Periodic Ground States in the Neutral Falicov-Kimball Model in Two Dimensions

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Abstract

We consider the Falicov-Kimball model in two dimensions in the neutral case, i.e., the number of mobile electrons is equal to the number of ions. For rational densities between 1/3 and 2/5 we prove that the ground state is periodic if the strength of the attraction between the ions and electrons is large enough. The periodic ground state is given by taking the one dimensional periodic ground state found by Lemberger and then extending it into two dimensions in such a way that the configuration is constant along lines at a 45 degree angle to the lattice directions.

Keywords: Falicov-Kimball model, ground state, periodic.
In the Falicov-Kimball model there are two types of particles. The electrons hop between nearest neighbor sites of the lattice while the ions do not hop at all. The electrons do not interact with each other except for the fact that they obey Fermi statistics. At most one ion is allowed at each lattice site and there is an on-site attraction between the ions and electrons. The Hamiltonian is

\[ H = \sum_{<x,y>} c_x^\dagger c_y - 4U \sum_x c_x^\dagger c_x V_x \]  \hspace{1cm} (1)

where \( c_x^\dagger \) and \( c_x \) are creation and annihilation operators for the electrons. \( V_x \) is the occupation number for the ions, i.e., \( V_x = 1 \) if there is a ion at \( x \) and \( V_x = 0 \) if there is not. The sum over \( <x,y> \) is over nearest neighbor bonds in the lattice. (The factor of 4 in front of the \( U \) is included for latter convenience.) The interaction of the electrons with the ions and the kinetic energy and fermi statistics of the electrons lead to an effective interaction for the ions. We will study the ground states of this model on the square lattice in the neutral case (the numbers of ions and electrons are equal) when the parameter \( U \) is large and positive. A review of rigorous work on the Falicov-Kimball model may be found in [4].

In any number of dimensions the ground state for density 1/2 is the checkerboard configuration for all \( U > 0 \) [1, 2]. In two dimensions with large \( U \) the ground states for densities 1/3, 1/4 and 1/5 are known rigorously and are periodic [3, 7]. Numerical results indicate periodic ground states for numerous other values of the density [13]. In one dimension Lemberger [10] showed that for any rational density, the ground state is the periodic arrangement of the ions which is “most homogeneous.” (He gives an explicit algorithm for determining the most homogeneous configuration.) His proof applies when \( U > U_0 \) where \( U_0 \) depends on the denominator of the rational density.

The above results suggest that in two dimensions in the neutral case with large \( U \) the ground state is always periodic. However, Kennedy [8] proved that for densities between 1/5 and 1/4 other than 2/9 the ground state is not periodic. In this density range there is phase separation. The ground states for densities 1/5, 2/9 and 1/4 are periodic, but in between the ground state is formed by adjoining regions with these periodic configurations so as to produce the desired density. Haller [6] proved that the same phenomenon occurs for densities between 1/6 and 2/11. So an analog of Lemberger’s result cannot hold in two dimensions for all densities. In this paper we prove that an analog of his result does hold for rational densities between 1/3 and 2/5. In this interval of densities the ground state is periodic. It is constant along lines of slope ±1 (one choice for the entire ground state), and its restriction to any horizontal or vertical line is the same as the one dimensional ground state found by Lemberger. We expect that this result is true for the entire density interval 1/3 to 1/2, but for technical reasons we cannot prove it for densities between 2/5 and 1/2.

There is a physical argument that suggests that the neutral model with rational density wants to spread out the ions as much as possible. Each electron spends most of its time at
a site with an ion since the attraction between electrons and ions is large. Now consider
the effect of the kinetic energy of an electron. If nearby sites have ions, then the electrons
at those sites will restrict the movement of the electron we are considering. So its kinetic
energy is minimized by arranging the ions so as to maximize the space that each electron
has to move in. However, Watson [14] emphasized that there will typically be a mismatch
between the lattice and the ion configuration that maximizes this space in the absence
of a lattice. Thus the lattice structure can frustrate the exclusion principle’s attempts to
put the ions in the “most homogeneous” configuration. In one dimension this frustration
does not occur. In two dimensions Kennedy’s and Haller’s results on phase separation
show it does occur for some intervals of density, while this paper shows it does not for
other intervals. There does not appear to be any easy way to predict which intervals of
density will have periodic ground states and which ones will have phase separation.

The results for the neutral model on the square lattice with large $U$ are summarized
in figure 1. The obvious open problem is to determine whether there is a periodic ground
state or phase separation in these intervals where the behavior is unknown. However, the
more important problem is to understand when there is phase separation vs. periodicity
without the extensive calculations required in this paper or the papers on phase separation.

The results on periodicity and phase separation in the square lattice depend on ex-
tensive calculations of an effective Hamiltonian and thus depend heavily on the lattice
structure. One possible way to gain some insight into the general periodic vs. phase
separation question is to investigate other lattices. There are rigorous results on periodic
ground states on the triangular lattice for some densities [5]. In three dimensions, nothing
is known with the exception of density 1/2.

