

RENORMALIZATION GROUP AND PHASE TRANSITIONS IN SPIN,
GAUGE, AND QCD LIKE THEORIES

by

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CERTIFICATE OF APPROVAL

PH.D. THESIS

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To the memory of my father, Deliang Liu.

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ABSTRACT

In this thesis, we study several different renormalization group (RG) methods, including the conventional Wilson renormalization group, Monte Carlo renormalization group (MCRG), exact renormalization group (ERG, or sometimes called functional RG), and tensor renormalization group (TRG).

We use the two dimensional nearest neighbor Ising model to introduce many conventional yet important concepts. We then generalize the model to Dyson's hierarchical model (HM), which has rich phase properties depending on the strength of the interaction. The partition function zeros (Fisher zeros) of the HM model in the complex temperature plane is calculated and their connection with the complex RG flows is discussed. The two lattice matching method is used to construct both the complex RG flows and calculate the discrete β functions. The motivation of calculating the discrete β functions for various HM models is to test the matching method and to show how physically relevant fixed points emerge from the complex domain.

We notice that the critical exponents calculated from the HM depend on the blocking parameter b . This motivated us to analyze the connection between the discrete and continuous RG transformation. We demonstrate numerical calculations of the ERG equations. We discuss the relation between Litim and Wilson-Polchinski equation and the effect of the cut-off functions in the ERG calculation.

We then apply methods developed in the spin models to more complicated and more physically relevant lattice gauge theories and lattice quantum chromodynamics (QCD) like theories. Finite size scaling (FSS) technique is used to analyze the Binder cumulant of the $SU(2)$ lattice gauge model. We calculate the critical exponent ν and ω of the model and show that it is in the same universality class as

the three dimensional Ising model.

Motivated by the walking technicolor theory, we study the strongly coupled gauge theories with conformal or near conformal properties. We compare the distribution of Fisher zeros for lattice gauge models with four and twelve light fermion flavors. We also briefly discuss the scaling of the zeros and its connection with the infrared fixed point (IRFP) and the mass anomalous dimension.

Conventional numerical simulations suffer from the critical slowing down at the critical region, which prevents one from simulating large system. In order to reach the continuum limit in the lattice gauge theories, one needs either large volume or clever extrapolations. TRG is a new computational method that may calculate exponentially large system and works well even at the critical region. We formulate the TRG blocking procedure for the two dimensional $O(2)$ (or XY) and $O(3)$ spin models and discuss possible applications and generalizations of the method to other spin and lattice gauge models.

We start the thesis with the introduction and historical background of the RG in general.

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CHAPTER 1

INTRODUCTION

1.1 Introduction

The renormalization group (RG) has played an important role in the development of quantum field theories in a variety of settings. It provides an effective way of analyzing which short-distance feature effects long-distance physics. On the other hand, renormalization group methods have also been developed for statistical mechanical systems in order to deal with systems with many degrees of freedom and of strong correlations. It provides a universal language that can be applied over a broad range of areas, i.e., particle physics, condensed matter physics, atomic physics, etc.

The basic idea of RG was introduced by Stueckelberg and Petermann [4] in 1951 in the form of the group transformation of the S -matrix S

$$\mathbf{P}_{\mathbf{M}\alpha}S = \frac{\partial S}{\partial \log \epsilon_M}, \quad (1.1)$$

where ϵ_M is the coupling constant and $\mathbf{P}_{\mathbf{M}\alpha}$ is the infinitesimal operator. Two years later, in 1953, the same authors [5] (see also [6]) formulated their ideas more explicitly and introduced the function $h(x, \mu, e)$ in quantum electrodynamics (QED), which can be considered as a first version of the β function

$$\mathbf{P}_{\mathbf{i}}S = h_{i_e}(x, \mu, e) \frac{\partial S(x, \mu, e, \dots)}{\partial e}. \quad (1.2)$$

They were able to relate different reparametrizations of QED mass (x, μ) and electric charge (e) via some transformations. They called them “group of normalization”, which is given by infinitesimal operations $\mathbf{P}_{\mathbf{i}}$. Gell-Mann and Low [7] in 1954 obtained the functional equations for QED propagators in the low wavelength/high-energy, or in another word the ultraviolet (UV) limit. They related the physical charge e with the bare charge e_0 via the family of parameters e_λ in different ($\lambda \rightarrow 0$,

$\lambda \rightarrow \infty$) limits. Bogoliubov and Shirkov [8] soon after in 1955 – 1956 gave a clearer picture on the connection between the above two groups work and they developed a simple way to analyze the UV and infrared (IR) asymptotic (see also [9]). Not many RG conceptional ideas on the field theory were developed until the beginning of the 1970s, Politzer [10] and Gross and Wilczek [11] showed that the non-Abelian gauge theories are asymptotically free by showing that the Callan [12] Symanzik [13, 14] β function has an opposite sign comparing to the QED case. This explained why quarks acted like free particles at small distances. The discovery of the asymptotic freedom strengthened the foundations of the quantum field theory (QFT) and solved the “inconsistency” of QED, as argued by Landau [15]. Those were the early development of the RG in particle physics. More references can be found in [16, 17, 18].

Wilson [19, 16, 20] generalized the idea of RG to the theory of the critical phenomena. It provided a general framework to deal with systems with many coupled degrees of freedom, including many interesting systems in the condensed matter physics, statistical physics, and many other different areas. Numerical methods coupled with RG makes the analysis of the properties of the systems very near to the critical points possible.

The Monte Carlo renormalization group (MCRG) method is one of the powerful numerical tools to study critical properties of the spin and gauge systems [21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45]. Applying Monte Carlo methods to the RG study was first suggested by Shang-Keng Ma [46] in the 1970s. The method was extensively used in the 1980s. The two lattice matching method has been proved to be a very useful technique to calculate the non-perturbative β function of gauge and spin models. Recently, Hasenfratz [40] has applied the two lattice matching method to many flavor $SU(3)$

gauge theories and analyzed critical properties of the theory. This motivated us to study more carefully the MCRG methods on both spin and lattice gauge models. Along this line, we carefully analyzed critical properties of Dyson's hierarchical model, discussed the connection between the continuous and discrete RG method, went through the functional RG, the tensor RG and finally the multi-flavor lattice gauge models.

The plan of the thesis is first to apply the two lattice matching method to the hierarchical model, where simple integral relation is exact. We will try to match a series of couplings and draw flow lines in the coupling space. Moreover, we will extend the flow to the complex coupling space and try to support the idea that Fisher Zeros, the partition function zeros in the complex temperature plane, (see section 2.2.4) form a boundary of the flow. We will eventually apply the technique developed to the lattice gauge models and lattice QCD like models. The structure of the thesis is the following: in the remaining of this chapter, I will briefly introduce some basic ideas used in the following chapters, and give a brief summary on what we have done so far. In Chapter 2, I will start with the conventional Ising model and then move on to Dyson's hierarchical Ising model. We will spend a certain amount of time on the hierarchical model. Most of the concepts and techniques used later on will be introduced here. Section 2.2 gives a brief introduction of the hierarchical model, shows the calculation of the critical properties, and discuss the linear analysis; section 2.2.4 summarizes the Fisher zeros calculations done in the hierarchical model; Finite size scaling techniques are discussed in section 2.2.8 and applied to the gauge model. In Chapter 3, I will show some exact renormalization group (ERG) results. The motivation for this chapter is to analyze the discrete and continuous RG transformations. In Chapter 4, I will switch our attention to the more complicated and more interesting lattice gauge models. I will first analyze

the connection between the Ising model and the $SU(2)$ lattice gauge model by calculating the critical exponents. I will then work on the $SU(3)$ lattice gauge model with many light flavors. Due to the existence of the matter field, the computation is more tricky and time consuming. I will not show the most up-to-date result but only some results shown in 2012 lattice conference. New results will be published soon later. Tensor renormalization group (TRG) method will be discussed in Chapter 5. The advantages of the TRG is that it can be done for arbitrary large volume as long as one can perform the RG transformation once and the accuracy can be systematically improved.

I have been focusing my work on the field of theoretical high energy physics, with particular emphasis on phase transitions of gauge and spin models, and physics beyond the standard model. One of the purpose of my research was to figure out the phase structure of various gauge models and to unambiguously determine the existence of nontrivial infrared fixed point (IRFP) for those models. In order to do so, we have developed new techniques to prove the existence/nonexistence of the IRFP by monitoring the complex zeros of the partition function, which is conventionally called Fisher zeros, as the volume increases. The method was first constructed and tested on spin models and then extended to lattice gauge models, such as pure gauge $SU(2)$ and $U(1)$ case (I was not directly involved in the $U(1)$ case though). During this process, I have been trained to solve problems both analytically and numerically. On the analytical side, RG and especially finite size scaling (FSS) technique has been used through out my whole research period. I have been able to estimate the leading and sub-leading critical exponents for particular spin and gauge models. Finite volume effect of Fisher zeros and discrete β functions has also been carefully analyzed. The most important part is that I managed to construct the RG flows in the complex plane and showed that Fisher zeros separate

flows going into different fixed points.

In the following, I will briefly present my current and further research. With the upcoming experimental effort at Large Hadron Collider (LHC), there has been a lot of recent activities in the lattice community regarding the existence of the non-trivial IRFP in lattice models. The analysis involves non-perturbative dynamics like in QCD. Therefore lattice calculations are essential. Despite the intrinsic theoretical interest of the realization of conformal symmetry near an IRFP, it seems that phenomenologically, the preferred option is a “walking” coupling constant for extended technicolor interactions [47, 48].

Recently, many groups have been doing simulations on different systems in order to locate possible IRFP. Most of the simulations were done on $SU(2)$ and $SU(3)$ with different numbers of fermion flavors N_f in the fundamental representation. It seems to be uncontroversial that the β function is always negative in the range of couplings measured for $SU(3)$ with $N_f \leq 8$. The corresponding continuum theory is asymptotically free, chirally broken and confining. For $N_f > 8$, the situation is somewhat controversial. The Yale collaboration claimed an IRFP for $SU(3)$, $N_f = 12$, which contradicts the Wuppertal-Budapest collaboration’s observation that the systems with $N_f = 8$ and 12 are chirally broken and confining. For $SU(3)$ with $N_f = 2$ flavors of sextet fermions, the Colorado-Tel Aviv collaboration first claimed indications for an IRFP but was then disproved by the same authors. For $SU(2)$ with 2 ad-joint representations, there are also indications for the existence of an IRFP in the strong coupling region, but it is not completely uncontroversial either.

Given this situation, it is necessary to develop new methods that would lead to an unambiguous determination of the existence of IRFP or of a non-perturbative “walking” coupling constant.

On the theoretical side, developing new methods and applying existing RG ideas, which are usually tested on relative simple spin models, to the complicated gauge models are essential. Functional RG has been very successfully applied to many branches of physics. It is demanding to seek its application to every field of physics, especially the newly emerged ultra-cold atoms (optical lattice).

1.1.1 Spin Models

The Hierarchical Ising model (HM) is an ideal laboratory to test various ideas before one applies them to the more complicated full QCD case. Depending on the dimensionality and the interaction strength among blocks, the HM can have second order phase transition, no phase transition, or have a Thouless effect like in the Anderson model [49]. Fisher zeros and critical exponents of the HM can be calculated with very high accuracy. Figure 1.1 (Left Panel) shows the real and imaginary part of the lowest Fisher zeros for $D = 3$ HM. The lowest Fisher zeros accumulate toward the critical value β_c as the volume increases. The departure from a linear behavior is significant and requires sub leading corrections. The dotted lines are the fittings with sub leading corrections included.

The role of Fisher zeros can be clearly seen from the construction of the RG flows in the complex parameter plane. Complex RG flows can be constructed by using a two-lattice matching method. This method is a way of measuring the running of the bare couplings based on the fact that all the observables will have the same value if the models have the same effective action. Figure 1.2 shows complex flows for different dimensions. It can be clearly seen that Fisher zeros form the basin of attraction of the IRFP [50, 51]. That means that one only needs to get Fisher zeros of the system in order to determine whether there is a IRFP or not [52]. Figure 1.2 (Right Panel) also shows that Fisher zeros for larger volumes move approximately “backward” along the separation flow lines. This behavior

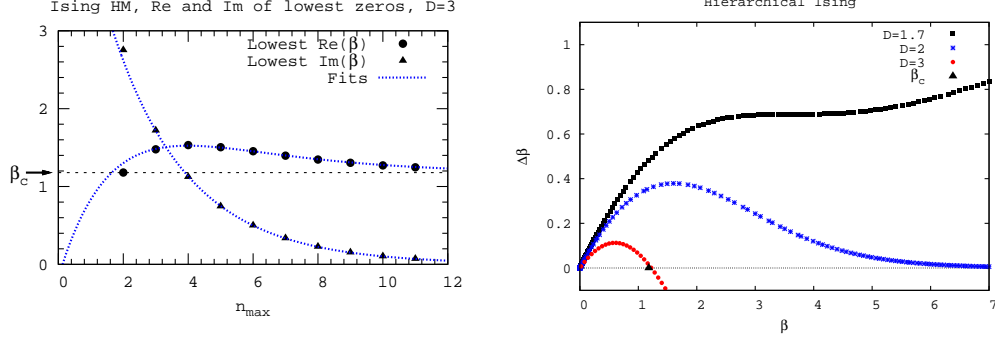


Figure 1.1: Real and imaginary parts of Fisher zeros for $D = 3$ and n_{max} going from 2 to 11 and discrete β functions for $D = 3, 2$, and 1.7 .

approximately agree with the argument that Fisher zeros for 2^n sites should map into Fisher zeros for 2^{n-1} sites.

Calculating the discrete β function is one of the methods used to look for or exclude IRFP. Discrete β function of the HM can be calculated numerically and semi-analytically. The zero of the discrete β function signals the appearance of a fixed point. However, sometimes finite volume effect may generate pseudo fixed point. This has been discussed extensively recently [49]. If one is interested in the corresponding continuous Callan-Symanzik β function, for example in the gauge model, one could use functional conjugation method to construct it from the discrete one [53, 49].

1.1.2 Gauge Models

In order to locate Fisher zeros for gauge models, we start from the inverse Laplace transform of the partition function, which is also called the density of state (DOS). The DOS can be constructed from plaquette configurations generated from the Monte Carlo simulation. The DOS of pure gauge $SU(2)$ on L^4 lattice has been successfully constructed [54, 55], see Fig. 1.3 (Left Panel). Once the DOS is known,

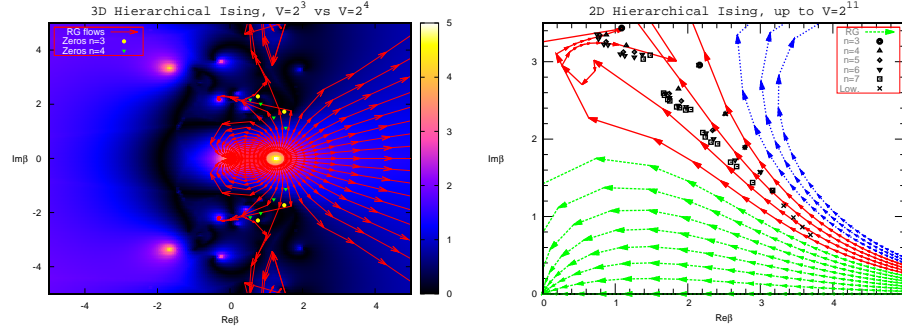


Figure 1.2: RG flows for the $D = 3$ hierarchical model in the complex β plane obtained by the two lattice method. and close up view of $D = 2$ RG flows and Fisher zeros for different volumes.

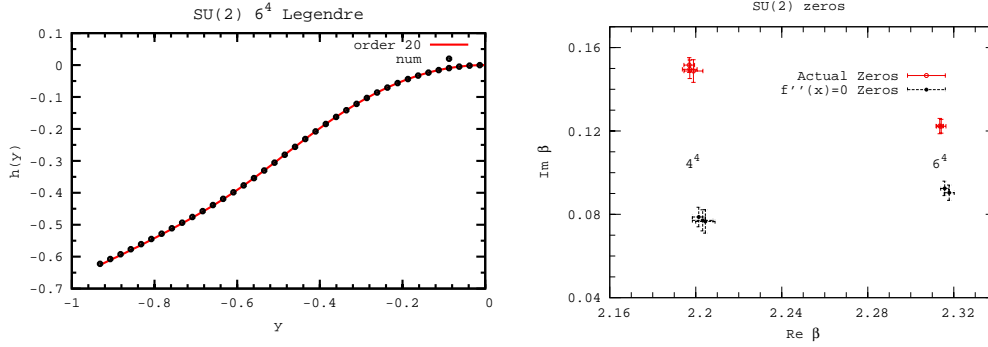


Figure 1.3: Density of state related function $h(y)$ together with the expansion in Legendre polynomials up to order 20 and Fisher zeros for the pure gauge $SU(2)$ without an ad-joint term for different volumes [1].

it is possible to calculate the partition function for any complex value of β , where $\beta = 2N/g^2$. Fisher zeros can then be calculated by using the re-weighting method. Figure 1.3 (Right Panel) shows Fisher zeros for pure $SU(2)$ gauge models on two different volumes. $SU(2)$ with ad-joint term and $U(1)$ cases have also been done [1]. The fact that Fisher zeros stabilizes away from the real axis for $SU(2)$ and pinches the real axis for $U(1)$ signals different cases of phase transitions. Similar phenomena have also been well tested in spin models [52].

Since we are ultimately interested in the thermodynamic limit, finite size scaling (FSS) technique is essential in analyzing the data and extrapolating relevant quantities to the infinite volume limit. Considerable experience on FSS technique has been acquired and already been applied to the analysis of the spin and gauge models [52, 56]. Leading and sub leading critical exponents of the gauge model have been extrapolated from the data by calculating the 4th order Binder cumulant.

1.2 Research Results

- Gauge Models

I have applied the techniques discussed above to $SU(3)$ lattice gauge model with various fermion flavor numbers and further located the partition function zeros of these models. This will provide an alternative way to locate the conformal window non-pertubatively.

We have a working parallel code with Dr. Don Sinclair to simulate $SU(3)$ with 2 and 3 staggered fermions. I have modified this existing code to simulate with N_f going from 4 to 16 and possibly non integer N_f by using rational hybrid Monte Carlo (RHMC) algorithm. The $N_f = 8$ and 16 case will be dealt with HMC algorithm separately. I started from small volume 4^4 with $N_f = 3$, where the rough location of β_c is known, and gradually increased the number of flavors to monitor how the β_c changes with the number of flavors. I have generated data for higher volumes. All the codes developed for this purpose are working well now and have adaptive routines. It would be good if we could compare results with other fermion actions. Therefore, we also use Professor Anna Hasenfratz group's code, which is based on the publicly available MILC code, to investigate the role of nHYP and APE spearing on the effects of the phase transitions.

- Spin Models

I have two undergoing projects in hand. One is the “numerical instabilities associated with block spinning non-integer numbers of sites”, where we block spin non-integer number of sites at each step and try to push the block spinning number to the continuum limit 1, where Functional RG methods work. We found out that the result is numerically unstable which may due to the logarithmic divergent term. Another one relates to the “exact renormalization group” (ERG) equations. I found out that some ERG equation are invariant under some particular rescaling.

Up to now, we have constructed complex RG flows for models with second order phase transition ($D = 3$ HM) and with no phase transition ($D = 2$ HM). I would like to find a suitable model with first order phase transition and see how the flows behave.

- Tensor Renormalization Group

The RG is a beautiful theory in dealing with systems with different length scales. It has inspired new concepts and computational methods in many branches of physics. In the lattice gauge theory, however, we recently found out that the conventional Migdal-Kadanoff (MK) method does not serve the purpose of high accuracy calculations due to its crude “bond sliding” approximation [57]. This inspired us to develop new method to “block spin” the system accurately and provide a tool to study lattice models near conformality. We started out noticing the density matrix renormalization group (DMRG) [58] methods which are widely used in the condensed matter community. We realized that despite its success in dealing with quantum systems in one space and one time dimension, the method quickly fails in dealing with higher dimensions. Recently, we developed the tensor renormalization group

(TRG) [59, 60] to deal with classical discrete spin models away from criticality. This inspired us to develop new methods to deal with continuous spin and lattice gauge models based on TRG. A variant of that, second renormalization of tensor-network state (SRG), which takes into account the effect of the environment and improves the accuracy at the criticality is also under development. Our goal is to apply the method to four dimensional lattice gauge models with fermions.

This is a long term project partially due to its complexity in four space-time dimension. Up to now, we have successfully written down the tensor state and figured out how to perform contractions for the two and three dimensional $O(N)$ models, and three dimensional $U(1)$ model. New mathematical and computational tools may be needed to deal with such complicated cases.

CHAPTER 2

RENORMALIZATION GROUP METHODS ON SPIN MODELS IN THE HIERARCHICAL APPROXIMATION

2.1 Ising Model

Before discussing about the hierarchical model, let us start with the usual Ising model. The Ising model was first proposed by Lenz [61] to his student Ising [62] in the early 1920s in order to analyze ferromagnetism. The hope was to devise a model with inter-particle interaction to display a magnetic phase transition. Ising managed to solve the model on a one-dimensional lattice and found that the model actually had no phase transition at finite temperature. We now sometimes say that the one dimensional Ising chain has a phase transition at zero temperature since the system only has long range order at $T_c = 0$. The ‘failure’ of describing the phase transition made Ising conclude that the model would not have any phase transition in higher dimension either. Of course, we now know that the Ising model has phase transitions in two and higher dimensions. In 1936, Peierls [63] proved that the Ising model should have phase transitions in two and three dimension at sufficiently low temperatures. Later in 1941, Kramers and Wannier [64] successfully located the exact critical temperature of the two dimensional Ising model. They argued that the nearest neighbor Ising model partition function at temperature T_1 could be transformed into another Ising model at a different temperature T_2 . They identified the critical temperature T_c by assuming that there is only one critical temperature. In 1944, Onsager[65] , derived the explicit expression of the free energy without external magnetic field in two dimension and further got the singularity behavior of the specific heat, which put Van der Waals mean field theories [66, 67, 68] into question and led to the discovery of the λ transition of the liquid helium [69] and the liquid vapor critical point [70] in Argon. After Onsager’s tremendous success,

many other two dimensional lattice models have been solved exactly. However, none of those different approaches can be generalized to the more physically interesting three dimensional case. In 1952, Lee and Yang [71, 72] gave a rigorous mathematical description of the phenomena of the phase transition. They studied the lattice gas model, which can be mapped into Ising model, in an external magnetic field. They discovered that the complex zeros of the partition function lie on a unit circle in the complex activity plane, or in another word on the imaginary external magnetic field h plane. They could relate the phase transition to the pinching of the zeros onto the real h axis. It was natural to generalize Lee-Yang's theorem to the complex temperature plane. There is no known general theorem to constrain the distribution of the partition function zeros in the complex temperature plane. However, Fisher [73] first observed that the zeros of the two dimensional Ising model on the square lattice distribute on two circles in the complex $\tanh(\beta)$ plane. We usually call the the zeros in the h plane Lee-Yang zeros and the ones in the β plane Fisher zeros. In the following, we will briefly discuss the Ising model and use that to introduce important concepts and quantities discussed later in the thesis.

The Hamiltonian of the nearest neighbor Ising model on a regular lattice with external magnetic field h is defined as

$$H = -J \sum_{\langle i,j \rangle} \sigma_i \sigma_j - h \sum_i \sigma_i, \quad (2.1)$$

where $\sigma_i = \pm 1$, and the first sum is over all the nearest neighbor pair $\langle i, j \rangle$. J is the coupling constant, which controls the interaction strength of the neighboring sites. We usually consider $J > 0$ case and the interaction is ferromagnetic. The

partition function can then be written as

$$Z[\beta, h] = \sum_{\{\sigma_i\}} e^{-\beta H} \quad (2.2)$$

$$= \sum_{E=0}^{N_1} \sum_{M=0}^{N_2} \alpha(E, M) x^E y^M, \quad (2.3)$$

where β is the inverse temperature T , $x = e^{-2\beta J}$ and $y = e^{-2\beta h}$. N_1 and N_2 are total number of sites and bonds of the lattice, and $\alpha(E, M)$ is the number of the configuration with E sites and M bonds [74].

The lattice gas Hamiltonian is the following

$$H = -4J \sum_{\langle i, j \rangle} \rho_i \rho_j - \mu \sum_i \rho_i, \quad (2.4)$$

where $\rho_i = 0, 1$, and μ is the chemical potential. The lattice gas model can be transformed into the Ising model by the following variable changes [71] :

$$\rho_i = \frac{1}{2}(\sigma_i + 1), \quad (2.5)$$

$$\mu = 2h - 4qJ, \quad (2.6)$$

where q is the total lattice site number.

In the zero magnetic field case, $h = 0$, the Ising model is in the ferromagnetic phase for $T < T_c$, where there is a spontaneous symmetry breaking, and in the paramagnetic phase for $T > T_c$. T_c is the so-called critical temperature of the system. For the lattice gas model, there are also gas and liquid phases for different temperatures for $\mu = -4qJ$. I will not go through the exact solution of the two dimensional Ising model here. They can be found in original papers [65, 75] and text books [76, 77, 78, 79]. Despite the tremendous success of Onsager's exact solution [65], the Ising model and its variants were still considered as the pure mathematical trick which had no physical meaning. Until the 1960s, people realized that the lattice gas model can reasonably well describe the critical behavior of the real liquid gas transition. Combining with the “universality class” and finite size scaling (FSS)

concepts, Ising model and other ‘simple’ models have since been considered seriously as approximation of the real world.

2.2 Dyson’s Hierarchical Model

In the previous section, we introduced the Ising model and explained that the model can only have phase transitions in the two or higher dimensions. From the universality point of view, the properties of the transition is only related to the dimensionality of the system and the internal symmetries. but not to the detailed structure of the model. Keep in mind that the model we introduced has only nearest neighbor interactions. It has been proved by Peierls [63] that for the ferromagnetic system with short range interactions between the spins, the order-disorder transition requires at least two dimensions. Colin Thompson and Mark Kac conjectured in 1967-1968 that there could be a phase transition for the one dimensional system with interaction strength proportional to $d^{-\alpha}$ with d to be the separation between two spins and $1 < \alpha \leq 2$. The conjecture has been proved by Frohlich and Spencer [80] in 1987. But at that time it was still a mystery [81]. Dyson happened to know the conjecture and wanted to prove it. To achieve that, he constructed a special model with long range interaction. He was able to prove that with the interaction strength $d^{-\alpha}$, the system could have a phase transition for $1 < \alpha < 2$ but he could not prove the $\alpha = 2$ case [82, 83, 84]. The $\alpha = 2$ case is in fact very interesting because it corresponds to the Anderson model with inverse square interaction [85, 86, 87]. Thouless [88] predicted that the Anderson model should have a very unusual phase transition with discontinuous spontaneous magnetization, which is usually called “Thouless effect”. I will discuss about that later in this chapter.

The tool Dyson used was called the hierarchical model (HM). It was first introduced by Freeman J. Dyson [89] in order to mimic a one dimensional Ising model with long-range interactions. It has been proved that the model has a second order

phase transition under certain conditions. Related models were re-investigated by George A. Baker [90]. His motivation was to find a model where Kenneth G. Wilson's [91] approximate recursion formula was exact. He also extended the model to higher dimensions. Because of its complexity (it has phase transition) and simplicity (it has an exact recursion formula), the model has been investigated intensively by P. M. Bleher and Ja. G. Sinai [92, 93], G. Gallavotti etc [94, 95], P. Collet and J. -P. Eckmann [96, 97, 98], and some others [99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 114, 115, 116, 117] in the past forty years. For a recent review, see Ref. [118] and the references therein.

2.2.1 Definition Of The Model

The partition function of the HM is defined as:

$$Z = \int e^{-\beta H}, \quad (2.7)$$

where H is the Hamiltonian of the model [82, 90]

$$H = -\frac{1}{2} \sum_{m=1}^N \left(\frac{c}{4}\right)^m \sum_{p_N, \dots, p_{m+1}} \left(\sum_{p_m, \dots, p_1} \phi_{(p_N, \dots, p_1)} \right)^2 \quad (2.8)$$

The meaning of the parameters and the notations used in the following sections are defined as the following:

D : Dimension of the system.

ϕ : Scalar field in the configuration space.

$\frac{c}{4} = b^{-2-D}$: Control the decay of the interaction.

b^D : Number of sites blocked in each RG iteration.

a : Lattice Spacing.

Λ : UV cutoff.

where ϕ is the scalar field in the configuration space. For the Ising model

discussed below, ϕ can only be ± 1 . The index p_i is set to 0 or 1, depending on which block we are referring to, and the summation is over all the combination of the 0 and 1 [118]. The parameter c controls the strength of the interaction. In order to have a phase transition, we require the parameter c to be positive and smaller than 2. That means that the interaction strength decreases with the distance. The model was initially developed as a one dimensional chain model. It is very easy to extend the original model to higher dimensions. We can include the dimension D to the strength parameter c : $c/4 = b^{-2-D}$, where b is the rescaling factor. For reasons why we can do so, please refer to [118]. Usually we will take $b^D = 2$ without further notice. One of the properties of the model is that the partition function $Z = \sum e^{-\beta H}$ can be calculated iteratively. In order to get the action, we need to sum up all the spin configuration of the system, or equivalently integrate over a local measure. For the Ising case, the spins can only have the value ± 1 . Therefore, the local measure for the Ising case is

$$W_0(\phi) = \delta(\phi^2 - 1). \quad (2.9)$$

In the following, we always work with the Ising measure. The model can also be constructed with Landau-Ginsburg measure

$$W_0(\phi) = \exp(-\frac{1}{2}m^2\phi^2 - \lambda\phi^4). \quad (2.10)$$

In the rest of this section, we will discuss the local measure in both the configuration ϕ space and the Fourier transformed k space. Suppose we start from the initial Ising measure in ϕ space

$$W_0(\phi) = \delta(\phi^2 - 1) \quad (2.11)$$

$$= \frac{1}{2}(\delta(\phi - 1) + \delta(\phi + 1)). \quad (2.12)$$

When we do the renormalization group (RG) transformation, we keep the sum of the spin in one block constant. We can easily get the new measure from the old one

due to the hierarchical structure of the model:

$$W_{n+1}(\phi) = [\int d\phi_1 d\phi_2 \delta(\phi_1 + \phi_2 - \phi) * W_n(\phi_1) W_n(\phi_2)] * e^{\frac{\beta}{2}(\frac{\epsilon}{4})^{n+1} \phi^2}, \quad (2.13)$$

where W_n is the unnormalized probability distribution of ϕ in a block of size 2^n .

In order to get rid of the delta function in equation 2.13, we introduce a new variable

$$\xi = \frac{\phi_1 - \phi_2}{2}, \quad (2.14)$$

and write

$$\phi_1 = \frac{\phi + 2\xi}{2} \quad (2.15)$$

$$\phi_2 = \frac{\phi - 2\xi}{2}. \quad (2.16)$$

Then we can change variable from ϕ_1 and ϕ_2 to ϕ and ξ . Equation 2.13 becomes

$$W_{n+1}(\phi) = [\int d\xi W_n(\frac{\phi}{2} + \xi) W_n(\frac{\phi}{2} - \xi)] * e^{\frac{\beta}{2}(\frac{\epsilon}{4})^{n+1} \phi^2}. \quad (2.17)$$

This is the iteration formula for the local measure in ϕ space. If we introduce the Fourier transform of the measure $W_n(\phi)$

$$W_n(\phi) = \int \frac{dk}{2\pi} e^{ik\phi} R_n(k), \quad (2.18)$$

we will get

$$\begin{aligned} R_{n+1}(k) &= \int d\phi e^{-ik\phi} W_{n+1}(\phi) \\ &= \int d\phi e^{-ik\phi} e^{\frac{\beta}{2}(\frac{\epsilon}{4})^{n+1} \phi^2} * \\ &\quad [\int d\xi \int \frac{dk_1 dk_2}{(2\pi)^2} e^{ik_1(\frac{\phi}{2} + \xi)} R_n(k_1) \\ &\quad e^{ik_2(\frac{\phi}{2} - \xi)} R_n(k_2)]. \end{aligned} \quad (2.19)$$

Equation 2.19 can be rewritten as

$$\begin{aligned} & \int d\phi e^{-ik\phi} e^{\frac{\beta}{2}(\frac{\xi}{4})^{n+1}\phi^2} \dots \\ &= \int d\phi e^{-\frac{\beta}{2}(\frac{\xi}{4})^{n+1}\nabla_k^2} e^{-ik\phi} \dots \end{aligned} \quad (2.20)$$

Integrating out ξ in equation 2.19, the rest becomes

$$\begin{aligned} & \int d\xi \int \frac{dk_1 dk_2}{(2\pi)^2} e^{ik_1(\frac{\phi}{2}+\xi)} R_n(k_1) e^{ik_2(\frac{\phi}{2}-\xi)} R_n(k_2) \\ &= \int \frac{dk_1 dk_2}{(2\pi)^2} 2\pi \delta(k_1 - k_2) e^{i(k_1+k_2)\frac{\phi}{2}} R_n(k_1) R_n(k_2) \\ &= \int \frac{dk_1}{2\pi} R_n^2(k_1) e^{ik_1\phi}. \end{aligned} \quad (2.21)$$

Combining the above two equations, the recursion formula for the Fourier transform of the measure can be written as

$$R_{n+1}(k) = e^{-\frac{\beta}{2}(\frac{\xi}{4})^{n+1}\nabla_k^2} R_n^2(k). \quad (2.22)$$

And the Fourier transform of the initial measure W_0 (Eq: 2.12) is

$$R_0(k) = \cos(k). \quad (2.23)$$

We can get the measure for volume $V = 2^N$ by starting from $R_0(k)$ and applying the recursion formula 2.22 N times. The nice property of the HM is that the block spin transformation only changes the local measure:

$$W_{n+1}(\phi) = C_{n+1} e^{\frac{\beta}{2}(\frac{\xi}{4})^{n+1}\phi^2} \int d\phi' W_n\left(\frac{(\phi - \phi')}{2}\right) W_n\left(\frac{(\phi + \phi')}{2}\right). \quad (2.24)$$

And the recursion formula for the $R_n(k)$ is then

$$R_{n+1}(k) = C_{n+1} e^{-\frac{1}{2}\beta\frac{\partial^2}{\partial k^2}} \left(R_n\left(\sqrt{\frac{c}{4}} k\right) \right)^2. \quad (2.25)$$

2.2.2 Partition Function Of The hierarchical Model

In the following, we will give an example on how to construct the partition function with volume $V = 2^2$ from that with volume $V = 2^1$. The Hamiltonian in

equation 2.8 with $N = 2$ is

$$H_{N=2} = -\frac{1}{2} \left[\frac{c}{4} \sum_{p_2} (\phi(p_2, 0) + \phi(p_2, 1))^2 \right. \quad (2.26)$$

$$\left. + \left(\frac{c}{4}\right)^2 (\phi(0, 0) + \phi(0, 1) + \phi(1, 0) + \phi(1, 1))^2 \right] \quad (2.27)$$

$$= -\frac{1}{2} \left[\frac{c}{4} ((\phi(0, 0) + \phi(0, 1))^2 + (\phi(1, 0) + \phi(1, 1))^2) \right. \quad (2.28)$$

$$\left. + \left(\frac{c}{4}\right)^2 (\phi(0, 0) + \phi(0, 1) + \phi(1, 0) + \phi(1, 1))^2 \right], \quad (2.29)$$

and the corresponding partition function is

$$\begin{aligned} Z_{N=2} &= \sum e^{-\beta H_{N=2}} \\ &= \sum \exp \left[\frac{\beta c}{2 \cdot 4} (\phi(0, 0) + \phi(0, 1))^2 \right] * \end{aligned} \quad (2.30)$$

$$\exp \left[\frac{\beta c}{2 \cdot 4} (\phi(1, 0) + \phi(1, 1))^2 \right] * \quad (2.31)$$

$$\exp \left[\frac{\beta}{2} \left(\frac{c}{4}\right)^2 (\phi(0, 0) + \phi(0, 1) + \phi(1, 0) + \phi(1, 1))^2 \right], \quad (2.32)$$

where the summation is over all configurations with $\phi(p_2, p_1) = \pm 1$. The partition function for $N = 1$ is

$$\begin{aligned} Z_{N=1} &= \sum e^{-\beta H_{N=1}} \\ &= \sum_{\phi(0, p_1) = \pm 1} \exp \left[\frac{\beta c}{2 \cdot 4} (\phi(0, 0) + \phi(0, 1))^2 \right]. \end{aligned} \quad (2.33)$$

We notice that equation 2.33 has the same form as equation 2.30 and 2.31. Suppose we have $Z_{N=1}$ already, we only need to calculate equation 2.32 in order to get $Z_{N=2}$. Furthermore, we can construct $Z_{N=i+1}$ from $Z_{N=i}$. That enables us to calculate the partition function exactly for relatively large volume system without running over all the possible configurations.

