Linear free energy relationships in electrostatic catalysis

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The use of electric fields to modify chemical reactions is a promising, emerging technique in catalysis. However, there exist few guiding principles, and rational design requires assumptions about the transition state or explicit atomistic calculations. Here, we present a linear free energy relationship, familiar in other areas of physical organic chemistry, that microscopically relates field-induced changes in the activation energy to those in the reaction energy, connecting kinetic and thermodynamic behaviors. We verify our theory using first-principles electronic structure calculations of a symmetric SN2 reaction and the dehalogenation of an aryl halide on gold surfaces and observe hallmarks of linear free energy relationships, such as the shifting to early and late transition states. We also report and explain a counterintuitive case, where the constant of proportionality relating the activation and reaction energies is negative, such that stabilizing the product increases the activation energy, i.e., opposite of the Bell-Evans-Polanyi principle.

Recent years have seen an increase in the study of electric field effects on chemical reactions [1-10], motivated in part by the large electric fields that have been implicated in the catalytic performance of biological enzymes [11-14]. Chemical reactions, whose transitions states have large dipole moments are natural targets for electrostatic catalysis, because an external electric field will modify the activation energy and thus the reaction kinetics. However, as a rational design principle, this requires knowledge of the transition state geometry and electronic structure, which is nontrivial to obtain. In contrast, the reactant and product, which define the reaction thermodynamics, are much more easily characterized. Here, we derive and computationally test a linear free energy relationship, which relates field-induced changes in the activation energy to those in the reaction energy, providing a powerful new framework for design and characterization of electrostatic catalysis.

In the presence of a uniform electric field \( F \), the change in free energy \( G \) of a molecular system is approximately given by \( \Delta G = -\mu \cdot F \), where \( \mu \) is the component of the molecular dipole moment in the direction of the field. We consider a chemical reaction progressing through a reactant (R), transition state (TS), and product (P) geometry. Assuming that the field does not qualitatively alter the reaction pathway, the activation energy \( \Delta G^\dagger = G_{\text{TS}} - G_{\text{R}} \) and reaction energy \( \Delta G_{\text{rxn}} = G_{\text{P}} - G_{\text{R}} \) are modified according to

\[
\Delta G^\dagger = - (\mu_{\text{TS}} - \mu_{\text{R}}) F \equiv - \Delta \mu^\dagger F \quad (1)
\]

\[
\Delta G_{\text{rxn}} \equiv - (\mu_{\text{P}} - \mu_{\text{R}}) F \equiv - \Delta \mu_{\text{rxn}} F \quad (2)
\]

Combining Eqs. (1) and (2) gives the linear free energy relationship (LFER)

\[
\Delta G^\dagger = m \Delta G_{\text{rxn}} \quad (3a)
\]

\[
m = \frac{\Delta \mu^\dagger}{\Delta \mu_{\text{rxn}}} \quad (3b)
\]

Assuming an Arrhenius form to the reaction rate with a prefactor that is independent of the field yields the relationship

\[
\log \frac{k_F}{k_0} = m \log \frac{K_F}{K_0} \quad (4)
\]

where \( k_F, K_F \) (\( k_0, K_0 \)) are the reaction rate constant and equilibrium constant in the presence (absence) of a field. Therefore, the LFER (3) quantitatively relates the kinetics and thermodynamics of chemical reactions in electric fields.

Unlike most LFERs that are empirical, the LFER (3) provides a microscopic expression for the constant of proportionality or slope, \( m = \Delta \mu^\dagger/\Delta \mu_{\text{rxn}} \), whose magnitude determines the sensitivity of the activation energy to changes in the reaction energy. In particular, it encodes the common wisdom that reactions, whose transition state has a large dipole moment are especially susceptible to catalysis by an electric field, because in that case \( |\Delta \mu^\dagger| \gg 0 \). However, the slope \( m \) can have positive or negative sign. The case with positive slope signals that an electric field that stabilizes the product with respect to the reactant will also lower the activation energy; this behavior is conventional and forms the basis of the Bell-Evans-Polanyi principle [15]. The case with negative slope is unconventional and predicts the opposite behavior: applying an electric field that lowers the reaction energy will increase the activation energy.

