Commensurate-Incommensurate Transition and Domain-Wall Dynamics of Adsorbed Overlayers on a Honeycomb Substrate

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Abstract – We introduce an effective one-mode Phase-field Crystal model for studying the commensurate-incommensurate transition and domain wall dynamics of the $(\sqrt{3} \times \sqrt{3})R30^{\circ}$ phase found in systems such as Xe/Pt(111), or Xe and Kr on Graphite. The model allows us to study large systems where the domain walls can be separated over large macroscopic distances and at same time incorporate the microscopic details of the domain wall structures. The resulting phase diagram shows that an intermediate stripe incommensurate phase always separates the commensurate phase from the honeycomb incommensurate phases. The energy of the domain wall crossing is investigated. We also find that near a step edge, the domain walls tend to align perpendicular to the step edge, in agreement with recent experimental observations.

In many adsorption systems the maxima (minima) of 1 the adsorption potential due to the substrate form a hon-2 eycomb (triangular) lattice [1]. This has been referred to previously as a honeycomb substrate [2]. Examples of such systems, which have been investigated in detail, are 5 Xe on graphite [3], Kr on graphite [4] and Xe/Pt(111) [5]. 6 For these adsorption systems, the ideal separation between the adsorbate atoms is close to $\sqrt{3}a$, where a is the distance between neighboring adsorption potential minimum. 9 For small strain energies, the lowest energy commensurate 10 states is the one that adatoms occupy only a third of the 11 minima, forming the commensurate $(\sqrt{3} \times \sqrt{3})R30^{\circ}$ phase. 12 Experimentally, an apparent continuous phase transition 13 from the commensurate $(\sqrt{3} \times \sqrt{3})R30^\circ$ to an incommen-14 surate phase with a honeycomb network of domain walls 15 (HoI phase) has been observed for Kr and Xe on Graphite 16 as function of temperature and coverage. In addition, for 17 Xe/Pt(111) another uniaxial symmetry incommensurate 18 phase with a striped array of domain walls has been ob-19 served [3, 5]. 20

²¹ Several theoretical works [6–8], however, have deter-

mined that the transition from the commensurate ($\sqrt{3}$ × 22 $\sqrt{3}R30^{\circ}$ to the HoI phase should be first order. Bak 23 et al. [7] argued that the symmetry of the domain wall 24 patterns in the incommensurate phase is determined by 25 the energy of the domain wall crossing. For a negative 26 value of domain wall crossing energy, the HoI phase is fa-27 vored rather than a parallel set of striped domain wall 28 (SI) phase. This theory predicts that in this case a dis-29 continuous transition from commensurate phase to a HoI 30 phase should take place. For positive domain-wall crossing 31 energy, the number of crossings must be as small as pos-32 sible and the transition of the commensurate phase into 33 SI phase is favored. This transition is predicted to be 34 continuous with the distance between the walls diverging 35 logarithmically as one approaches the critical point. This 36 is followed by a first-order transition from the SI phase to 37 the HoI phase. Note that the terminology of stripe, hon-38 eycomb used here refers to the symmetry of the domain 39 wall pattern separating the commensurate domains and 40 not the lattice joining the center of domains that has also 41 been used in the literature [9]. 42

Although these theoretical arguments about the rela-43 tion of energy of the domain wall crossing to the nature 44 of the actual incommensurate phases are quite general, 45 they are based on a mean field theory that ignore ther-46 mal fluctuations and wall thickness. Even within mean 47 field theory, only the SI phase is well understood, while 48 the domain wall crossing energy and the details of the two 49 dimensional honeycomb network of domain walls and its 50 energetics have not been fully investigated analytically. 51 In addition computational efforts using molecular dynam-52 ics have been hindered by the large length scales required 53 for examining the commensurate-incommensurate transi-54 tion. In this work, we will examine these issues using 55 an amplitude representation of the phase field crystal for-56 malism [11] which allows us to study the microscopic de-57 tails of the domain wall crossings and the HoI and SI 58 phases, yet allowing the sample size to be macroscopic 59 [9, 10] which is particularly important near the onset of 60 the commensurate-incommensurate transition. 61