2.2.3 Extension Of The Recursion Formula

The recursion formula 2.25 can be extended for an arbitrary scale transformation. The number of sites integrated for the HM, namely 2, appears as the exponent in the recursion formula. In order to generalize the formula to arbitrary dimension

D , we only need to do the following replacements

$$2 \rightarrow b^D \quad (2.34)$$

$$\frac{c}{4} \rightarrow b^{-2-D}. \quad (2.35)$$

And under the renormalization group (RG) transformation, the lattice spacing a and the momentum cutoff becomes

$$a \rightarrow ba \quad (2.36)$$

$$\Lambda \rightarrow \frac{\Lambda}{b}. \quad (2.37)$$

The recursion formula 2.25 becomes

$$R_{n+1}(k) = C_{n+1} e^{-\frac{1}{2}\beta \frac{\partial^2}{\partial k^2}} \left(R_n(b^{-(D+2)/2} k) \right)^{b^D}. \quad (2.38)$$

The usual recursion formula is obtained for $b = 2^{1/3}$ and $D = 3$.

2.2.4 Zeros Of The Partition Function

Before starting to address the technical details of how to locate the zeros of the partition function, we would like to discuss the importance of finding the zeros of the partition function, or more generally the zeros of some analytic function.

The first thing coming out of my mind when talking about the zeros of the analytic function is Riemann hypothesis. It states that all the nontrivial zeros of the Riemann zeta function $\zeta(z)$ lies on the critical line $Re(z) = 1/2$. Although trillions of zeros have been calculated and all of them lie in the critical line, the hypothesis is still one of the most important unsolved mathematical problems. Another example relating to the zeros of the analytic function is the famous Lee-Yang theorem [119, 120]. The theorem states that the partition function zeros of the Ising ferromagnet are purely imaginary in the complex magnetic field plane, or lie on the unit circle in the complex e^h plane. They also pointed out the relationship between the distribution of the zeros and the order of the phase transition. The theorem has

been tested on numerous models by different authors. Years later in 1964, M. E. Fisher studied the zeros of the partition function in the complex temperature plane [121]. As mentioned in the previous section, we called these two types of partition function zeros Lee-Yang zeros and Fisher zeros. In this section, we will discuss the later one.

In general, the partition function of the Dyson's hierarchical model (HM) can not be expressed as a finite order polynomial of the variable $e^{-2\beta}$ or others as can be done in the regular Ising model. This property makes the analysis of distribution and scaling of the partition function zeros more difficult. Whether there is a regular distribution function $g(\theta)$ or not is still an open question.

The partition function of the hierarchical Ising model has the following form

$$Z[\beta] = \sum_{k=0}^l a_k e^{\lambda_k \beta}. \quad (2.39)$$

The coefficient a_k are non-negative integers and λ_k are non-negative real numbers. Equation (2.39) is not an ordinary polynomial but a quasi-polynomial [122]. The general form of the quasi-polynomial is

$$Z[\beta] = \sum_{j=0}^m \sum_{k=0}^l a_{jk} \beta^j e^{\lambda_k \beta}. \quad (2.40)$$

Equation (2.39) is a special case with $m = 0$. Obviously, there is no real solution for the equation $Z[\beta] = 0$ because all the coefficients a_k in equation 2.39 are non-negative. In order to find the partition function zeros, it is necessary to extend β to the complex plane. These zeros are the Fisher zeros mentioned above.

We have discussed how to get the exact expression for the partition function in section 2.2.2. In principle, the problem of finding the Fisher zeros is just the problem of finding the roots of the quasi-polynomial 2.39. If all the λ_k were integers, we could write the partition function 2.39 as a finite order polynomial with the variable $e^{a\beta}$, where a is an integer depending on the actual form of the partition. Many system

studied have the above properties, i. e. [74]. Finding the roots of the finite order polynomial is not too difficult as long as the order is not too large. The model we are interested in here, however, is not so simple. Most of the times λ_k is not integer and not even an rational number for the HM. This makes the calculation of the Fisher zeros not so trivial. Even though we could get the expression of the partition function, it is impractical to calculate its roots analytically. We developed two methods to locate the Fisher zeros. The first method is by calculating both the real and imaginary part of the partition function, finding the zeros of each part, and locating their crossing. The second one is the “logarithmic residue” method which is based on the Cauchy’s Residue Theorem. The method relies on the following fact

$$\frac{1}{2\pi i} \oint_C \beta^n \frac{Z'(\beta)}{Z(\beta)} = \sum_{i=1}^k \beta_i^n, \quad (2.41)$$

where β_i are all the zeros in the integration region C . It should be noted that Riemann used similar techniques to estimate the number of zeros in the particular region in his original paper on Riemann hypothesis. The “logarithmic residue” method works well if we know the expression of the function and if we only need to find one or several roots in a specified region, which is just the case we studied here. We usually start from $n = 0$ and the right hand side of the equation 2.41 gives the number of roots located in the specified region C

$$\frac{1}{2\pi i} \oint_C \frac{Z'(\beta)}{Z(\beta)} = k. \quad (2.42)$$

We can always adjust the integration region C and make sure that there is only one zero in the region. And then we can set $n = 1$ to calculate the location of the zero.

For the lattice gauge models that will be discussed in the following chapters, one usually can not get the exact expression of the partition function. In this case, one can first construct the density of state and get the zeros of the partition function out of the numerical Laplace transformation.

Let us look at some examples of the first zero searching method. In order to locate the complex zeros of the partition function, we extend the parameter β , which corresponds to the inverse temperature of the system, to the complex plane. We scan the the complex β plane in some region. For every complex β , we calculate the real and imaginary part of the partition function Z . We record the points where the partition function changes sign. These points correspond to the zeros of the real or imaginary part of the partition function. Only the points with zeros in both real and imaginary parts of the partition function are the complex Fisher zeros. Because of the special structure of the equation 2.8, we do not need to sum up all the spin configuration of the system. We can build the Hamiltonian iteratively and calculate the partition function precisely.

Figure 2.1 shows the zeros for $D = 3$ and the volume $V = 2^4$. The Fisher zeros correspond to the crossing of the red and the blue lines. In the region we plotted, there are four Fisher zeros. They are not scattered everywhere into the complex plane. Instead, there is a clear tendency that they follow in some curve. As we increase the volume to 2^5 , the Fisher zeros in the same region double (see Fig. 2.1) and the lowest zero moves closer to the real axis. This is in agreement with the fact that there will be no zero in the real axis for a finite volume system. When we go to the infinite volume limit, the zeros pinch to the real temperature axis, which is the indication of the phase transition.

In order to see all the the zeros in the complex temperature plane, we also search the partition function zeros in the complex $\frac{1}{\beta}$ plane. The result for $V = 2^4$ is shown in Figure 2.3, 2.4. Clearly, all the Fisher zeros accumulate in the small region near the origin. From the complex inverse mapping on the extended complex plane, we know that the line in the complex $\frac{1}{\beta}$ plane corresponds to a circle in the complex β plane. The accumulation in the complex $\frac{1}{\beta}$ plane means there is no zero

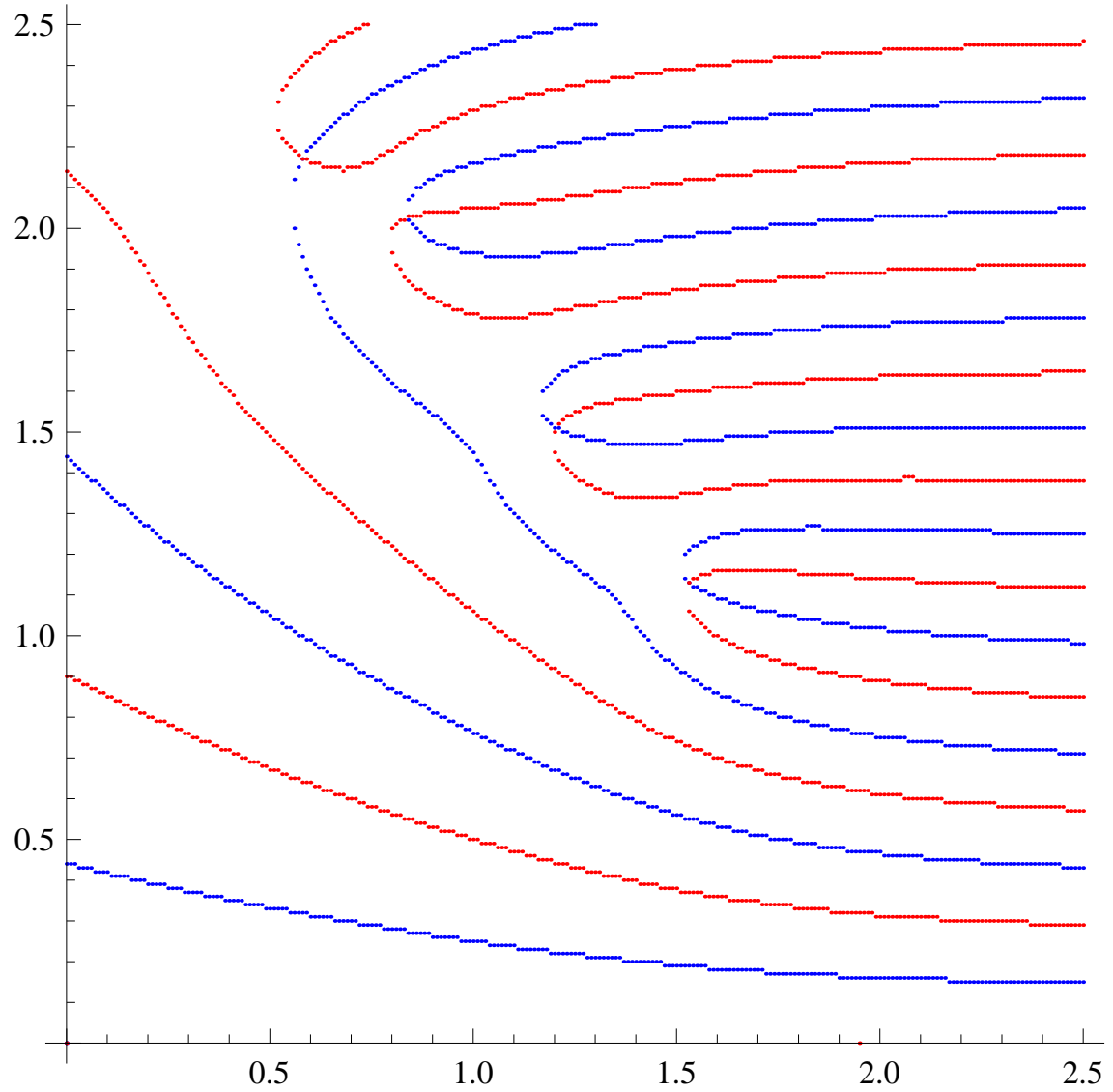


Figure 2.1: The partition zeros in the complex β plane for the HM with volume 2^4 in 3 dimension.

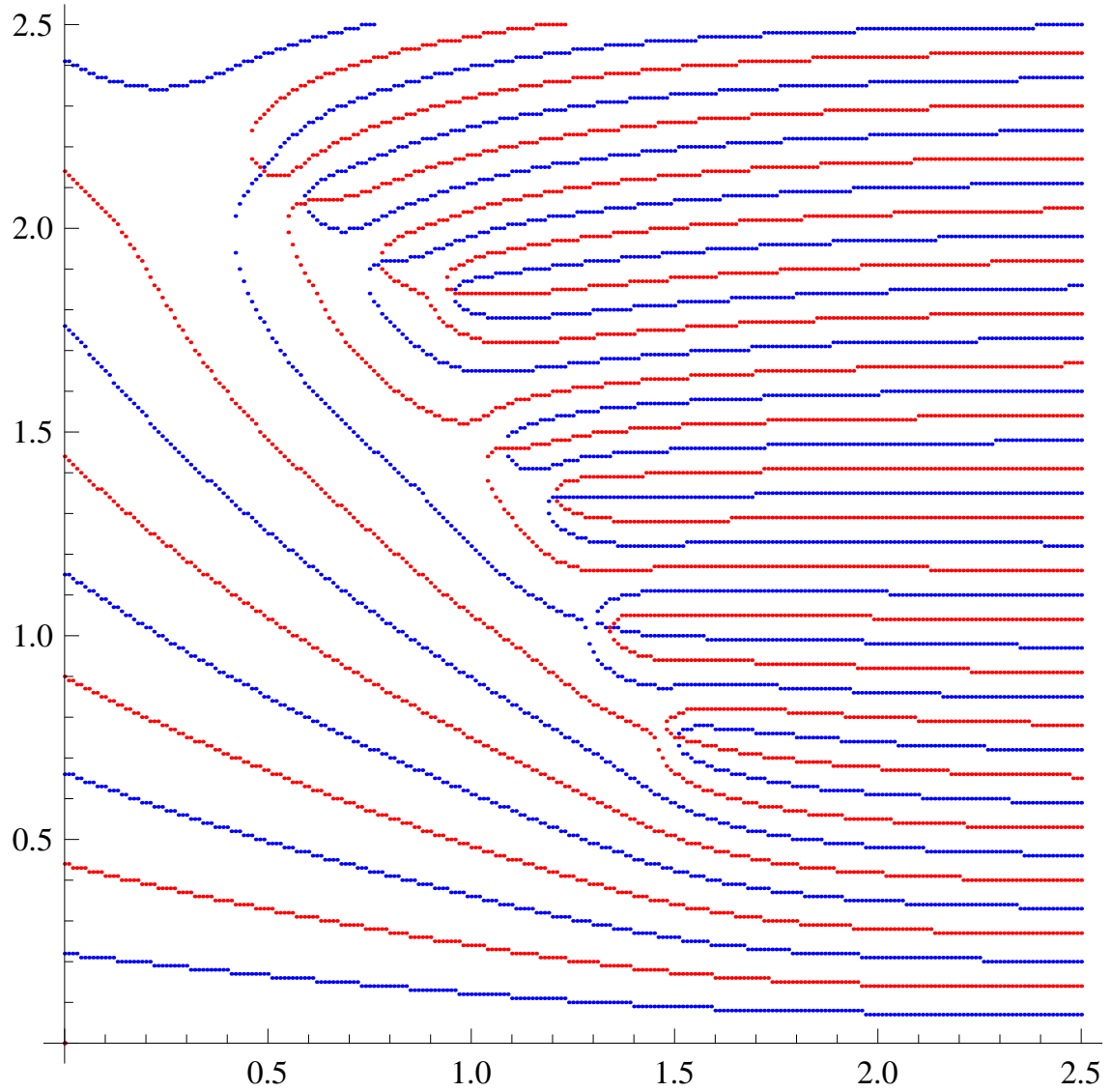


Figure 2.2: The partition zeros in the complex β plane for the HM with volume 2^5 in 3 dimension.

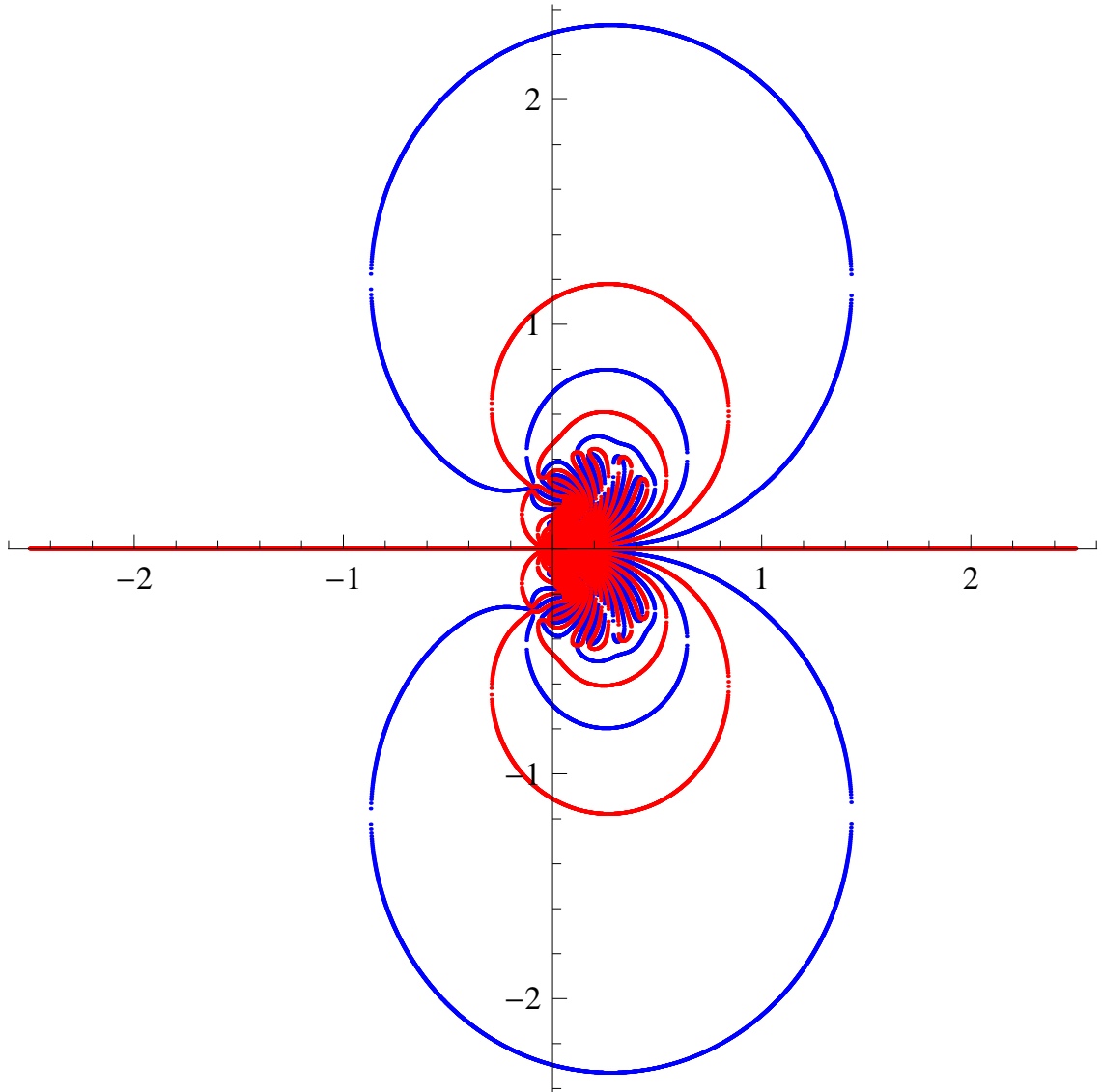


Figure 2.3: Same system as in Fig. 2.1 but in the complex inverse β plane.

near the origin in the complex β plane. We can also see the corresponding zeros of Figure 2.1 from the Figure 2.4.

2.2.5 Distribution Of The Zeros For Different Variables

At the end of this section, we want to briefly discuss the distribution of the Fisher zeros. From figure 2.14, we can clearly see that the number of zeros in the

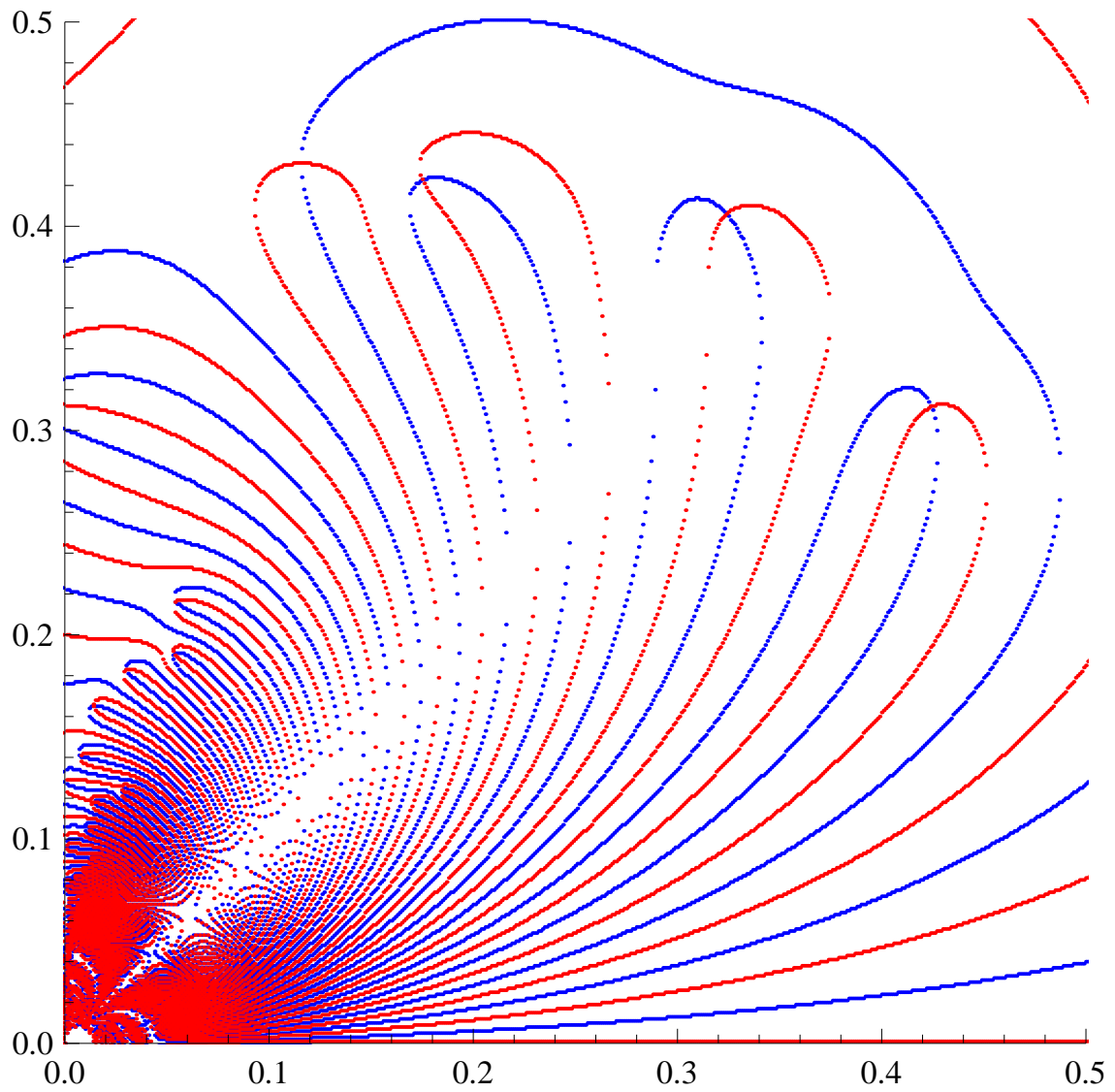


Figure 2.4: Same system as in Fig. 2.1 but in the complex inverse β plane.

specific region is proportional to the logarithm of the volume.

Sometimes the distribution of the partition function zeros is considered in the complex $\tanh \beta$ plane or the complex $e^{-2\beta}$ plane. The zeros is usually distributed on the circles. In the following, we will try to connect different distribution patterns with different choices of variables.

Suppose the Fisher zeros in the complex β plane has the form $\beta = \beta_r + i\beta_i$. Then if we express these zeros in the complex $\tanh \beta$ plane, we will get

$$\begin{aligned}\tanh \beta &= \tanh(\beta_r + i\beta_i) \\ &= \frac{\sinh(2\beta_r)}{\cos(2\beta_i) + \cosh(2\beta_r)} + i \frac{\sin(2\beta_i)}{\cos(2\beta_i) + \cosh(2\beta_r)}.\end{aligned}\tag{2.43}$$

We further assume β_r is a constant, which was observed in the hierarchical Ising model in the infinite volume limit. That means the distribution of the Fisher zeros form a line which is parallel to the imaginary β plane (Actually there are two lines, one crosses the positive β and another crosses the negative β . We are only interested in the real β or ferromagnetic region.). Then in the $e^{-2\beta}$ or $\tanh \beta$ plane, the Fisher zeros will distribute in a circle. The reason is that both $f(\beta) = e^{-2\beta}$ and $f(\beta) = \tanh \beta$ are conformal mapping and they will map the lines parallel to the imaginary axis in the β plane to circles in the β and $\tanh \beta$ plane.

The following are examples of the distributions of zeros in different variables. The partition function for $D = 2$ and $V = 2^3$ HM is

$$\begin{aligned}Z(\beta) &= 64e^{21\beta/32} + 16e^\beta + 32e^{5\beta/4} + 32e^{11\beta/8} \\ &\quad + 16e^{13\beta/8} + 32e^{53\beta/32} + 4e^{2\beta} + 16e^{69\beta/32} + \\ &\quad 16e^{77\beta/32} + 8e^{21\beta/8} + 2e^{3\beta} + 2e^{7\beta/2} + 16.\end{aligned}\tag{2.44}$$

It is easy to change the variable from e^β to x^{32} and then write the partition as a polynomial of variable x . We can then get all the roots of the polynomial by

conventional methods. Figure 2.6 is a plot of all the roots in the complex $x = e^{\beta/32}$ plane for $V = 2^4$ case. We can see “clear” pattern that all the roots are located around the unit circle. However, if we plot the roots in the complex $\log(x)$ plane, then the roots will scatter around the plane and no obvious pattern persists.

Things become more difficult when we switch to $D = 3$ case. The following is the partition function for $D = 3$ and $V = 2^3$,

$$\begin{aligned}
Z(\beta) = & 16e^{\sqrt[3]{2}\beta} + 4e^{2\sqrt[3]{2}\beta} + 64e^{\frac{\beta}{4\sqrt[3]{2}} + \frac{\beta}{2^{2/3}} + \frac{\beta}{16}} \\
& + 32e^{\sqrt[3]{2}\beta + \frac{\beta}{4\sqrt[3]{2}} + \frac{\beta}{2^{2/3}} + \frac{\beta}{16}} \\
& + 32e^{\sqrt[3]{2}\beta + \frac{\beta}{2\sqrt[3]{2}}} + 32e^{\sqrt[3]{2}\beta + \frac{\beta}{2\sqrt[3]{2}} + \frac{\beta}{4}} \\
& + 16e^{\sqrt[3]{2}\beta + \frac{\beta}{\sqrt[3]{2}} + \frac{\beta}{4}} + 16e^{\sqrt[3]{2}\beta + \frac{5\beta}{4\sqrt[3]{2}} + \frac{\beta}{2^{2/3}} + \frac{\beta}{16}} \\
& + 16e^{\sqrt[3]{2}\beta + \frac{5\beta}{4\sqrt[3]{2}} + \frac{\beta}{2^{2/3}} + \frac{9\beta}{16}} + 8e^{2\sqrt[3]{2}\beta + \frac{\beta}{\sqrt[3]{2}} + \frac{\beta}{4}} \\
& + 2e^{2^{2/3}\beta + 2\sqrt[3]{2}\beta} + 2e^{2^{2/3}\beta + 2\sqrt[3]{2}\beta + \beta} + 16.
\end{aligned} \tag{2.45}$$

2.2.6 $D = 2$

In Ref. [123], we try to argue that the Fisher zeros form the “gate” to control the RG flows. In the following, we will consider two different cases: the first case has no phase transition and the other one has a second order phase transition. We will see how we can extract the information about the phase transition from the location of the Fisher zeros.

In the $D = 2$ case, the HM has no phase transition at finite temperature [82]. That means that the partition function should not be zero on the real axis even in the infinite volume limit. In figure (2.7) we plot the lowest Fisher zeros of the HM with different volume. The distribution of the Fisher zeros is symmetric about the real β plane so that we only need to show the zeros in the upper half plane.

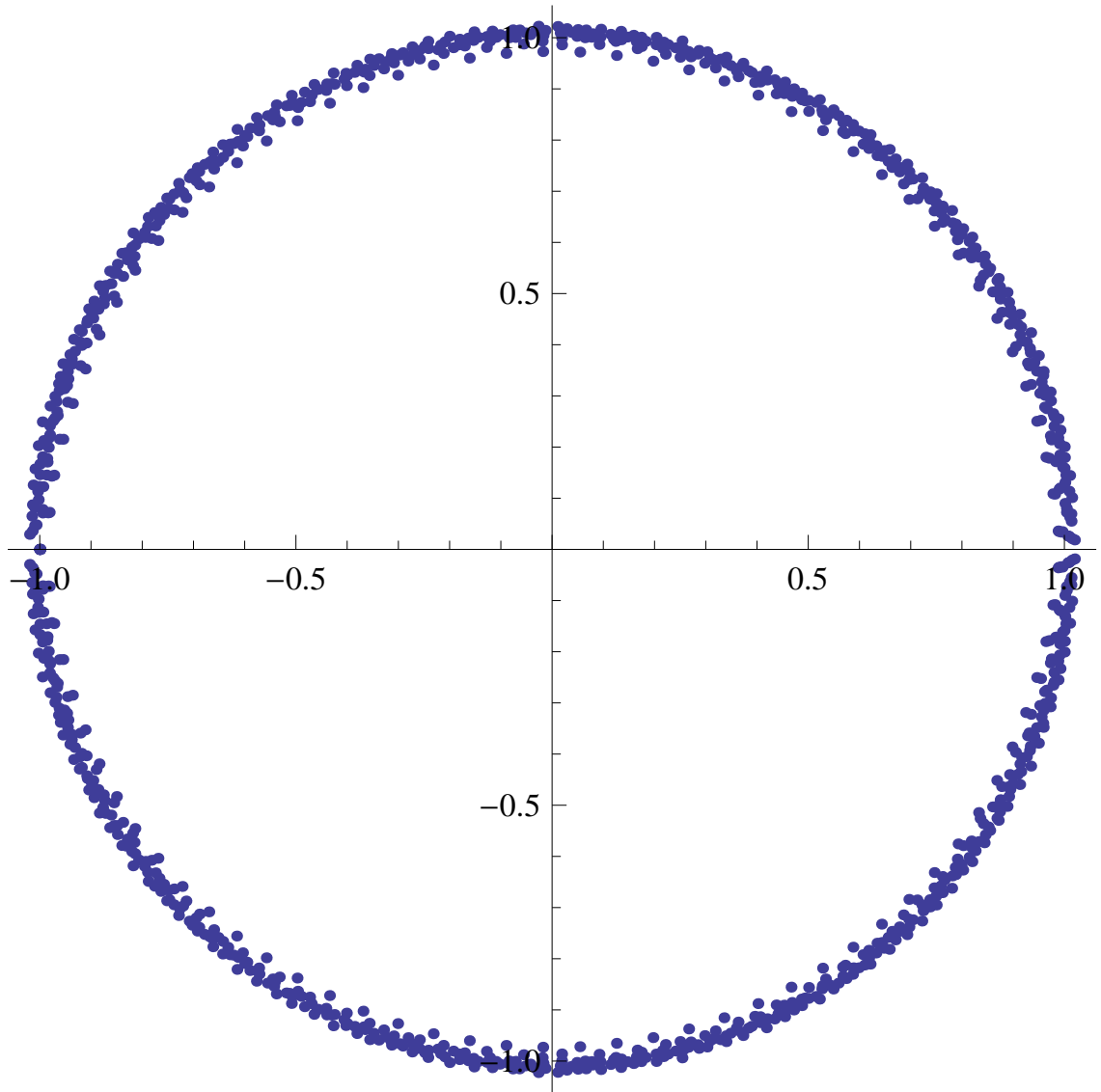


Figure 2.5: The roots of the hierarchical model partition for $D = 2$ and $V = 2^4$ in the $x = e^\beta$ plane.

