surface structure. Surface Science Reports, 1985, 5, 1-85. | 3.8 | 260 |
| 2 | Perturbation treatment of correlations in transition metals. Journal De Physique, 1980, 41, 281-289. | 1.8 | 200 |
| 3 | Segregation and Ordering at Surfaces of Transition Metal Alloys: The Tight-Binding Ising Model. Europhysics Letters, 1988, 7, 575-580. | 0.7 | 165 |
| 4 | Effect of coulomb correlations on energy bands in ferromagnetic transition metals: Ni, Co and Fe. Journal De Physique, 1982, 43, 341-353. | 1.8 | 152 |
| 5 | Band gaps and asymptotic behaviour of continued fraction coefficients. Journal of Physics C: Solid State Physics, 1982, 15, 2891-2924. | 1.5 | 144 |
| 6 | Surface-sandwich segregation in Pt-Ni and Ag-Ni alloys: Two different physical origins for the same phenomenon. Physical Review B, 1987, 35, 4338-4344. | 1.1 | 141 |
| 7 | Comments on the electronic structure of nickel. Physical Review B, 1980, 21, 3729-3733. | 1.1 | 135 |
| 8 | Structures of a Ag monolayer deposited on Cu(111), Cu(100), and Cu(110) substrates: An extended tight-binding quenched-molecular-dynamics study. Physical Review B, 1992, 46, 16018-16030. | 1.1 | 129 |
| 9 | Electronic structure, pairwise interactions and ordering energies in binary f.c.c. transition metal alloys. Solid State Communications, 1981, 39, 149-153. | 0.9 | 126 |
| 10 | Alloy surfaces: segregation, reconstruction and phase transitions. Computational Materials Science, 1999, 15, 196-235. | 1.4 | 118 |
| 11 | Phase transitions in surface segregation of PtcNi1â^calloys from tight-binding Ising-model calculations. Physical Review B, 1990, 41, 4422-4434. | 1.1 | 104 |
| 12 | New magic numbers in metallic clusters: an unexpected metal dependence. Surface Science, 1997, 383, L719-L727. | 0.8 | 102 |
| 13 | Ag/Cu(111) structure revisited through an extended mechanism for stress relaxation. Physical Review B, 1999, 59, 10910-10917. | 1.1 | 99 |
| 14 | Correlation effects on Auger spectra in unfilled d band metals. Journal of Physics C: Solid State Physics, 1981, 14, 4347-4355. | 1.5 | 92 |
| 15 | Ge/Ag(111) semiconductor-on-metal growth: Formation of anAg2Gesurface alloy. Physical Review B, 2000, 62, 16653-16656. | 1.1 | 86 |
| 16 | Modeling free and supported metallic nanoclusters: structure and dynamics. Phase Transitions, 2004, 77, 101-113. | 0.6 | 83 |
| 17 | Surface segregation near the temperature of bulk phase separation: Incomplete wetting in Cu(Ag) alloys. Physical Review B, 1991, 44, 5842-5854. | 1.1 | 77 |
| 18 | Experimental and Theoretical Evidence for a Strong Anisotropy of the Surface Debye-Waller Factor as Determined for a Monolayer of Cobalt on Copper (111) by Surface Extended X-Ray-Absorption Fine Structure. Physical Review Letters, 1986, 56, 1272-1275. | 2.9 | 73 |

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| 19 | Is the segregation-dissolution kinetics driven by a surface local equilibrium? An answer via the kinetic tight-binding Ising model. Surface Science, 1992, 274, 297-305. | 0.8 | 69 |
| 20 | Anisotropy of diffusion along steps on the (111) faces of gold and silver. Physical Review B, 1994, 50, 12104-12117. | 1.1 | 68 |
| 21 | How to compare superficial and intergranular segregation? A new analysis within the mixed SMA–TBIM approach. Acta Materialia, 1999, 47, 2705-2715. | 3.8 | 63 |
| 22 | Study of the W (Ta) core level shifts induced by the adsorption of oxygen on tungsten (tantalum) (110). Journal of Physics C: Solid State Physics, 1981, 14, 3463-3473. | 1.5 | 61 |
| 23 | Electronic structure and relative stabilities of L12 and D022 ordered structures occuring in transition metal alloys. Solid State Communications, 1983, 45, 585-590. | 0.9 | 60 |