It has been shown [2, 9–11] that instead of the fully microscopic atomic model, a useful theoretical formalism for studying the different phases in adsorption systems and the transition between them is the phase field crystal (PFC) model. In this model, the free energy of the adsorption system is described by the functional

$$F[n(\vec{r})] = \int d\vec{r} \left\{ \frac{n(\vec{r})}{2} \left[\Delta B + B_x (q_0^2 + \nabla^2)^2 \right] n(\vec{r})(1) - \frac{t}{3} n(\vec{r})^3 + \frac{v}{4} n(\vec{r})^4 + V(\vec{r}) n(\vec{r}) \right\}.$$

where $V(\vec{r})$ is the periodic adsorption potential due to the 68 adatom substrate interaction with a lattice constant a as 69 depicted in Fig. 1a, and n(r) is a coarse grained dimen-70 sionless density field measured from the average density 71 \bar{n} characterizing the real space configuration of the ad-72 sorbate film. The adatoms form a hexagonal phase in 73 the absence of the substrate potential with lattice con-74 stant $a_f = 4\pi/\sqrt{3}q_0$ and the commensurate $\sqrt{3} \times \sqrt{3}R30^\circ$ 75 phase has lattice constant $a_c = \sqrt{3}a$. The lattice misfit 76 parameter is defined as $\epsilon = (a_c - a_f)/a_c$. We are inter-77 ested in the various phases which can occur due to the 78 competition of the strain energy of the film resulting from 79 the lattice mismatch and the potential coupling between 80 substrate and film. The equilibrium density $n(\vec{r})$ is deter-81 mined by minimizing the free-energy functional in Eq.(1). 82 However, solving for the minimum free energy phase using 83 the full PFC model is limited to relative small size sys-84 tems because of the fine-grid mesh required to resolve all 85 the atomic scale details of the various phases. In the past, 86 an alternate amplitude expansion has been developed to 87 overcome this problem [9, 10, 12-17]. 88

When applying this formalism to the adsorption system, it is useful to expand the density in terms of the commensurate sublattices states that are depicted in Fig. 1b, i.e.,

$$n(\vec{r}) = \sum_{kl} \left[\eta_{kl} e^{i\vec{G}_{kl}\cdot\vec{r}} + c.c. \right], \qquad (2)$$

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where η_{kl} are complex amplitudes, $\vec{G}_{kl} = k\vec{q}_1 + l\vec{q}_2$, (\vec{q}_1, \vec{q}_2) are the principal reciprocal lattice vectors and klare the Miller indices for a film of two dimensional triangular symmetry the reciprocal vectors are $\vec{q}_1 = q_c(1,0)$ and $\vec{q}_2 = q_c(-1,\sqrt{3})/2$, where $q_c = 4\pi/\sqrt{3}a_c$. For the commensurate $(\sqrt{3} \times \sqrt{3})R30^\circ$ phase, there are three equivalent sublattices with the lattice constant $a_c = \sqrt{3}a$ as shown in Fig. 1b. A given sublattice is a triangular lattice with lattice constant a_c and displaced from the other sublattices by $\vec{\Delta}_1 = [0,0], \vec{\Delta}_2 = [a,0]$ and $\vec{\Delta}_3 = [2a,0]$.



Fig. 1: Schematic of a) graphite surface and b) possible commensurate sublattices. In a) the grey circles and black points represent potential maxima and minima, respectively. In b) the red, blue and green points (or three shades of grey in monochrome rendering) represent the three equivalent $(\sqrt{3} \times \sqrt{3})R30^{\circ}$ commensurate sublattices.



Fig. 2: The phases found numerically for $\epsilon = 3\%$. The colors correspond to the closest sublattice (shown in Fig. 1b) that the adsorbed atoms sit at for the a) HoI and b) SI $(\sqrt{3} \times \sqrt{3})R30^{\circ}$ phases.

