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From adsorption to dissipation: insights from computer simulations of solid H₂O and CO

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From Adsorption to Dissipation: Insights from Computer Simulations of Solid H₂O and CO

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Cover description: Mountains and valleys symbolize energy peaks and troughs on solid surfaces. Figures climbing rock walls symbolize adsorbed surface species. Pebbles fall from the rock walls into the water below to create ripples, symbolizing energy dissipation. Water molecules are referenced via the sea in the cove; CO molecules are referenced via the *CO-conuts* in the palm tree.

Contents

1	Introduction	1
1.1	Computer Simulations	2
1.2	Adsorption	4
1.3	Vibrational Energy Relaxation	5
1.4	Aims and Scope	7
1.5	Main Results	8
1.6	Outlook	9
1.A	Bibliography	10
2	Methods	19
2.1	Force Fields	19
2.1.1	General Considerations	19
2.1.2	Force Fields for H ₂ O	22
2.1.3	Force Fields for CO	26
2.2	Geometry Optimization	29
2.3	Vibrational Analysis	30
2.3.1	Harmonic Approximation	30
2.3.2	Velocity Autocorrelation Function	33
2.4	Molecular Dynamics	36
2.4.1	Microcanonical Ensemble	38
2.4.2	Canonical Ensemble	39
2.A	Bibliography	40
3	YetAnotherSimulationSuite.jl: An Atomic Simulation suite in Julia	45
3.1	Features	46
3.2	Installation	46
3.3	Benchmarks	47
3.3.1	Geometry Optimization	47
3.3.2	Harmonic Frequencies	47
3.3.3	Molecular Dynamics	48
3.4	Memory Considerations	48
3.4.1	Geometry Optimization	49
3.4.2	Harmonic Frequency Calculation	49

3.4.3	Molecular Dynamics Simulation	49
3.5	Examples	49
3.A	Bibliography	51
4	Abundance of Exceptionally Strong Binding Sites for H₂O Adsorption on the Ice Ih(0001) Surface	53
4.1	Abstract	54
4.2	Introduction	54
4.3	Methods	56
4.4	Results and Discussion	58
4.5	Conclusion	63
4.A	Appendix	64
4.A.1	Optimized Lattice Vectors and Cohesive Energy	64
4.A.2	Order Parameter 2.0 Surfaces	64
4.A.3	Binding Energy Correlations	64
4.A.4	Bond Angle Analysis	65
4.B	Bibliography	70
5	Floating in Space: How to Treat the Weak Interaction between CO Molecules in Interstellar Ices	75
5.1	Abstract	76
5.2	Introduction	76
5.3	Computational Details	78
5.3.1	Force Field	78
5.3.2	Density Functional Theory	80
5.4	Results and Discussion	82
5.4.1	DFT Benchmark	82
5.4.2	Amorphous CO	83
5.4.3	Crystalline CO	90
5.5	Astrophysical Implications	92
5.6	Conclusion	93
5.7	Data Availability Statement	94
5.A	Appendix	94
5.A.1	Pair Interaction Validation	94
5.A.2	Geometry Optimization Threshold	94
5.A.3	Total Interaction Energy	96
5.A.4	DBSCAN	98

5.A.5	Dimer Geometries	98
5.B	Bibliography	101
6	Vibrational Energy Relaxation in Solid Carbon Monoxide	109
6.1	Abstract	110
6.2	Introduction	110
6.3	Methods	113
6.4	Structural and Vibrational Properties	116
6.5	Analyzing Vibrational Energy Relaxation with Classical Dynamics . .	120
6.5.1	Trajectory Ensembles	120
6.5.2	Frequency Shift	122
6.5.3	Frequency-Gap Dependent Decay Rate	124
6.6	Results and Discussion	125
6.6.1	Amorphous Cluster	125
6.6.2	Crystalline Cluster	127
6.6.3	Analysis of Energy Redistribution	128
6.6.4	Non-Resonant Energy Transfer	130
6.6.5	Resonant Energy Transfer	132
6.7	Summary and Outlook	133
6.8	Data Availability Statement	135
6.A	Appendix	135
6.A.1	Size Convergence Tests	135
6.A.2	Geometric and Vibrational Properties for the PIP–NN Potential	136
6.A.3	Decomposition of Phonon DOS	137
6.A.4	Trajectory Ensemble for Crystalline Clusters	138
6.A.5	Decomposition of Bi-Exponential Fits for Non-Resonant Energy Transfer	138
6.B	Bibliography	145
	Summary	151
	Samenvatting	155
	Publications	159
	Curriculum vitae	161
	Acknowledgments	163

Data Availability

165