| 24 | Electronic structure and phase stability of A15 transition metals and alloys. Journal of Physics F: Metal Physics, 1983, 13, 2543-2567. | 1.6 | 60 |
| 25 | Bulk and surface vibrational and thermodynamical properties of fcc transition and noble metals: a systematic study by the continued fraction technique. Journal De Physique, 1985, 46, 987-1000. | 1.8 | 60 |
| 26 | Generalised perturbation theory in disordered transition metal alloys: application to the self-consistent calculation of ordering energies. Journal of Physics F: Metal Physics, 1978, 8, 1437-1456. | 1.6 | 58 |
| 27 | Pt/Co(0001) superstructures in the submonolayer range: A tight-binding quenched-molecular-dynamics study. Physical Review B, 1999, 60, 2781-2788. | 1.1 | 58 |
| 28 | Surface Segregation in Transition Metal Alloys. Progress of Theoretical Physics Supplement, 1990, 101, 159-180. | 0.2 | 58 |
| 29 | New opportunities to understand heterogeneous catalysis processes on nanoscale bimetallic particles through synchrotron radiation and theoretical studies. Applied Catalysis A: General, 2000, 200, 47-54. | 2.2 | 52 |
| 30 | Surface segregation in PtRh alloys revisited in the framework of the tight-binding Ising model. Surface Science, 1990, 236, 398-408. | 0.8 | 50 |
| 31 | A 'quenched molecular dynamics' approach to the atomic stability of the (100) face of BCC transition metals. Journal of Physics C: Solid State Physics, 1986, 19, 4463-4472. | 1.5 | 49 |
| 32 | Equilibrium and kinetics in the (111) surface of Cu-Ag alloys: Comparison between mean-field and Monte Carlo calculations. Physical Review B, 1994, 50, 1912-1921. | 1.1 | 48 |
| 33 | Surface segregation in CuNi and AgNi alloys formulated as an area-preserving map. Surface Science, 1990, 225, 319-330. | 0.8 | 44 |
| 34 | W4f core level shift study on unreconstructed and hydrogen reconstructed W(100) faces. Journal of Physics C: Solid State Physics, 1982, 15, 4023-4032. | 1.5 | 43 |
| 35 | A theoretical inquiry into the question of W and Ta (100) atomic structures. Journal of Physics C: Solid State Physics, 1983, 16, 2407-2419. | 1.5 | 43 |
| 36 | W(100) reconstruction studied by core level spectroscopy. Solid State Communications, 1984, 50, 393-396. | 0.9 | 43 |

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| 37 | Electronic structure and pairwise interactions in substoichiometric transition metal carbides and nitrides. Journal De Physique, 1985, 46, 1001-1015. | 1.8 | 43 |
| 38 | Tight-binding molecular dynamics study of diffusion on Au and Ag(111). Surface Science, 1995, 331-333, 920-924. | 0.8 | 42 |
| 39 | Core-level spectroscopy of clean and adsorbate-covered Ta(100). Physical Review B, 1984, 30, 5487-5493. | 1.1 | 40 |
| 40 | Microstructure of the surfactantlike effect in Ni/Ag(100) and (111). Physical Review B, 1997, 55, $10931-10937$. | 1.1 | 38 |
| 41 | Site segregation in size-mismatched nanoalloys: Application to Cu–Ag. Surface Science, 2006, 600, 5011-5020. | 0.8 | 38 |
| 42 | Theoretical investigation of chemical and morphological ordering inPdcCu1â^'cclusters. Physical Review B, 2002, 66, . | 1.1 | 37 |
| 43 | High-Temperature Study of the Schwoebel Effect in Au(111). Physical Review Letters, 1996, 76, 2109-2112. | 2.9 | 36 |
| 44 | Thermodynamic derivation of the coherent potential approximation and ordering processes in transition alloys. Journal of Physics F: Metal Physics, 1980, 10, 2137-2146. | 1.6 | 34 |
| 45 | Influence of surface stress in the missing row reconstruction of fcc transition metals. Surface Science, 2006, 600, 5131-5135. | 0.8 | 34 |