Note that the potential for the Xe/Pt(111) has exactly 103 the same symmetry shown in Fig. 1 since the adsorp-104 tion sites for this systems are the on-top positions for the 105 Pt(111) substrate [1] with triangular symmetry, and the 106 density field $n(\vec{r})$ for the $(\sqrt{3} \times \sqrt{3})R30^\circ$ phase of this 107 system can be described by the same Fourier expansion 108 as the Kr/Graphite and Xe/Graphite systems. Through-109 out the present study, we will adopt the simplest peri-110 odic adsorption potential with the right triangular sym-111 metry containing the lowest-order harmonics as given by $V(\vec{r}) = -V_0 \sum_{kl} e^{i\vec{G}_{kl}^s \cdot \vec{r}}$, where \vec{G}_{kl}^s are the same as \vec{G}_{kl} 112 113 given following Eq. (2) except rotated by 30° and larger 114 by a factor of $\sqrt{3}$. In the commensurate phase, the ampli-115 tudes, η_{kl} , in Eq. (2) are constant. To allow for the devel-116



Fig. 3: Phase diagram from the effective single mode model, defined by Eq. (4). The red (upper) and blue (lower) points correspond to the C-SI and SI-HoI transitions, while the lines are guides the eye. The dashed black line corresponds to the Sine-Gordon prediction [31] for the C-SI transition, i.e., $f^{\rm ad}/K = (\pi^2/16)\epsilon^2$. Note in the inset the blue and red points almost overlap.

opment of domain walls and the incommensurate phases,
these amplitudes are allowed to be a function of position
but only slowly varying in space, i.e., containing only small
wave vectors (compared to the reciprocal lattice vectors
for the commensurate phase).

From a practical point of view, one needs to limit the 122 number of amplitudes or modes η_{kl} in the expansion. Pre-123 viously, it has been demonstrated that even a single mode 124 approximation including all the amplitudes corresponding 125 to the reciprocal lattice vectors of the smallest magnitude 126 (i.e., the set η_{10} , η_{01} and $\eta_{\overline{1}\overline{1}}$) works well for the (1×1) 127 phase and yields all the interesting qualitative results for 128 the phase diagram [2, 9-11]. For the present commensu-129 rate $(\sqrt{3} \times \sqrt{3})R30^{\circ}$ phase of interest here, the smallest 130 amplitudes do not couple directly to the substrate adsorp-131 tion potential and the essential physics would be lost in 132 this "one" mode approximation. The minimal number of 133 sets of modes of the same magnitude that is required for 134 coupling to the substrate potential is two. 135

For the purpose of illustration, it is useful to derive an explicit expression for the free energy in the commensurate $(\sqrt{3} \times \sqrt{3})R30^{\circ}$ phase when only the first two sets of modes (corresponding to the set first set above and the second set $\eta_{12}, \eta_{1\bar{1}}$ and $\eta_{\bar{2}\bar{1}}$). Setting the magnitude of the first and second sets to be the constants Φ and Ω , respectively, gives a free energy per unit area A as,

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$$F_{c}[\Phi,\Omega]/A = F_{L} + 3\Delta B \left(\Omega^{2} + \Phi^{2}\right) - 6V_{0}\Omega \qquad (3)$$

+ $3B_{x} \left(\Omega^{2} \left(q_{0}^{2} - 3q_{c}^{2}\right)^{2} + \Phi^{2} \left(q_{0}^{2} - q_{c}^{2}\right)^{2}\right)$
- $4t \left(\Omega^{3} + 3\Omega\Phi^{2} + \Phi^{3}\right)$
+ $v \left(\frac{45}{2}\Omega^{4} + 90\Omega^{2}\Phi^{2} + 36\Omega\Phi^{3} + \frac{45}{2}\Phi^{4}\right),$

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where F_L is a constant independent of Φ and Ω . Note that 143 in the $(\sqrt{3} \times \sqrt{3})R30^{\circ}$ phase, the adsorption potential $V(\vec{r})$ 144 only couples to the second mode amplitude Ω . Minimizing 145 the free energy with respect to Ω first, we can solve for Ω 146 as a function of Φ . Substitution of this back into the free 147 energy then yields a free energy functional $F(\Phi)$ that de-148 pends only on the amplitude of the first mode Φ . This free 149 energy describes a first order phase transition from a dis-150 ordered or liquid state (i.e., Φ) to an ordered or crystalline 151 state (Φ finite) as a function of ΔB as illustrated in Fig. 4. 152 This procedure can obviously be generalized to eliminate



Fig. 4: Free energy density as a function of Φ for (from top to bottom) $\Delta B = 0.05, 0.047$ and 0.02. For $\Delta B < 0.047$ the free energy density is minimized by a crystalline state (finite Φ) and a liquid state for $\Delta B > 0.047$.

all higher modes to yield an effective one-mode free energy $F(\Phi)$. Note that aside from the modification of all the coefficients of the various powers of the first mode amplitude Φ in the free energy, the crucial modification is in the coupling term to the adsorption potential, which in this case will yield nonlinear terms of the form $V_0(a\Phi^2 + b\Phi^3 + ...)$.