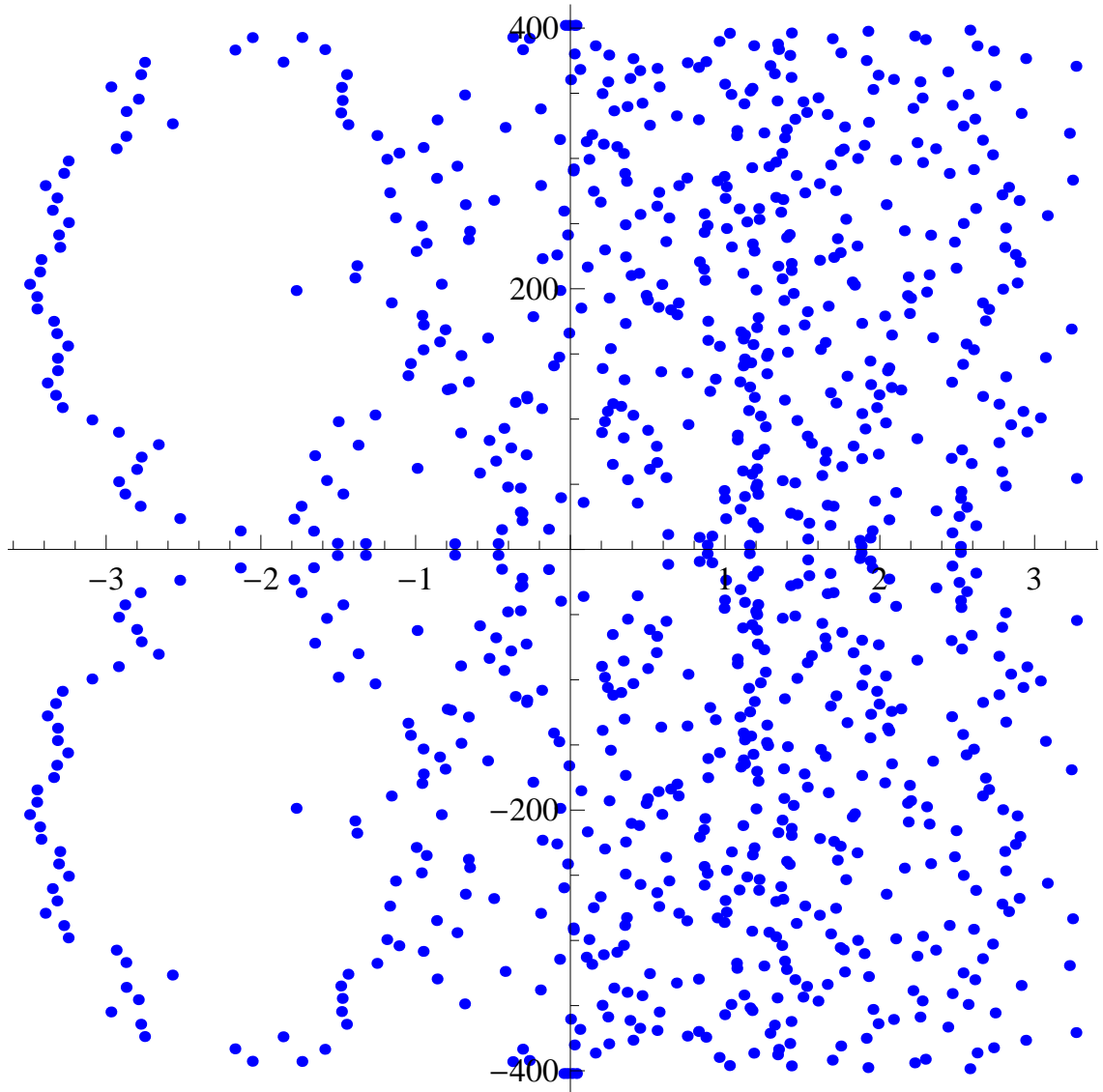


Figure 2.6: The roots of the HM partition for $D = 2$ and $V = 2^4$ in the $\log(x) = \beta/128$ plane.

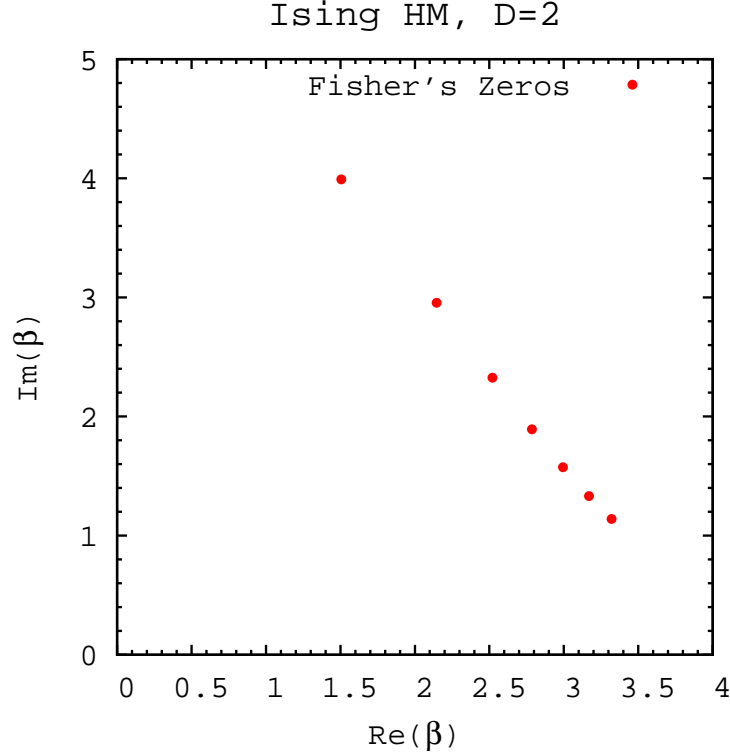


Figure 2.7: The Fisher zeros for the two dimensional HM with different volume $V = 2^N$; from up to down: $N = 2, 3, 4, 5, 6, 7, 8$.

The partition function has infinite number of complex zeros due to the periodicity of the exponential function. Here we are only interested in the zero that is closest to the real axis, or the “lowest” zero in the point of view of the absolute value. Because only through these zeros we can decide the order of the phase transition. For $D = 2$, the lowest Fisher zeros get closer and closer to the real axis as the volume increases and at the same time, they move to the right with a slower rate. There is no tendency for the zeros to pinch the real axis. In figure (2.8), we plot the real and imaginary part of the lowest zeros with the same data as in Fig. (2.7).

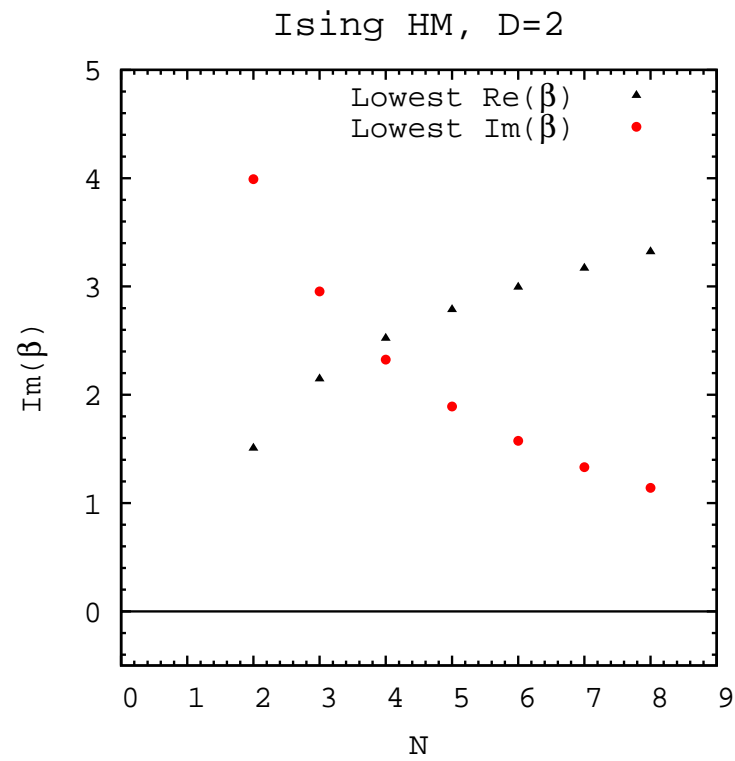


Figure 2.8: The real and imaginary part of the Fisher zeros for the two dimensional HM with different volume $V = 2^N$.

2.2.7 $D = 3$

For $D = 3$ case, the system has a second order phase transition. The second derivative of the free energy diverges at the some particular temperature (or inverse temperature β) in the thermodynamic limit. This temperature is called the critical temperature. The partition function must be zero at the critical temperature. We know that the critical β for the three dimensional HM is around 1.179 [118]. In figure (2.9), we can see that the lowest Fisher zeros really accumulate around 1.179 for the large volume and they move faster to the real axis than the two dimensional case. Another interesting phenomena can be observed if we compare Fig. 2.7 with Fig. 2.9 is that the lowest zeros go all the way to the right for the two dimensional case but they curve back to the critical β for the three dimensional case. That is a characteristic to distinguish the order of the phase transition. That means that we might only need to calculate the lowest Fisher zeros for several small volumes and “guess” the order of the phase transition by looking at the tendency of the zeros to move. Figure 2.10 again is the real and imaginary part of the lowest zeros with the same data as in Fig. 2.9.

2.2.8 Finite Size Scaling

Up to now, we only have Fisher zeros up the volume $V = 2^{11}$, which is still a small system compared to the Avogadro constant 2^{23} . For a system with a second order phase transition, the phase transition only occurs in the thermodynamic limit, in which the volume(V) and the number of particles(N) are infinite while the density $\rho = V/N$ stays finite [124, 125]. For the finite volume system, the intensive quantities are continuous functions of the temperature(T), magnetic field(H), etc. That means that there is no true critical point for a finite system. While in the

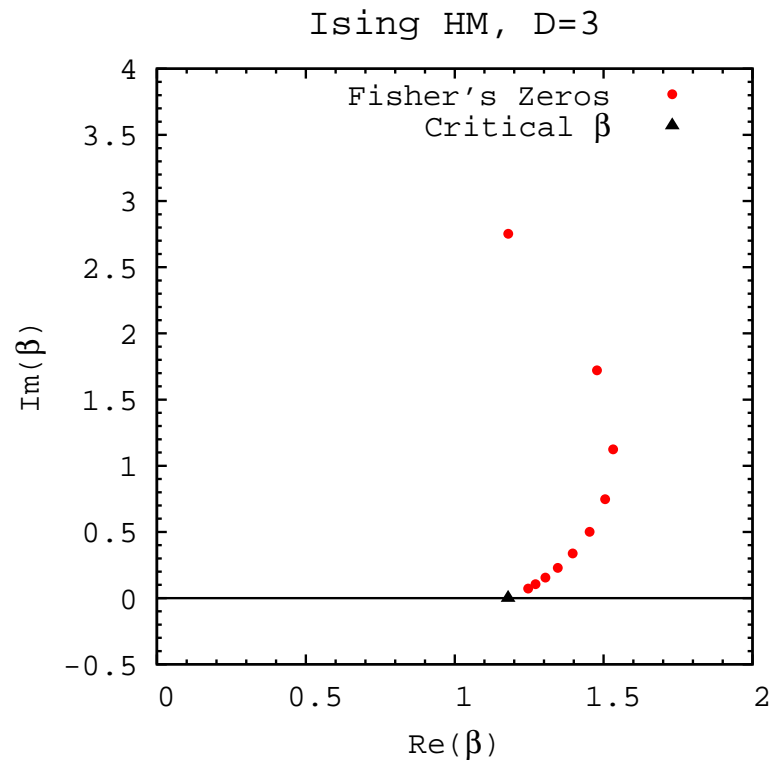


Figure 2.9: The Fisher zeros and the critical β for the 3 dimensional HM with different volume $V = 2^N$; from up to down: $N=2, 3, 4, 5, 6, 7, 8, 9, 10, 11$.

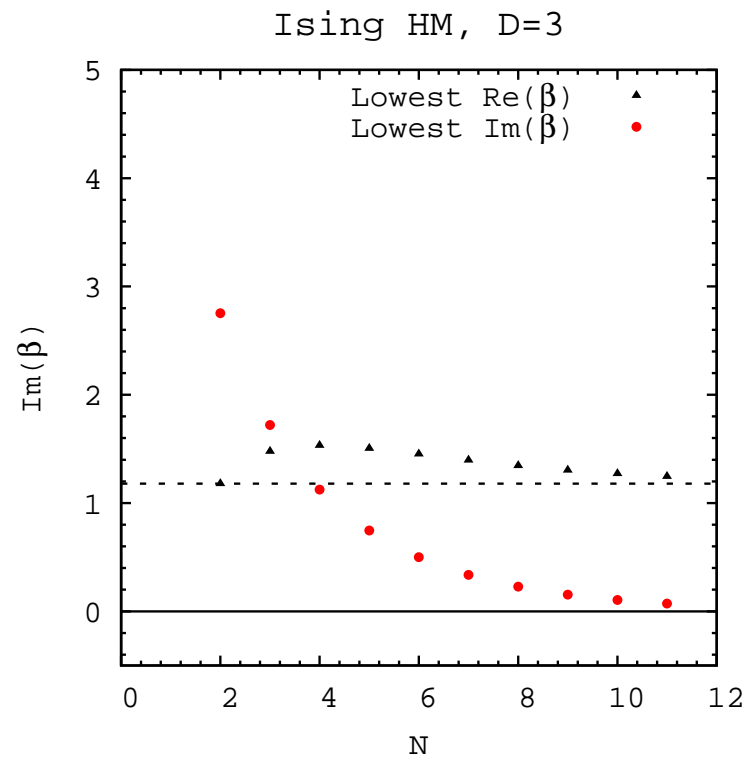


Figure 2.10: The real and imaginary part of the Fisher zeros for the 3 dimensional HM with different volume $V = 2^N$.

thermodynamic limit, the divergence may occur. In order to obtain the thermodynamic limit information from the finite volume system, we will need finite size scaling (FSS) technique.

In finite size system, there are three length scales: a , L and ξ . Near the critical point, a dependence is very weak. The thermodynamic quantities depend on $\frac{\xi}{L}$. We can treat L^{-1} as a scaling variable, with eigenvalue $y = 1$. The FSS assumes that: $\langle O \rangle_L = L^{\alpha/\nu} \phi(L^{\frac{1}{\nu}} t)$ if $\langle O \rangle_\infty(t) \propto |t|^{-\alpha}$ when $t \rightarrow \infty$, where $t = (T - T_c)/T_c$. The above relation can be proved by using renormalization group (RG) transformation.

Itzykson et al. [74] pointed out that the leading zeros ($\beta_1(L)$) of the partition function obey the finite size scaling relation. One could extract the critical exponents from the leading zeros. More precisely, one would have

$$\beta_1(L) - \beta_c \approx AL^{-\frac{1}{\nu}}, \quad (2.46)$$

where $L = V^{\frac{1}{D}}$ is the linear size of the system and ν is the critical exponent associated with the correlation length. We note that L here might not be integers in the system studied, although the total number of the sites V is always integer. For example, we could have $V = 2^2$, $D = 3$ and $L = V^{\frac{1}{D}} = 4^{\frac{1}{3}}$. As mentioned above, the critical temperature β_c has been known for a while [118]: $\beta_c = 1.17903017 \dots$ and in principle could be calculated to any significant digits. If the relation 2.46 holds, it is straight forward to get the critical exponent ν by performing a linear fit of the leading zeros (See Fig. 2.11). Comparing with the known value $\nu = 0.64957$ [126, 127], the critical exponent we extracted from the Fisher zeros is 1.3% larger. It has been suggested that the scaling of the imaginary part of the Fisher zeros corresponds to the critical exponent ν and the real part of the lowest zeros corresponds to the shift exponent λ [128], where the shift exponent is defined as $|\beta_L - \beta_c| \approx L^{-\lambda}$. However, we found out that the distance scaling, where the

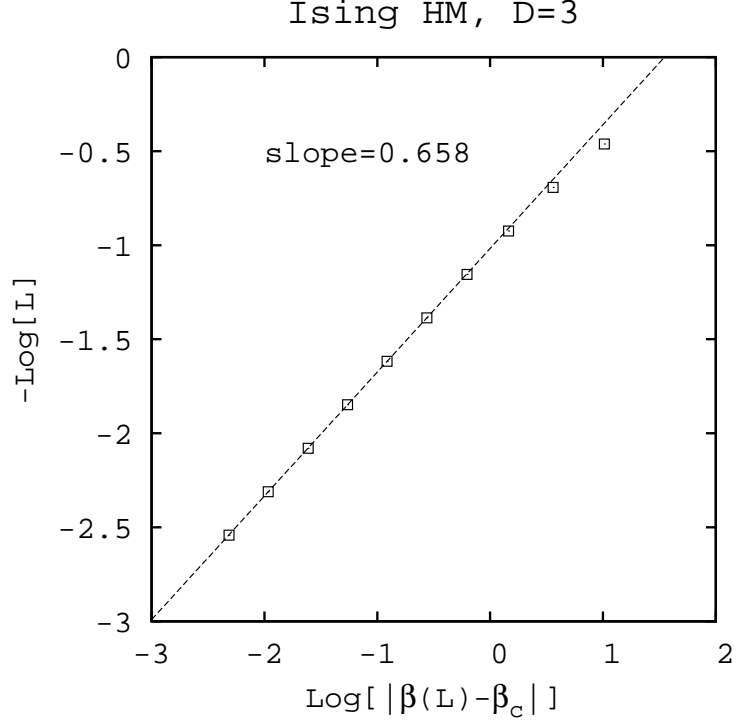


Figure 2.11: $-\ln L$ versus $\ln|\beta(L) - \beta_c|$ for $n = 2 - 11$; the fitting was done with $n = 8 - 11$.

information of both the real and imaginary part of the zeros is used, is optimum for the extraction of the critical exponent ν . See Fig. 2.12 and 2.13 for the scaling of the imaginary and real part of the zeros. We notice that the exponent extracted from the imaginary part of the zeros is smaller than ν while the corresponding real part is larger.

Following Itzykson et al. [74], we could also build up the relationship between the zeros of the partition function and the sub-leading critical exponent ω . We know that the zeros of the partition function is the same as the zeros of the scaling function Q

$$Z_{1/\lambda}(K_1 L^{y_1} \lambda^{y_1}, \dots) = Q(x_1, x_2, \dots), \quad (2.47)$$

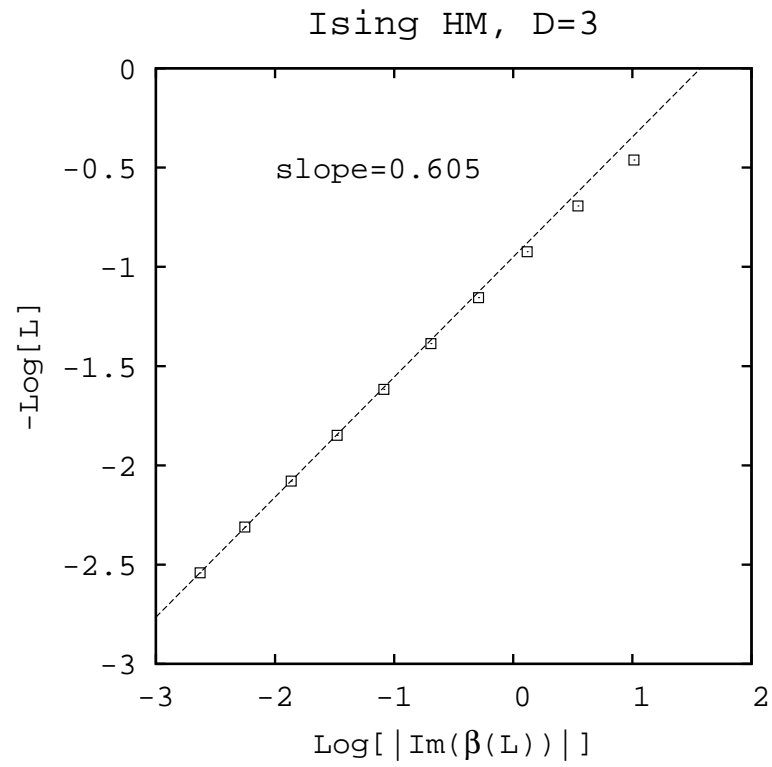


Figure 2.12: $-\ln L$ versus $\ln|\text{Im}\beta(L)|$ for $n = 2 - 11$; the fitting was done with $n = 8 - 11$.

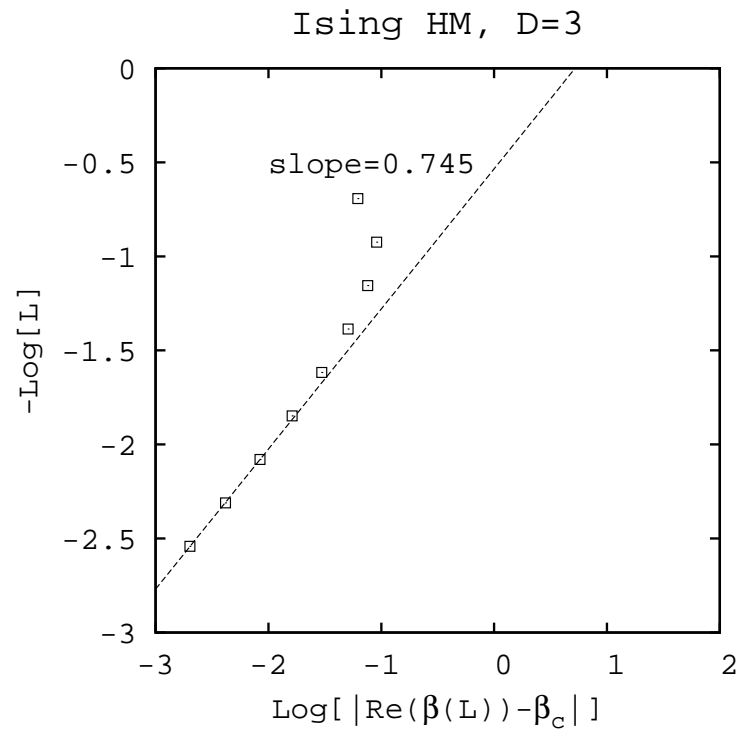


Figure 2.13: $-\ln L$ versus $\ln|\text{Re}\beta(L) - \beta_c|$ for $n = 3 - 11$; the fitting was done with $n = 8 - 11$.

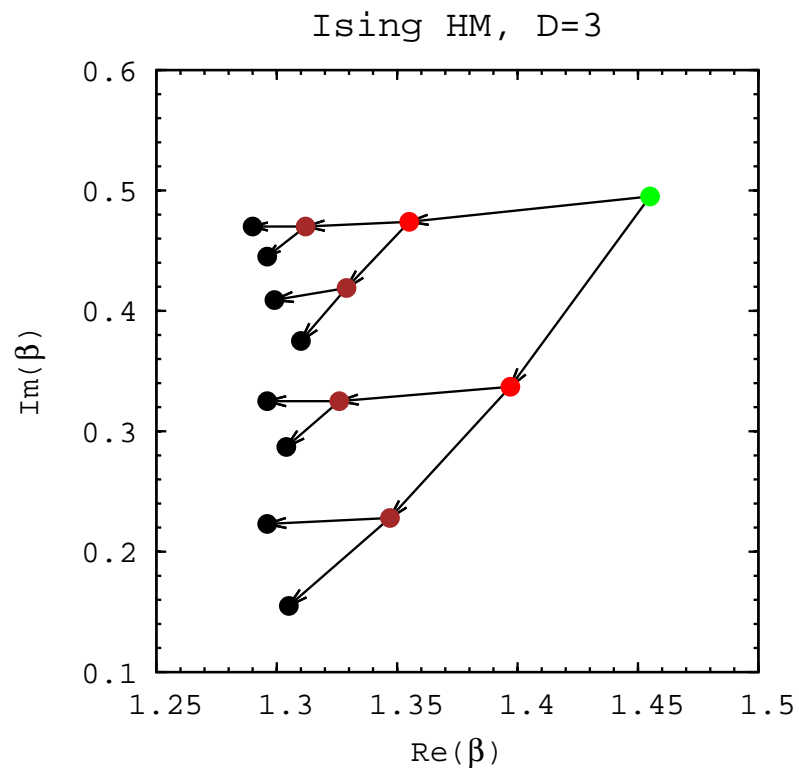


Figure 2.14: The density of the zeros is proportional to $\ln V$.

where $\lambda = L/b$ and $x_i = K_i L^{y_i}$ are the scaling variables. Here we will consider the zero field case ($h = 0$). In this case, if we consider the leading thermal scaling variable and the sub-leading variable, we will have the following scaling relation for the zeros of the Q

$$Q(x_t, x_\omega) = Q(KL^{1/\nu}, UL^{-\omega}), \quad (2.48)$$

where $K = \beta - \beta_c$ and ω corresponds to the exponent of the sub-leading correction. We further assume that we could solve the above equation and get

$$KL^{1/\nu} = f^{-1}(UL^{-\omega}), \quad (2.49)$$

If we expand the right hand side of the equation 2.49 and only keep the linear term in $L^{-\omega}$, we will get

$$KL^{1/\nu} = (\beta - \beta_c)L^{1/\nu} \quad (2.50)$$

$$= f^{-1}(UL^{-\omega}) \quad (2.51)$$

$$= A + BL^{-\omega} + O(L^{-2\omega}), \quad (2.52)$$

where A and B are in general both complex numbers. Notice that we have absorbed the coefficient U into B .

From equation 2.52, we can get the scaling relation of both real and imaginary part of the partition function zeros

$$Re(\beta(L)) - \beta_c = Re(A)L^{-1/\nu} + Re(B)L^{-1/\nu-\omega}, \quad (2.53)$$

$$Im(\beta(L)) = Im(A)L^{-1/\nu} + Im(B)L^{-1/\nu-\omega}. \quad (2.54)$$

We have the values for β_c , ν , and ω already so that we could use these values to test the above scaling relations. In figure 2.15, the real and imaginary part of the zeros with volume ranging from $V = 2^2$ to $V = 2^{11}$ are shown along with the fitting curves based on the equation 2.53 and 2.54.

The fittings agree with the data except for the smallest volume. This opens

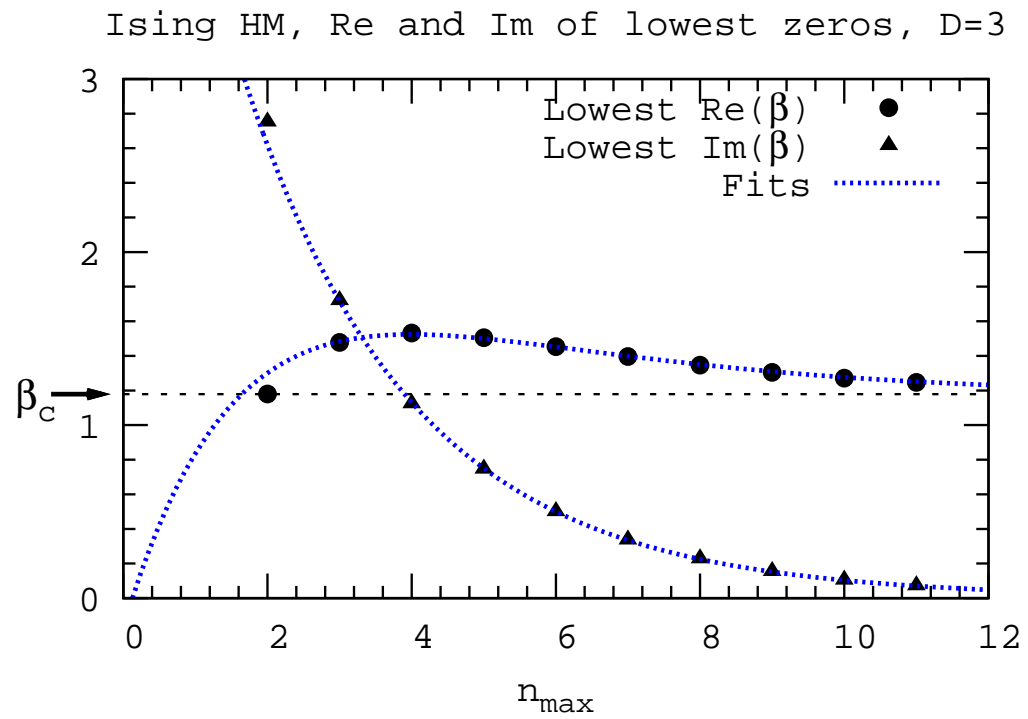


Figure 2.15: The real and imaginary parts of the Fisher zeros for $n=2-11$.

up the possibility of extracting the sub-leading exponent from the Fisher zeros.

2.2.9 From Discrete To Continuous RG transformation

Up to now, we only considered $b^D = 2$ case. Actually, we are also interested in the limit $b \rightarrow 1$ for the following reasons. In recent years, the local potential approximation (LPA) has been widely used in the context of Exact Renormalization Group Equations (ERGE), see next section, and has generated a lot of interest. We recommend references [129, 130] for reviews of the recent progress. ERGE allow in principle the study of global aspects of the RG flows of field theoretical or statistical models. However, truncation methods are necessary in order to make practical calculations. One particularly popular choice is to combine Polchinski's ERGE [131] with the LPA [132]. This results in a simple partial differential equation (called the “Polchinski's equation” below) for the effective potential. Polchinski's equation can also be obtained [133] as an infinitesimal version of the RG equation for the HM. This suggests that the linearized theories should be close to each other. Many accurate calculations of the critical exponents [134, 135, 136, 137, 138] show that the exponents differ only in the fifth significant digit. This small difference is significant. We find that the critical exponents of the HM depends on the blocking scheme. There is also a logarithmic term preventing one going from discrete RG transformation to the continuous one. These will be shown in section 2.2.10 and 2.2.11.

For $b = 2$ and $D = 3$, the value $\gamma = 1.30033\dots$ was obtained numerically in Ref. [139]. This value is significantly different from the value $\gamma = 1.2991407\dots$ obtained in references [126, 127] for $b = 2^{1/3}$ and $D = 3$. For $D = 3$, the limit $b \rightarrow 1$ was studied in Ref. [140] with the result $\gamma = 1.299124$. The difference in the fifth digit is significant and was confirmed by new calculations [138]. This shows that again different values of b correspond to different classes of universality and also

that for a given $b \neq 2^{1/D}$, the extensions given in equations are in-equivalent. In addition we see that the b dependence is much weaker and the slope is apparently opposite to the slope found analytically. We calculated the critical exponent γ for integer b^D up to 8. We try to extend the calculation to non-integer b^D . However, we found there is a stability window where everything converges only within some window of l_{\max} (see section 2.2.11 for definition of l_{\max}). This prevents us from making connections between discrete block spinning(integer b^D) transformation and the continuous one($b^D \rightarrow 1$).

2.2.10 Linear Analysis

2.2.10.1 Fixed Point Equation

Usually we normalize the recursion formula (2.38) by requiring the constant term in $R_n(k)$ to be 1 in order for $R_n(k)$ to have a direct probabilistic interpretation. In this case, the fixed point equation is :

$$R^*(k) = C_{n+1} e^{-\frac{1}{2}\beta_c \frac{\partial^2}{\partial k^2}} \left(R^*(b^{-(D+2)/2} k) \right)^{b^D}. \quad (2.55)$$

We can get the stability matrix L by linearizing the RG transformation right about the fixed point R^* . In the above case, we will only get one eigenvalue that is larger than 1 which corresponds to the relevant direction. The critical exponent γ can be obtained from $\gamma = \ln(2/c)/\ln\lambda_1$, where λ_1 is the largest eigenvalue.

However, we can also absorb the normalization coefficient into the $R^*(k)$ by multiplying both sides of (2.55) by $C_{n+1}^{1/(b^D-1)}$. The new fixed point equation becomes

$$R_0^*(k) = e^{-\frac{\beta_c}{2} \frac{\partial^2}{\partial k^2}} \left(R_0^*(b^{-(D+2)/2} k) \right)^{b^D}. \quad (2.56)$$

The corresponding polynomial expansion becomes

$$R_n(k) = a_{n,0} + a_{n,1}k^2 + a_{n,2}k^4 + \dots + a_{n,l_{\max}}k^{2l_{\max}}, \quad (2.57)$$

or equivalently (only even power of k is involved due to $Z(2)$ symmetry of the ϕ

field),

$$R_n(u) = a_{n,0} + a_{n,1}u + a_{n,2}u^2 + \dots + a_{n,l_{max}}u^{l_{max}}. \quad (2.58)$$

The $_0$ under the R^* in (2.56) means the 0th order in the following small ζ expansion about integer b^D . If we use the above unnormalized fixed point to get the stability matrix, we will get two eigenvalues larger than 1. The first one is just $\lambda_0 = b^D$. The reason we use the unnormalized fixed point equation is that we do not need to expand the coefficient C_{n+1} when we do the small ζ expansion which considerably simplifies the calculation.