| 46 | Electronic structure of Pd clusters in the tight-binding approximation: influence of spd-hybridization. Surface Science, 1996, 352-354, 675-679. | 0.8 | 32 |
| 47 | Charge redistribution at Pd surfaces:Ab initiogrounds for tight-binding interatomic potentials. Physical Review B, 1997, 56, 12161-12166. | 1.1 | 32 |
| 48 | Ordered surface alloy formation of immiscible metals: The case of Pb deposited on Ag(111). Physical Review B, 2005, 72, . | 1.1 | 32 |
| 49 | Ordering and surface segregation in <mml:math xmlns:mml="http://www.w3.org/1998/Math/MathML"><mml:mrow><mml:msub><mml:mi>Co</mml:mi><mml:r A theoretical study from surface alloys to nanoalloys. Physical Review B, 2015, 91, .</mml:r </mml:msub></mml:mrow></mml:math> | nr dwl> <mr< td=""><td>ml:31n>1</td></mr<> | ml: 31 n>1 |
| 50 | Alloy surfaces and surface alloys: from equilibrium to kinetics. Surface Science, 1993, 287-288, 371-376. | 0.8 | 30 |
| 51 | Is ordering in PtNi alloys induced by spin-orbit interactions?. Journal of Physics F: Metal Physics, 1987, 17, 1935-1944. | 1.6 | 29 |
| 52 | Multilayer relaxation and reconstruction in bcc and fcc transition and noble metals. Vacuum, 1990, 41, 311-314. | 1.6 | 29 |
| 53 | Model of surface segregation driving forces and their coupling. Physical Review B, 2008, 78, . | 1.1 | 29 |
| 54 | On a "surfactant-like―behaviour of deposit. Surface Science, 1994, 307-309, 531-537. | 0.8 | 28 |

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| 55 | Core level spectroscopy of the low index faces of Tantalum. Surface Science, 1985, 162, 46-50. | 0.8 | 27 |
| 56 | Theoretical study of the Debye-Waller factor in surface extended x-ray-absorption fine structure: Influence of an adsorbed monolayer. Physical Review B, 1986, 34, 6662-6668. | 1.1 | 26 |
| 57 | Molecular dynamics simulations for the Ag/Cu (111) system: from segregated to constitutive interfacial vacancies. Applied Surface Science, 2000, $162-163$, $219-226$. | 3.1 | 26 |
| 58 | Relation between surface stress and $(1\tilde{A}-2)$ reconstruction for $(1\ 1\ 0)$ fcc transition metal surfaces. Applied Surface Science, 2003, 212-213, 866-871. | 3.1 | 26 |
| 59 | Link between the surface wetting in Cu(Ag) and the layer-by-layer dissolution mode of a thick Ag deposit on a Cu substrate. Surface Science, 1995, 331-333, 805-810. | 0.8 | 25 |
| 60 | Vacancy generation at steps and the kinetics of surface alloy formation. Surface Science, 1996, 364, 453-466. | 0.8 | 25 |
| 61 | Clean surface studies by photoelectron diffraction analysed within a single scattering theory. Journal De Physique, 1988, 49, 227-236. | 1.8 | 25 |
| 62 | Further comments on the electronic structure of nickel. Physical Review B, 1980, 22, 6472-6473. | 1.1 | 23 |
| 63 | An atomistic modelling of the porosity impact on UO2 matrix macroscopic properties. Journal of Nuclear Materials, 2011, 415, 210-216. | 1.3 | 23 |
| 64 | Layer-by-layer versus surfactant dissolution modes in heteroepitaxy. Physical Review B, 1999, 60, 13890-13901. | 1.1 | 22 |
| 65 | Intergranular segregation and vibrational effects: A local analysis. Physical Review B, 2000, 61, 14470-14480. | 1.1 | 22 |
| 66 | Theoretical determination of two critical sizes for strain relaxation during Co/Pt(111) heteroepitaxy. Surface Science, 2000, 446, 272-282. | 0.8 | 22 |
| 67 | Cu-Ag (111) Polymorphism Induced by Segregation and Advacancies. Physical Review Letters, 2003, 91, 176103. | 2.9 | 22 |
| 68 | Surface core level spectroscopy of the stepped surface. Solid State Communications, 1984, 52, 635-639. | 0.9 | 21 |