The preceding discussion and indeed all of this paper is only concerned with ground
states. For density 1/2 and any $U > 0$ it is known that there are at least two Gibbs states
if the temperature is sufficiently low [1, 9] and the number of dimensions is at least two.
For densities 1/3, 1/4 and 1/5, a “quantum” Pirogov-Sinai theory must be used to prove
that for large $U$ there are multiple Gibbs states at low temperature $[2, 11, 12, 13]$. The key ingredient in these approaches is a Peierls estimate. One can prove such an estimate for densities $1/6, 2/11$ and $2/9$ and so the methods should apply to these densities as well. For densities in $(1/3, 2/5)$ we do not know how to prove the needed Peierls estimate, and so the existence of multiple Gibbs states at low temperatures is an open problem for these densities.

We now turn to our main result. The precise statement is as follows.

**Theorem 1** Suppose $\rho = p/q \in [1/3, 2/5]$ where $p$ and $q$ are relatively prime. We assume the lattice $\Lambda$ is $L \times L$ where $L$ is a multiple of $4q$, and we impose periodic boundary conditions. Then there exists $U(q) > 0$ such that for all $U \geq U(q)$, the ground states are translations, reflections, and rotations of the configuration $S$ defined by the following properties: $S$ is constant along lines of slope $+1$, and the restriction of $S$ to any vertical or horizontal line is the same as the one dimensional large $U$ ground state with density $\rho$.

First we give an overview of the proof. When $U$ is large one may do perturbation theory in $1/U$. We start by considering $3 \times 3$ blocks of sites and consider the perturbation series for the energy up to fourth order. Kennedy has shown that at this order, each block which looks like one of those shown in figure 3 contributes the same amount of energy, while any other block contributes a higher amount of energy. So if a configuration exists in which every $3 \times 3$ block looks like one of those from figure 3, it minimizes $H$ through fourth order. It turns out that such configurations do exist. In fact, Watson [14] has shown that they correspond to tilings by squares and parallelograms as shown in figure 4. In such a tiling the configuration must be constant on lines of slope $\pm 1$, where the sign for the entire configuration is determined by the slope of the short sides of one of the parallelograms.

Up to rotations and reflections, the ions in such a tiling must be one of the three types shown in figure 4. We may write the energy through eighth order in terms of the number of each vertex type. Through sixth order we find that every tiling has the same energy. At eighth order, tilings with only type B and C vertices have the same energy, but type A vertices increase the energy. So if there exists configurations that correspond to square-parallelogram tilings with no type A vertices, then they minimize $H$ through eighth order. Such configurations do exist for densities in $[1/3, 2/5]$.

When we include the higher order terms, the most we can say at this point is that the ground state consists primarily of large regions corresponding to a square-parallelogram tiling with no type A vertices. Because of the lack of type A vertices, these regions must contain parallelograms. Furthermore, the slopes of the short sides of these parallelograms must be the same throughout each individual region. This means that within each region, the configuration must be constant on lines of slope $+1$ or $-1$, but not both, as it would in a region consisting entirely of type A vertices.
On each of these “+” and “−” regions, the problem essentially becomes one-dimensional, since we are now only concerned with the spacing between these “stripes” of occupied sites (see figure 2). It is tempting to try to apply a procedure to each region, similar to that of Lemberger for the one-dimensional case, in order to determine the cheapest stripe arrangement. However, there are two difficulties that arise. First there is the problem of the shape of these regions. Their boundaries may be very irregular whereas an extension of Lemberger’s method would require applying periodic boundary conditions to a fairly regular shape such as a square or rectangle. Second there is a density constraint. The density inside the + and − regions need not be the same as the overall density. This is a problem because how large $U$ needs to be in Lemberger’s argument depends on the density. Even if we could say something about the configuration in each + and − region, this would not take into account the ions lying outside these regions, so we effectively lose track of the original ion density. We must therefore find more regular regions to work with that allow us to keep track of the original density.

The first step in overcoming these difficulties is to partition the lattice into diamonds containing 8 sites, as shown in figure 3. If one of these diamonds intersects a $3 \times 3$ block not shown in figure 3 or it contains a type A ion, it may be associated with an increase in energy at eighth order. We call such diamonds “bad.” The eighth order terms actually occur with $U^{-7}$, so the total energy cost of the bad diamonds is at least $O(U^{-7})N$, where $N$ is the number of bad diamonds. Each of the remaining “good” diamonds is contained in one of the + or − regions mentioned above. In a + region (− region), we group the diamonds into $-45^\circ$ strips ($+45^\circ$ strips) in such a way that the strips are bounded on each of the short sides by a bad diamond. This is shown in figure 4.