2.2.10.2 Stability Matrix

Let us make a small variation δR around fixed point R^*

$$R_0^* + \delta R' = RG_0[R_0^* + \delta R] = R_0^* + L[\delta R], \quad (2.59)$$

where RG_0 stands for the 0th order RG transformation. If we only keep the first order in δR , we will get the stability matrix L

$$L \approx e^{-\frac{1}{2}\beta_c \frac{\partial^2}{\partial k^2}} [2R_0^*(b^{-(D+2)/2} k)] \quad (2.60)$$

In general, the stability matrix is not symmetric and involves very large and very small numbers. Fig.2.16 shows a typical stability matrix. The matrix elements range from 10^{-109} to 10^{+42} for $l_{max} = 40$. That requires us to use very high precision for the calculations. The non-symmetric property also requires the consideration of the left and right eigenvectors separately (see section 2.2.10.3 for details).

For the special $b^D = 2$ case, the stability matrix can be calculated in the following way

$$\delta a_{n+1,\sigma} = \sum_{\nu=0}^{l_{max}} L_{\sigma,\nu} \delta a_{n,\nu} \quad (2.61)$$

$$L_{\sigma,\nu} = \frac{\partial a_{n+1,\sigma}}{\partial a_{n,\nu}} \quad (2.62)$$

$$a_{n+1,\sigma} = \Gamma_{\sigma}^{\mu\nu} a_{n,\mu} a_{n,\nu} \quad (2.63)$$

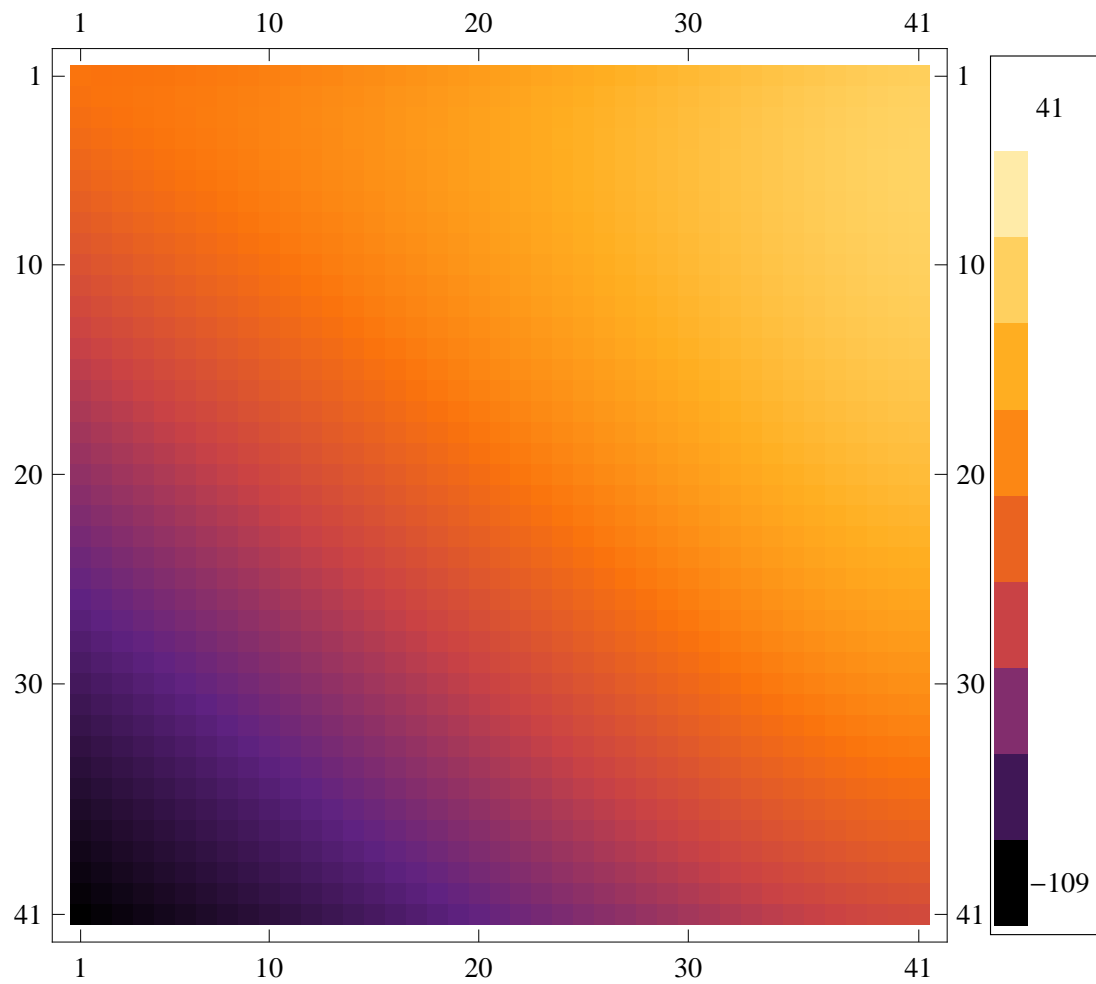


Figure 2.16: Stability matrix for $l_{\max} = 40$.

$$\Gamma_{\sigma}^{\mu\nu} = (c/4)^{\mu+\nu} \frac{(-1/2)^{\mu+\nu-\sigma} (2(\mu+\nu))!}{(\mu+\nu-\sigma)!(2\sigma)!}. \quad (2.64)$$

2.2.10.3 Small ζ Expansion

In order to investigate the non-integer b^D behavior of the recursion formula, we expand b^D right around integer values. This is similar to the ϵ expansion extensively used in statistical mechanics and quantum field theory. Often, people expand dimension D around integer values, usually 4 or 2. It has been proved that this theoretical method is quite useful although there is no significant physical meaning. In the following, we will show that we can do a similar expansion for b^D in our system instead of D . We will fix the dimension of the system to be $D = 3$. The physical meaning of b^D in our system is the number of sites blocked in each RG transformation. For example, for the system with $b^D = 2$ and $D = 3$, we block 2 sites in each iteration. One should notice that the “number of sites” blocked in each direction is $2^{1/3}$. Blocking non-integer number of sites seems impossible in the position space. However, this is very natural if we consider it in the k space. One should also notice that we not only change the power in the formula (2.56) but also the rescaling factor when we do the b^D expansion

$$b^D = 2 + \zeta, \quad (2.65)$$

$$\frac{c}{4} = b^{-(D+2)} = \frac{c_0}{4} \left(1 - \frac{5}{6}\zeta\right), \quad (2.66)$$

where c_0 corresponds to the $\zeta = 0$ case. We suppose the first order correction to the fixed point R_0^* is R_1^* . The new fixed point equation to the first order in ζ becomes

$$RG_{\zeta}[R_0^* + \zeta R_1^*] = R_0^* + \zeta R_1^*. \quad (2.67)$$

We can again make a small variation around the new fixed point (set $\delta R = -\zeta R_1^*$ in equation 2.59)

$$RG_{\zeta}[R_0^*] = R_0^* + \zeta R_1^* - \zeta L'[R_1^*]. \quad (2.68)$$

In the limit $\zeta \rightarrow 0$, L' approaches L ,

$$\frac{RG_\zeta[R_0^*] - R_0^*}{\zeta} \rightarrow R_1^* - L[R_1^*]. \quad (2.69)$$

On the other hand, if we expand the left hand side of equation (2.67) to first order in ζ and compare with the right hand side, we obtain

$$R_1^* = L[R_1^* + G], \quad (2.70)$$

where

$$G = -\frac{5}{6}k^2 \frac{\partial R_0^*}{\partial k^2} + \frac{1}{2}R_0^* \ln R_0^*. \quad (2.71)$$

From (2.69) and (2.70), we obtain

$$\frac{RG_\zeta[R_0^*] - R_0^*}{\zeta} \rightarrow L[G] \quad (2.72)$$

in the $\zeta \rightarrow 0$ limit. The above relation has been tested numerically for $b^D = 2$ with different l_{\max} .

2.2.10.4 Linearization

If we can construct R_1^* numerically, we can construct the fixed point for non-integer b^D from R_0^* , which interpolate among the fixed points on Fig. 2.17 for $b^D = 2, \dots, 8$. One might have already noticed that everything we have done is very similar to the variational method used in Quantum Mechanics. The only difference is that we are dealing with non-Hermitian matrix here. Actually the matrix L is not symmetric (see Fig. 2.16). For the non symmetric matrix, the left and right eigenvectors are not each others transpose. In the following, we will choose the right eigenvectors as the basis to expand R_1^* and G . We will also normalize left and right eigenvectors individually. Suppose the right eigenvectors of L are v_n and the corresponding eigenvalues are λ_n . Then we can write

$$R_1^* = \sum \alpha_n v_n \quad (2.73)$$

$$G = \sum \beta_n v_n. \quad (2.74)$$

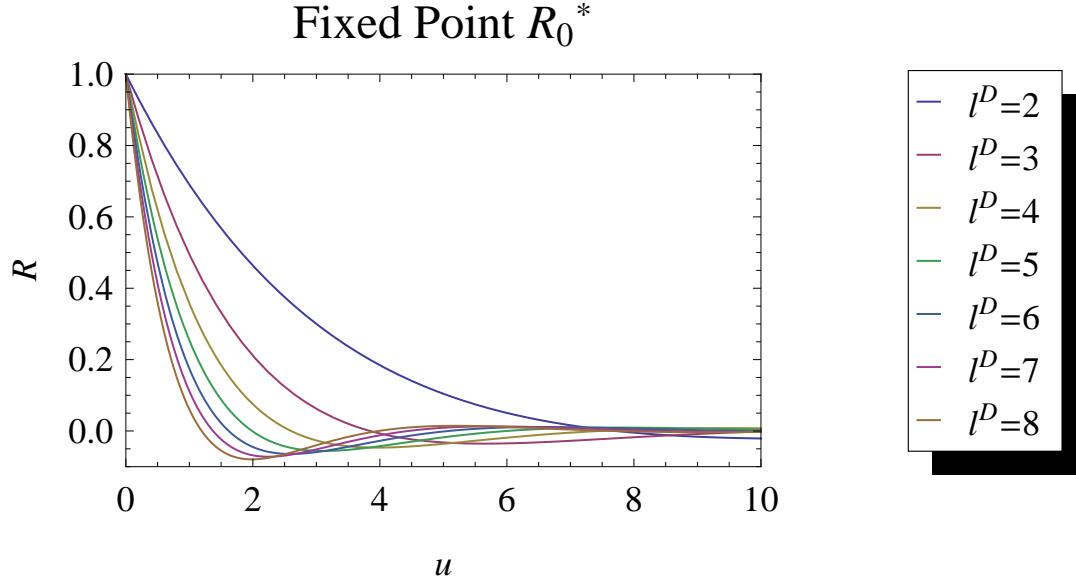


Figure 2.17: The fixed point polynomial R_0^* for different b^D .

By using the orthogonality relations among the left and right eigenvectors, we multiply the left eigenvectors on both sides of equation (2.70) and then normalize. We obtain

$$\alpha_n = \frac{\beta_n \lambda_n}{1 - \lambda_n}. \quad (2.75)$$

We have already have G (see equation (2.71)). We can easily get β_n by multiplying both sides of equation (2.74) by the corresponding left eigenvectors. Then $\alpha_n(R_1^*)$ is obtained from equation (2.75).

2.2.11 Window Of Stability

From Eq. 2.71 and 2.74, we obtain

$$G = -\frac{5}{6}k^2 \frac{\partial R_0^*}{\partial k^2} + \frac{1}{2}R_0^* \ln R_0^* = \sum \beta_n v_n. \quad (2.76)$$

Therefore,

$$\beta_n = \frac{\langle v_L | G \rangle}{\langle v_L | v_R \rangle}, \quad (2.77)$$

where v_{Ln} and v_{Rn} are the left and right eigenvectors of L .

In order for β_n to be stable with increasing l_{\max} (although there is no physical reason to require β_n to be stable, we should at least avoid singularity.), we require both $\langle v_L|G \rangle$ and $\langle v_L|v_R \rangle$ decrease or increase with the same rate as we increase l_{\max} . However, this is not the case. We observe that when we increase l_{\max} , $\langle v_L|G \rangle$ and $\langle v_L|v_R \rangle$ are only parallel within some range. There is a maximum l_{\max} where everything is normal within this range. If we further increase l_{\max} , $\langle v_L|G \rangle$ decreases much slower than $\langle v_L|v_R \rangle$. See Fig. 2.18 for details. That is to say, when we increase l_{\max} , which in principle is equivalent to getting better approximation to the original measure, we will inevitably get very large (if not infinity) β_n . That prevents us from getting accurate critical temperature and critical exponents. We also notice that the stability window is smaller with smaller b^D . The reason for the diverging property of β_n is that there is a logarithm term $\ln R_0^*$ in the expression G . The right eigenvectors are not good basis for this logarithm term.

In Fig. 2.19, we show the result β_c obtained from different b^D . The red dots correspond to the values for integer b^D . We can calculate them up to arbitrary precision, which is equivalent to say that we can use arbitrary l_{\max} to approximate the initial measure. The blue dots are for non-integer b^D . The results for non-integer b^D are unstable. That confirms our analysis above.

In Fig. 2.20, the critical exponent γ is showed with different b^D and l_{\max} . All the calculations were performed with maximum l_{\max} in the stability window. We can see that the stability window decreases with decreasing b^D . The γ we can get for non-integer b^D can not reach the accuracy acquired for our comparison with LPA (5th digit).

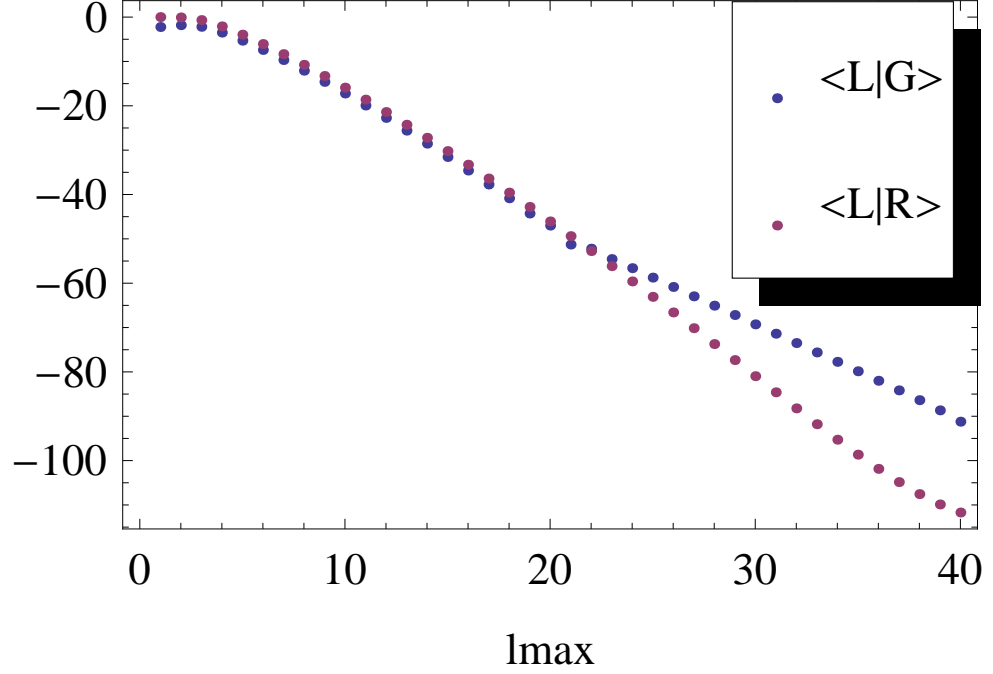


Figure 2.18: Red dots: $\langle L|R \rangle$; Blue dots: $\langle L|G \rangle$.

2.2.12 Two Lattice Matching

In this section, we will discuss how to construct the renormalization group (RG) flow in the complex β plane by using the two lattice matching method.

Now, let me briefly introduce the basic idea of the two lattice matching method. For simplicity, we assume there is only one relevant direction for the system studied. We start the RG transformation at one coupling β_1 near the phase transition, after several transformations, the system will go along the relevant direction and follow the flow thereafter. We can try to search for another coupling β_2 , where we only need to do one less time the transformation to reach the relevant direction and match the same point as the previous case in the parameter space. Suppose the scaling factor for the transformation is b , lattice spacing $a \rightarrow ba$, the

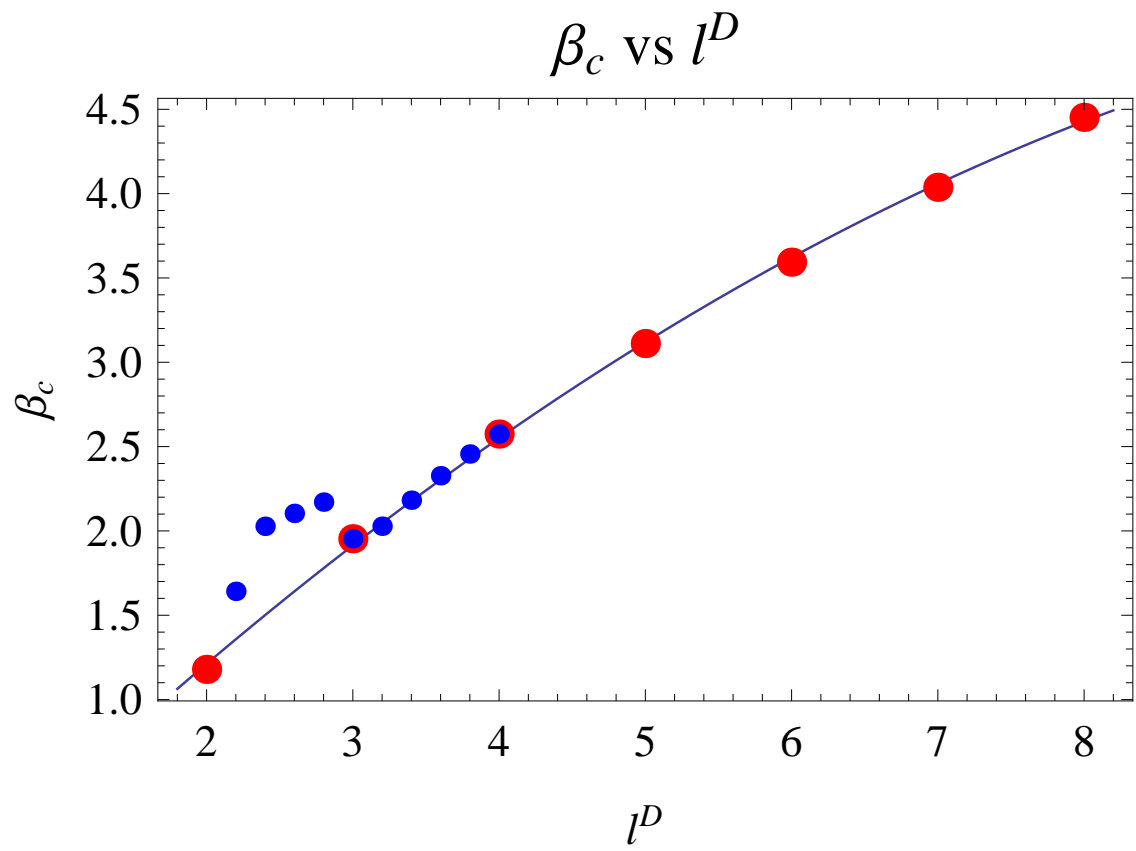


Figure 2.19: The critical temperature β_c changes with b^D .

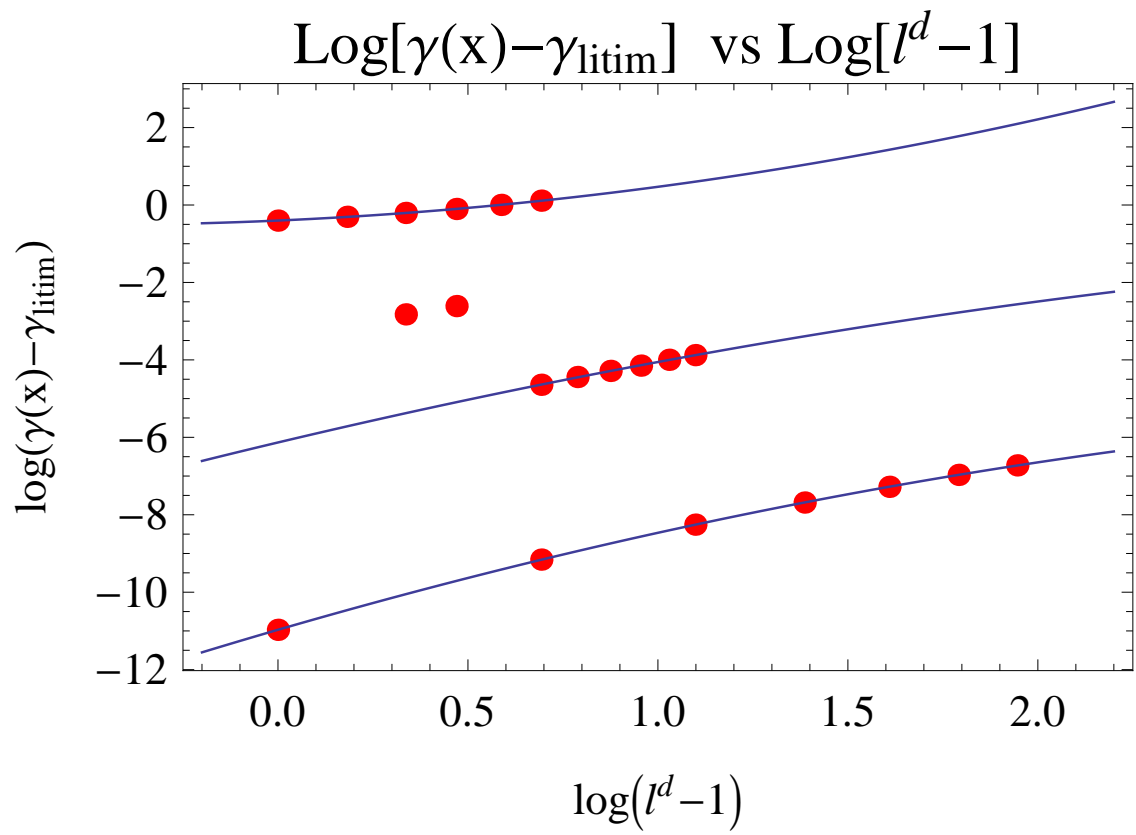


Figure 2.20: The critical exponent γ changes with b^D .

two β 's have the following relationship

$$\beta_2 - \beta_c = b^{\frac{1}{\nu}}(\beta_1 - \beta_c). \quad (2.78)$$

Usually, one will choose the scaling variable $b = 2$. In the hierarchical model described below, b can be non-integer, i.e., $2^{\frac{1}{3}}$. In practice, we will choose two systems with different volume, where the linear size of the two systems differ by a factor of b . We want these two systems describe the same long distance physics at β_1 and β_2 . For example, we can keep the correlation length ξ in physical units to be constant. We start with one system with volume L^D , lattice spacing a , and coupling β_1 . After n iteration, the lattice spacing changes from a to $b^n a$. The correlation length in lattice units changes from $\frac{\xi}{a}$ to $\frac{\xi}{b^n a}$. ξ is the physical length divided by a , $\xi = \frac{1}{ma}$ where m is a dimensionful physical mass. We can simulate another system with volume $\frac{L^D}{b}$, lattice spacing ba , and coupling β_2 . This time we iterate $n - 1$ times. The lattice spacing changes from ba to $b^{n-1}ba = b^n a$. The correlation length in the lattice unit changes from $\frac{\xi}{ba}$ to $\frac{\xi}{b^n a}$. They both end up with the same correlation length if we measure them in the lattice unit. We further assume they have the same action in the blocked lattice. Usually the blocked action will involve a large number of couplings and it is impractical to calculate it. However, we do not need to calculate it directly. We use the fact that when the action of the system is the same, all the observables should also be the same. We only need to measure a set of different observables. For details, please see [141].

The observable we use to match is the spin-spin correlation, which is a little different from that of Ref. [23, 25]. We will see it in the following that the observable $R(\beta, 2^N)$ defined in 2.79 does not depend on the lattice rescaling.

Suppose we have 2^N sites in the whole volume. We can split the system into two blocks B_1 and B_2 . Each of them has 2^{N-1} sites. Figure (2.21) is an illustration

for $D = 1$ case. The observable we considered is

$$R(\beta, \mathcal{V}/a^D) \equiv \frac{\left\langle (\sum_{x \in B_1} \phi_x)(\sum_{y \in B_2} \phi_y) \right\rangle_\beta}{\left\langle (\sum_{x \in B_1} \phi_x)(\sum_{y \in B_1} \phi_y) \right\rangle_\beta}, \quad (2.79)$$

where \mathcal{V} is the physical volume of the system and a is the lattice spacing in the physical unit (i.e., in the unit of meter, centimeter, etc.). The observable R has the following advantages: 1), it is relatively easy to calculate for the system considered; 2), it does not require to calculate the partition function; 3), we do not need to consider the field rescaling associated with the full RG transformation. Both 2) and 3) are due to the rational form of the observable R . The observable can be calculated both in ϕ space and k space. In k space, the model can be very easily extended to the $O(N)$ model with arbitrary N and arbitrary dimension D . This enables us to extend the technique developed here over a wide range of interesting questions.

In the following, we will give the details on how to calculate R . From the relationship between the measures in the successive hierarchy (see Equation 2.13), we can immediately get the following formula

$$\begin{aligned} & \left\langle \left(\sum_{x \in B_1} \phi_x \right) \left(\sum_{y \in B_2} \phi_y \right) \right\rangle_\beta \\ &= \frac{\int d\phi_1 d\phi_2 e^{\frac{\beta}{2}(\frac{\epsilon}{4})^N (\phi_1 + \phi_2)^2} \phi_1 \phi_2 W_{N-1}(\phi_1) W_{N-1}(\phi_2)}{Z_N}, \end{aligned} \quad (2.80)$$

and

$$\begin{aligned} & \left\langle \left(\sum_{x \in B_1} \phi_x \right) \left(\sum_{y \in B_1} \phi_y \right) \right\rangle_\beta \\ &= \frac{\int d\phi_1 d\phi_2 e^{\frac{\beta}{2}(\frac{\epsilon}{4})^N (\phi_1 + \phi_2)^2} \phi_1^2 W_{N-1}(\phi_1) W_{N-1}(\phi_2)}{Z_N}. \end{aligned} \quad (2.81)$$

As mentioned above, we do not need to calculate the partition function Z_N in the denominator to calculate the ratio in Eq. 2.79. Only the numerators matter. The local measure W_{N-1} can be constructed easily from W_0 and hence the

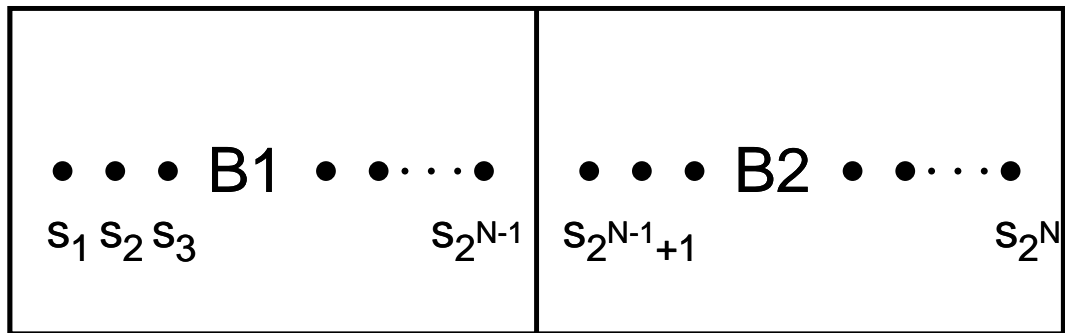


Figure 2.21: An example of blocking for the one dimensional hierarchical model.

observable R .

In the k space, the calculation is a little tricky. We notice that the calculation of $\left\langle (\sum_{x \in B_1} \phi_x)(\sum_{y \in B_2} \phi_y) \right\rangle_\beta$ is similar to that of the partition function except that we need to replace $W(\phi)$ by $\phi W(\phi)$ or $\phi^2 W(\phi)$.

$$\begin{aligned} \phi W(\phi) &= \phi \int \frac{dk}{2\pi} e^{ik\phi} R(k) \\ &= \int \frac{dk}{2\pi} (-i \frac{\partial}{\partial k} e^{ik\phi}) R(k) \\ &= \int \frac{dk}{2\pi} e^{ik\phi} (i \frac{\partial}{\partial k} R(k)) + \int \frac{dk}{2\pi} (-i \frac{\partial}{\partial k} (e^{ik\phi} R(k))) \end{aligned} \quad (2.82)$$

where the last term can be set to 0 if $R(k)$ decays fast enough at $\pm\infty$. We can perform the same trick to the $\phi^2 W(\phi)$ term. After several lines of simple algebra, we will get

$$R(\beta, 2^N) = \frac{e^{-\frac{\beta}{2}(\frac{c}{4})^N \nabla_k^2} (-(R'_{N-1}(k))^2)}{e^{-\frac{\beta}{2}(\frac{c}{4})^N \nabla_k^2} (-R''_{N-1}(k) R_{N-1}(k))} \quad (2.83)$$

In practical calculations, we expand the operator $e^{-\frac{\beta}{2}(\frac{c}{4})^N \nabla_k^2}$ up to certain orders and approximate the local measure by polynomials. We need to increase the order of the polynomial until the result of the observable stabilizes.

For the $O(N)$ model, we only need to change the initial measure $W_0(\phi)$ from $\delta(\phi^2 - 1)$ to $\delta(\phi^2 - 1)$. We have already mentioned that changing the dimensionality D is equivalent to changing the strength parameter c : $c/4 = b^{-2-D}$.

Having figured out how to calculate the observable, we use the two lattice matching method to construct the RG flows. The flows are constructed by the following procedure. We first choose an initial β_0 and then calculate the observable at β_0 with the larger volume, say 2^5 . We then use the Newton's method or the "logarithmic residue" method to find the corresponding β_1 which gives the same observable at smaller volume, say 2^4 . We can further use β_1 to the larger volume and try to match another β_2 to the smaller volume \dots . By doing this, we can get a sequence of β s starting from β_0 and ending to β_n . By choosing different initial

starting points, we can get different flow lines(connecting β_i and β_{i+1} with arrows). Figure (2.22) shows several flow lines starting from $\text{Re}\beta = 5$ and different $\text{Im}\beta$. We can see clearly that all the flows stay in the boundary formed by the Fisher zeros and they also all flow to the trivial strong coupling fixed point.

The above matching procedure can be written as

$$R(\beta, 2^5) = R(\beta', 2^4). \quad (2.84)$$

We should notice that there are usually more than one β' satisfying the matching condition 2.86 for one single β . This can be easily seen from the case we will discuss in the following.

2.2.12.1 $D = 2$

For $D = 2$ and $b^D = 2$, we have $c = 1$. In this case, all the λ_k in equation (2.39) are non-negative rational numbers. In general, the least common multiple of the denominator of λ_k is $V^2/2$, where V is the volume of the system. The matching condition 2.86 also has the similar property. This nice property allows us to write the matching condition as a polynomial equation by changing the variable β' to $V^2/2 \log[n]$. Searching for the matching point β' reduces to solving for the roots of the polynomial equation. The number of the solutions depends on the order of the polynomial. That explains why there are more than one β' for a fixed β . In order to get unambiguous flows, we need to develop some strategy to pick the “right” matching β' . A natural way is to compare the distance between β and the β' and choose the closest one to the β as the matching point. We found out that most of the time it is easy to determine the closest β' because the other matching candidates are all very far away from the initial β . In order to quantify the ambiguity of choosing the closest matching β' , we define the following factor

$$f(\beta) = \log |\beta - \beta'_2| - \log |\beta - \beta'_1|, \quad (2.85)$$

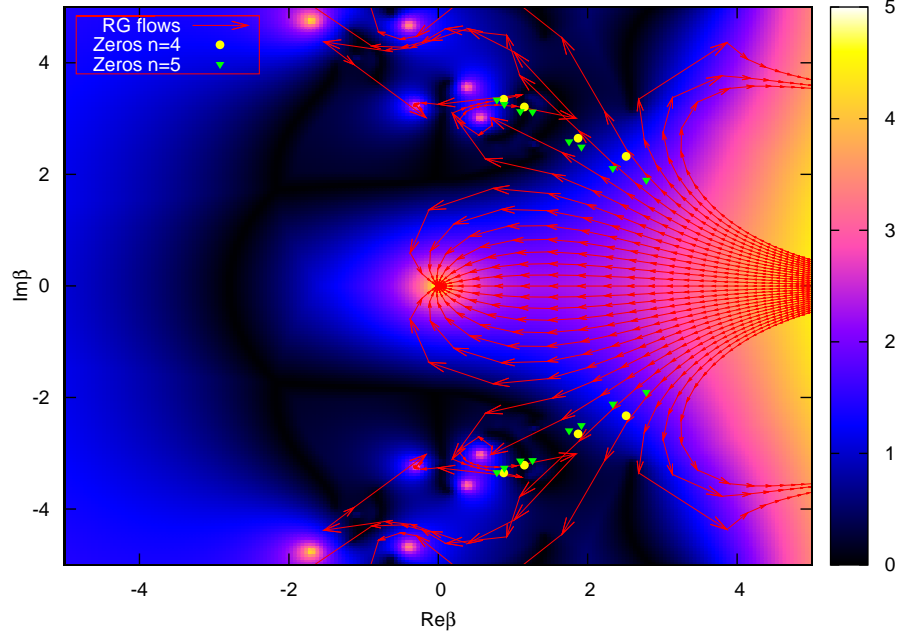


Figure 2.22: RG flows obtained by the two lattice matching methods for hierarchical Model.

where β'_1 and β'_2 are the closest and next closest β' to β . If $f(\beta)$ is very close to zero, β'_1 and β'_2 are very close to each other. That means that it is more ambiguous to choose the closest β'_1 as the matching point. On the other hand, it is less ambiguous if $f(\beta)$ is large. In figure 2.22, we make a contour plot of $f(\beta)$. The darker the color, the closer the β'_1 to β'_2 , in another word, the more ambiguous to pick β' . Notice that the flows near the Fisher zeros are more ambiguous and unpredictable.