Similar non-linear coupling terms would also appear if 160 a non-linear coupling of the density, n, to the surface potential, V were included in Eq. (1.) In this spirit we will 162

consider the lowest-order non-linear coupling term that couples the first set of modes to the potential. Replacing nV with $\lambda n^2 V$ in Eq. (1) gives for the first set of amplitudes,

$$F_{\eta} = \int d\vec{r} \left[\sum_{kl} \left(B^{x} |\mathcal{G}_{kl}\eta_{kl}|^{2} - \frac{3v}{2} |\eta_{kl}|^{4} \right) + \frac{\Delta B}{2} A^{2} + \frac{3v}{4} A^{4} - 2t \left(\prod_{kl} \eta_{kl} + \text{c.c.} \right) + 2\lambda V_{0} \left(\eta_{10}\eta_{01} + \eta_{01}\eta_{\bar{1}\bar{1}} + \eta_{\bar{1}\bar{1}}\eta_{10} + \text{c.c.} \right) \right], (4)$$

where the sums and products are over only the first set of modes, $A^2 \equiv 2 \sum_{kl} |\eta_{kl}|^2$, $\mathcal{G}_{kl} \equiv \nabla^2 + 2i\alpha \vec{G}_{kl}^f \cdot \vec{\nabla} + q_0^2 - q_c^2$ and c.c. is the complex conjugate. As can be seen in the 167 168 169 last line of Eq. (4) the density is now explicitly coupled 170 to the potential, showing that this approach is an effective 171 one-mode model for studying the $(\sqrt{3} \times \sqrt{3})R30^\circ$ phase 172 and its transition into an incommensurate phase. In fact, 173 all other similar higher-order commensurate states such 174 as the $(\sqrt{7} \times \sqrt{7})R19.1^{\circ} p(2 \times 2)$ phases can be studied 175 with a similar effective one-mode model, in spite of the 176 much more complicated domain wall structures and larger 177 number of equivalent sublattices [28]. 178

In the commensurate $(\sqrt{3} \times \sqrt{3})R30^{\circ}$ phase, one of the 179 three sublattices shown in Fig. 1b of the adsorption sites 180 is occupied. As the amplitude of adsorption potential V_0 181 decreases and/or the misfit ϵ increases, the adsorbate sys-182 tem undergoes a transition to an incommensurate phase, 183 consisting of commensurate regions occupying one of the 184 sublattices, separated by domain walls. Depending on the 185 values of the misfit parameter ϵ and the coupling strength 186 of the adsorption potential V_0 , such domain-wall structure 187 can form a HoI phase (Fig. 2a) or a SI phase (Fig. 2b). 188

¹⁸⁹ We have determined the phase diagram as a function of ¹⁹⁰ the lattice mismatch and strength of the adsorption poten-¹⁹¹ tial by minimizing the free-energy functional using the ef-¹⁹² fective single mode model of Eq. (4). The patterns emerge ¹⁹³ essentially due to a competition between the elastic and ¹⁹⁴ adhesion energies. For easy comparison with experiment ¹⁹⁵ we define the adhesion free energy per unit length, $f^{\rm ad}$ as

$$f^{\rm ad} = f^{\rm com} - f^{\rm elas},\tag{5}$$

where f^{com} and f^{elas} are the free energies of the commensurate and incommensurate states at the lattice spacing a_c per unit area, respectively. For Eq. (4), $f^{\text{ad}} = 8\lambda V_0 \Phi^2$. For the elastic contribution the relevant elastic modulus K or a displacement from one sublattice to another is

$$K = (C_{11} + C_{12})^2 / C_{11}, (6)$$

which can be measured experimentally. For the free energy given in Eq. (4), $K = 16\Phi^2 B^x$. Thus for easy comparison we plot the phase diagram in the plane of two dimensionless variables, the ratio of adhesion to elastic free energy densities $f^{\rm ad}/K$ and the mismatch parameter ε . The result is shown in Fig. 3.