If we were to apply periodic boundary conditions to a particular strip and compare the resulting energy with that of the energy of the strip in the original configuration, it would
seem that this difference would be proportional to the size of the boundary of the strip. However, consider one of the 45° strips of diamonds. Inside the strip, the configuration is constant on lines of slope −1. These lines of constant configuration persist through the long sides of the strip, and continue in S until they encounter a bad diamond. So if there are no bad diamonds near the long sides, then imposing periodic boundary conditions on the strip is effectively the same as the boundary conditions the strip had to begin with. This allows us to show that the total difference between the energies of the strips in the original configuration and their energies with periodic boundary conditions is $O(U^{-9})N$.

We then glue all the strips of good diamonds together with the bad diamonds end to end to form one long thin strip. We apply periodic boundary conditions to the configuration on this strip. The difference between the energy of this long strip with periodic boundary conditions and the energy of the original configuration is $O(U^{-9})N$. Each bad diamond costs an energy $O(U^{-7})$, so the energy of the original configuration is at least as large as that of our long strip. Finally, in the single long strip we can apply Lemberger’s argument to conclude that the best arrangement is the most homogeneous one.

**Proof of Theorem 1**: We denote an ion configuration by $S$. We let $H(S)$ be the minimum energy of $H$ with this ion configuration, i.e., we minimize the Hamiltonian with respect to the electronic wave function. If $U$ is sufficiently large, then the function $H(S)$ may be expanded in powers of $U^{-1}$.

$$H(S) = \sum_{m=2}^{\infty} U^{1-m} h_m(S)$$

(Details of this perturbation theory may be found in [3, 10].) Because we are on a square lattice, only the even values of $m$ have a nonzero $h_m$. $h_m(S)$ is a local function. It is a sum of terms, each of whose support is contained in the support of a closed nearest neighbor walk with $m$ steps. We will let $H_m(S)$ denote $U^{1-m} h_m(S)$.

Let $M$ be the number of $3 \times 3$ blocks in a configuration $S$ which are not a reflection or rotation of one of those shown in figure 3. Kennedy [7] showed that

$$\alpha MU^{-3} \leq H_2(S) + H_4(S) + U^{-3}L^2(a + b \rho) \leq \gamma MU^{-1}$$

where $\alpha > 0$, $\gamma > 0$, $a$, and $b$ are constants. So after adding the constant $U^{-3}L^2(a + b \rho)$ to the Hamiltonian, we see that every $3 \times 3$ block which is not shown in figure 3 contributes a positive amount of energy at fourth order (or lower), and the configurations in which every $3 \times 3$ block is one of those in figure 3 have zero energy through fourth order. The following lemma gives us a geometric interpretation of a configuration in which every $3 \times 3$ block is one of those shown in figure 3.

**Lemma 1 (Watson [14])** Let $S$ be a configuration in which every $3 \times 3$ block is one of those shown in figure 3. Then there exists a tiling by squares and parallelograms as shown in figure 4, in which every vertex corresponds to an ion in $S$. 

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Proof: Let S be a configuration satisfying the hypothesis of the lemma. Consider any occupied site. It is the center of a type 1, 2, or 3 block from figure 3. Now consider a 3 by 3 block which shares six sites with the original 3 by 3 block we considered. Since the configuration at six of its sites is known and it must be one of the blocks in figure 3, there are only a few possibilities for the configuration on the three unknown sites. By continuing in this manner and keeping track of all the cases one may determine the allowable local ion arrangements. These turn out to be rotations and reflections of the three arrangements in figure 4.

Figure 3: Configurations which minimize the energy locally.

![Figure 3](image)

From the lemma and equation (3), we see that the configurations which minimize the energy through fourth order correspond to tilings by squares and parallelograms. However, the densities one can obtain from such tilings is restricted. A configuration in which every ion is type C has density 1/3. One in which every ion is type A has density 1/2. We may construct a configuration having any rational density between 1/3 and 1/2 by placing an interface of slope ±1 between a region of type A ions and a region of type C ions, the size of each region being chosen to give the desired density. The ions on the interface will then be type B. So we see that from these square-parallelogram tilings, one may obtain any rational density in [1/3, 1/2]. (Of course, there are many other ways to do so besides the construction we have given.)

![Figure 4](image)
Consider a square-diamond tiling and let \( n_A, n_B, n_C \) be the number of type \( A, B, C \) ions, respectively. These three integers do not determine the configuration, but they do determine enough of the local structure of the configuration that the energy \( H(S) \) may be computed through eighth order from \( n_A, n_B \) and \( n_C \). We show in the appendix that for \( m = 6 \) and \( 8 \)

\[
H_m(S) = U^{1-m}(n_A e_m^A + n_B e_m^B + n_C e_m^C)
\]

where

\[
egin{align*}
e_6^A &= -3200, & e_8^A &= 98000, & e_6^B &= -1520, \\
e_8^B &= 33600, & e_6^C &= 160, & e_8^C &= -15120.
\end{align*}
\]

In a square diamond tiling, the numbers \( n_A, n_B, \) and \( n_C \) are not independent. Since the area is \( L^2 \), we have

\[
2n_A + \frac{5}{2}n_B + 3n_C = L^2
\]

We are also keeping the number of ions fixed at \( \rho L^2 \), so

\[
n_A + n_B + n_C = \rho L^2.
\]

Using eqs. (4) and (8) to solve for \( n_B \) and \( n_C \) in terms of \( n_A \), we have

\[
H_6(S) + H_8(S) - L^2 f(U, \rho) = 15680U^{-7}n_A.
\]

\( f(U, \rho) \) is a function of \( U \) and \( \rho \), but there is no need to write it out explicitly.