To elucidate the theory, we first consider a simple, symmetric SN2 reaction of I\(^-\) with methyl iodide [Fig. 1]. Using density functional theory (DFT) with the B3LYP functional, we calculated the minimum energy reaction pathway in the absence and presence of an external electric field directed along the I-C-I bond axis [Fig. 1(b)]. First we observe that, over the range of field strengths -1 to 1 V/nm, the activation energy can be tuned by over 8 kcal/mol, corresponding to a factor of \( 10^5 \) in the reaction rate at room temperature and highlighting the potential power of electrostatic catalysis. More important for the present work, we observe that the reaction energy and activation energy are found to vary roughly linearly with the field strength [Fig. 1(d)], in agreement with Eqs. (1) and (2) (deviations from linearity at larger field strengths are due to molecular polarizabilities and changes in the geometries). Therefore, the LFER (3) holds to an excellent approximation, with \( \Delta G^\dagger \approx (1/2) \Delta \Delta G_{\text{rxn}} \) [Fig. 1(e)].

The observed slope \( m \approx +1/2 \) is readily explained by an analysis of the evolution of the dipole moment over the course of the reaction in the absence of a field [Fig. 1(c)]. Importantly, the dipole moment decreases monotonically, from \( \mu = \ldots \)

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FIG. 1. Symmetric $S_\sigma^2$ reaction of I$^-$ with methyl iodide. (a) Geometries of reactant (R), transition state (TS), and product (P) in the absence of a field. (b) Reaction energy in the absence and presence of a field of strengths indicated. (c) Dipole moment evolution in the absence of a field. (d) Reaction energy (orange, left axis) and activation energy (blue, right axis) as a function of field strength. (e) Comparison of the LFER prediction to the DFT simulation results.

8.4 D (R) to 0.0 D (TS) to −8.4 D (P). Therefore, the dipole moment differences $\Delta \mu R = 8.4$ D and $\Delta \mu_{TS} = 16.8$ D have the same (negative) sign, and their ratio, the slope $m$, is positive. Moreover, the magnitude of the slope is explained by the ratio $m = \Delta \mu R / \Delta \mu_{TS} = (-8.4)/(-16.8) = +1/2$, in agreement with the computational observation.

The field-dependent reaction profiles in Fig. 1(b) also show the behavior of “early” and “late” TSs, consistent with Hammond’s postulate. For example, in the presence of a field with $F = -1$ V/nm, the reaction is more exothermic, has a lower reaction energy, and has an early TS. The change in the TS geometry can be quantified: the bond length between the attacking iodine and the carbon is found to be 2.60 Å, 2.72 Å, and 2.85 Å and the angle made by the attacking iodine, carbon, and hydrogen is found to be 94°, 98°, and 86°, in the early, middle, and late TS geometries, respectively. We conclude that simple reactions influenced by electric fields obey the LFER (3) to a good approximation and exhibit many of its hallmarks known from synthetic organic chemistry.

Next, we test our theory on a more complicated example from heterogeneous catalysis. Specifically, we computationally study the dehalogenation of an iodobenzene molecule on a gold surface. This reaction is a key step in the metal-catalyzed Ullmann coupling reaction of aryl halides [16]. Participation of the metal is crucial, because the aryl halide bond dissociation energy is about +70 kcal/mol in the gas phase. All calculations are performed with periodic DFT with the optB86b-vdW functional, whose accuracy has been established for noncovalent interactions [17], and we identify minimum-energy reaction pathways using the climbing-image nudged elastic band approach.

We apply an electric field in the direction perpendicular to the gold surface with both positive and negative sign. This electric field may be internal, due to the electrostatics of a solid-liquid interface or local field effects. The field may also be external and applied controllably using the gold tip of a scanning tunneling microscope (STM), which has recently been used in a break-junction arrangement to experimentally demonstrate electrostatic catalysis [18, 19]. Again, we consider a maximum field strength of 1 V/nm, which is appropriate for these experimental realizations. We emphasize that quantitative properties of the electric field will depend on atomistic details of the surface and solvent as well as their dynamics, and our study is only qualitative.