| 69 | Physical origin of thickness-controlled sequential phase formation during reactive diffusion: Atomistic modeling. Physical Review B, 2010, 82, . | 1.1 | 21 |
| 70 | Photoelectron spectroscopy study of $Pb/Ag(111)$ in the submonolayer range. Surface Science, 2006, 600, 1227-1230. | 0.8 | 20 |
| 71 | Atomistic modeling of strain and diffusion at the <mml:math display="inline" xmlns:mml="http://www.w3.org/1998/Math/MathML"><mml:mrow><mml:mtext>Si</mml:mtext><mml:mo>/</mml:mo><mml:msub><mml:mrow><mpl:mrow><mpl:mrow><mpl:mtext>Si</mpl:mtext></mpl:mrow></mpl:mrow></mml:mrow></mml:msub></mml:mrow></mml:math> | ıml <mark>:m</mark> text: | ∙Sið ^Q /mml:m |
| 72 | Step-driven molecular adsorption of Sb on Si(111). Surface Science, 1998, 395, 317-325. | 0.8 | 18 |

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| 73 | New trends in heterogeneous catalysis processes on metallic clusters from synchrotron radiation and theoretical studies. Applied Surface Science, 2000, 164, 140-146. | 3.1 | 18 |
| 74 | Electronic structure of NiZr2C16 compound. Journal of Physics F: Metal Physics, 1982, 12, 441-447. | 1.6 | 17 |
| 75 | One- and two-hole excitation spectra of the Hubbard model in the non-half-filled case. Physical Review B, 1986, 34, 5101-5110. | 1.1 | 17 |
| 76 | On the influence of topology on the energy profile in metallic Pd clusters. Surface Science, 1994, 307-309, 735-740. | 0.8 | 17 |
| 77 | Kinetics of segregation and dissolution in Cu1â^'cAgc and surface phase transition: comparison between mean field and Monte Carlo calculations. Surface Science, 1994, 307-309, 804-809. | 0.8 | 17 |
| 78 | Surface-induced ordering in phase separation systems: influence of concentration and orientation. Surface Science, 1999, 441, 225-239. | 0.8 | 17 |
| 79 | New Structures and Atomistic Analysis of the Polymorphism for the $\hat{a}^* = 5$ (210) [001] Tilt Boundary. Journal of Materials Science, 2000, 8, 55-69. | 1.2 | 17 |
| 80 | Incomplete wetting of very dilute Cu(Ag) alloys by surface segregation. Surface Science, 1991, 251-252, 664-669. | 0.8 | 16 |
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| 82 | Topology of a Ag monolayer on a $Cu(111)$ substrate: a tight-binding quenched molecular dynamics study. Surface Science, 1993, 287-288, 476-481. | 0.8 | 16 |
| 83 | On the various terminations occurring in CuPt-ordered alloys: the TBIM approach. Surface Science, 1994, 307-309, 440-444. | 0.8 | 16 |
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| 85 | One-hole excitation spectra of the one-dimensional Hubbard model. Physical Review B, 1985, 32, 2167-2177. | 1.1 | 15 |
| 86 | Experimental study of the growth and dissolution of Re/W(110) by core level photoemission spectroscopy: formation of a surface alloy. Surface Science, 1993, 286, 150-167. | 0.8 | 15 |
| 87 | On a surprising anisotropy of surface segregation in CuPt alloys. Surface Science, 1993, 287-288, 851-856. | 0.8 | 15 |
| 88 | An unusual composition profile: a LEED–TBIM study of Pt25Cu75(). Surface Science, 2003, 527, 71-79. | 0.8 | 15 |
| 89 | Unified picture of </a href=" mailto:="" mml:math="">-band"><b .<="" 2011,="" 83,="" alloys.="" b,="" in="" metal="" physical="" review="" shifts="" td="" transition=""><td>1.1</td><td>15</td> | 1.1 | 15 |
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| 91 | Influence of Coulomb correlations on the electronic excitations of transition metal surfaces: application to Mo(100) and Ni(100), (110) and (111). Journal of Physics C: Solid State Physics, 1980, 13, 6063-6072. | 1.5 | 14 |