The preceding paragraph assumed that \( S \) was a tiling. In general, this will not be the case. Suppose \( S \) is not a tiling and \( M \) is the number of \( 3 \times 3 \) blocks which are not rotations or reflections of those in figure 3. Then the number of sites in the region where \( S \) is not a tiling is at most \( 9M \). So the terms intersecting this region contribute at most \( O(U^{-5})M \) to \( H_6 + H_8 \). Equations (7) and (8) will no longer hold, but the difference between the right and left hand sides will be bounded by a constant times \( M \). Thus for a general \( S \), we have

\[
H_6(S) + H_8(S) - L^2 f(U, \rho) = 15680U^{-7}n_A + O(U^{-5})M.
\]

We may now determine the ground states through order 8. From inequality (3) we see that any configuration with \( M = 0 \) will minimize the energy through fourth order. In equation (9), \( M \) appears at order 6. So for large enough \( U \) we want \( M = 0 \). Since the coefficient of \( n_A \) is positive in equation (9), we also want \( n_A = 0 \). So the configurations which minimize the energy through order 8 are those where every ion is type \( B \) or \( C \). These configurations correspond to square-parallelogram tilings with no adjacent \( \pm 45^\circ \) strips of squares. They have zero energy through order 8 after we have subtracted \( L^2 f(U, \rho) \). All we have shown so far is that zero is a lower bound for the energy through eighth order for densities in \([1/3, 2/5]\). We can construct a configuration that attains this lower bound as follows. A region with only type \( B \) vertices has density \( 2/5 \), while a region with only type
C has density $1/3$. By adjoining a type B region with a type C region with an interface between them at $45^\circ$ we can obtain any density in $[2/5, 1/3]$ and attain our lower bound.

If we just consider the energy through eighth order, then any configuration that has a “bad” $3 \times 3$ block or a type A ion will have energy higher than the ground state energy. The crucial point is that there exists a positive constant $c$ so that difference of the energy and the ground state energy is at least $cU^{-7}$ times the number of these “local defects” in the configuration. We now move on to the higher order terms.

Recall that $\rho = p/q \in [1/3, 2/5]$ where $p$ and $q$ are relatively prime, and the lattice $\Lambda$ is $L \times L$ where $L$ is a multiple of $4q$. We partition $\Lambda$ into diamonds containing 8 sites as shown in figure 5.

![Figure 5: Partitioning of the lattice into diamonds containing 8 sites.](image)

We define a diamond $b$ to be bad if it contains a type A ion or intersects a bad $3 \times 3$ block. Otherwise, $b$ is good. We let $N$ denote the number of bad diamonds. A good diamond is one which is contained in a region of the configuration corresponding to a square-parallelogram tiling with no type A ions. We know from the above that the energy through eighth order is bounded below by a term proportional to the number of bad $3 \times 3$ blocks plus the number of type A ions, which is proportional to $N$. So inequality (3) and equation (9) tell us that

$$H_2 + H_4 + H_6 + H_8 + U^{-3}L^2(a + b\rho) - L^2f(U, \rho) \geq cNU^{-7}$$

where $c > 0$ is a constant. For future convenience, we define

$$\tilde{H} = H + U^{-3}L^2(a + b\rho) - L^2f(U, \rho)$$
Let $S$ be a configuration with density $\rho(S) \in [\frac{1}{3}, \frac{2}{5}]$. We label a parallelogram in $S$ with a $+$ if its short sides have slope $+1$, and a $-$ if its short sides have slope $-1$. Now, every good diamond must contain a type B or C ion, both of which are the vertex of a parallelogram. Any other parallelogram intersecting this diamond must have the same sign. Thus the good diamonds may be labelled $+$ or $-$ in a natural way. Furthermore, since all the ions in a good diamond are type B or C, two diamonds that share an edge must have the same sign. One may think of $S$ as being divided into regions where the diamonds are $+$, regions where the diamonds are $-$, and regions of bad diamonds. A $+$ region and a $-$ region must be separated by a region of bad diamonds. Inside every $+$ region, $S$ is constant on lines of slope $+1$, and inside every $-$ region it is constant on lines of slope $-1$. We call this the stripe property. More, however, may be said about the arrangement of the occupied stripes inside each good region. Since every ion must be type B or C, each pair of occupied stripes must be separated by one or two unoccupied stripes. Also, since there are no type A ions, we may not have consecutive stripes corresponding to the following occupation sequence: occupied, unoccupied, occupied, unoccupied, occupied. This rules out the checkerboard configuration, the sign of which is ambiguous. When a region has the stripe property with no type A ions, we say that the stripes are correctly spaced.