For a large misfit there is a SI phase between the com-207 mensurate and HoI phases. It can be shown [28] in the 208 small-displacement limit that Eq. (4) can be mapped 209 into a Sine-Gordon model for the C-SI transition (in the 210 mean field limit) and a continuous transition occurs when 211 $f^{\rm ad}/K = (\pi^2/16)\varepsilon^2$. As can be seen in Fig. 3 this pre-212 diction works very well at small strains. It appears that 213 the C-SI and SI-HoI transition lines become very close at 214 small strains, but to the limit of the numerical simula-215 tions they never merge indicating that the SI state region 216 is present at all strains. It is important to note that it is 217 very difficult to examine the strains less than 1% as the 218 length scale of the patterns scale as the inverse of the mis-219 fit strain. For the smallest strain examined, $\varepsilon = \pm 1\%$ the 220 SI phase was present for some values of $f^{\rm ad}/K$. Finally, we 221 have also evaluated the phase diagram using the full PFC 222 model of Eq. (1) without using the effective one-mode 223 approximation and using the latter approximation with a 224 Vn^3 non-linear coupling instead of the Vn^2 coupling used 225 here. Although there are small quantitative changes in 226 the location of the phase boundaries, the topology of the 227 phase diagrams obtained from these models are essentially 228 the same as that shown in Fig. 3 for the effective one mode 229 model. 230

Close to the phase transition we found that for large 231 misfit parameters, $|\epsilon|$, the free energy of the wall crossing 232 is greater than that of the walls themselves. This can be 233 inferred from the local free energy (i.e., the integrand of 234 Eq. (4)) plot shown in Figs. 5a and b. For the case $\varepsilon = 8\%$ 235 it can be seen that the free energy density is higher at the 236 point where three domain walls cross. On the other hand, 237 for lower misfits the free energy density reduces slightly 238 at the crossing. For strong adsorption potential and large 239 misfit, we find wide regions where the SI phase (Fig. 2 b)) 240 is the lowest energy. Surprisingly, the SI to commensurate 241 transition in this limit is of first order. For lower values of 242 pinning the lowest energy phase is the HoI phase (Fig. 2 243 a)). The transition from SI to HoI is discontinuous. When 244 the misfit parameter $|\epsilon|$ is decreased the stability range of 245 the SI phase (Fig. 3) decreases quickly. For misfit parame-246 ters between 1% and -1%, it is numerically very costly to 247 analyze the stability of the SI phase, but it reasonable to 248 assume that the SI phase is present for all misfit param-249 eters in the absence of fluctuations. The narrow region 250 of the stability of the stripe phase in the phase diagram 251 for decreasing misfit is consistent with the corresponding 252 decrease of the wall crossing energy. In fact, according 253 to the domain-wall arguments of Bak et al. [7], for any 254 negative value of the domain wall crossing, a HoI struc-255 ture would be favoured rather than a SI phase. Our re-256 sults for the phase diagram imply that the commensurate-257 incommensurate transition proceeds first via a transition 258 from the commensurate phase into an SI phase and then 259 from a SI phase into a HoI phase. For small misfit param-260 eters and weak coupling to the substrate, the stripe phase 261 occurs in such a narrow region of the phase diagram that it 262 may not be experimentally observable. It should be kept 263 in mind, however, that our results are at the level of mean
field theory since it does not take into account the effects
of thermal fluctuations, which may eliminate the SI phase.



Fig. 5: In a) the local free energy density is displayed as a function of space for $\varepsilon = 1\%$ very close to the HoI/SI transition for a system of size $111a_f \times 111a_f$. In this figure the dark and light shades correspond to high and low free energy densities, respectively. In b) the normalized local free energy is plotted as function of x along the grey line shown in a).