2.2.12.2 $D = 3$

As stated in the previous section, there is a phase transition for the $D = 3$ case and the critical β is around 1.179 in the infinite volume limit. In Figure 2.23, we can clearly see that the flow goes from the nontrivial fixed point to the trivial fixed points 0 and ∞ .

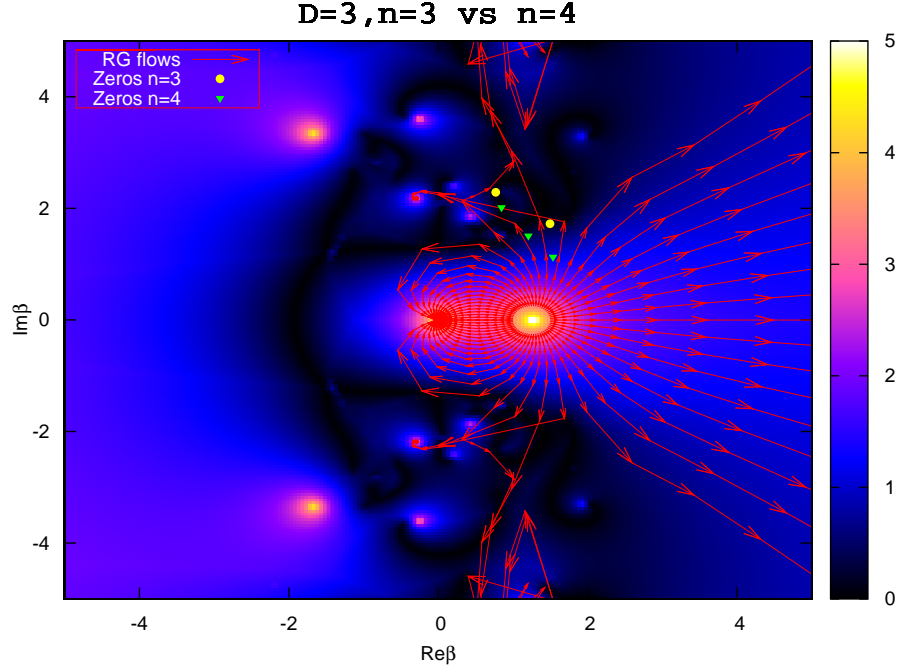


Figure 2.23: Unambiguous RG flows for the hierarchical model in the complex β plane obtained by the two lattice method.

One of the motivations of doing the two lattice matching for the hierarchical model was to construct the RG flow in the complex parameter plane and find connections between the flow and the Fisher zeros. One can further construct the β function, which is similar to the Callan-Symanzik β function used in the field theory language. We will discuss more about that in the lattice gauge theories chapters later.

The matching condition can be chosen as

$$R(\beta, L^D) = R(\beta', (L/b)^D). \quad (2.86)$$

We can then start from the bare value β_0 and get the new coupling from Eq. 2.86

$$\beta = \beta_0 \implies \beta'_1, \quad (2.87)$$

$$\beta = \beta'_1 \implies \beta''_1, \quad (2.88)$$

$$\vdots \tag{2.89}$$

The discrete β function is then defined as

$$\Delta\beta(\beta, L^D \rightarrow (L/b)^D) = \beta - \beta'. \tag{2.90}$$

In the following, we will consider the further generalized hierarchical model with $2^{n_{max}}$ lattice sites

$$H = -\frac{1}{2} \sum_{n=1}^{n_{max}} \left(\frac{c}{4}\right)^n f(n) \sum_{B^{(n)}} \left(\sum_{x \in B^{(n)}} \phi_x \right)^2. \tag{2.91}$$

All the other parameters are the same as the conventional one except that there is an extra $f(n)$ factor, which further controls the interaction strength. The generalized model has many nice properties and is an ideal laboratory to test various ideas before one applies them to the more complicated full QCD case. Here is a list of properties we used [82, 84, 142]

- For $D > 2$ and $f(m) = 1$, the model has a second order phase transition, which is similar to D=3 regular Ising model.
- For $D \leq 2$ and $f(m) = 1$, the model has no phase transition at finite temperature, which is different from D=2 regular Ising model.
- For $D = 2$ and $f(m) = \log(m)$, the model has Thouless effect, which is equivalent to Anderson model.

In the following, we will consider D=3, 2, $f(m) = 1$ and various dimensions for $f(m) = \log(m)$. One of the reasons to investigate systems with different parameters is to get better understanding on how infrared fixed point (IRFP) emerges, which plays a key role in understanding the techni-color related theories in the high energy physics field. Fixed points correspond to zeros of the β function. It has been pointed out that zeros of the β function can disappear in three ways as one or several

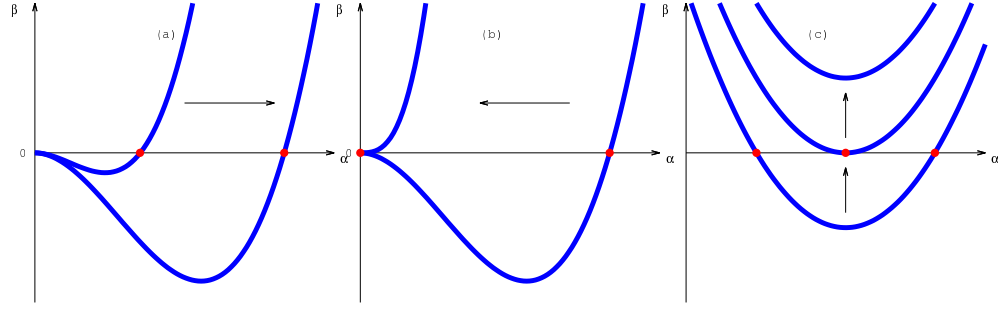


Figure 2.24: Three mechanisms for the loss of the fixed point(s) described in [2].

parameters change [2]. They are schematically described in Fig. 2.24. The first two ways are interchangeable depending on what parameters are used ($\alpha \leftrightarrow 1/\alpha$). For the third way, the fixed points that disappeared can be recovered in the complex parameter plane [143]. This third way can be easily seen for the hierarchical model by tuning the dimensionality D . Figure 2.27 shows discrete β functions for $f(n) = \log(n+1)$, $D = 1.9, 1.994$, and 2. The corresponding complex RG flows are shown in Fig. 2.28, 2.29, 2.30. Figure 2.31 shows that Fisher zeros appear to accumulate along lines that separate the flows ending on different fixed points.

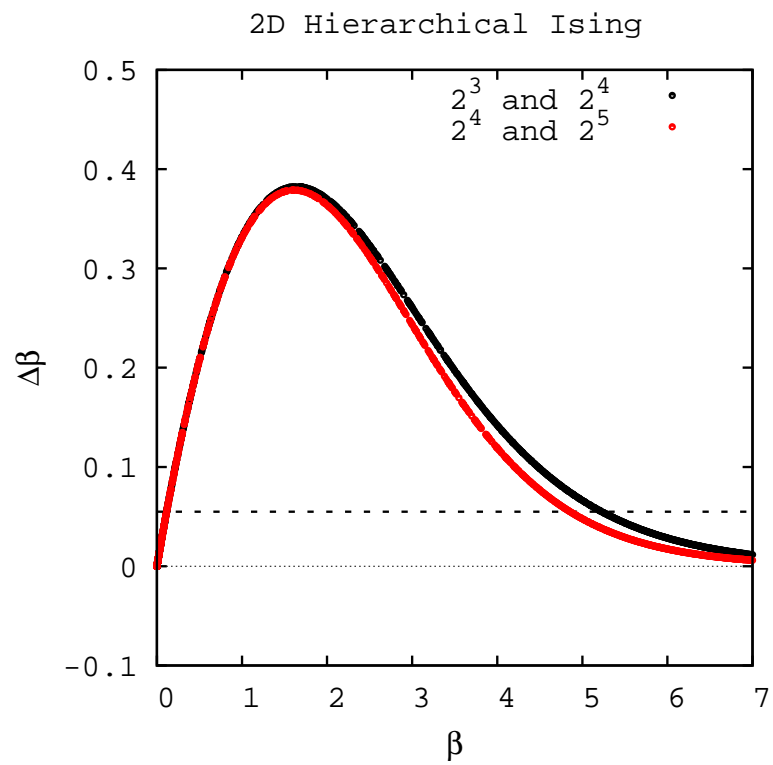


Figure 2.25: Discrete β function for the hierarchical model with $D = 2$ and $f(m) = 1$.

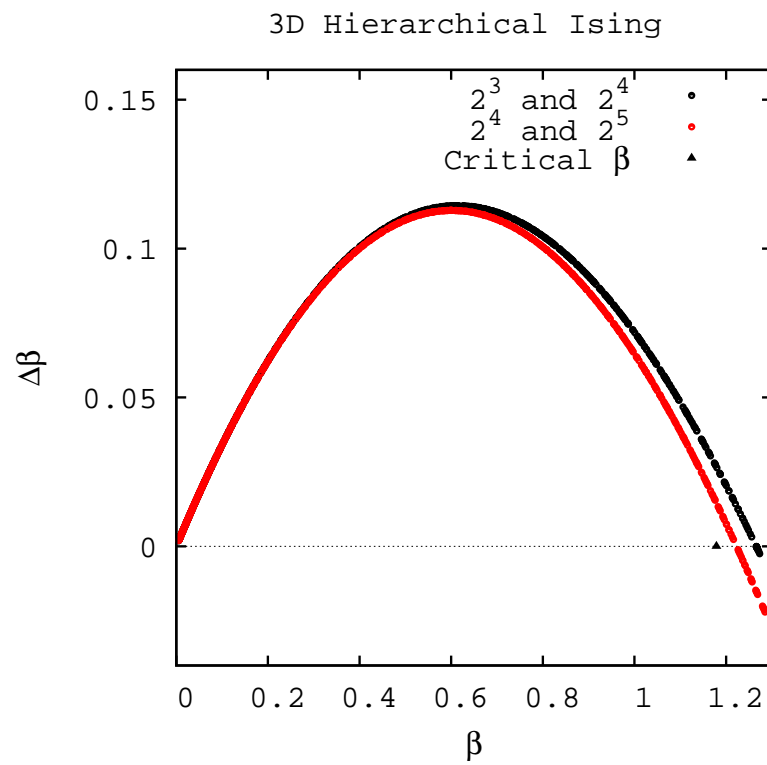


Figure 2.26: Discrete β function for the hierarchical model with $D = 3$ and $f(m) = 1$.

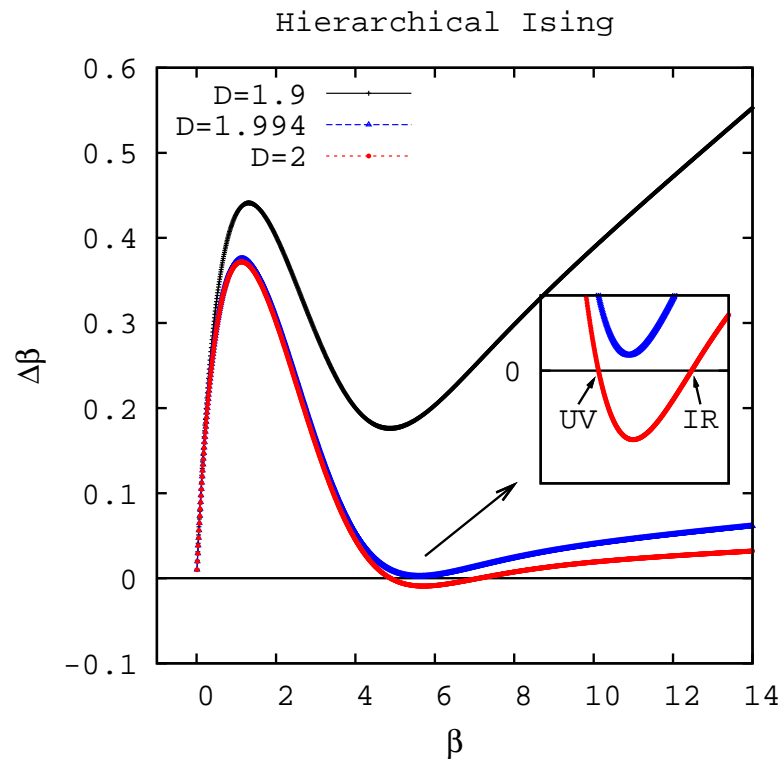


Figure 2.27: Discrete β function for the hierarchical model with $D = 1.9, 1.994$, and 2 and $f(m) = \log(m)$.

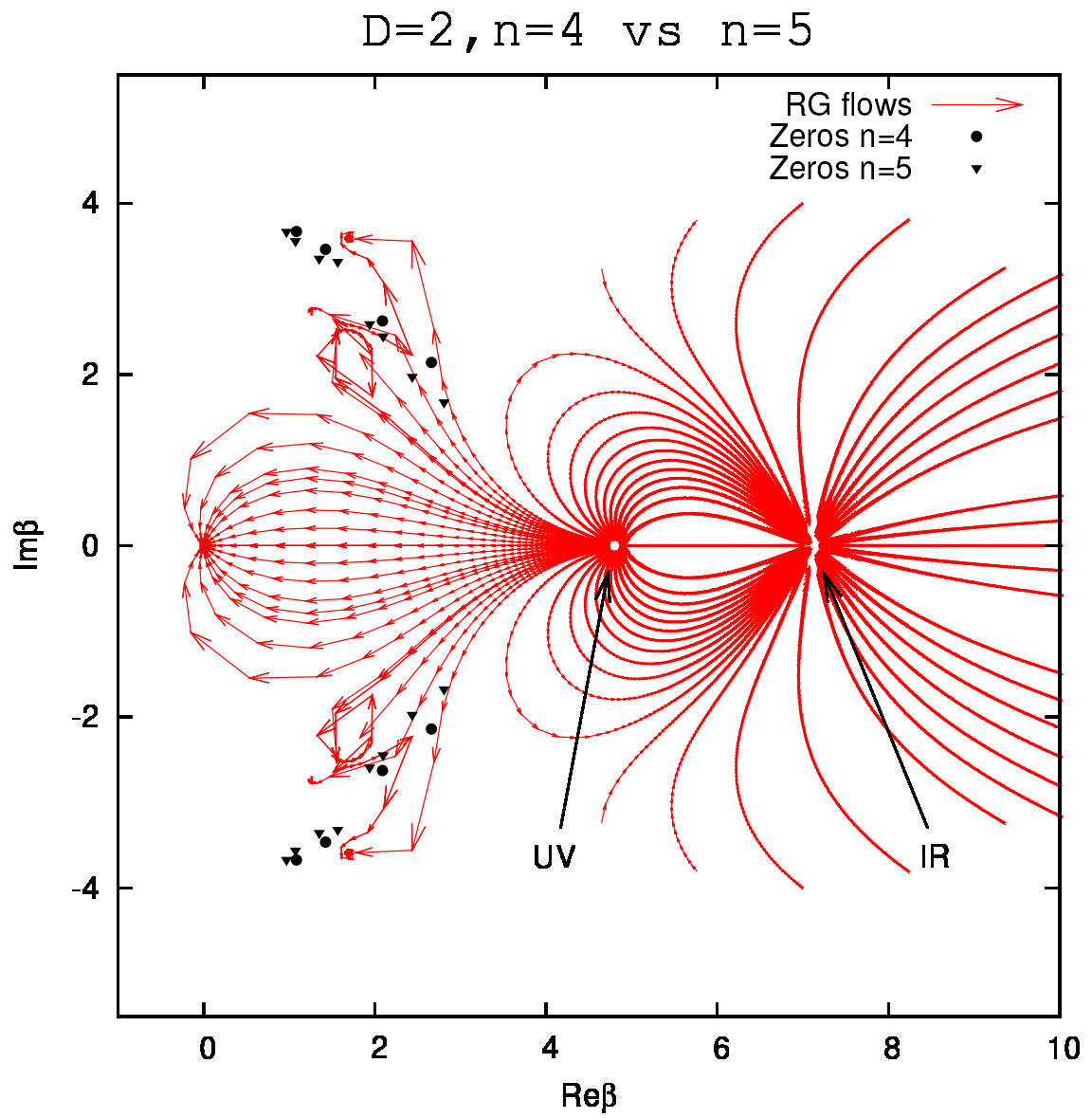


Figure 2.28: Complex RG flows for $D = 2$.

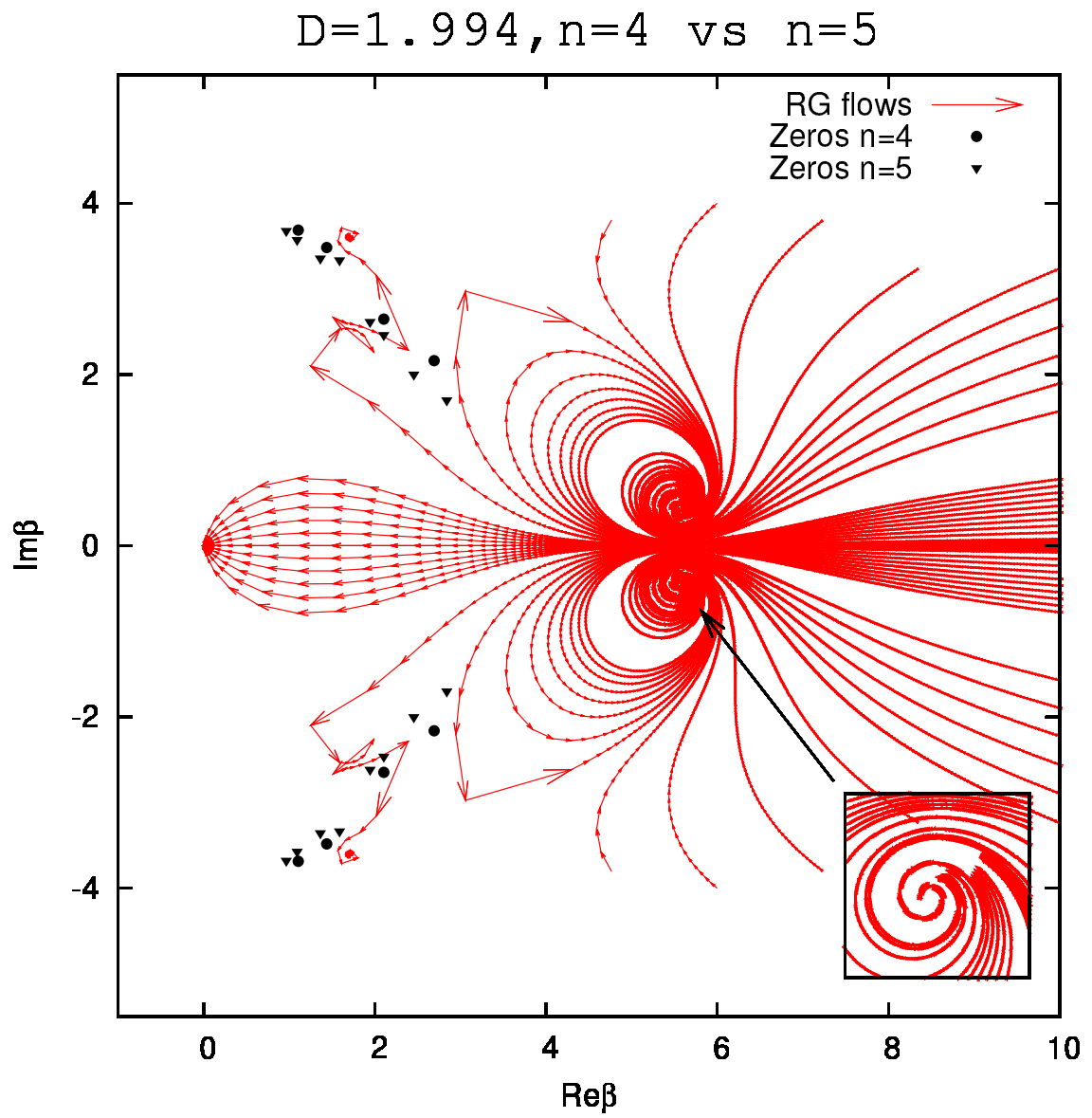


Figure 2.29: Complex RG flows for $D = 1.994$.

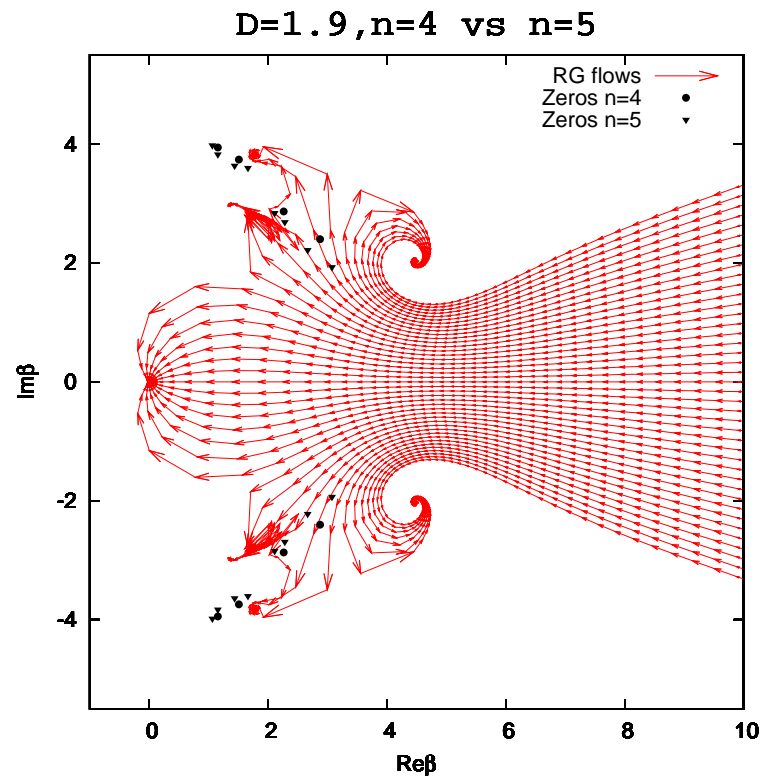


Figure 2.30: Complex RG flows for $D = 1.9$.

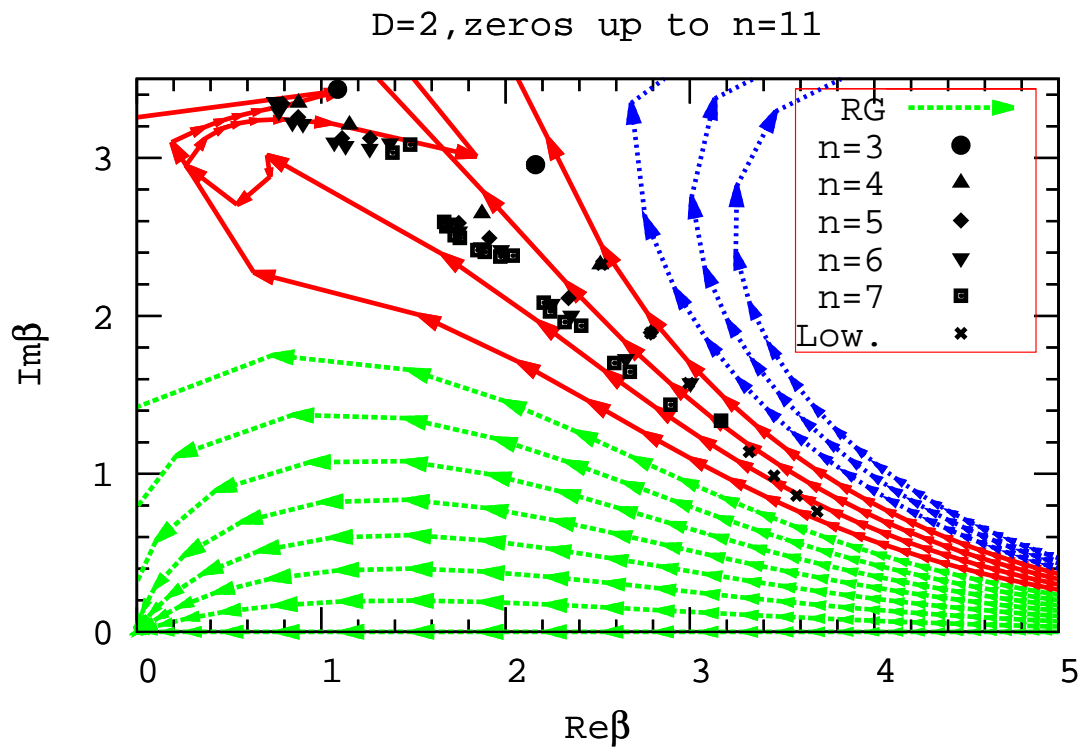


Figure 2.31: The density of the zeros is proportional to the $\ln V$.

CHAPTER 3

EXACT RENORMALIZATION GROUP

3.1 Introduction

In section 2.2, we see that the critical exponent γ of the hierarchical model depends on the blocking scheme. This is a very unpleasant result. In general, the critical exponent should be independent of the blocking. We plan to find methods to improve the hierarchical model. We will start with the hierarchical model and find a scheme independent model which has the same nice properties as the hierarchical model and at the same time can approximate the nearest neighbor model in D dimensions.

On the other hand, Daniel Litim [3] pointed out the critical exponents ν and ω calculated by the exact renormalization group (ERGE) method under local potential approximation (LPA) are highly correlated. He showed that the critical exponents of the hierarchical model with $b^D = 2$ also fell into the same curve (Fig. 3.1). We further show that all the other results with integer b^D also nicely fit into this curve. We are interested in knowing why this is the case. In the following section of this chapter, we will discuss the technique of the ERGE and its connection with the RG discussed in the previous chapter.

The ERGE was first introduced by Wilson [144], and then Wegner and Houghton [145] in the 1970s. Later on, it was reformulated by Polchinski [131] in 1984. ERGE is a continuous realization of the Wilson renormalization group. “Exact” comes from the fact that there is no approximation or expansion involved in the renormalization group (RG) transformation of the action. However, due to the complexity of the formulation, approximation or truncation are usually needed in order to perform practical calculations. Therefore, ERGE calculations are usually not “exact” and some people prefer the term “functional renormalization group”.

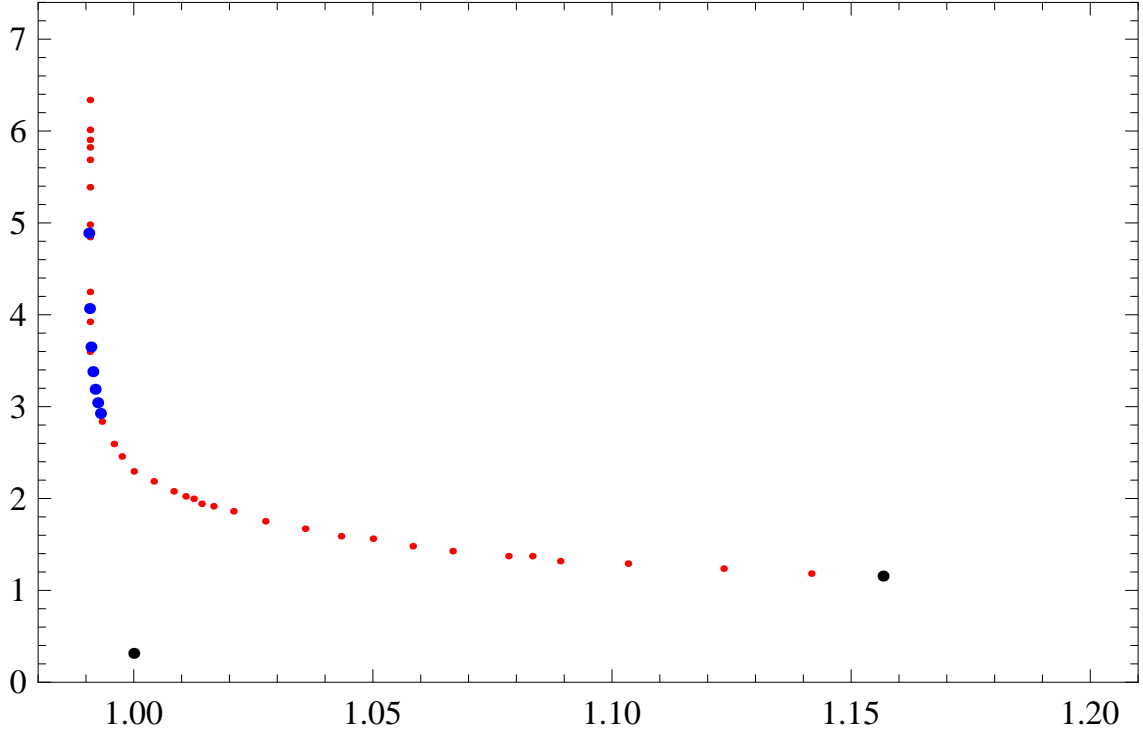


Figure 3.1: The red dots data were obtained from Fig. 4 in Ref. [3].

ERGE has been used in many sub-field of the physics. References can be found in the review articles [129, 146, 147, 148] and textbooks [149, 78, 150, 151].

In this chapter, we will mainly focus on the Wilson-Polchinski ERGE [144, 131], where the cutoff is smooth, under the local potential approximation (LPA) of the derivative expansion. We will also show the equivalence between the Wilson-Polchinski equation and the Litim optimized renormalization group equation [152, 153, 154, 155].

3.2 Overview

Renormalization group method has been used successfully in quantum field theory and condensed matter fields to calculate the critical exponents and other observables near the critical region. Wilson renormalization group (RG) is based on the idea of integrating out the high momentum degrees of freedom successively and

therefore interpolating between the low momentum and high momentum physics. There are many different realization of this process and the obvious advantage is that one can do various flexible approximations. Furthermore, optimization technique can be used to increase the reliability of the calculation.

For completeness, let us re-derive the formula we are interested in. The partition function with classical action S and a source term J can be written as

$$Z_k[J] = \int \exp(-S[\phi] + \int d^d x J(x)\phi(x)), \quad (3.1)$$

where d is the dimensionality of the system and ϕ is the scalar field. Let us then add a cutoff term $\Delta S_k[\phi]$, which takes into account of the rescaling of the field ϕ and keeping the original partition function invariant. One usually only consider the one-loop structure of the flow equation because higher loops will introduced complicated loop integration overlaps. The one-loop requirement constrains the cutoff term to be at most quadratic. Therefore, we can write

$$\delta S_k[\phi] = \frac{1}{2} \int d^d x \phi R \phi. \quad (3.2)$$

Here R is the cutoff function depending on the infrared scale k . The role of R is to suppress the low momentum mode while keep the high momentum mode intact. It also has to keep the correct initial condition $\Gamma_k|_{k \rightarrow \Lambda} = S_\Lambda$. Therefore, we will need R to satisfy the following condition

$$\lim_{q^2/k^2 \rightarrow 0} R_k(q^2) > 0, \quad (3.3)$$

$$\lim_{k^2/q^2 \rightarrow 0} R_k(q^2) = 0, \quad (3.4)$$

$$\lim_{k \rightarrow \Lambda} R_k(q^2) \rightarrow \infty. \quad (3.5)$$

The flow equation for the Legendre transformed effective action $\Gamma_K[\phi] = \int d^d x J(x)\phi(x) - \ln Z_k[\phi] - \Delta S_k[\phi]$ is [156, 157, 158, 159]

$$\partial_t \Gamma_k[\phi] = \frac{1}{2Tr(\frac{\partial^2 \Gamma_k[\phi]}{\partial \delta \phi(p) \partial \delta \phi(q)} + R_k)} \partial_t R_k, \quad (3.6)$$

where $t = \ln k$ is the logarithmic scale parameter, Tr means integrating over momentum and summing over indices.

The flow is linear and infrared and ultraviolet finite. Using the derivative expansion [160], to leading order in other words the local potential approximation (LPA) [144, 161, 162], the effective action of the $O(N)$ symmetric scalar theory has the following form

$$\Gamma_k = \int d^d x (V_k(\phi) + \frac{1}{2} \partial_\nu \phi \partial_\nu \phi), \quad (3.7)$$

where $V_k(\phi)$ is the local potential. From Eq. 3.6 and 3.7, we can get the flow equation for V_k field. If we rewrite the flow equation in terms of the dimensionless variables $u(\rho) = V_k/k^d$ and $\rho = 1/2\phi^2 k^{2-d}$ and taking the proper trace [163], we obtain

$$\partial_t u(\rho) = -du(\rho) + (d-2-\eta)\rho u' + \frac{K_d}{2}(N-1)l(u') + \frac{K_d}{2}l(u' + 2\rho u''), \quad (3.8)$$

where $K_d = \frac{2\pi^{d/2}}{(2\pi)^d \Gamma(d/2)}$ is the surface of the d dimensional unit sphere divided by $(2\pi)^d$. The function $l(\omega)$ is defined as

$$l(\omega) = \frac{1}{2} \int_0^\infty dy y^{d/2} \frac{\partial_t r(y)}{y(1+r) + \omega}, \quad (3.9)$$

where $y \equiv q^2/k^2$ and $\partial_t r(y) = -2yr'(y)$. Here $r(y)$ is the dimensionless regulator function

$$R(q^2) = q^2 r\left(\frac{q^2}{k^2}\right). \quad (3.10)$$

As stated before, one can choose different cutoff functions $r(y)$ as long as they satisfy the proper conditions. Different cutoff functions correspond to different coarse-graining process. Unfortunately, solutions to the truncated flow equation 3.6 depend on the choice of the cutoff function. This is similar to the perturbative QCD [164] case, where at finite order in perturbation theory different regularization

schemes gives different physical observables. It was found that there is an optimized cutoff function which leads to faster convergence and stable flow [165]. The function has the following form

$$r(y)_{\text{opt}} = \left(\frac{1}{y} - 1\right)\theta(1 - y). \quad (3.11)$$

In the following sections, we will mainly focus on the optimized cutoff function 3.11 and its variants.