We now divide up the $-$ regions into $45^\circ$ strips as follows. Choose any good diamond whose sign is $-$. As shown in figure 6, we extend this set in the directions $\pm (1, 1)$ until we hit a bad diamond. Every diamond in this $45^\circ$ strip is good, and, since one of them is $-$, they must all be $-$. We may then carry our labelling one step further and label this strip with a $-$. Note that every good $-$ diamond is contained in exactly one such $-$ strip.

![good "-" diamond](image)

Figure 6: A good "$-$" diamond is chosen. The sites in the shaded region are contained in good "$-$" diamonds.

We divide up the $+$ region into $-45^\circ$ strips by changing all the signs in this construction. We label these strips with a $+$. See figure 7.
Figure 7: Left: a − strip. Right: a + strip. The configuration inside the − strip (+ strip) is constant along lines of slope −1 (+1).

We now take the strips and rotate them so they are all horizontal and then glue them together to form one long strip. Now rotate the bad diamonds and glue them onto one end to form an even longer strip. This strip will contain all the sites of the original Λ. This is shown in figure 8. The strips have the stripe property. After rotation this means that in figure 8 the portion of the configuration that came from the strips is constant along vertical lines of lattice sites. In figure 8, vertical lines of lattice sites only contain two sites, so this simply means that either both of these sites are occupied or both are empty. In the portion of figure 8 which contains the bad diamonds, the configuration may not satisfy this property. However, in the statement of the theorem L was assumed to be a multiple of 4q. Thus the total number of ions is a multiple of 4p. There are an even number of ions in each + and − strip, so there must be an even number of ions in the bad region of our thin strip. Therefore we may rearrange the configuration in this portion of the strip so it also has either zero or two ions in each vertical line of lattice sites. We now glue the short sides together and the long sides together to form a torus, call it T. Let S1 denote the resulting configuration on T. We want to bound $|\tilde{H}(S) - \tilde{H}(S_1)|$. As we shall see, we will only need to estimate this difference at orders $\geq 10$.

| strips          | bad diamonds |
|-----------------|---------------|

Figure 8: The thin strip obtained by gluing together the + and − strips as well as the bad diamonds.

The perturbation theory leads naturally to an expression for $H$ as a sum over nearest neighbor closed walks $\gamma$.

$$H = \sum_\gamma w(\gamma)$$
Lemberger gave a nice formula for the contribution of a walk:

\[ w(\gamma) = \frac{(-1)^{m(S, \gamma)}}{|\gamma|} (2U)^{1-|\gamma|} \left( \frac{|\gamma| - 2}{m(S, \gamma) - 1} \right) \]  

(11)

where \(|\gamma|\) is the number of steps in \(\gamma\) and \(m(S, \gamma)\) is the number of ions at sites in the walk \(\gamma\). Obviously, \(H_n\) is equal to the sum over walks with \(n\) steps.

We want to set up a correspondence between the terms in the perturbation series of \(\tilde{H}(S)\) and \(\tilde{H}(S_1)\). So we need to set up a correspondence between walks on the torus \(T\) and the original square \(\Lambda\). We think of \(Z^2\) as the covering space of the torus \(T\). Then a walk \(\gamma_1\) on \(T\) has a natural lift to a walk \(\hat{\gamma}_1\) on \(Z^2\). Each site in \(T\) comes from a site in \(\Lambda\). We let \(\gamma\) be the translate of \(\hat{\gamma}_1\) such that \(\gamma(0)\) is the site in \(\Lambda\) that corresponds to the site \(\gamma_1(0)\) in \(T\). The map \(\gamma_1 \rightarrow \gamma\) takes walks on \(T\) to walks on \(\Lambda\).

There is a problem with this correspondence. The lifted walk \(\hat{\gamma}_1\) need not return to its starting point. In fact the lifted walk will return to its starting point if and only if the original walk \(\gamma_1\) is homotopic to zero. So we redefine \(\tilde{H}(S_1)\) and \(\tilde{H}(S)\) to be the sum of all the terms in the perturbation series that come from walks homotopic to zero. If a walk on \(\Lambda\) is not homotopic to zero, then it has at least \(L\) steps. \((L\) is the length of the square \(\Lambda\).) Thus this changes \(\tilde{H}(S)\) by terms of order \(U^{-L}\). So the total change in \(\tilde{H}(S)\) is of order \(L^2U^{-L}\). This is tiny even compared to the energy cost of a single bad diamond. The change in \(\tilde{H}(S_1)\) is not so small since the torus \(T\) is quite narrow in one direction. But we are free to define \(\tilde{H}(S_1)\) any way we wish. It merely serves to organize our proof.