To model the surface roughness, we first study the dehalogenation reaction on a gold (111) surface with an additional gold adatom at a hollow site (Fig. 2). In the absence of a field, we find a reaction energy of $\Delta G_{TS} = -13.4$ kcal/mol and an activation energy of $\Delta G^\ddagger = +4.1$ kcal/mol. Notably, this activation energy is significantly less than the one previously calculated for the same reaction on a clean gold surface, i.e., without an adatom, which was +16.9 kcal/mol [16]. This large reduction in the activation energy agrees with the common wisdom that rough metal surfaces are better catalysts than clean metal surfaces.

Assuming that the reaction pathway does not change in the presence of the field, we recalculate the electronic energy in the presence of an external electric field in the range $F = -1$ V/nm to $F = +1$ V/nm [Fig. 2(b)], where a posi-
tive field points away from the surface, i.e., in the positive z-direction. (Preliminary testing indicates that reoptimizing the geometries in the presence of the field does not significantly alter our findings over the range of fields considered here.) We find that application of an electric field indeed catalyzes the reaction, albeit only slightly: a field of $F = +1 \text{ V/nm}$ lowers the activation energy by 0.45 kcal/mol, which would increase the reaction rate by a factor of two at room temperature. Reversing the direction of the field raises the activation energy by a similar amount and thus suppresses the reaction. More importantly, the reaction energy and activation energy are again observed to be roughly linear in the field strength [Fig. 2(d)] and the TS shifts to an earlier one as the reaction becomes more favorable [inset of Fig. 2(b)]. Similar to the $S_N(2)$ behavior, the LFER (3) holds approximately with a positive slope less than one, $\Delta G^\ddagger \approx (+0.3)\Delta G_{\text{rxn}}$ [Fig. 2(e)].

The sign and magnitude of the slope can again be explained by analyzing the evolution of the dipole moment (in the direction of the field) over the course of the reaction in the absence of the field from $\mu = 4.4$ D (R) to 3.4 D (TS) to 0.75 D (P) [Fig. 2(c)]. Therefore, the dipole moment differences $\Delta \mu^\ddagger = -1.0$ D and $\Delta \mu_{\text{rxn}} = -3.6$ D have the same (negative) sign, and their ratio, the slope $m$, is positive. Moreover, the magnitude of the slope is explained by the ratio $\Delta \mu^\ddagger / \Delta \mu_{\text{rxn}} \approx 0.3$, which is why the change in the activation energy is about three times smaller than the change in the reaction energy.

Unlike for the simple $S_N(2)$ reaction, the behavior of the dipole moment for this surface-mediated dehalogenation is not easily rationalized, because the largest changes to the electronic structure occur in the direction parallel to the surface, i.e., perpendicular to the field. However, analysis of our DFT calculations in terms of electron densities and Löwdin charges (not shown) can provide some insight into the evolution of the dipole moment in the z direction. In the chemisorbed reactant, there is significant electron transfer from the molecule to the gold, establishing a dipole moment in the positive z direction. The reaction occurs via an oxidative addition-like pathway, whereby electron transfer from the undercoordinated gold adatom onto the aryl halide facilitates bond breaking and slightly reduces the dipole moment of the TS. In the product geometry, the aryl ring has a greater electron density and thus a less positive total charge, which we attribute to improved resonance with the gold orbitals. Overall, the charge separation between molecule and gold is smallest in the product, consistent with its small dipole moment.

In our opinion, the previous two computational studies confirm the most important result of the present work, i.e., that a LFER holds for electrostatic catalysis and that, if its slope is positive, changes to the activation energy are positively proportional to changes in the reaction energy. For the endeavor of electrostatic catalysis, this finding shifts focus from the activation energy—and its associated challenge of transition state design—to the much more manageable reaction energy. Alternatively, an experimental measurement of the slope $m$ appearing in the LFER (3) can be used to infer details about the reaction pathway, including its TS and associated dipole moment differences. We expect the positivity of the slope to hold in many, but not all, reactions. Specifically, a sufficient, but not necessary condition for the positivity of the slope is that the dipole moment evolution is monotonic (increasing or decreasing) over the course of the reaction, which is a common and intuitive behavior. However, many important reactions have a transition-state dipole moment that is larger than that of the reactant and product. In this case, the slope may be negative, which we demonstrate now with a closely related computational example.