3.3 Numerical Methods To Find The Fixed Points

Let us start from the flow equation 3.8 and 3.9 For $d = 3$, $\eta = 0$ and $N = 1$, we obtain

$$-3u + \rho u' + \int_0^\infty dy \frac{-y^{\frac{5}{2}} r'(y)}{y(1+r) + u' + 2\rho u''} = \partial_t u. \quad (3.12)$$

Note that in Eq. (2) of [166] we believe that it should be $y^{5/2}$ instead of $y^{3/2}$. If we choose the cutoff function to be $r(y) = b(\frac{1}{y} - 1)\theta(1 - y)$, where $\theta(x)$ is Heaviside step function, after proper rescaling of the field we obtain

$$-3u + \rho u' + \frac{1}{4\pi^2} \frac{2b(\sqrt{\frac{b+w}{b-1}} \text{ArcTanh} \sqrt{\frac{b-1}{b+w}} - 1)}{b-1} = \partial_t u \quad \text{for } b > 1, \quad (3.13)$$

and

$$-3u + \rho u' + \frac{1}{1 + u' + 2\rho u''} = \partial_t u \quad \text{for } b = 1, \quad (3.14)$$

with $w = u' + 2\rho u''$.

Let us focus on the $b = 1$ case first and the $b > 1$ case can be treated with the same technique after taking the series expansion.

The fixed point equation corresponds to $\partial_t u' = 0$. Since the integration constant does not affect the fixed point properties, we can equivalently write

$$-3u + \rho u' + \frac{1}{1 + u' + 2\rho u''} = 0, \quad (3.15)$$

or equivalently

$$(-3u + \rho u') * (1 + u' + 2\rho u'') + 1 = 0. \quad (3.16)$$

Equation (3.14) has two fixed points, one is the trivial Gaussian fixed point $u_* = \text{Constant}$ and another one is the interesting non-trivial Wilson-Fisher fixed point. In this section, we will describe different numerical methods we used to explore the non-trivial fixed point solution of equation (3.12), including the special case $b = 1$ (3.14) and several other expansions with different parameter b .

3.3.1 Polynomial Expansion of the Potential $u(\rho)$ About the Origin

It has been suggested [167, 168, 169, 170] that one can expand the effective potential as polynomials [155]. We first Taylor expand the potential $u(\rho)$ about the origin to order m

$$u(\rho) = \sum_{n=0}^m \frac{1}{n!} \lambda_n \rho^n. \quad (3.17)$$

Inserting equation (3.17) into left hand side of equation (3.16), we will get a polynomial of order $2m$ in terms of ρ

$$\begin{aligned} & f_0(\lambda_0, \lambda_1, \dots, \lambda_m) + f_1(\lambda_0, \lambda_1, \dots, \lambda_m)\rho, \\ & + f_2(\lambda_0, \lambda_1, \dots, \lambda_m)\rho^2 + \dots + f_{2m}(\lambda_0, \lambda_1, \dots, \lambda_m)\rho^{2m} = 0, \end{aligned} \quad (3.18)$$

where all the coefficients f_i s are in general functions of $\lambda_0, \dots, \lambda_m$. The fixed point condition corresponds to setting all the coefficients of ρ to zero. Since we only expand $u(\rho)$ to order m and the u' and $\rho u''$ terms bring the power down by 1, we can only impose the condition that $f_0 = 0, f_1 = 0, \dots, f_{m-1} = 0$. There are $m + 1$ variables and m independent equations so that we need to impose one more condition in order to solve the system. In practice, we notice that f_i only depends on $\lambda_0, \dots, \lambda_{i+1}$ (this is very easy to understand by looking at equation (3.16): the u' and $\rho u''$ bring the power down by 1). This nice property enable us to solve the equation iteratively and express all the λ s in terms of λ_1 . We can get λ_0 as a function of λ_1 from the condition $f_0 = 0$. And then we can substitute λ_0 by a

function of λ_1 in f_1, \dots, f_{m-1} . We can further get λ_2 as a function of λ_1 from $f_1 = 0$ condition. By doing so, we get the following relations

$$\begin{aligned}
\lambda_0 &= \frac{1}{3\lambda_1 + 3}, \\
\lambda_2 &= -\frac{2}{3}\lambda_1(\lambda_1 + 1)^2, \\
\lambda_3 &= \frac{2}{15}\lambda_1(\lambda_1 + 1)^3(13\lambda_1 + 1), \\
\lambda_4 &= -\frac{8}{7}\lambda_1^2(\lambda_1 + 1)^4(7\lambda_1 + 1), \\
\lambda_5 &= \frac{16}{189}\lambda_1^2(\lambda_1 + 1)^5(623\lambda_1^2 + 121\lambda_1 + 2), \\
&\vdots
\end{aligned} \tag{3.19}$$

We further apply the boundary condition $\lambda_m = 0$ in order to get the value of λ_1 . Since λ_m is a polynomial of λ_1 , we will get multiple solutions from $\lambda_m = 0$ condition. The fixed point value corresponds to the stable one with increasing m . See Figure (3.2).

Once we get the fixed point solution, we perturb around the fixed point and get the stability matrix and its the eigenvalues. The negative eigenvalue λ_1 relates to the critical exponent ν , $\nu = -1/\lambda_1$, and the smallest positive eigenvalue corresponds to the first sub-leading critical exponent ω .

The stability matrix M_{ij} is defined as

$$M_{ij} = \frac{\partial f_i}{\partial \lambda_j}, \tag{3.20}$$

where i and j run from 0 to m . The matrix M can be calculated from numerical derivative for large m . The method described above (expanding around the origin) is not the most efficient way to get the fixed point since the critical values or the fixed point values oscillate around the asymptotic value when we increase the expansion order m . This can be seen from Figure (3.3). In fact, we can only get 8 significant digits with order up to $m = 80$. This is clearly not practical for high precision calculations. Therefore, a new method must be introduced.

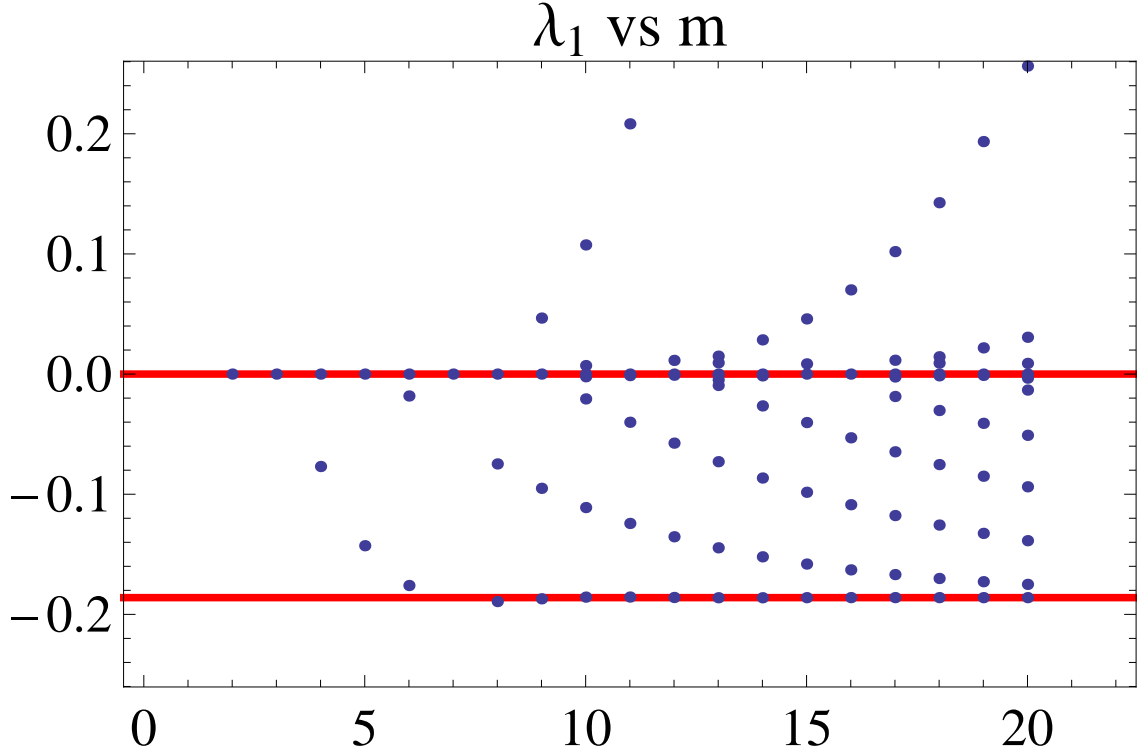


Figure 3.2: Two λ_1 s are stable with increasing polynomial order m .

3.3.2 Polynomial Expansion of the Potential $u(\rho)$ About the Minimum

It turns out that the convergence is much faster if we use the following expansion Equation (3.21)

$$u(\rho) = \sum_{n=1}^m \frac{1}{n!} \lambda'_n (\rho - \rho_{min})^n, \quad (3.21)$$

where ρ_{min} corresponds to the minimum of $u(\rho)$ (See Figure (3.4)) and the expansion coefficients λ'_n are in general different from the coefficients used in Eq. (3.17).

The ρ_{min} in Eq. 3.21 can be calculated from the method described in the previous section since we do not need very high accuracy for it. The boundary condition can be chosen such that $\lambda'_{m-1} = 0$ and $\lambda'_m = 0$. All the λ' s are implicitly

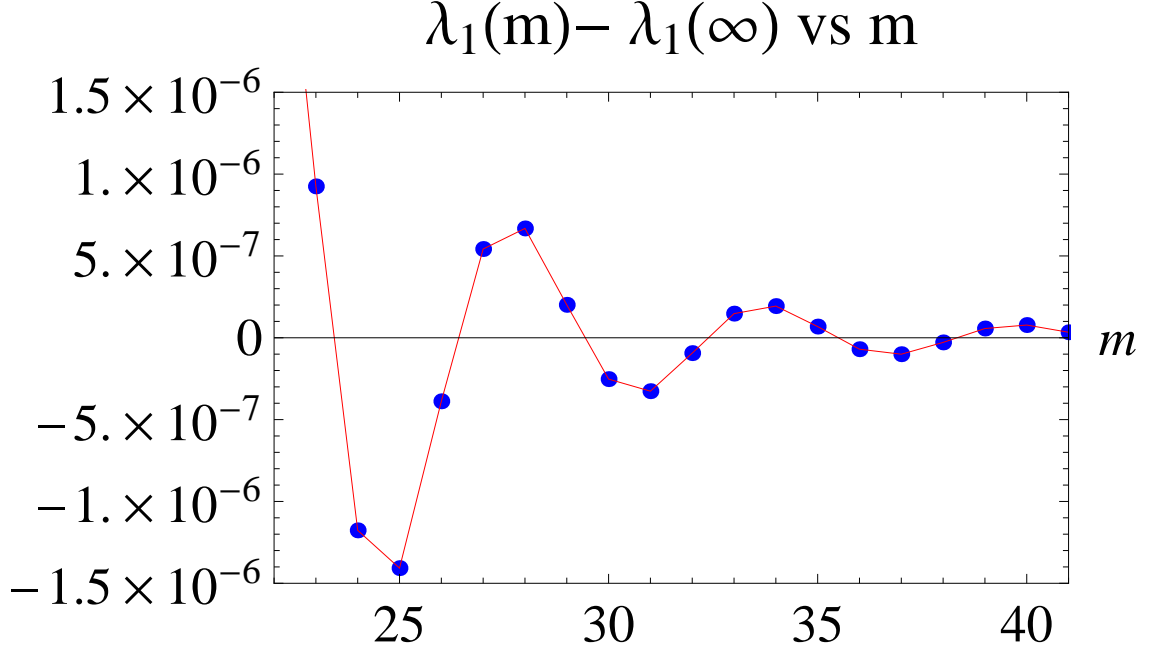


Figure 3.3: λ_1 approaches the asymptotic value $\lambda_1(\infty) = -0.18606424947 \dots$ slowly with increasing polynomial order m by using the expansion around the origin method.

dependent on λ_1 and λ_2 . We can express λ'_{m-1} and λ'_m as functions of λ_1 and λ_2

$$\lambda'_{m-1} = f_1(\lambda'_1, \lambda'_2) = 0, \quad (3.22)$$

$$\lambda'_m = f_2(\lambda'_1, \lambda'_2) = 0.$$

The solution of Eq. (3.22) can be obtained from the two dimensional Newton method. The result is shown in Table (3.1). The method converges fast with increasing polynomial order m , see Fig. (3.5). The first six critical exponents are listed in Table 3.2.

We also tried the expansion slightly away from the potential minimum ρ_{min} and get similar results

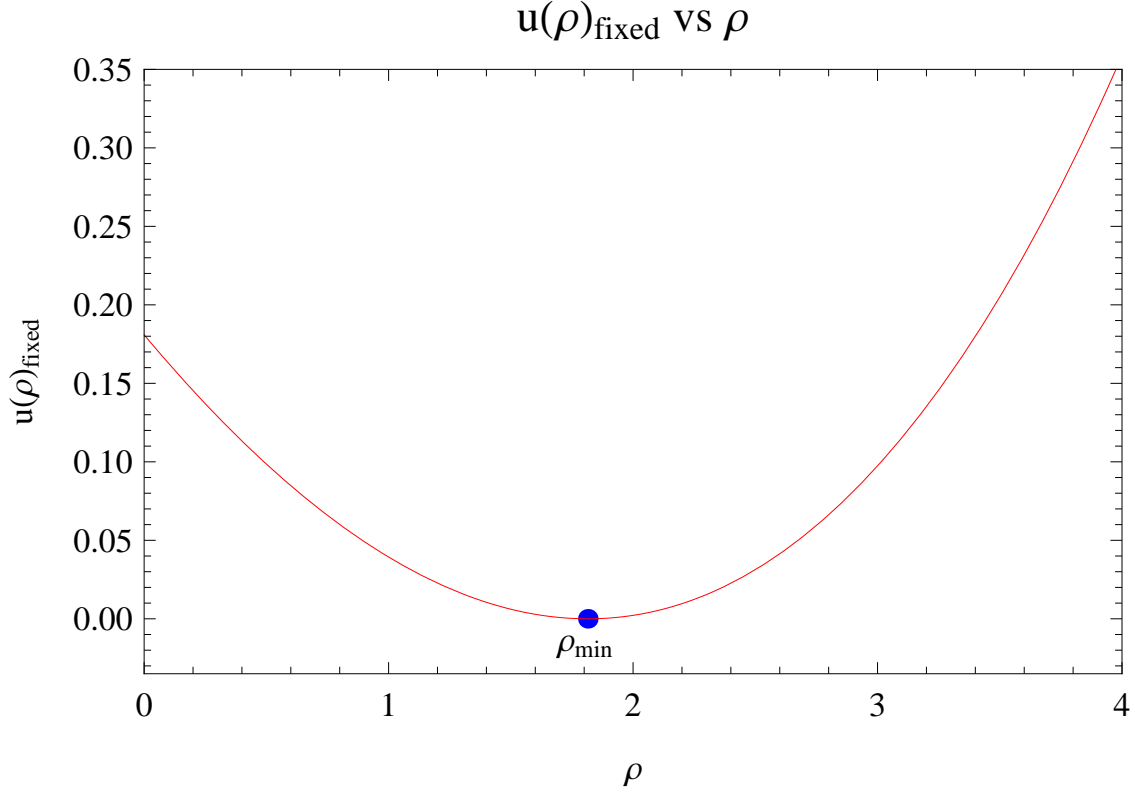


Figure 3.4: $u(\rho)$ versus ρ . ρ_{min} corresponds to the minimum of the potential $u(\rho)$.

Fixed point couplings	
λ'_1	$-3.92436863865 * 10^{-14}$
λ'_2	0.126164421218
λ'_3	0.0298149647677
λ'_4	0.00626281638385
λ'_5	-0.000275905517003
λ'_6	-0.00112762344235

Table 3.1: The first six couplings at the fixed point of Eq. (3.14) for the expansion of Eq. (3.21) with $m = 45$ and $\rho_{min} = 1.814898403687$.

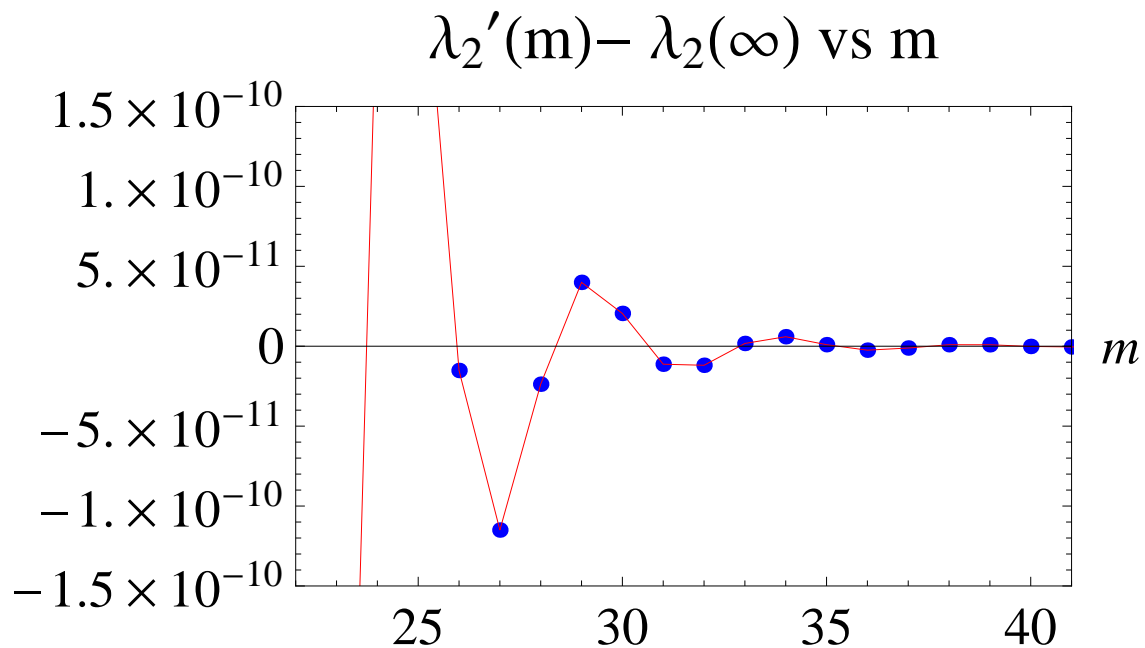


Figure 3.5: λ_2' approaches the asymptotic value very fast with increasing polynomial order m by using the expansion around the potential minimum method.

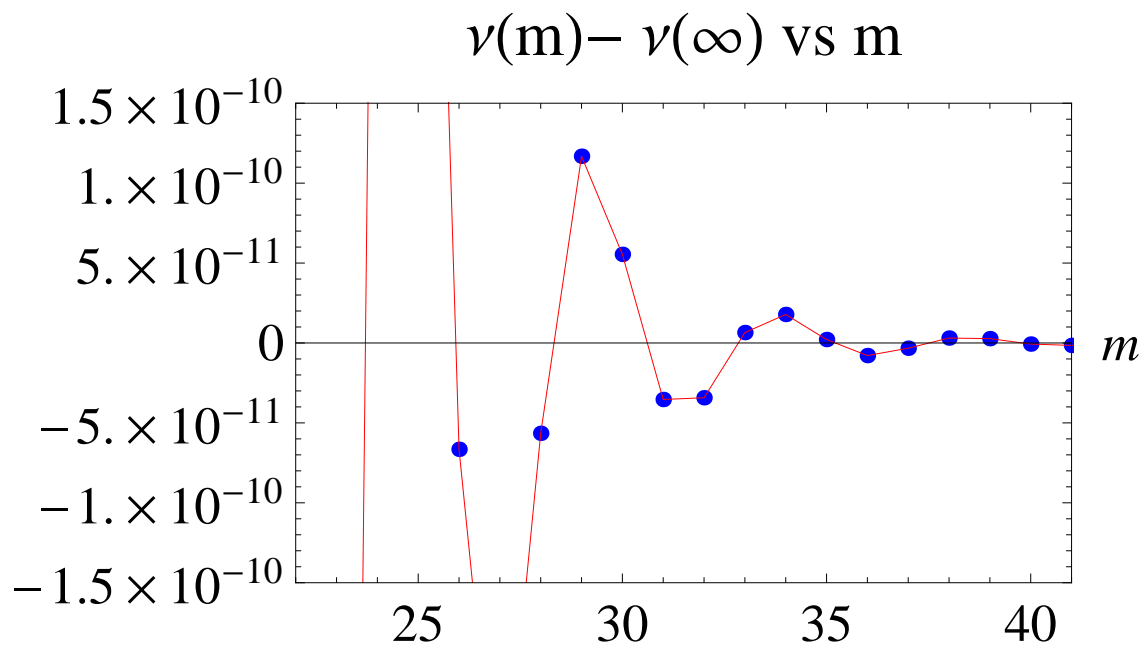


Figure 3.6: The convergence of the critical exponent ν .

Critical exponents	
ν	0.64956177388
ω	0.6557459391
ω_2	3.18000651
ω_3	5.9122305
ω_4	8.796093
ω_5	11.79808

Table 3.2: The first six critical exponents calculated from the expansion of Eq. (3.21) with $m = 42$ and $\rho_{\min} = 1.814898403687$.

3.4 Equivalence between Litim and Wilson-Polchinski equation

Before discussing the critical properties of the flow equation under other different cutoff functions, let us look more closely at the Wilson-Polchinski flow equation. Felder [133] has shown that the hierarchical RG is equivalent to the Wilson-Polchinski RG in the continuous blocking spin limit, i.e., $l^D \rightarrow 1$ limit. Moreover, Morris [154] proved that the ERGE with optimized cutoff function is also equivalent to the Wilson-Polchinski RG.

If we rewrite the Eq. 3.8 in terms of the variable $v(\phi) = V_k(\phi)/k^d$ and $\psi = \phi/k^{(2-d)/2}$, we obtain the following flow equation for $d = 3$

$$\partial_t v = -3v + \frac{1}{2}\varphi v' + \frac{1}{1+v''}. \quad (3.23)$$

The flow equation for the Wilson-Polchinski flow equation in terms of $v(\psi)_w$ is

$$\partial_t v_w = -3v_w + \frac{1}{2}\varphi_w v'_w - v''_w + v'^2_w. \quad (3.24)$$

In this section, I will show how the above two equations 3.23 and 3.24 are equivalent.

The potential $v(\varphi)$ in (3.23) is related to $v_w\varphi_w$ in (3.24) by a Legendre transformation

$$v(\varphi) = v_w(\varphi_w) - \frac{1}{2}(\varphi_w - \varphi)^2. \quad (3.25)$$

$$\varphi = \varphi_w - v'_w(\varphi_w). \quad (3.26)$$

Equation 3.25 corrects a typo in Eq. (9) of [155].

We can further get the relationship between the derivative of v and the that of v_w

$$\begin{aligned} v' &= \frac{\partial v}{\partial \varphi} \\ &= \frac{\partial v_w}{\partial \varphi_w} \frac{\partial \varphi_w}{\partial \varphi} - (\varphi_w - \varphi) \left(\frac{\partial \varphi_w}{\partial \varphi} - 1 \right) \\ &= v'_w \frac{1}{1 - v''_w} - v'_w \frac{v''_w}{1 - v''_w} \\ &= v'_w, \end{aligned} \quad (3.27)$$

$$v'' = \frac{\partial v'_w}{\partial \varphi} = v''_w \frac{\partial \varphi_w}{\partial \varphi} = v''_w \frac{1}{1 - v''_w}, \quad (3.28)$$

and further

$$v = v_w - \frac{1}{2}v'^2_w, \quad (3.29)$$

$$\frac{1}{1 + v''} = \frac{1}{1 + \frac{v''_w}{1 - v''_w}} = 1 - v''_w. \quad (3.30)$$

Finally,

$$\begin{aligned} & -3v + \frac{1}{2}\varphi v' + \frac{1}{1 + v''} \\ &= -3(v_w - \frac{1}{2}v'^2_w) + \frac{1}{2}(\varphi_w - v'_w)v'_w + (1 - v''_w) \\ &= -3v_w + \frac{1}{2}\varphi_w v'_w + v'^2_w - v''_w + 1 \\ &= -3v_w + \frac{1}{2}\varphi_w v'_w - v''_w + v'^2_w + 1. \end{aligned} \quad (3.31)$$

Since at the fixed point $\partial_t v' = 0$, the extra constant 1 plays no role on the fixed point equation.

3.5 Effects Of The Cutoff Functions

In Sec. 3.3, Eq. 3.13 and 3.14 are derived after proper field rescaling. This can be understood from App. B. In this section, we will use the properties derived in App. B to expand the flow equation and analyze the cutoff dependence of the critical exponents.

The “wrong” flow equation stated in [166] Eq. (2) is

$$-3u + \rho u' + \int_0^\infty dy \frac{-y^{\frac{3}{2}} r'(y)}{y(1+r) + u' + 2\rho u''} = 0, \quad (3.32)$$

with $r(y) = b(\frac{1}{y} - 1)\theta(1 - y)$, where $\theta(x)$ is Heaviside step function.

After integrating over y , Eq. (3.32) becomes

$$-3u + \rho u' + \frac{2b}{\sqrt{(b-1)(b+w)}} \text{ArcTanh} \sqrt{\frac{b-1}{b+w}} = 0 \quad \text{for } b > 1. \quad (3.33)$$

By applying App. B “No A Dependence” Theorem, we can remove the prefactor

$$\frac{2b}{\sqrt{b-1}} \Rightarrow$$

$$-3u + \rho u' + \frac{1}{\sqrt{b+w}} \text{ArcTanh} \sqrt{\frac{b-1}{b+w}} = 0. \quad (3.34)$$

Expanding $\text{ArcTanh}(x) = x + \frac{x^3}{3} + \frac{x^5}{5} + \dots$, the ArcTanh dependent term will become

$$\begin{aligned} & \frac{1}{\sqrt{b+w}} \left(\sqrt{\frac{b-1}{b+w}} + \frac{1}{3} \left(\sqrt{\frac{b-1}{b+w}} \right)^3 + \dots \right) \\ &= \sqrt{b-1} \left(\frac{1}{b+w} + \frac{1}{3} \frac{b-1}{(b+w)^2} + \frac{1}{5} \frac{(b-1)^2}{(b+w)^3} + \dots \right) \\ & \xrightarrow{\text{No A dependence}} \frac{1}{b+w} + \frac{1}{3} \frac{b-1}{(b+w)^2} + \frac{1}{5} \frac{(b-1)^2}{(b+w)^3} + \dots \end{aligned} \quad (3.35)$$

$$\begin{aligned} & \Rightarrow \frac{1}{b(1+\frac{w}{b})} + \frac{1}{3} \frac{b-1}{b^2(1+\frac{w}{b})^2} + \dots \\ & \xrightarrow{\text{No A dependence}} \frac{1}{1+\frac{w}{b}} + \frac{1}{3} \frac{b-1}{b} \frac{1}{(1+\frac{w}{b})^2} + \frac{1}{5} \left(\frac{b-1}{b} \right)^2 \frac{1}{(1+\frac{w}{b})^3} \dots \\ & \xrightarrow{\text{No B dependence}} \frac{1}{1+w} + \frac{1}{3} \frac{b-1}{b} \frac{1}{(1+w)^2} + \frac{1}{5} \left(\frac{b-1}{b} \right)^2 \frac{1}{(1+w)^3} + \dots \end{aligned} \quad (3.36)$$

Note that if one “accidentally” used $y^{5/2}$ instead of $y^{3/2}$, i.e.,

$$-3u + \rho u' + \int_0^\infty dy \frac{-y^{\frac{5}{2}} r'(y)}{y(1+r) + u' + 2\rho u''} = 0, \quad (3.37)$$

then after integration one will get

$$-3u + \rho u' + \frac{2b(\sqrt{\frac{b+w}{b-1}} \text{ArcTanh} \sqrt{\frac{b-1}{b+w}} - 1)}{b-1} = 0 \quad \text{for } b > 1. \quad (3.38)$$

For $y^{3/2}$ we will have

$$-3u + \rho u' + \frac{2b}{b-1} x(x + \frac{x^3}{3} + \frac{x^5}{5} + \dots) = -3u + \rho u' + \frac{2b}{b-1} (x^2 + \frac{x^4}{3} + \frac{x^6}{5} + \dots), \quad (3.39)$$

and for $y^{5/2}$ we will have

$$-3u + \rho u' + \frac{2b}{b-1} [\frac{1}{x} (x + \frac{x^3}{3} + \frac{x^5}{5} + \dots) - 1] = -3u + \rho u' + \frac{2b}{b-1} (\frac{x^2}{3} + \frac{x^4}{5} + \dots), \quad (3.40)$$

where $x = \sqrt{\frac{b-1}{b+w}}$.

Therefore, we can see that the difference between $y^{5/2}$ and $y^{3/2}$ is just in the coefficients of the x^{2n} . From Eq. 3.39, one can also see that one can start with Eq. 3.16 and consider higher orders of $x^2 = \frac{b-1}{b+w}$ as perturbations. The first step is to add a ϵx^4 to Eq. 3.16. For $b = 1$ case, we add $\epsilon \frac{1}{1+w}$. One will find that the critical exponents ν and ω change with different ϵ , see Fig. 3.7. If one plot them in the $\omega - \frac{1}{\nu}$ plane, one would get Fig. 3.8

We can also calculate critical exponents from Eq. 3.34 and 3.38 directly for different values of parameter b . Parts of the results are shown in Fig. 3.9 and 3.10. From Fig. 3.9 we can see that the $y^{3/2}$ results, which corresponds to the Eq. 3.32, converges to the “exact” Wilson-Polchinski RG result but not the $y^{5/2}$. The possible explanations are that either $y^{5/2}$ result “jump” to the “exact” result in the $b \rightarrow 1$ limit or $y^{3/2}$ is the proper form for the optimized cutoff ERG equation. I have not re-derived the Eq. 3.8 and 3.9 but prefer the later explanation.

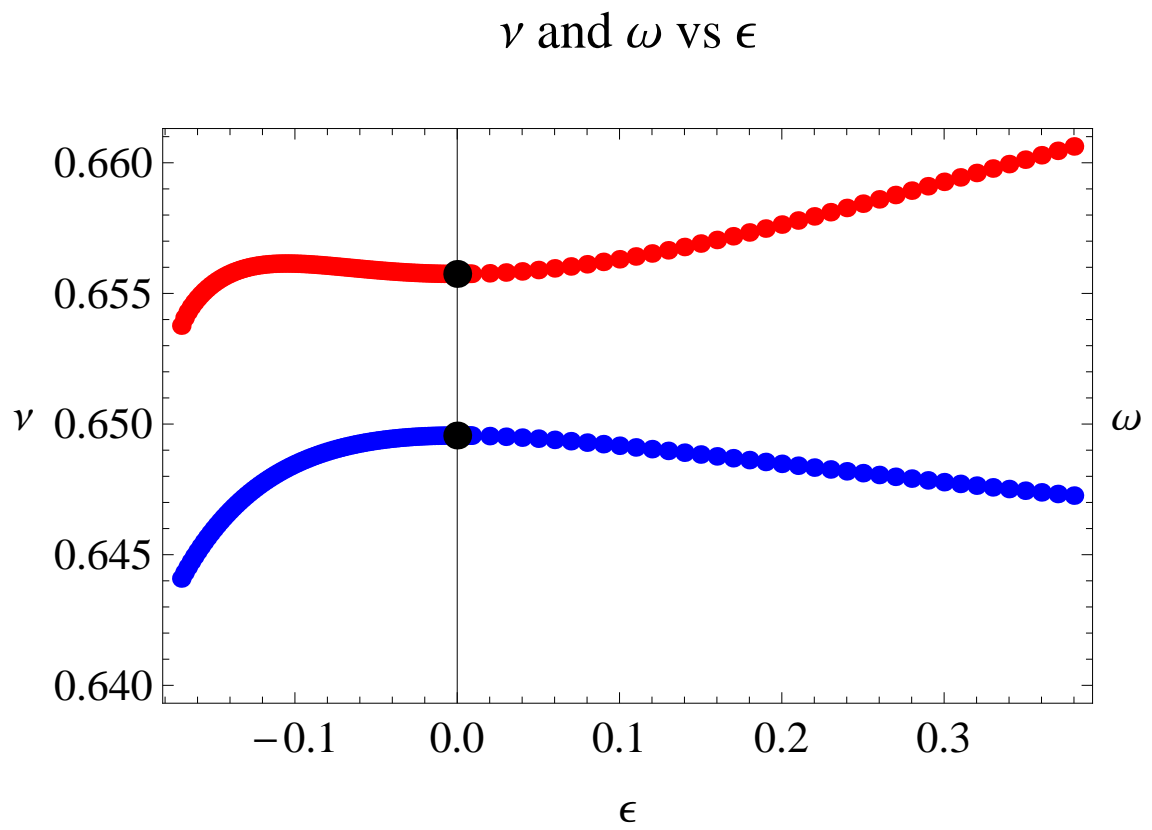


Figure 3.7: Critical exponents ν and ω change as function of ϵ .