Now suppose that \(\gamma_1 \rightarrow \gamma\) and \(\gamma\) does not encounter a bad diamond in \(S\). Then \(\gamma\) lies entirely in either a region of + diamonds or a region of − diamonds. Consider the first case. So the strips have slope −1 and the configuration is constant along lines with slope +1 in the vicinity of the walk \(\gamma\). It follows that \(\gamma\) sees the same configuration in \(S\) that \(\gamma_1\) sees in \(S_1\), and so \(\omega(\gamma) = \omega(\gamma_1)\). Thus the terms in \(\tilde{H}_n(S) - \tilde{H}_n(S_1)\) that do not cancel come from walks \(\gamma\) in \(S\) that do not encounter a bad diamond. So the number of terms that do not cancel is of order \(n^2N\). Hence we have

\[ |\tilde{H}_n(S) - \tilde{H}_n(S_1)| \leq U^{1-n}c^n n^2N \]  

(12)

Note that although \(S_1\) satisfies the stripe property, its stripes may not be correctly spaced. The idea now is to form a finite sequence of configurations, \(\{S_i\}_{i=1}^K\), such that

\[ |\tilde{H}_n(S_{i+1}) - \tilde{H}_n(S_i)| \leq U^{1-n}c^n, \]

\(K = O(N)\), and the stripes in the final configuration \(S_K\) are correctly spaced. The strategy is as follows. The reason why the stripes in \(S_1\) may not be correctly spaced is because of the introduction of the bad diamonds, as well as the cuts. There are \(O(N)\) stripes in the bad region of \(S_1\) and \(O(N)\) cuts, so we may hope to correct the spacings with \(O(N)\) modifications, each modification costing at most \(U^{1-n}c^n\) at order \(n\).
In making these modifications we must be sure that each one fixes a “fault” in $S_i$ without creating a similar fault. All the modifications follow the same basic procedure. We illustrate this by considering a pair of adjacent occupied stripes in $S_1$. Since the density of $S_1$ is $\leq 2/5$, if there is a pair of adjacent occupied stripes, there must be a pair of adjacent unoccupied stripes. As shown in figure 9, we introduce two “markers,” one between the pair of adjacent occupied stripes and one between the pair of adjacent unoccupied stripes. We let $W$ be the portion of the configuration between the markers. With the two axes of rotation $a$ and $b$ as shown in figure 9, we form $W^{-1}$ by rotating $W$ about the axis $a$. We then replace $W$ by $W^{-1}$ and, if necessary, rotate $W^{-1}$ about the axis $b$ so it fits into place. (Whether or not we need to rotate about $b$ depends on the number of sites in $W$.) $S_2$ is then the modified configuration. Any walk which gives a non zero contribution to $\tilde{H}_n(S_2) - \tilde{H}_n(S_1)$ must cross one of the markers, and so must visit one of the 8 sites adjacent to the markers. So we have that $|\tilde{H}_n(S_2) - \tilde{H}_n(S_1)| \leq U^{1-n} c^n$.

Our modification has allowed us to separate a neighboring pair of occupied stripes without creating a new pair. So we see that we can get rid of all such pairs with $O(N)$ modifications.

The next problem we focus on is the possibility of more than two consecutive unoccu-
pied stripes. If this occurs, since the density is $\geq 1/3$ there must be at least one pair of occupied stripes separated by a single unoccupied stripe. Figure 10 shows how we may reduce the number of adjacent unoccupied stripes by one by a procedure similar to the above.

![Figure 10: Reducing the number of consecutive unoccupied stripes.](image)

After $O(N)$ repetitions of the above procedures we reach a configuration in which every pair of occupied stripes is separated by one or two unoccupied stripes. We now face the possibility that our configuration contains type A ions. These ions could have come from $S_1$ or they could have been formed by the modifications above. But we only made $O(N)$ modifications, and $S_1$ contained at most $O(N)$ type A ions. So our configuration can contain at most $O(N)$ type A ions. We let a “1” denote an occupied stripe and a “0” denote an unoccupied stripe. The stripe sequence signifying a type A ion is then: 1, 0, 1, 0, 1. If this occurs, then since the density is $\leq 2/5$ there must exist the following stripe sequence: 1, 0, 0, 1, 0, 0, 1. In figure 11 we see how the procedure used above can get rid of these type A ions. At first glance, it may seem that this last modification, although removing a type A ion at the left end of $W$ in $S_i$, may have created a type A ion at the right end of $W^{-1}$ in $S_{i+1}$. However, if we do get a type A ion in $S_{i+1}$ after the inversion of $W$, it must have been there before the inversion, in $S_i$. Thus the number of type A ions is indeed reduced.