From the phase diagram of Fig. 3, the intermediate SI 267 phase is more pronounced for large coupling parameter 268 $f^{\rm ad}/K$. For lower couplings, the SI phase exist only in a 269 very narrow region. Experimentally it could appear as a 270 direct transition from commensurate to HoI phase when 271 the temperature and/or the coverage is varied. This be-272 havior is consistent with the existing experimental data 273 for Xe and Kr on Graphite [1]. Since for Xe/Pt(111) the 274 corrugation potential is much larger, corresponding to a 275 large coupling parameter, the transition from commensu-276 rate to HoI phase should clearly involve an intermediate 277 SI phase, as observed experimentally [1,5]. Of course, for 278 this interpretation to hold, we have to identify qualita-279 tively the change of temperature as effectively changing 280 the coupling parameter, and the change of coverage as ef-281 fectively changing the mismatch parameter ϵ . 282

The effective single mode model allows us to study the 283 domain wall pattern in the incommensurate phase at large 284 length scales, but still retaining atomistic resolution. One 285 interesting question that can be addressed with this ap-286 proach is the influence of adsorbate step edges on the ori-287 entation of the domain-wall structure. Recently, it has 288 been observed experimentally by STM imaging of Xe over-289 layers on Graphite [27] that the domain walls orient per-290 pendicularly to Xe island step edges on the surface. We 291 can verify this behavior with the present model. To mimic 292 a step edge on the overlayer, we set the pinning potential 293 to a constant value on one side of the step edge and zero 294 on the other size, and then follow the time development 295 from an initial condition of a uniformly strained film (cor-296 responding to the average strain of the equilibrium state) 297 assuming non-conserved dissipative dynamics [9, 10], given 298 by the equation of motion 299

$$\frac{\partial \eta_j}{\partial t} = -\frac{\partial F_\eta}{\partial \eta_j *}.$$
(7)



Fig. 6: Snapshots of a portion of a domain pattern at times t = 50 in a) and c) and t = 50000 in b) and d) for a strain of $\varepsilon = 7\%$. In a) and b) the colors correspond to the three sublattices and in c) and d) the color is proportional to the energy as in Fig. 5. Inside the central region $f^{\rm ad} = 0$ and $f^{\rm ad} = 0.27 \times 10^{-2}$ outside.

Fig. 6 shows the pattern in the HoI phase for increasing times near a step of a curved edge. The domain wall structure undergoes a reorientation near the step edge such that domain walls are preferentially perpendicular to the step edge, as observed experimentally [27].

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In this work, we have introduced an effective one mode 305 PFC model designed for studying higher order commensu-306 rate states such as the $(\sqrt{3} \times \sqrt{3})R30^\circ$ phase found in sys-307 tems such as Xe/Pt(111), Xe/Graphite, and Kr/Graphite, 308 and the corresponding phase diagrams as the temperature 309 and/or lattice mismatch is varied. Our model allows us 310 to study systems at macroscopic scales where the domain 311 walls in the incommensurate phases can be separated over 312 large distance scale. It also allows us to study the micro-313 scopic details of the domain wall structures both in the 314 honeycomb incommensurate (HoI) phase and the stripe 315 incommensurate phase (SI) phases. In particular, we are 316 able to evaluate the domain wall crossing energy as a func-317 tion of the adsorption potential strength and lattice mis-318 match. New results such as the general persistence of a 319 SI phase in between the commensurate phase and the HoI 320 phase is found. Our results are qualitatively in agree-321 ment with the experimental observation where both the 322 HoI and SI phases have been found in the Xe/Pt(111)323 system whereas only the HoI phase have been observed 324 for the Xe/Graphite and Kr/Graphite systems. We have 325 also found that near a step edge of the adsorbate, the do-326 main walls tend to align perpendicular to the step edge, in 327 agreement with recent experimental observation [27]. In 328

the future, it would be interesting to follow up the present 329 study with inclusion of thermal fluctuation effects. We 330 have demonstrated before [30] that thermal fluctuations 331 can be effectively incorporated in the PFC model by treat-332 ing the PFC free energy functional as an effective Hamil-333 tonian. It is known that thermal fluctuations can lead 334 to generation of dislocations in the domain wall structure, 335 possibly resulting in a fluid phase in between the commen-336 surate and incommensurate phases [29]. 337

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