| 92 | Interpretation of Cr (001) photoemission spectra: influence of correlations. Journal of Physics F: Metal Physics, 1984, 14, 1317-1324. | 1.6 | 14 |
| 93 | Is the (110) face of tungsten reconstructed?. Solid State Communications, 1983, 47, 279-282. | 0.9 | 13 |
| 94 | Competition or synergy between surface segregation and bulk ordering: the Cuî—,Pd system. Surface Science, 1996, 352-354, 588-591. | 0.8 | 12 |
| 95 | On the "exotic―behaviour of the PtSn system. Surface Science, 1997, 377-379, 1033-1037. | 0.8 | 12 |
| 96 | Misfit dislocation loops or incommensurate structure at an interface: Vibrational and anharmonic effects. Physical Review B, 2002, 66, . | 1.1 | 12 |
| 97 | Surface and bulk core-level lineshapes in tungsten and tantalum. I. Theory. Journal of Physics C: Solid State Physics, 1987, 20, 2647-2657. | 1.5 | 11 |
| 98 | Photoelectron diffraction study of the unreconstructed and the hydrogen reconstructed W(110) surfaces. Solid State Communications, 1989, 69, 1015-1018. | 0.9 | 11 |
| 99 | Local-density approximation study of semiconductor/metal adsorption characteristics: Ge/Ag(100). Physical Review B, 1999, 59, 15337-15345. | 1.1 | 11 |
| 100 | Compressive strain versus tensile strain. Applied Surface Science, 2001, 177, 238-242. | 3.1 | 11 |
| 101 | Linear time dependence of the surfactant effect: A local equilibrium under flux. Physical Review B, 2004, 69, . | 1.1 | 11 |
| 102 | Self-organization of Ge tetramers on Ag(001) surface: A 2D realization of unusual substrate mediated interactions. Surface Science, 2008, 602, 506-510. | 0.8 | 11 |
| 103 | Strain effect on self-diffusion in silicon: Numerical study. Physical Review B, 2009, 79, . | 1.1 | 11 |
| 104 | Theoretical investigation of the influence of reaction and diffusion kinetics upon thin-film reactive diffusion. Physical Review B, 2012, 85, . | 1.1 | 11 |
| 105 | Theoretical investigation of Cottrell atmosphere in silicon. Acta Materialia, 2014, 65, 1-9. | 3.8 | 11 |
| 106 | Surface and bulk core-level lineshapes in tantalum. II. Experiments. Journal of Physics C: Solid State Physics, 1988, 21, 287-295. | 1.5 | 10 |
| 107 | Ge deposition on Ag surfaces: Dependence of the adsorption characteristics on the surface orientation. Physical Review B, 2000, 61, 8469-8474. | 1.1 | 10 |
| 108 | Comment on some â€~â€~new'' results on the multilayer reconstruction of W(100). Physical Review B, 1940, 6440-6441. | 989 1.1 | 9 |

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| 109 | Flux dependence of the surfactant effect in : a theoretical study. Surface Science, 1996, 352-354, 562-566. | 0.8 | 9 |
| 110 | Step-descent mechanisms on Ag and Au(111). Surface Science, 1997, 377-379, 843-846. | 0.8 | 9 |
| 111 | Rules for tight-binding calculations in bi-metallic compounds based on density functional theory: the case of Co–Au. Journal of Physics Condensed Matter, 2010, 22, 505503. | 0.7 | 9 |
| 112 | Coulomb correlations in nickel within the alloy analogy of the Hubbard model. Solid State Communications, 1981, 39, 1113-1116. | 0.9 | 8 |
| 113 | Hydrogen chemisorption on tungsten (110) studied by core level spectroscopy. Journal of Physics C: Solid State Physics, 1983, 16, 1555-1566. | 1.5 | 8 |
| 114 | Variation at the surface of mean correlated displacements in fcc transition and noble metals studied by a continued fraction technique. Surface Science, 1985, 162, 126-131. | 0.8 | 8 |
| 115 | The face dependence of the effective electron mean free path derived from spherical-wave corrections in photoelectron diffraction of $W(110)$ and $W(100)$ surfaces. Journal of Physics Condensed Matter, 1989, 1, 1879-1888. | 0.7 | 8 |