The last configuration in this sequence of modifications, $S_K$, has correctly spaced stripes. Furthermore, $K = O(N)$ and $|\tilde{H}_n(S_{i+1}) - \tilde{H}_n(S_i)| \leq U^{1-n}c^n$ for each consecutive
Figure 11: Removing the type A ions.

pair in our sequence. This gives us the following telescoping sum

\[ \tilde{H}(S) - \tilde{H}(S_K) = [\tilde{H}(S) - \tilde{H}(S_1)] + \sum_{i=1}^{K-1} [\tilde{H}(S_i) - \tilde{H}(S_{i+1})]. \] (13)

The construction of our sequence above and inequality (12) imply that

\[ \sum_{\text{even } n \geq 10} |\tilde{H}_n(S) - \tilde{H}_n(S_K)| \leq K \sum_{\text{even } n \geq 10} U^{1-n} c^nn^2 \leq cNU^{-9} \] (14)

In the last inequality we used the fact that \( K \) is of order \( N \). Since \( S_K \) satisfies the stripe property and its stripes are correctly spaced, we know that

\[ \sum_{\text{even } n = 2}^{8} \tilde{H}_n(S_K) = 0 \]

Equation (13) and the bounds (10) and (14) imply

\[ \tilde{H}(S) \geq \tilde{H}(S_K) + \alpha NU^{-7} - cNU^{-9}. \] (15)

Consider the torus \( T \) on which \( S_K \) is defined. Let \( S_L \) be a configuration on \( T \) which satisfies the stripe property and such that the restriction of the configuration to a line in one of the lattice directions is the one-dimensional ground state for density \( \rho(S) \) when \( U \) is
large as determined by Lemberger. \((S_L \text{ is unique up to lattice symmetries.})\) We claim that among all the configuration on \(T\) having density \(\rho(S)\) and satisfying the stripe property, the energy is minimized by \(S_L\). In particular, the claim implies \(H(S_K) \geq H(S_L)\).

To establish the claim we show that there is a close relationship between the perturbation series for the two-dimensional model and that of the one-dimensional model. The terms in the Hamiltonian at order \(2m\) may be written in terms of the walks as

\[
H_{2m} = \sum_{\gamma: |\gamma|=2m} w(\gamma)
\]

with \(w(\gamma)\) given by eq. (11). For the one-dimensional model the only change that needs to be made in this formula is that \(\gamma\) is only summed over walks in the one-dimensional lattice. We will show that if the configuration has the stripe property then

\[
H_{2m} = c(m) \sum_{\gamma: |\gamma|=2m, \gamma \text{ hor}} w(\gamma)
\]

where \(\gamma \text{ hor}\) means that the sum is over walks which only take steps in the horizontal direction. So these walks are one-dimensional although they may start at any point in the two-dimensional lattice. The combinatorial factor \(c(m)\) is given by

\[
c(m) = \sum_{k=1}^{m} \left( \begin{array}{c} m \\ k \end{array} \right)^2
\]

To show (17) we think of a walk \(\omega\) as a sequence of \(2m\) letters chosen from L (left), R (right), U (up) and D (down). The number of L’s must equal the number of R’s, and the number of U’s must equal the number of D’s. (This is because of our redefinition of the Hamiltonian so that the perturbation series only contains terms coming from walks that are homotopic to zero.) Let \(P\omega\) be the projection of the walk that is obtained by replacing each U by an R and each D by an L. (This assumes that we are in the case where the slopes are \(-1\). For slope +1, we replace U by L and D by R.) Then for configurations that satisfy the stripe property, \(H(\omega) = H(P\omega)\). Now let \(\omega'\) be a walk with only L and R steps. Since the walk is closed there are \(m\) R’s and \(m\) L’s. We obtain all walks \(\omega\) with \(P\omega = \omega'\) by replacing some number of L’s with D’s and the same number of R’s with U’s. If \(k\) is the number of L’s that we replace, then there are \(\binom{m}{k}\) ways to make the L \(\rightarrow\) D replacements and an equal number of ways to make the R \(\rightarrow\) U replacements. So the number of \(\omega\) with \(P\omega = \omega'\) is \(c(m)\) as defined above. Equation (17) says that for stripe configurations the two-dimensional Hamiltonian is the same as the one-dimensional Hamiltonian order by order except for a multiplicative factor at each order. It is straightforward to check that Lemberger’s argument applies unchanged when the \(c(m)\) factors are included.

The stripe arrangement in \(S_L\) follows the Farey tree, just as the ion arrangement in the 1-d case. So \(S_L\) consists of a unit cell of length \(q\), which is periodically extended to
cover the torus. Our original square lattice $\Lambda$ has sides whose lengths are multiples of $4q$. So we may build a configuration $S^\Lambda_L$ on $\Lambda$ which has the same stripe arrangement as $S_L$, and satisfies $H(S^\Lambda_L) = H(S_L)$. We may then rewrite inequality (15) as

$$H(S) \geq H(S^\Lambda_L) + \alpha NU^{-7} - cNU^{-9}. \quad (19)$$

Assuming $U \geq \sqrt{c/\alpha}$, this shows that $S^\Lambda_L$ is a ground state. To see that these are the only ground states, we consider two cases. If $N > 0$ then the above bound says $H(S) > H(S^\Lambda_L)$. If $N = 0$, then $S$ satisfies the stripe property with the stripes correctly spaced. So we know from Lemberger’s argument that $H(S) \geq H(S^\Lambda_L)$, with equality holding only when $S = S^\Lambda_L$ up to a lattice symmetry. $\blacksquare$
Appendix

In this appendix we derive eq. (4) which expresses the sixth and eighth order terms in the Hamiltonian in terms of the number of ions of types A, B and C. Recall that the Hamiltonian at a given order is given in terms of walks by

\[ H_{2m} = \sum_{\gamma: |\gamma|=2m} w(\gamma) \]  

(20)

This may be rewritten as

\[ H_{2m} = \sum_X c_{m,X} S(X) \]  

(21)

where the sum is over subsets of the lattice and \( S(X) \) is 1 if the configuration \( S \) has an ion at every site in \( X \) and 0 otherwise.