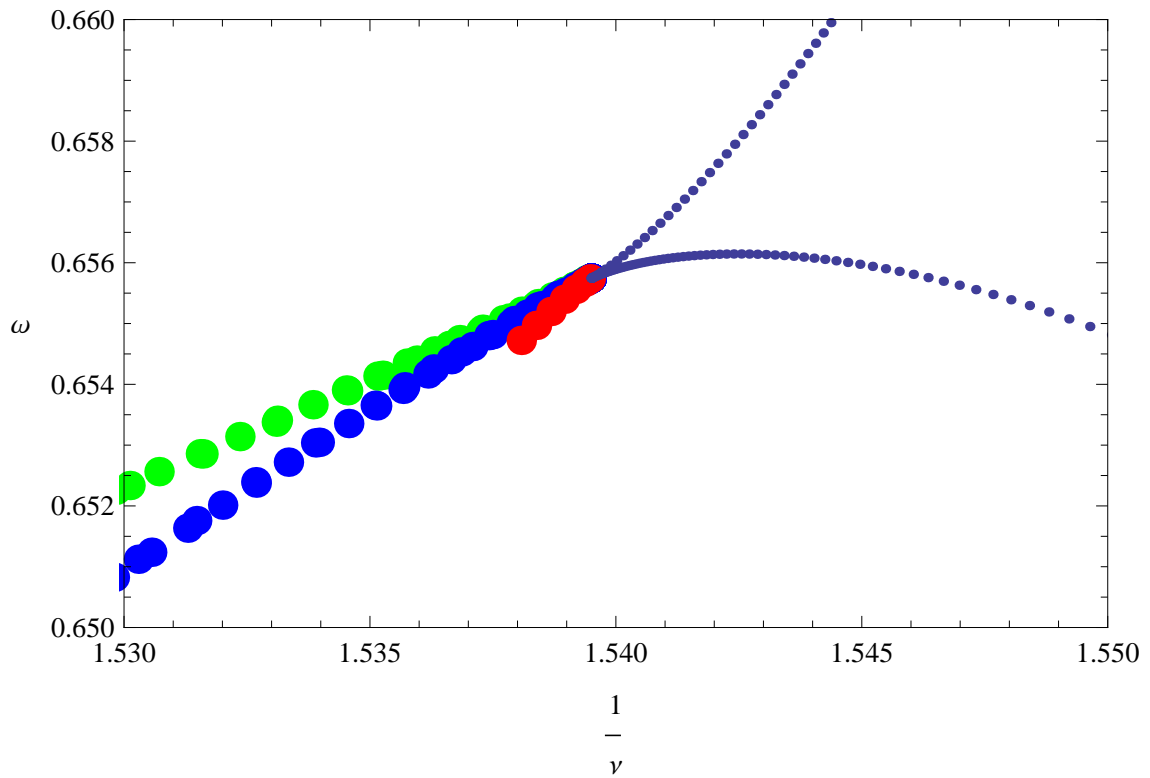


Figure 3.8: Critical exponents ν and ω for different ϵ .

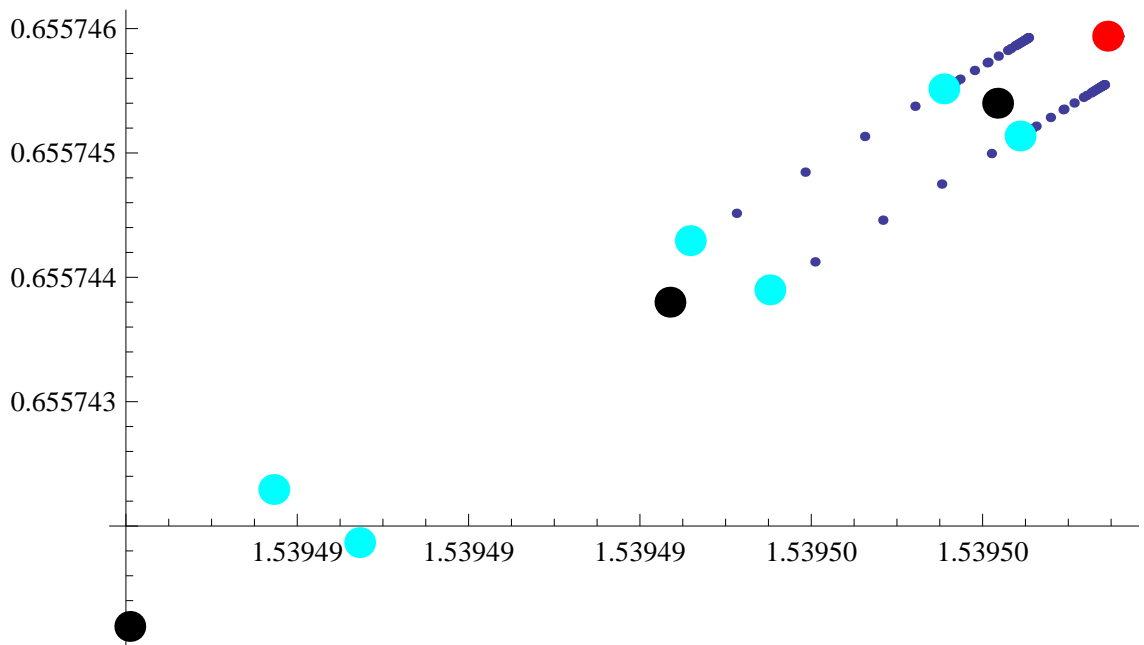


Figure 3.9: Show that Litim used formula corresponding to $y^{5/2}$ instead of $y^{3/2}$ to get the data “DataNuOmega.m”.

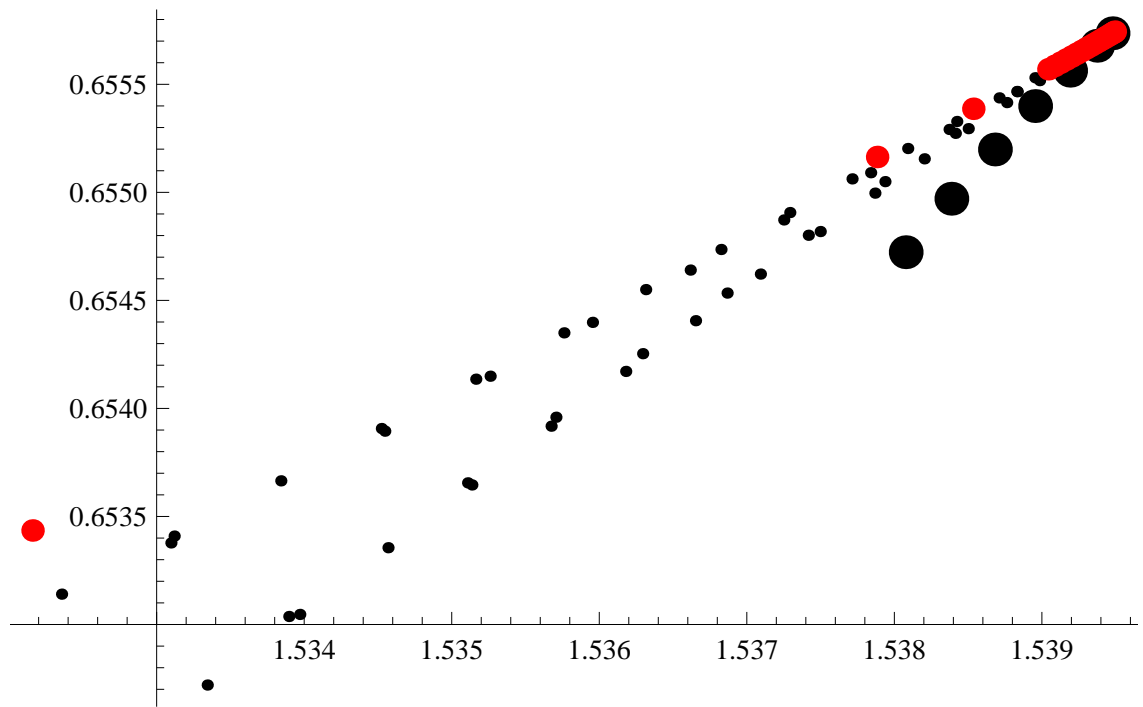


Figure 3.10: y axis is the ω and x axis is the $1/\nu$.

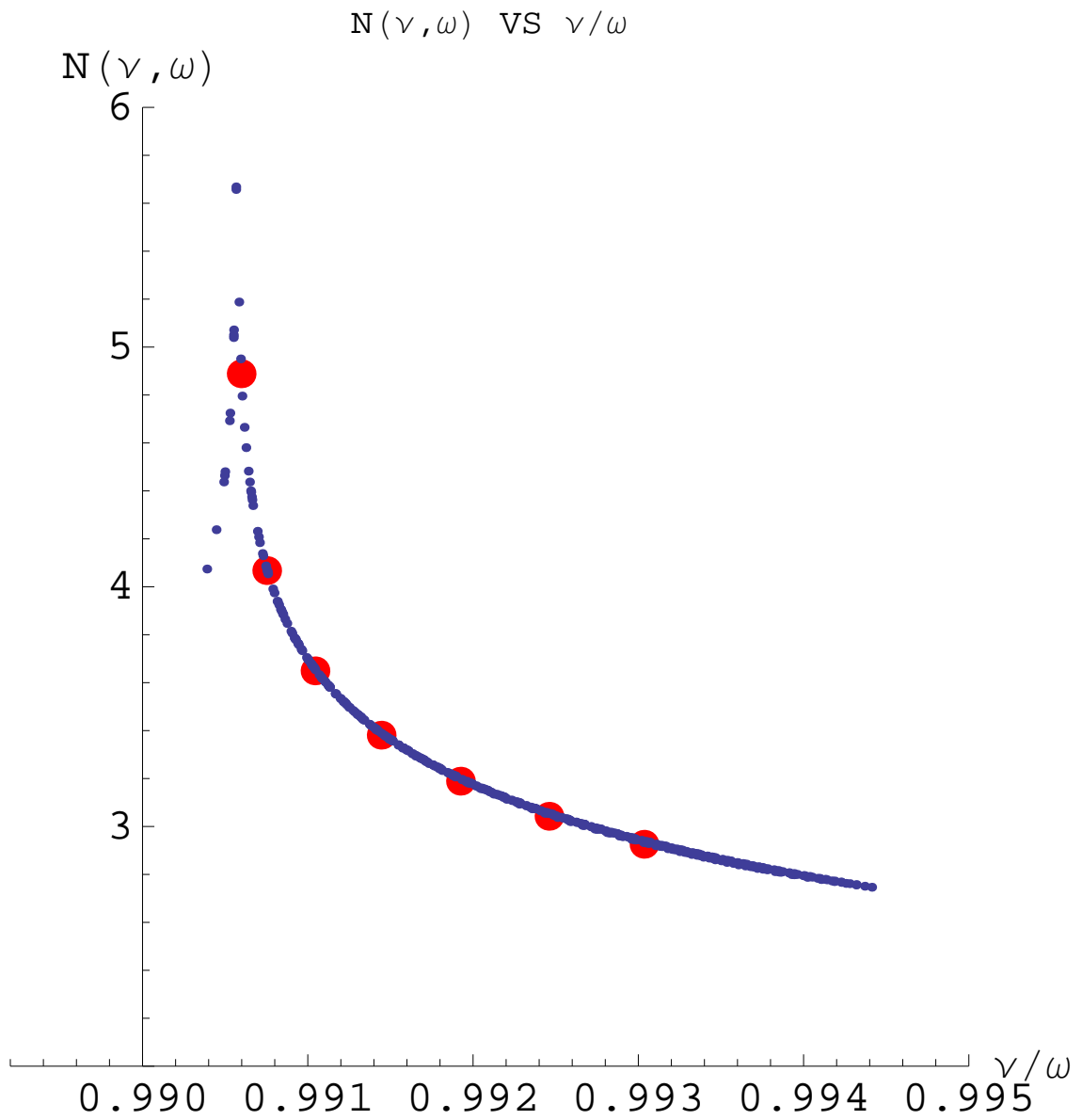


Figure 3.11: Blue points calculated with different ϵ_1 and ϵ_2 ; red points are calculated from the hierarchical model with different b^D .

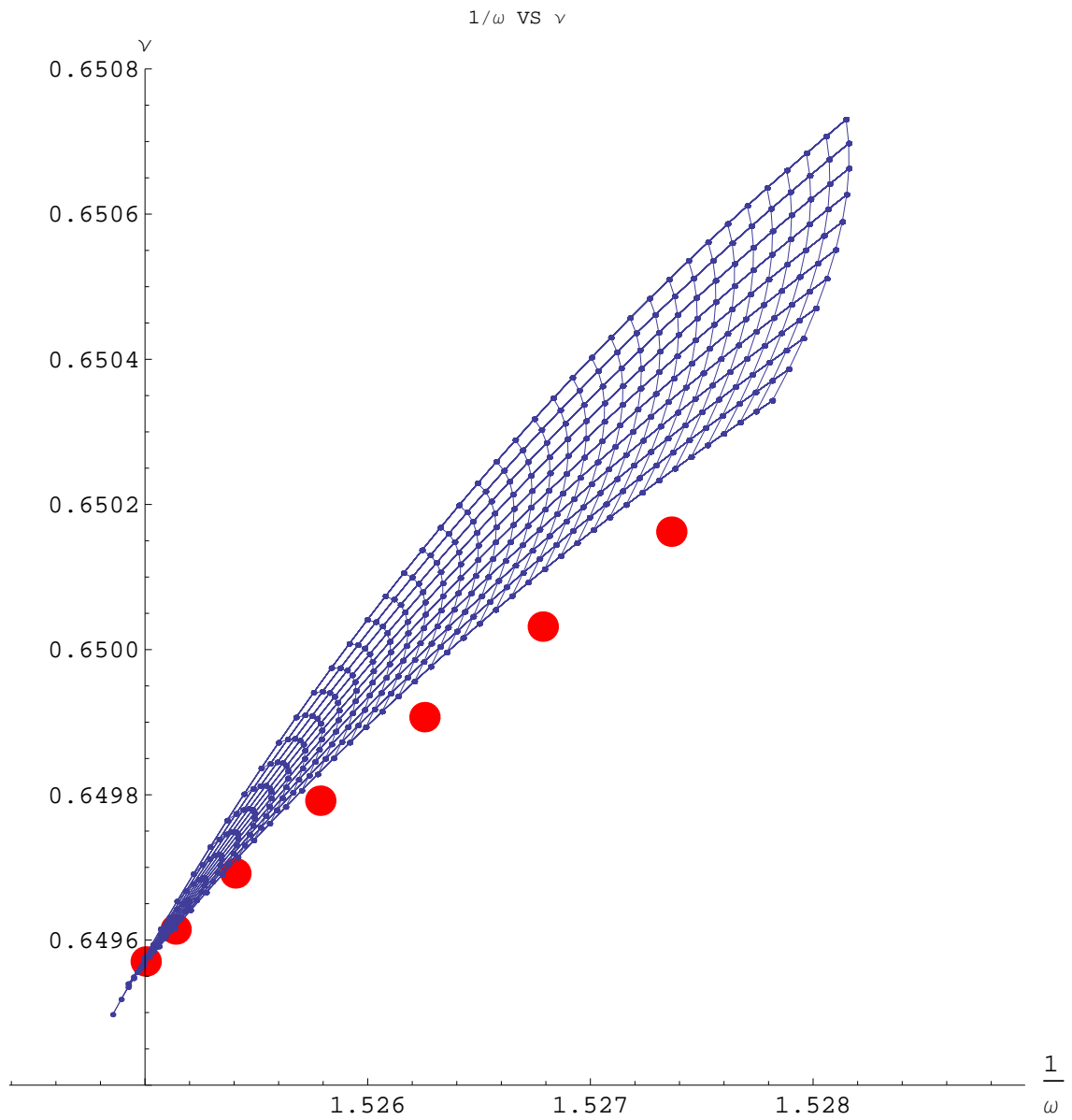


Figure 3.12: Blue points calculated with different ϵ_1 and ϵ_2 ; red points are calculated from the hierarchical model with different b^D .

CHAPTER 4

RENORMALIZATION GROUP AND LATTICE GAUGE THEORIES

4.1 Motivations

In the previous chapters, we have been focusing on the classical spin models. We detailed the properties of the partition function zeros, Lee-Yang zeros and Fisher zeros, the finite size scaling (FSS) technique, their relations to the critical exponents, discrete and continuous block-spinning, two lattice matching, β functions, complex RG flows, and ERGE. In the following two chapters, we will apply the technique developed and switch our attention to the more complicated and more physically important lattice gauge models. In this chapter, I will focus on the pure gauge $SU(2)$ lattice gauge model.

As stated in Sec. 2.2.8, FSS is a powerful tool to extrapolate information from finite system to the thermodynamic limit. It has also be used to study the critical behavior of the system, such as calculating critical exponents. The essence of FSS is the following: when the system is close to the second order phase transition, long distance properties are only determined by some global features of the system, such as the internal symmetries, the dimensionality, and the range of the interaction. The details of the Hamiltonian interactions is not so important compared to those global feature. Universality further enables us to classify seemingly different systems into certain classes. Connection between spin models and lattice gauge models can be seen from the conjecture of Svetitsky and Yaffe [171]. They pointed out that, for $d + 1$ dimensional $SU(N)$ pure gauge model, “if any portion of the boundary is second-order, then the critical behavior will be described by some fixed point of d -dimensional, $Z(N)$ invariant spin systems.”

For a pure gauge $SU(2)$ theory in 3+1 dimensions, we expect the universality class of the finite temperature transition to be the same as the 3 dimensional Ising

model. Existing results on FSS for $SU(2)$ [172, 173, 174, 175, 176] agree well with this expectation. The above conjecture was based on the assumption that the effective spin system of the gauge theory has only short range interactions. However, there were indications that the effective interaction of the Polyakov loops might be long range [177]. The simulation of $SU(2)$ gauge theory in 2+1 dimensions [178] also showed that the critical exponent $\beta(0.2 \pm 0.04)$ deviates from the corresponding 2D Ising exponent (0.125). In this chapter, we would like use FSS to analyze the lattice gauge model and try to extrapolate the critical exponents ν and ω and compare those with the three dimensional nearest neighbor Ising model.

4.2 Introduction To Lattice Gauge Theory

Classical lattice spin models are familiar to physicists because they are reflections of the lattice structures in materials in the laboratory. Some spin models have been extensively discussed in the previous chapters. Here we are going to discuss some more abstracted systems. In particular, we are going to use space-time lattice to describe the gauge field theory. The goal is to construct solutions of cutoff theories which can be mapped onto the real continuous Minkowski space-time field theory. The trick is to avoid integrating over infinite number of degrees of freedom. One can convert the original field theory problems to statistical mechanics ones.

Non-Abelian gauge theories in four dimensions were discovered to be asymptotically free (AF) [10, 11]. Roughly speaking, AF means that the coupling “constant” $g(\Lambda)$ decreases with increasing Λ and vanishes as $\Lambda \rightarrow \infty$. In another word, the effective coupling g goes to zero at short distance. This has been tested extremely well in the deep-inelastic scattering of electrons or neutrinos off strongly interacting protons and neutrons. This supports the conjecture that the strong interaction can be described by the $SU(3)$ “color” gauge group with quarks in the fundamental representations. Later on, people conjectured that the quarks should

be confined in such theories, which is consistent with the observation that no free quark has ever been caught in the experiment. Confinement is very natural in the lattice gauge theory in the strong coupling limit [179]. Strong coupling limit in the lattice gauge theory corresponds to the high temperature expansion in the spin system. In the lattice gauge theory, the effective bare coupling will go to zero as one takes the lattice spacing to the continuum, which corresponds to the low temperature domain. The task then would be to obtain a theory that has no phase transition when one goes from the high temperature to low temperature region. This way, one could then construct a computable theory that has the properties of both confinement and AF. However, we will see later on that many lattice gauge models have non-trivial phase transition in the intermediate region. On the one hand, one should make sure that one is in the ‘right’ phase if one wants to simulate the continuous physics. On the other hand, the phase structure of different lattice gauge theories are interesting on their own. In the following, we will see that the statistical mechanics of lattice gauge theories share many similarities with the classical spin models. More details about the lattice gauge theories and lattice QCD like theories are discussed in the App. C.

4.3 Finite Size Scaling of The $SU(2)$ Lattice Gauge Theory

In this section, we work on $SU(2)$ lattice gauge theory in 3+1 dimensions: $N_\tau \times N_\sigma^3$. We fix the number of site in the time direction N_τ to be 4, and vary the number of site in the space dimension N_σ from 2, 4, 6, 8, \dots 16. We use periodic boundary conditions in all the directions. We focus on direct estimations of the exponents ν and ω . For $N_\tau = 4$, the only direct estimate of ν we are aware of is 0.65(4), which was obtained in Ref. [172]. This is compatible with the accurate value 0.6298(5) obtained in Ref. [180] for the 3-dimensional Ising model. In this

section, we would like to improve the accuracy of the estimate of Ref. [172] and resolve the corrections due to the irrelevant directions. The second aim is largely unexplored and a better understanding of these corrections could help us design methods to reduce these effects as done in Ref. [180] for the 3 dimensional Ising model. In the existing work for $N_\tau = 4$ in Refs. [172, 173, 176], a fixed β interval procedure was used. This means that the β interval is fixed for different volumes. In the following, we shrink the interval in order to reduce the nonlinear effects [181] and use a finer β resolution.

4.3.1 Binder cumulant and FSS

We define the 4th order Binder cumulant [182], g_4 , as

$$g_4 = 1 - \frac{\langle P^4 \rangle}{3\langle P^2 \rangle^2}, \quad P = \frac{1}{N_\sigma^3} \sum_{\mathbf{x}} \frac{1}{2} \text{Tr} \prod_{\tau=1}^{N_\tau} U_{\tau, \mathbf{x}; 0}. \quad (4.1)$$

Related definitions appear in the literature, such as $B_4 = \frac{\langle P^4 \rangle}{\langle P^2 \rangle^2}$. We assume that there is no external field and that the g_4 depends on the scaling variables as

$$g_4 = g_4(u_\kappa N_\sigma^{1/\nu}, u_1 N_\sigma^{-\omega}, \dots), \quad (4.2)$$

with

$$\begin{aligned} u_\kappa &= \kappa + u_\kappa^{(2)} \kappa^2 + \dots \\ u_1 &= u_1^{(0)} + u_1^{(1)} \kappa + \dots, \end{aligned}$$

where u_κ is the only relevant scaling variable, u_1 is the first irrelevant scaling variable, and κ is the reduced quantity $\kappa = (\beta - \beta_c)/\beta_c$. Figure 4.1 shows the Binder cumulant as function of β for different volumes. We can see that as the volume increases, all the curves cross around one β , which is just the critical β_c . This can be understood by expanding the Binder cumulant up to the first nonlinear corrections.

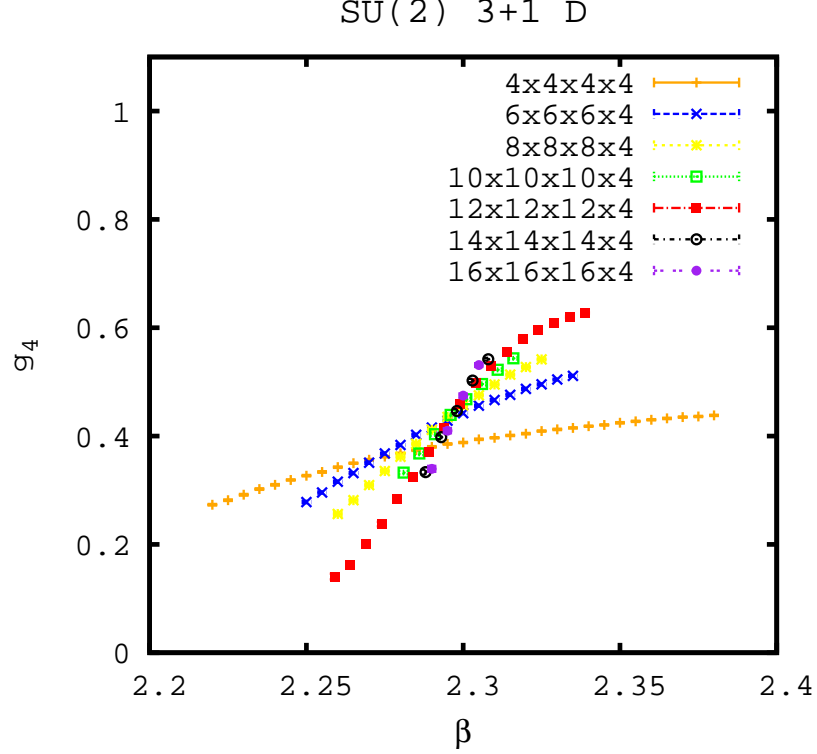


Figure 4.1: g_4 versus β for different volumes.

We obtain

$$g_4(\beta, N_\sigma) = g_4(\beta_c, \infty) + f_1 \kappa N_\sigma^{1/\nu} + f_2 \kappa^2 N_\sigma^{2/\nu} + (c_0 + c_1 \kappa N_\sigma^{1/\nu}) N_\sigma^{-\omega} + \dots \quad (4.3)$$

This expansion is accurate if $|\kappa| N_\sigma^{1/\nu}$ is small enough. In addition we would like the nonlinear effects of f_2 and c_1 to be negligible. It is easy to estimate f_2 from numerical data for intermediate values of κ [181]. However, c_1 is more difficult to resolve from an already small effect and its effect will be ignored. In the following, we will work with values of $|\kappa| N_\sigma^{1/\nu}$ such that the effects of f_2 are within the numerical errors of g_4 that we now proceed to discuss.

4.3.2 Determination of the critical exponent ν

We now focus on the estimation of ν . If we are reasonably close to β_c , we can use the linear form:

$$g_4(\beta, N_\sigma) \simeq g_4(\beta_c, \infty) + c_0 N_\sigma^{-\omega} + f_1 \kappa N_\sigma^{1/\nu}. \quad (4.4)$$

This expression contains 6 unknown parameters: $g_4(\beta_c, \infty)$, c_0 , ω , f_1 , β_c and $1/\nu$ and we will use a new strategy to attack this difficult problem. A first observation is that the dependence on ν can be isolated from the other parameters by studying the linear dependence in β . However, one should keep in mind that the slope of this linear fit depends implicitly on the choice of the center of the interval which should be as close as possible to β_c . In order to guarantee that nonlinear effects are under control, we will consider β intervals of the form

$$|\beta - (\beta_c)_{app}| < 0.015 \times (4/N_\sigma)^{(1/\nu)_{app}}. \quad (4.5)$$

We will start with reasonable values for $(\beta_c)_{app}$ and $(1/\nu)_{app}$ and then show that the effect of their variations is small. The factor 0.015 has been chosen following a procedure described in Ref. [181] and guarantees an approximate linear relation between g_4 and β at fixed N_σ :

$$g_4 \simeq a_{N_\sigma} + b_{N_\sigma} \times \beta. \quad (4.6)$$

This is illustrated in Fig. 4.2 for $(\beta_c)_{app} = 2.299$, $(1/\nu)_{app} = 1.6$ and $N_\sigma = 10$. We see that in the chosen interval the deviation from linearity seem mostly due to statistical errors rather than a systematic curvature that would be observed if we had chosen a broader interval. The volume dependence of the parameters of the

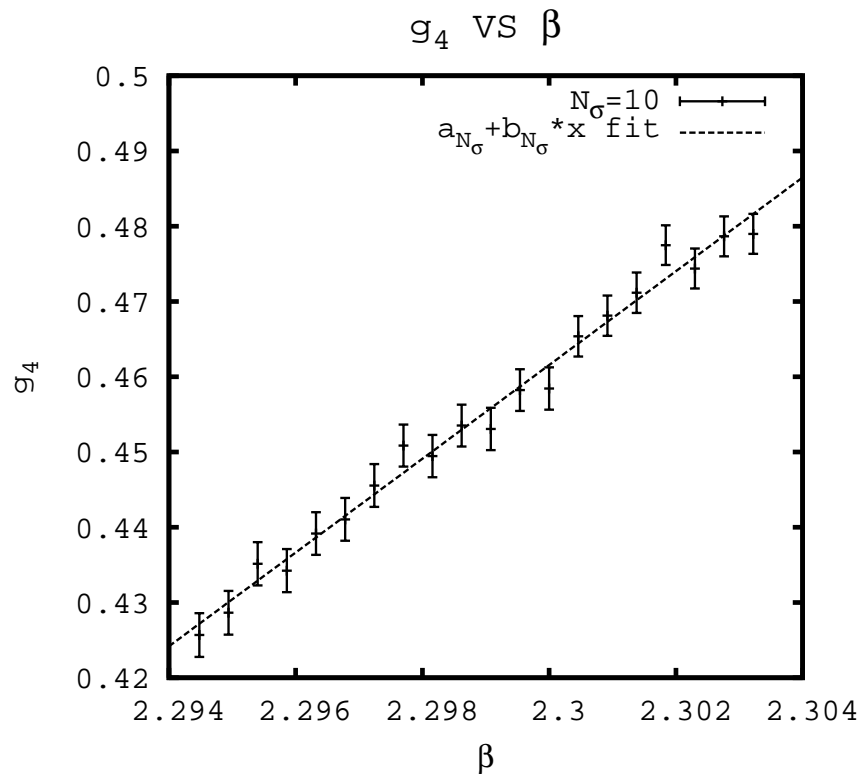


Figure 4.2: Linear fit of g_4 near β_c for $N_\sigma = 10$, $(\beta_c)_{app} = 2.299$ and $(1/\nu)_{app} = 1.6$.

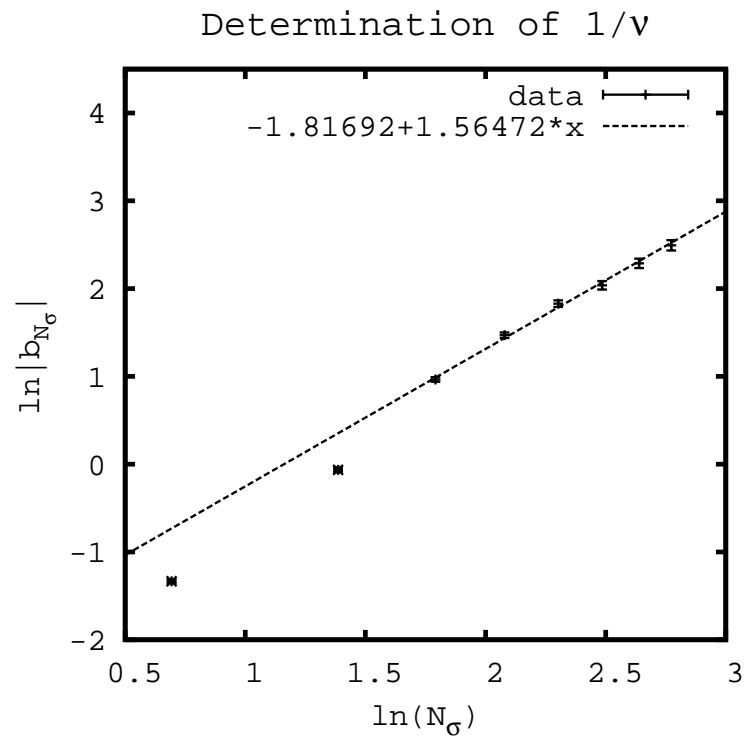


Figure 4.3: Determination of $1/\nu$ from the log-log fit discussed in the text.

linear fit are

$$a_{N_\sigma} \simeq g_4(\beta_c, \infty) + c_0 N_\sigma^{-\omega} - f_1 N_\sigma^{1/\nu} \quad (4.7)$$

$$b_{N_\sigma} \simeq f_1 N_\sigma^{1/\nu} / \beta_c. \quad (4.8)$$

Once we have b_{N_σ} for different N_σ , we can do a log-log fit to determine the inverse critical exponent $1/\nu$. One should note that the slope can be determined independently of β_c or $g_4(\beta_c, \infty)$. This is illustrated in Fig. 4.3. From the log-log fit with $N_\sigma \geq 6$, we obtain $1/\nu = 1.56(4)$. We plan to model and explain the deviations from linearity at low N_σ .

We need to address the dependence on $(\beta_c)_{app}$ and $(1/\nu)_{app}$. We used the central value $\beta_c = 2.2991$ from Ref. [176] as the critical value and changed the center of the interval $(\beta_c)_{app}$, between 2.297 and 2.301 and $(1/\nu)_{app}$ between 1.4 and 1.8. We calculated $1/\nu$ for a set of 189 values of $((\beta_c)_{app}, (1/\nu)_{app})$. The histogram is shown in Fig. 4.4. We can see that the values appear in a rather large range between 1.468 and 1.631. The average of this set of results is $\overline{1/\nu} = 1.570$ with $\sigma = 0.027$ which will be our preliminary result. In the meantime, we collected more data at large N_σ . If we choose the center of the data in a smaller range, namely with $(\beta_c)_{app}$ from 2.298 and 2.300 and the same range for $(1/\nu)_{app}$, we get a slightly different distribution shown in Fig. 4.5, but the average is essentially the same. It seems thus possible to average over $(\beta_c)_{app}$ and $(1/\nu)_{app}$ in order to get a more accurate value of $1/\nu$. For reference, $1/\nu_{Ising}$ is estimated as 1.5887(85) in Ref. [183] and 1.5878(12) in Ref. [180] and our preliminary result here is consistent with these more accurate values. This is preliminary, we plan to go to larger N_σ and to determine β_c independently using the method of Ref. [176].