| 116 | Atomistic study of porosity impact on phonon driven thermal conductivity: Application to uranium dioxide. Journal of Applied Physics, 2014, 115, . | 1.1 | 8 |
| 117 | Influence of spin-orbit coupling on the atomic stability of the (100) face of 5d BCC transition metals. Solid State Communications, 1985, 55, 961-965. | 0.9 | 7 |
| 118 | Theoretical study of surface alloy formation through generation and annihilation of vacancies. Surface Science, 1996, 352-354, 552-556. | 0.8 | 7 |
| 119 | Atomistic simulations of relaxation and reconstruction phenomena in heteroepitaxy: $Co/Au(1\ 1\ 1)$. Applied Surface Science, 2002, 188, 134-139. | 3.1 | 7 |
| 120 | Kinetics study of antimony adsorption on Si(1 1 1). Applied Surface Science, 2003, 212-213, 715-723. | 3.1 | 7 |
| 121 | Ordering trends in transition metal alloys from tight-binding electronic structure calculations. Physical Review B, 2011, 84, . | 1.1 | 7 |
| 122 | Surface Segregation Maps Derived from Tight-Binding Ising Model. Solid State Phenomena, 0, 172-174, 1008-1015. | 0.3 | 7 |
| 123 | Surface segregation trends in transition metal alloys. Physical Review B, 2013, 88, . | 1.1 | 7 |
| 124 | Theoretical study of xenon adsorption in UO ₂ nanoporous matrices. Journal of Physics Condensed Matter, 2014, 26, 485015. | 0.7 | 7 |
| 125 | Environment dependence of magnetic moment and atomic level shifts within tight-binding approximation: An illustration in the case of cobalt. Surface Science, 2016, 646, 261-268. | 0.8 | 7 |
| 126 | Computation of interatomic Green functions for transition metals using continued fraction techniques. Journal De Physique, 1984, 45, 283-290. | 1.8 | 7 |

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| 128 | Sb/Si(111)Adsorption: Hidden Phase Transitions Behind Langmuir-Like Isotherms. Physical Review Letters, 2005, 94, 076101. | 2.9 | 6 |
| 129 | display="inline"> <mml:mrow><mml:mi mathvariant="normal">Sb</mml:mi><mml:mo><mml:mo><mml:mi mathvariant="normal">Si</mml:mi><mml:mi><mml:mo><mml:mo><mml:mo><mml:mn>111</mml:mn><mml:mo>)</mml:mo>)</mml:mo> (*/mml:mo> (*/mm</mml:mo></mml:mo></mml:mi></mml:mo></mml:mo></mml:mrow> | mml:mo>< | /mml:mrow |
| 130 | 2008, 77, Experimental and theoretical investigation of correlation effects in aggregates of Ni and Pd as a function of their size. Journal of Physics Condensed Matter, 1989, 1, 5875-5885. | 0.7 | 5 |
| 131 | Comparison of an experimental and a theoretical study of the surface alloy Pt/Cu(111). Journal of Electron Spectroscopy and Related Phenomena, 1994, 68, 357-362. | 0.8 | 5 |
| 132 | Sb doping of Si molecular-beam epitaxial layers: $\hat{a} \in f$ Influence of the substrate misorientation. Physical Review B, 1997, 56, 7615-7622. | 1.1 | 5 |
| 133 | Formation of an unexpected ordered two-dimensional Ag2Pb surfacealloy on Ag(111): A SXRD and STM study. Journal of Physics and Chemistry of Solids, 2006, 67, 601-604. | 1.9 | 5 |
| 134 | Molecular dynamics simulation of silicon oxidization. Thin Solid Films, 2010, 518, 2422-2426. | 0.8 | 5 |
| 135 | Disentangling coordination and alloy effects in transition-metal nanoalloys from their electronic structure. Physical Review B, 2013, 88, . | 1.1 | 5 |
| 136 | Electronic structure of CoPt based systems: from bulk to nanoalloys. Journal of Physics Condensed Matter, 2015, 27, 455503. | 0.7 | 5 |
| 137 | How to derive tight-binding <i>spd </i> potentials? Application to zirconium. Journal of Physics Condensed Matter, 2015, 27, 336301. | 0.7 | 5 |