Recall that \( I(S) \) denotes the sets of sites at which there is an ion for the configuration \( S \). If \( S(X) \) is to give a non zero contribution to \( H_6(S) + H_8(S) \), \( X \) must be a subset of \( I(S) \) which is contained in a closed nearest neighbor walk of six or eight steps. Let \( S \) correspond to a tiling by squares and parallelograms. Then the only subsets \( X \subset I(S) \) which can contribute at orders 6 and 8 are those shown, up to rotations and reflections, in figure 12. We will refer to an ion and the eight ions arranged about it as in figure 12.

Figure 12: All possible subsets of \( I(S) \) (up to lattice symmetries) which contribute to the energy at orders 6 and 8 when \( S \) corresponds to a square-parallelogram tiling. The lines serve to distinguish the subsets.
as the “local arrangement” of the ion. Recall that the local arrangement about an ion in $I(S)$ must look like one of the three shown in figure 4. So for any $X$ which gives a nonzero contribution to $H_6(S) + H_8(S)$, there is at least one ion $x$ such that $X$ is a subset of the local arrangement about $x$. There may be more than one such $x$. For example, if $X$ consists of a single ion, $X$ will be contained in 9 local arrangements. To compensate for this over counting, we define $\tilde{c}_{m,X} = \frac{c_{m,X}}{\kappa(X)}$ where $\kappa(X)$ is the number of ions whose local arrangement contains $X$. Now fix a type A ion and let $L$ denote the local arrangement about it. Then for even $m \leq 8$ we have

$$e^A_m = \sum_{X \subseteq L} \tilde{c}_{m,X}$$

This may be written as

$$e^A_m = \sum_{X \in \text{figure 12}} \tau_{A,X} \tilde{c}_{m,X}$$

where “$X \in \text{figure 12}” means that $X$ is a rotation or reflection of one of the arrangements from figure 12, and $\tau_{A,X}$ is the number of translates of $X$ appearing in the local configuration about a type A ion. $e^B_m$ and $e^C_m$ are given by the same formula with $L$ taken to be the local arrangement of a type B or C ion respectively.

Table 1 gives the coefficients $c_{6,X}$ and $c_{8,X}$ for each $X$ in figure 12. The last three columns give the numbers $\frac{\tau_{A,X}}{\kappa(X)}$, $\frac{\tau_{B,X}}{\kappa(X)}$, and $\frac{\tau_{C,X}}{\kappa(X)}$. So we see, for example, that $e^A_6 = 64 \times 1 - 384 \times 2 + 96 \times 2 - 674 \times 4 = -3200$.

We should note that one can use the above type of argument to show that an equation of the form (4) must hold. The values of $e^A_m$, $e^B_m$, and $e^C_m$ can then be computed by carrying out the perturbation theory for $H_m(S)$ for several different periodic configurations $S$ and then using the results to set up a linear system of equations for $e^A_m$, $e^B_m$, and $e^C_m$. We have done this as an independent check on the calculation above.

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| $X$ | $c_{6,X}$ | $c_{8,X}$ | $A$ | $B$ | $C$ |
|-----|-----------|-----------|-----|-----|-----|
| 1   | 64        | 112       | 1   | 1   | 1   |
| 2   | -384      | 768       | 2   | 3/2 | 1   |
| 3   | 96        | -1360     | 2   | 1   | 0   |
| 4   | 216       | -768      | 0   | 1   | 2   |
| 5   | 24        | 448       | 0   | 1   | 2   |
| 6   | -672      | 11520     | 4   | 2   | 0   |
| 7   |           | 512       | 4   | 1   | 0   |
| 8   |           | 32        | 2   | 0   | 0   |
| 9   |           | 192       | 2   | 0   | 0   |
| 10  |           | 1152      | 2   | 1   | 1   |
| 11  |           | 768       | 2   | 1   | 1   |
| 12  | -1728     | 0         | 2   | 4   |     |
| 13  | -5184     | 0         | 1   | 2   |     |
| 14  | -384      | 8         | 2   | 0   |     |
| 15  | -1248     | 4         | 1   | 0   |     |
| 16  | 3840      | 4         | 1   | 0   |     |
| 17  | 23040     | 1         | 1/2 | 0   |     |
| 18  | 3840      | 4         | 1   | 0   |     |
| 19  | 960       | 1         | 0   | 0   |     |

Table 1: Values of $c_{m,X}$ used to calculate the energy of types A, B, and C ions at orders 6 and 8.
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