We study the same dehalogenation reaction on a clean gold (111) surface, i.e., without an adatom [Fig. 3]. In this case, oxidative addition by gold is sterically hindered and the activation energy is significantly higher, $\Delta G^\ddagger = 17.74$ kcal/mol. The activation energy and reaction energy are again linear in the strength of the field [Fig. 3(d)] and can be weakly tuned by about 0.3 kcal/mol and 0.4 kcal/mol, respectively. More interestingly, the LFER (3) holds, but with a negative slope [Fig. 3(e)]. In other words, the field orientation that lowers the reaction energy actually raises the activation energy and shifts the TS to an earlier one [Fig. 3(b)]. This negative slope is explained by the dipole moment, which is nonmonotonic over the course of the reaction, evolving from 2.8 D (R) to 3.2 D (TS) to 2.2 D (P), i.e., with $\Delta \mu^\ddagger \approx +0.4$ D and $\Delta \mu_{\text{rxn}} \approx -0.6$ D [Fig. 3(c)]. Thus the LFER (3) predicts a slope of $m \approx (+0.4)/(-0.6) = -(2/3)$, in good agreement with the computational result.

Again, analysis of our DFT calculations can rationalize the behavior of the dipole moment. We find that the charge assignments of the reactant and product are qualitatively similar to those in the presence of the adatom. However, we find that in the TS, rather than accepting electron density from the gold, the aryl iodide molecule donates additional electron density to the gold, increasing the extent of charge separation and thus
increasing the magnitude of the dipole moment. This example shows how the analysis of reactions in terms of the LFER (3) can be used to uncover unusual mechanistic details of complex chemical reactions.

The theory developed here is general and can be readily applied in other situations. For example, the electric field need not be externally applied, and could be self-induced, as was recently discussed for a molecule in a plasmonic nanocavity [20]. Specifically, a molecule with dipole moment $\mu$ sandwiched between two metals will induce a proportional reaction field $F \propto \mu$, which lowers the molecular energy by $\Delta G \propto \mu^2$. A LFER naturally follows with the form $\Delta G' = m'\Delta G_{\text{rxn}}$, where now the slope is given by the difference in the square of the dipole moments,

$$m' = \frac{\mu_{TS}^2 - \mu_R^2}{\mu_p^2 - \mu_R^2}. \quad (4)$$

Therefore, we expect the work presented here, which provided a general, but microscopic relationship between the kinetics and thermodynamics of chemical reactions in electric fields to have impact throughout the field of electrostatic catalysis, similar to how scaling relations have revolutionized heterogeneous catalysis [21–23].

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**COMPUTATIONAL DETAILS**

All DFT calculations for the $S_N2$ reaction were performed using the Orca [24] quantum chemistry package with the B3LYP functional [25, 26] and the def2-TZVP basis set [27, 28]. The climbing-image nudged elastic band method [29–33] followed by the eigenvalue-based TS searching algorithm [34, 35] (Orca keyword “NEB-TS”) was used to find the minimum energy path and the TS structure. For each applied field, the reaction path geometries were fully optimized.

All periodic DFT calculations for the dehalogenation reaction on a gold surface were performed using the Quantum Espresso package [36]. Following Ref. 16, we applied the optB86b-vdW functional [17], which provides an affordable description of dispersion interactions, yielding results comparable to those obtained by the random phase approximation [16, 17]. Core electrons were treated using the projector augmented wave method [37, 38]. We used a 500 eV kinetic energy cutoff and a 2×2×1 sampling of the Brillouin zone. Gold surfaces were modeled using 5×5 slabs, 15 Å of vacuum, and a lattice constant of 4.140 Å. The clean gold (111) surface was modeled by a four-layer slab, and the two uppermost layers were allowed to relax during geometry optimizations. The system with an extra adatom was modeled by a three-layer slab, and the adatom and uppermost layer were allowed to relax. The transition state calculations were performed with the climbing image nudged elastic band approach, using 12 images. The electric field was applied with a sawtooth potential [36] and all calculations were dipole-corrected [39] using a dipole length of 0.89 Å in the direction of the field.

[1] Sason Shaik, Debasish Mandal, and Rajeev Ramanan. Oriented electric fields as future smart reagents in chemistry. *Nat. Chem.*, 8(12):1091–1098, 2016.

[2] Fanglin Che, Jake T Gray, Su Ha, Norbert Kruse, Susannah L Scott, and Jean-Sabin McEwen. Elucidating the roles of electric fields in catalysis: A perspective. *ACS Catal.*, 8(6):5153–5174, 2018.