| 138 | Intergranular Segregation in Cu(Ag) and Ag(Cu) Systems: Analysis of the Driving Force Using a Tight-Binding Scheme. Materials Science Forum, 1996, 207-209, 701-704. | 0.3 | 4 |
| 139 | Thermal dependence of surface polymorphism: the Ag/Cu (111) case. Applied Surface Science, 2001, 177, 252-257. | 3.1 | 4 |
| 140 | Role of sp–d hybridization in the formation of stacking defects at metal surfaces. Surface Science, 2008, 602, 2681-2688. | 0.8 | 4 |
| 141 | SURFACE ALLOY FORMATION IN THE Cu–Pd(111) SYSTEM: A KTBIM APPROACH. Surface Review and Letters, 1997, 04, 1119-1122. | 0.5 | 3 |
| 142 | Tight-Bindingn-momentpotential for zirconium hydride atomistic modeling. Metallurgical Research and Technology, 2015, 112, 102. | 0.4 | 3 |
| 143 | Stress influence on substitutional impurity segregation on dislocation loops in IV–IV semiconductors. Computational Materials Science, 2016, 114, 23-32. | 1.4 | 3 |
| 144 | Alliage de surface et équilibre local dans le systÃ"me Pd/Cu (111). European Physical Journal Special Topics, 1996, 06, C7-155-C7-158. | 0.2 | 3 |

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| 145 | Combining Solid State Physics Concepts and X-Ray Absorption Spectroscopy to Understand DeNOx Catalysis. Oil and Gas Science and Technology, 2006, 61, 677-689. | 1.4 | 3 |
| 146 | On the temperature dependence of the surface sandwiches observed in PtNi and AgNi alloys. Vacuum, 1990, 41, 441-445. | 1.6 | 2 |
| 147 | Theoretical prediction of new dissolution modes during metal heteroepitaxy. Journal of Crystal Growth, 1999, 198-199, 83-88. | 0.7 | 2 |
| 148 | Effect of magnetism on surface segregation in FeNi alloys. Journal of Physics Condensed Matter, 2016, 28, 064003. | 0.7 | 2 |
| 149 | Atomistic modelling of residual stress at UO2surfaces. Journal of Physics Condensed Matter, 2016, 28, 015006. | 0.7 | 2 |
| 150 | Tight-binding modelling of ferromagnetic metals and alloys. Modelling and Simulation in Materials Science and Engineering, 2017, 25, 084004. | 0.8 | 2 |
| 151 | Tight-binding modeling of interstitial ordering processes in metals: Application to zirconium hydrides. Physical Review B, 2020, 101, . | 1.1 | 2 |
| 152 | An atomistic approach for stress relaxation in materials., 2001,, 119-150. | | 2 |
| 153 | Surface Segregation in Transition Metal Alloys: From Electronic Structure to Phase Portraits. Progress of Theoretical Physics Supplement, 2013, 101, 159-180. | 0.2 | 2 |
| 154 | One- and two-hole excitation spectra of the 1d Hubbard model. Journal of Magnetism and Magnetic Materials, 1986, 54-57, 1029-1030. | 1.0 | 1 |
| 155 | Structure and properties of nanoscale materials: theory and atomistic computer simulation. International Journal of Nanotechnology, 2012, 9, 576. | 0.1 | 1 |
| 156 | Atomistic Model for Ge Condensation under SiGe Oxidation. Defect and Diffusion Forum, 0, 363, 210-216. | 0.4 | 1 |
| 157 | Theoretical study of xenon adsorption on UO ₂ surfaces. Journal of Physics Communications, 2018, 2, 035041. | 0.5 | 1 |
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| 159 | Theoretical and Experimental Evidences of Sequential Phase Formation during Sub-Nanometric-Thick Film Reactive Diffusion. Solid State Phenomena, 0, 172-174, 633-639. | 0.3 | 0 |
| 160 | Nanometric-Size Effect upon Diffusion and Reaction in Semiconductors: Experimental and Theoretical Investigations. Defect and Diffusion Forum, 0, 323-325, 433-438. | 0.4 | 0 |
| 161 | Electronic Structure of Nanoalloys: A Guide of Useful Concepts and Tools. Engineering Materials, 2012, , 159-195. | 0.3 | 0 |
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