[3] Simone Ciampi, Nadim Darwish, Heather M Aitken, Ismael Díez-Pérez, and Michelle L Coote. Harnessing electrostatic catalysis in single molecule, electrochemical and chemical systems: a rapidly growing experimental tool box. *Chem. Soc. Rev.*, 47(14):5146–5164, 2018.

[4] Cina Foroutan-Nejad and Radek Marek. Potential energy surface and binding energy in the presence of an external electric field: modulation of anion–π interactions for graphene-based receptors. *Phys. Chem. Chem. Phys.*, 16(6):2508–2514, 2014.

[5] Vivian M Lau, William C Pfalzgra, Thomas E Markland, and Matthew W Kanan. Electrostatic control of regioselectivity in an (i)-catalyzed hydroarylation. *J. Am. Chem. Soc.*, 139(11):4035–4041, 2017.

[6] Craig F Gorin, Eugene S Beh, and Matthew W Kanan. An electric field–induced change in the selectivity of a metal oxide–catalyzed epoxide rearrangement. *J. Am. Chem. Soc.*, 134(1):186–189, 2012.

[7] Craig F Gorin, Eugene S Beh, Quan M Bui, Graham R Dick, and Matthew W Kanan. Interfacial electric field effects on a carbene reaction catalyzed by rh porphyrins. *J. Am. Chem. Soc.*, 135(30):11257–11265, 2013.

[8] Valerie Vaissier Welborn, Luis Ruiz Pestana, and Teresa Head-Gordon. Computational optimization of electric fields for better catalysis design. *Nat. Catal.*, 1(9):649–655, 2018.

[9] Sason Shaik, David Danovich, Jyothish Joy, Zhanfeng Wang, and Thijs Stuyver. Electric-field mediated chemistry: Uncovering and exploiting the potential of (oriented) electric fields to exert chemical catalysis and reaction control. *J. Am. Chem. Soc.*, 142(29):12551–12562, 2020.

[10] Jyothish Joy, Thijs Stuyver, and Sason Shaik. Oriented external electric fields and ionic additives elicit catalysis and mechanis-
tic crossover in oxidative addition reactions. J. Am. Chem. Soc., 142(8):3836–3850, 2020.

[11] Shina CL Kamertin, Pankaz K Sharma, Zhen T Chu, and Arieh Warshel. Ketosteroid isomerase provides further support for the idea that enzymes work by electrostatic preorganization. Proc. Natl. Acad. Sci., 107(9):4075–4080, 2010.

[12] Stephen D Fried, Sayan Bagchi, and Steven G Boxer. Extreme electric fields power catalysis in the active site of ketosteroid isomerase. Science, 346(6216):1510–1514, 2014.

[13] Arieh Warshel, Pankaz K Sharma, Mitsuonri Kato, Yun Xiang, Hanbin Liu, and Mats HM Olsson. Electrostatic basis for enzyme catalysis. Chem. Rev., 106(8):3210–3235, 2006.

[14] Valerie Vaissier Welborn and Teresa Head-Gordon. Fluctuations of electric fields in the active site of the enzyme ketosteroid isomerase. J. Am. Chem. Soc., 141(32):12487–12492, 2019.

[15] Rutger A Van Santen, Matthew Neurock, and Sharan G Shetty. Reactivity theory of transition-metal surfaces: a bornsted-evans–polanyi linear activation energy–free-energy analysis. Chem. Rev., 110(4):2005–2048, 2009.

[16] Jonas Bjork, Felix Hanke, and Sven Sstafstrom. Mechanisms of halogen-based covalent self-assembly on metal surfaces. J. Am. Chem. Soc., 135(15):5768–5775, 2013.

[17] Jiří Klimeš, David R Bowler, and Angelos Michaelides. Van der waals density functionals applied to solids. Phys. Rev. B, 83(19):195131, 2011.

[18] Rinat Meir, Hui Chen, Wenzhen Lai, and Sason Shaik. Oriented electrostatic catalysis. J. Phys. Chem. A, 125(13):3704–3705, mar 2003.

[19] Stefano Baroni, Nicola Bonini, Matteo Calandra, Rocco Carretta, Carlo Cavazzoni, Davide Ceresoli, Guido L Chiarotti, Matteo Cococcioni, Ismael Dabo, et al. Quantum espresso: a modular and open-source software project for quantum simulations of materials. J. Phys. Condens. Matter Phys., 21(39):395502, 2009.

[20] Claudia Climent, Javier Galego, Francisco J Garcia-Vidal, and Johannes Feist. Plasmonic nanocavities enable self-induced electrostatic catalysis. Angew. Chem. Int. Ed., 58(26):8698–8702, 2019.

[21] J.K. Nørskov, T. Bligaard, A. Logadottir, S. Bahn, L.B. Hansen, M. Bollinger, H. Bengaard, Z. Slijivanca, M. Mavrikakis, Y. Xu, S. Dahl, and C.J.H. Jacobsen. Universality in heterogeneous catalysis. J. Catal., 209(2):275–278, jul 2002.

[22] Francoise Jaccard, M. Dravitzki, and P. Gaspard. A quantum mechanical model for the reaction of hydrogen with nickel. J. Am. Chem. Soc., 59(19):12301, 1957.

[23] Frank Neese. Software update: the orca program system, version 4.0. WIREs Comput. Mol. Sci., 8(1):e1327, 2018.

[24] Axel D Becke. Density-functional exchange-energy approximation with correct asymptotic behavior. Phys. Rev. A, 38(6):3098, 1988.

[25] Axel D Becke. A new mixing of hartree–fock and local density-functional theories. J. Chem. Phys., 98(2):1377–1377, 1993.

[26] Angsar Schäfer, Christian Huber, and Reinhart Ahlrichs. Fully optimized contracted gaussian basis sets of triple zeta valence quality for atoms li to kr. J. Chem. Phys., 100(8):5829–5835, 1994.

[27] Florian Weigend and Reinhart Ahlrichs. Balanced basis sets of split valence, triple zeta valence and quadruple zeta valence quality for h to mo: Design and assessment of accuracy. Phys. Chem. Chem. Phys., 7(18):3297–3305, 2005.

[28] Greg Mills and Hannes Jönsson. Quantum and thermal effects in h2 dissociative adsorption: Evaluation of free energy barriers in multidimensional quantum systems. Phys. Rev. Lett., 72(7):1124, 1994.

[29] Gregory Mills, Hannes Jönsson, and Gregory K Schenter. Reversible work transition state theory: application to dissociative adsorption of hydrogen. Surf. Sci., 324(2-3):305–337, 1995.

[30] Bruce along, Giovanni Cicotti, and David F Coker. Classical and quantum dynamics in condensed phase simulations: Proceedings of the International School of Physics "Enrico Fermi", 1998.

[31] Graeme Henkelman, Blas P Uberuaga, and Hannes Jönsson. A climbing image nudged elastic band method for finding saddle points and minimum energy paths. J. Chem. Phys., 113(22):9901–9904, 2000.

[32] Graeme Henkelman and Hannes Jönsson. Improved tangent estimate in the nudged elastic band method for finding minimum energy paths and saddle points. J. Chem. Phys., 113(22):9978–9985, 2000.

[33] Ajit Barak, Noah Adams, Jack Simons, and Ron Shepard. Search for stationary points on surfaces. J. Phys. Chem., 89(1):52–57, 1985.

[34] Jon Baker. An algorithm for the location of transition states. J. Comput. Chem., 7(4):385–395, 1986.

[35] Paolo Giannozzi, Stefano Baroni, Nicola Bonini, Matteo Calandra, Roberto Caro, Carlo Cavazzoni, Davide Ceresoli, Guido L Chiarotti, Matteo Cococcioni, Ismael Dabo, et al. Quantum espresso: a modular and open-source software project for quantum simulations of materials. J. Condens. Matter Phys., 21(39):395502, 2009.

[36] Peter E Blöchl. Projector augmented-wave method. Phys. Rev. B, 50(24):17953, 1994.

[37] Peter E Blöchl, Ove Jepsen, and Ole Krogh Andersen. Improved tetrahedron method for brillouin-zone integrations. Phys. Rev. B, 49(23):16223, 1994.

[38] Lennart Bengtsson. Dipole correction for surface supercell calculations. Phys. Rev. B, 59(19):12301, 1999.