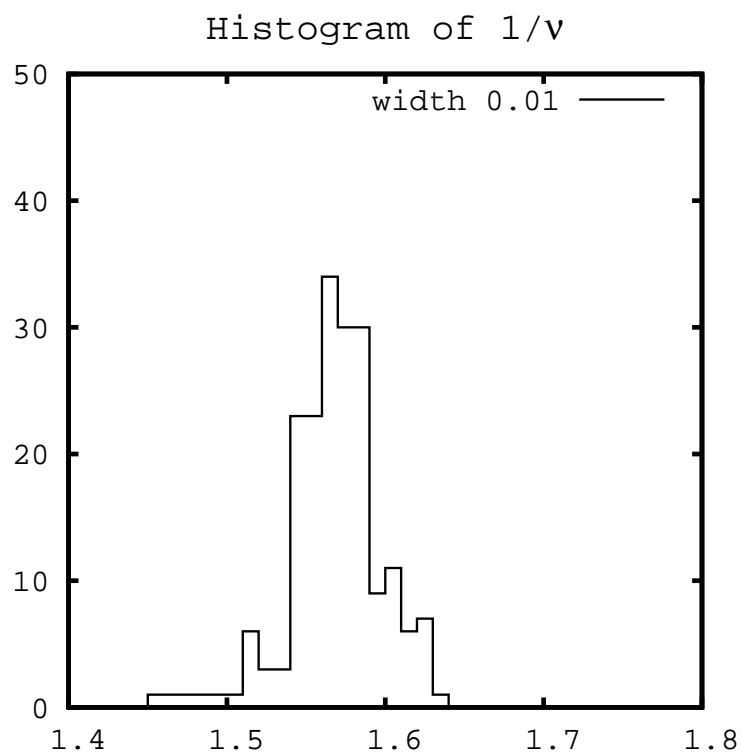


Figure 4.4: $(\beta_c)_{app}$ changes from 2.297 to 2.301; $(1/\nu)_{app}$ changes from 1.4 to 1.8. $\overline{1/\nu} = 1.570$; $\sigma = 0.027$.

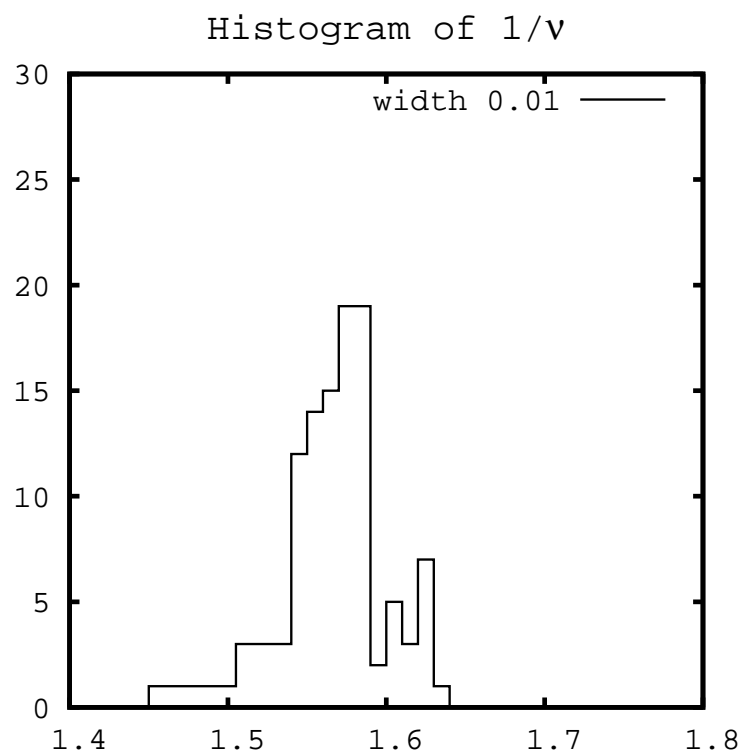


Figure 4.5: $(\beta_c)_{app}$ changes from 2.298 to 2.300; $(1/\nu)_{app}$ changes from 1.4 to 1.8. $\overline{1/\nu} = 1.571$; $\sigma = 0.028$.

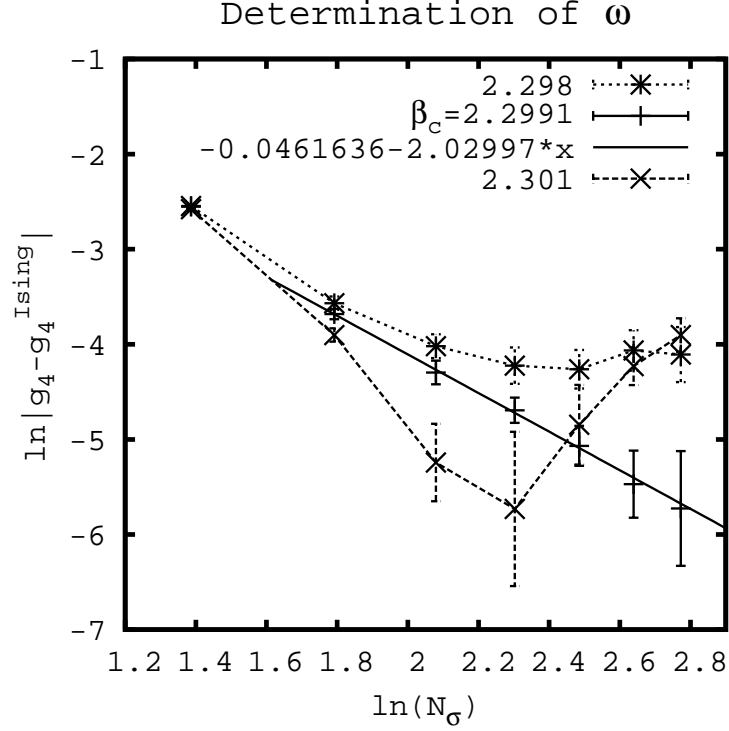


Figure 4.6: For $\beta_c = 2.2991$, the behavior is approximately linear: $g_4 \simeq g_4(\beta_c, \infty) + c_0 \times N_\sigma^{-\omega}$.

4.3.3 Determination of the critical exponent ω

Unless we determine β_c and $1/\nu$ very precisely, it is very difficult to subtract the effects of the third term of Eq. (4.4). If we can work at β_c , this term is absent:

$$g_4(\beta_c, N_\sigma) = g_4(\beta_c, \infty) + c_0 N_\sigma^{-\omega}. \quad (4.9)$$

Consistently with the previous section and the rest of the literature, we assume the universal value $g_4(\beta_c, \infty) = 0.46575$ as found in Ref. [180]. $\text{Log}[|g_4 - g_4(\beta_c, \infty)|]$ vs. $\text{Log}[N_\sigma]$ should be linear right at β_c and nonlinear for all the other β s. This is shown in Figure 4.6. At the same time, the slope is $-\omega$. The result we obtained from this analysis is $\omega = 2.030(36)$.

This is very different from $\omega_{Sing} = 0.812$ [183]. It is possible that the coefficient of the $N_\sigma^{-\omega}$ is very small and the exponent we extrapolated is a sub-subleading exponent. For a detail discussion of the sub-leading corrections, see Ref. [180]. The most plausible explanation seems that this exponent is related to the irrelevant direction associated with the breaking of rotational symmetry [184] and which is close to 2.

4.4 Infrared Fixed Point of the $SU(3)$ Multiflavor Lattice Gauge Theory

With the ongoing effort at the LHC, there has been a renewed interest in the phase diagram of lattice gauge theory models. Theories that have an infrared fixed point (IRFP) and are asymptotically free(AF) at high energies are called inside the “conformal window”. The location of the conformal windows for several families of models have triggered intense discussions. Different numerical and analytical techniques have been applied to QCD-like models with a large number of fermion flavors or with fermions in higher representations. It is important to understand the critical behavior of lattice models from various points of view. It was proposed [1, 123] to consider complex extensions of the picture of confinement proposed by Tomboulis. It was observed that the Fisher zeros, the zeros of the partition function in the complex β plane, determine the global properties of the complex RG flows. In the case where a phase transition is present, the scaling properties of the zeros allow us to distinguish between a first and second order phase transition. In the following, we briefly review the Finite Size Scaling (FSS) of the Fisher zeros. We then discuss numerical results for $SU(3)$ in 4D $N_f = 4$ and 12 flavors.

The Fisher zeros for $SU(3)$ with $N_f = 4$ and 12 quarks in the fundamental representation have been investigated. We used the standard Wilson gauge action and unimproved staggered fermion action with the Rational Hybrid Monte Carlo

(RHMC) algorithm [185, 186, 187]. We started with relatively small symmetric lattices and up to 50,000 configurations. The bare quark mass is set to be $m_q = 0.02$ for now. For $N_f = 12$, we found a discontinuity for the plaquette near $\beta \simeq 4.1$ for $V = 8^4$ lattices. The plaquette histories showed the characteristic hysteresis behavior. For comparison, we show the crossover for $N_f = 4$. Possible hypothesis to be tested are that for $N_f = 12$, the imaginary part scales like L^{-4} , which signals a first order phase transition and that the real part increases like $\log(L)$. Nonlinear effects seem important at small volume and larger volume calculations are in progress. The chiral condensate has a discontinuity near the same value of β as the plaquette. More recently, it was shown that improved actions can create a second discontinuity with a broken single-site shift symmetry between the two transitions.

We know that RG flows are defined in the space of couplings: (g_1, g_2, g_3, \dots) . β_{CS} functions describe how the couplings change with energy scales Λ :

$$\beta_{CS}(g_i) = \Lambda \frac{\partial}{\partial \Lambda} g_i. \quad (4.10)$$

In general, the beta function, $\beta(g)$, depends on the renormalization scheme. However, the first two terms are scheme independent under the perturbative expansion. The two loop beta function can be calculated for arbitrary N_c and N_f

$$\beta(a) = -\beta_0 a^2 - \beta_1 a^3 + \dots, \quad (4.11)$$

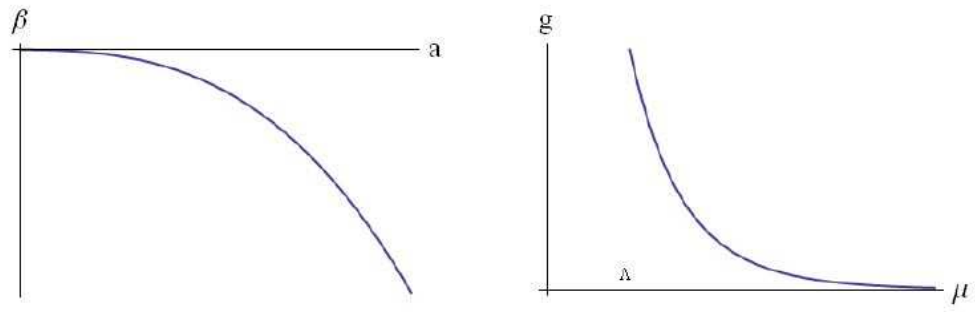
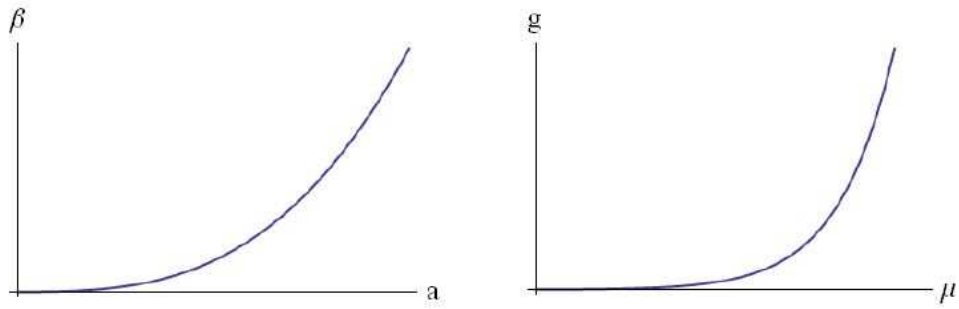
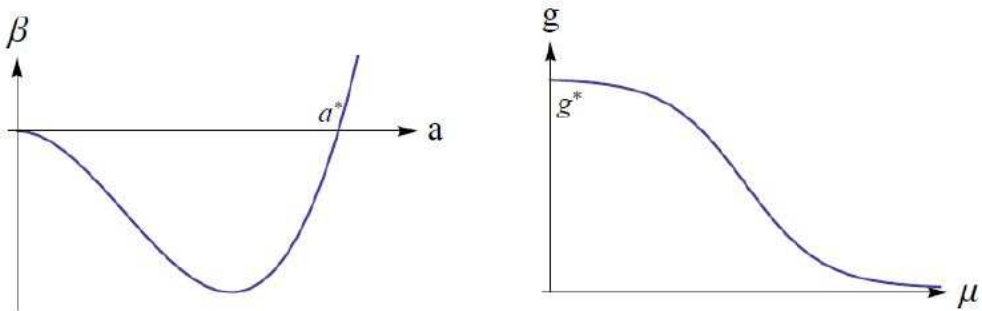
$$\beta_0 = \frac{11}{3} N_c - \frac{2}{3} N_f, \quad (4.12)$$

$$\beta_1 = \frac{34 N_c^2}{3} - \frac{10 N_c N_f}{3} - \frac{(-1 + N_c^2) N_f}{N_c}, \quad (4.13)$$

where $a \equiv \frac{g^2}{4\pi^2} = \frac{\alpha}{4\pi}$. The UV and IR properties of the theory depend on the signs of the coefficients. This is illustrated in Fig. 4.7, 4.8, and 4.9.

The partition function for full QCD with one fermion flavor is

$$Z = \int [dU][d\bar{\psi}][d\psi] \exp[-S_G(U) - \bar{\psi} M(U) \psi], \quad M = D + m. \quad (4.14)$$

Figure 4.7: $\beta_0 > 0$ and $\beta_1 > 0$.Figure 4.8: $\beta_0 < 0$ and $\beta_1 < 0$.Figure 4.9: $\beta_0 > 0$ and $\beta_1 < 0$.

N_f	Volume	Num. of β used	Num. of config. per β
4	4^4	21	25,000
4	6^4	35	8,000
4	8^4	36	25,000
4	12^4	21	25,000
4	16^4	5	2,500
12	4^4	31	50,000
12	6^4	41	50,000
12	8^4	15	8,000

Integrating over Grassmann variables

$$Z = \int [dU] \exp[-S_G(U)] \det M(U). \quad (4.15)$$

Introducing a pseudo fermion field Φ

$$Z = \int [dU][d\Phi^*][d\Phi] \exp[-S_G(U)] - \Phi^* M(U) \Phi. \quad (4.16)$$

For 2 flavors, one would have

$$Z = \int [dU] \exp[-S_G(U)] \det[M^\dagger M], \quad \det D = \det D^\dagger. \quad (4.17)$$

For staggered fermion

$$Z = \int [dU] \exp[-S_G(U)] \prod_f \det[M^\dagger M]^{N_f/8}. \quad (4.18)$$

We use standard Wilson gauge and naive staggered fermion action with the Hybrid Monte Carlo (HMC) algorithm. Focus on $N_f = 4$ and $N_f = 12$ with relatively small symmetric lattices. Bare quark mass is set to be $m_q = 0.02$ for now. The following is the configurations used to calculate Fisher zeros

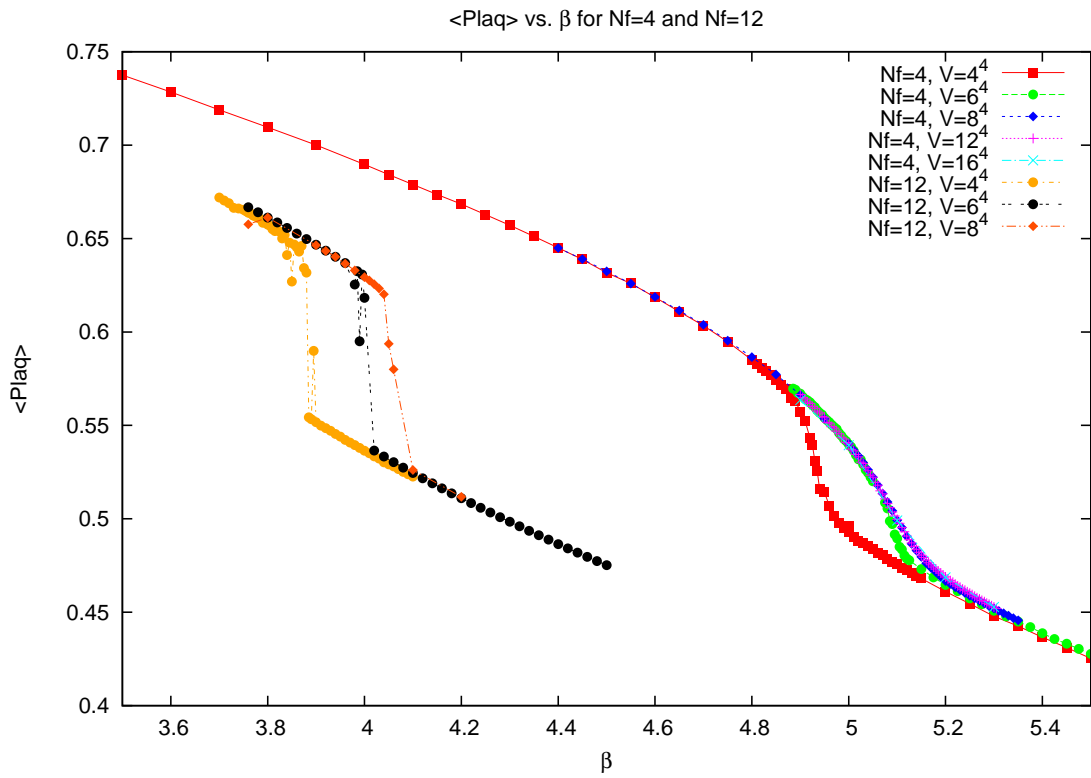


Figure 4.10: Average plaquette for $N_f = 4$ and $N_f = 12$ at different volumes.

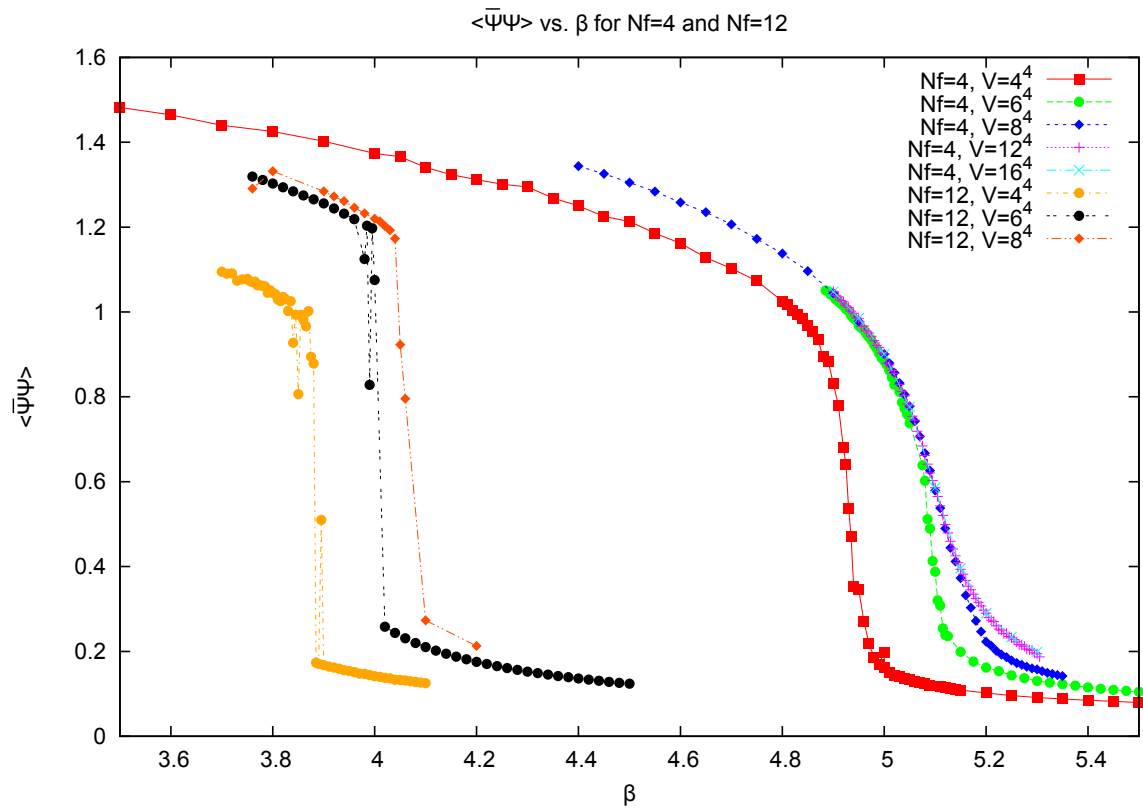


Figure 4.11: $\langle \bar{\Psi}\Psi \rangle$ for $N_f = 4$ and $N_f = 12$ at different volumes.

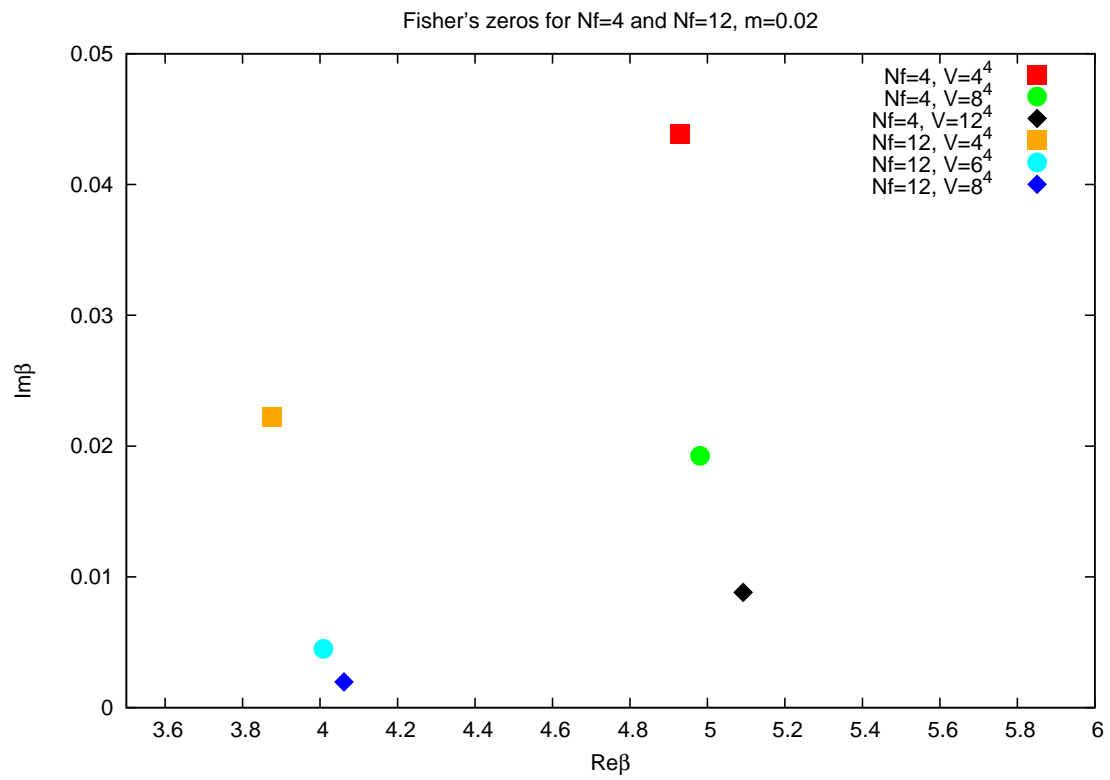


Figure 4.12: Fisher zeros for $N_f = 4$ and $N_f = 12$ at different volumes.

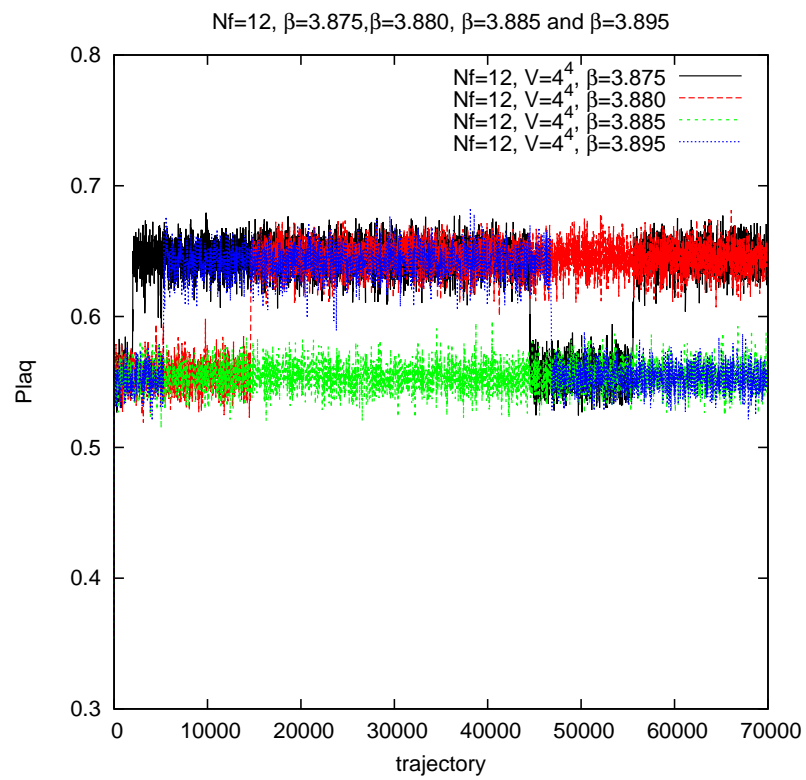


Figure 4.13: Typical plaquette history for the $V = 4^4$ system.

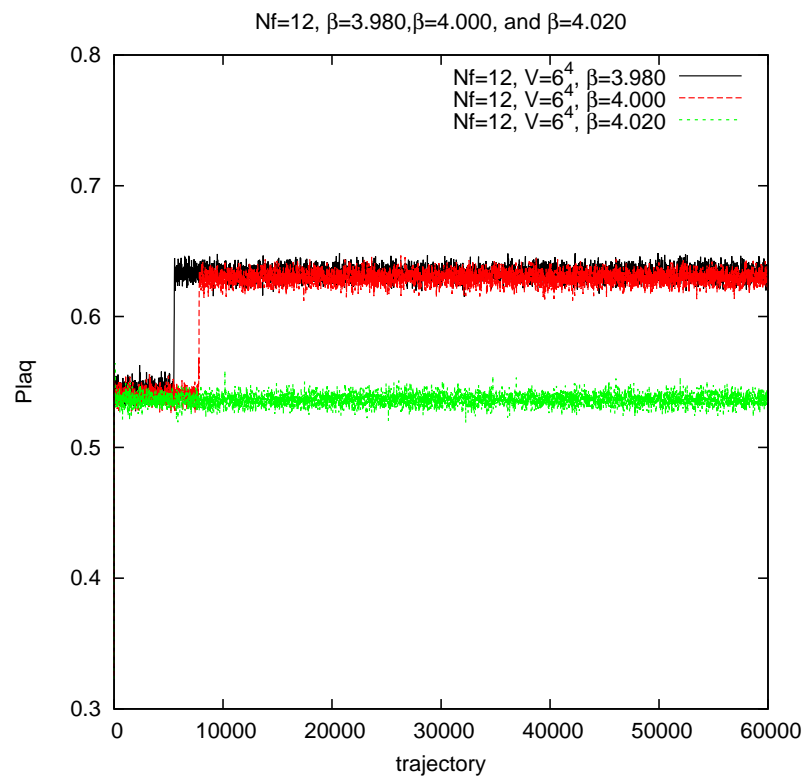


Figure 4.14: Typical plaquette history for the $V = 6^4$ system.

CHAPTER 5

TENSOR RENORMALIZATION GROUP

5.1 TRG on 2D $O(2)$ model

The Hamiltonian for the two dimensional $O(2)$ model can be written as

$$H = - \sum_{\langle ij \rangle} \mathbf{S}_i \cdot \mathbf{S}_j, \quad (5.1)$$

where \mathbf{S}_i is the unit vector staying at each site i : $(\cos(\theta), \sin(\theta))$.

H then can be written as

$$H = - \sum_{\langle ij \rangle} \cos(\theta_i - \theta_j) \quad (\pi < \theta_i < \pi). \quad (5.2)$$

The corresponding partition function is

$$Z = \int \prod_i \frac{d\theta_i}{2\pi} e^{\beta \sum_{\langle ij \rangle} \cos(\theta_i - \theta_j)}. \quad (5.3)$$

We can further expand the $e^{\beta \cos \theta}$ term as sum of modified Bessel function of the first kind

$$e^{\beta \cos \theta} = I_0(\beta) + \sum_{n=1}^{\infty} I_n(\beta) (e^{in\theta} + e^{-in\theta}) \quad (5.4)$$

$$= I_0(\beta) \left[1 + \sum_{n=-\infty, n \neq 0}^{\infty} \frac{I_n(\beta)}{I_0(\beta)} e^{in\theta} \right] \quad (5.5)$$

$$= I_0(\beta) \sum_{n=-\infty}^{+\infty} \frac{I_n(\beta)}{I_0(\beta)} e^{in\theta}. \quad (5.6)$$

Modified Bessel functions listed above have the following asymptotic properties

$$\frac{I_n(\beta)}{I_0(\beta)} = \frac{I_{-n}(\beta)}{I_0(\beta)} \sim \begin{cases} \frac{\beta^n}{2^n n!} & \beta \rightarrow 0 \\ e^{-\frac{n^2}{2\beta}} & \beta \rightarrow \infty. \end{cases} \quad (5.7)$$

By using Eq. (5.7), one could perform conventional high temperature expansion and so on. However, in the following, we would like to write down the partition function as product of matrices

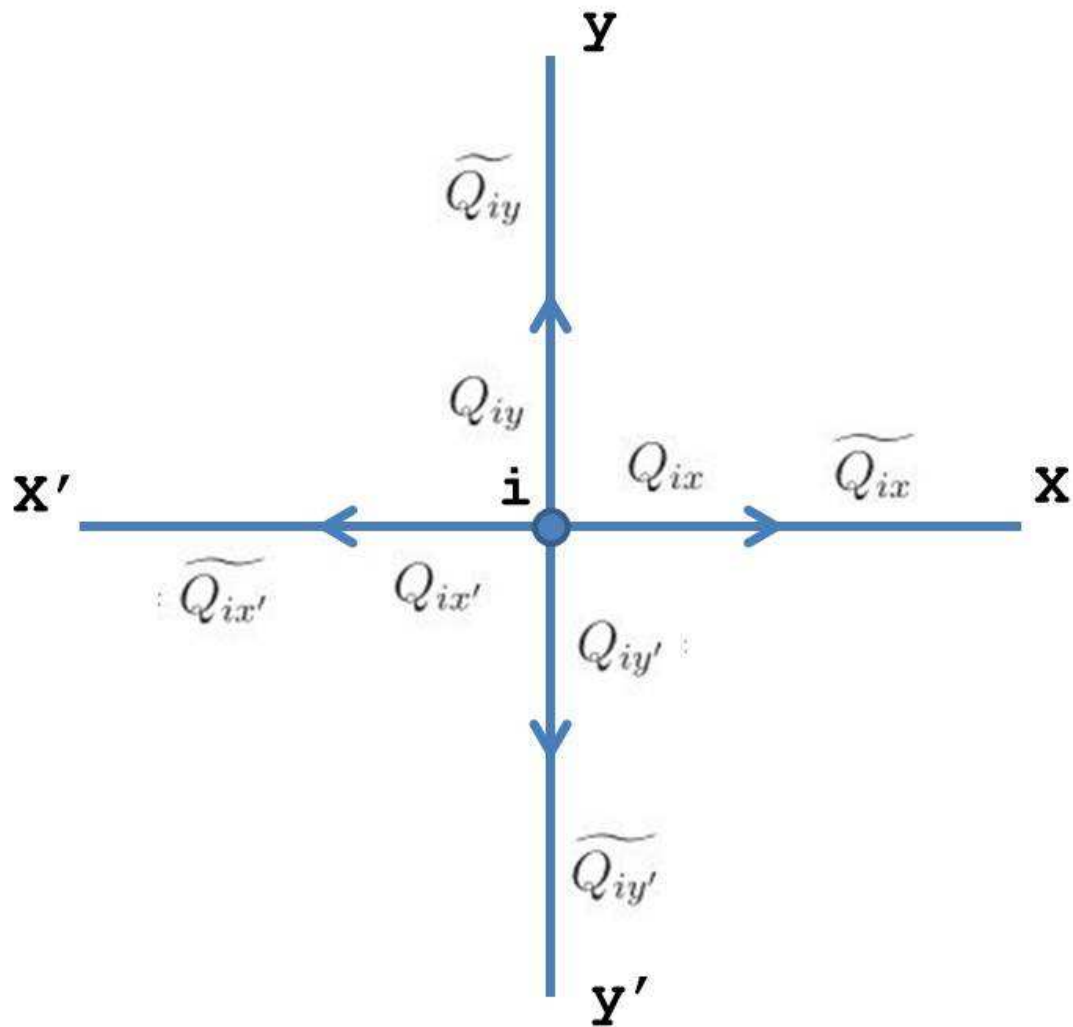


Figure 5.1: Tensor representation of the two dimensional $O(2)$ model.

$$\begin{aligned}
Z(\beta) &= \int \prod_k \frac{d\theta_k}{2\pi} \prod_{\langle ij \rangle} e^{\beta \cos(\theta_i - \theta_j)} \\
&= \int \prod_k \frac{d\theta_k}{2\pi} \prod_{\langle ij \rangle} \sum_{n_{ij}=-\infty}^{+\infty} I_{n_{ij}}(\beta) e^{in_{ij}(\theta_i - \theta_j)} \\
&= \int \prod_k \frac{d\theta_k}{2\pi} \prod_{\langle ij \rangle} W(\theta_i, \theta_j).
\end{aligned} \tag{5.8}$$

where $W(\theta_i, \theta_j) = e^{\beta \cos(\theta_i - \theta_j)} = \sum_{n_{ij}=-\infty}^{+\infty} e^{in_{ij}(\theta_i - \theta_j)} I_{n_{ij}}(\beta)$.

$W(\theta_i, \theta_j)$ can be further written as

$$W(\theta_i, \theta_j) = \sum_{n_{ij}} U(\theta_j, n_{ij}) \Lambda_{n_{ij}} V(\theta_i, n_{ij}), \tag{5.9}$$

where

$$U(\theta_j, n_{ij}) = e^{-in_{ij}\theta_j}, \tag{5.10}$$

$$\Lambda_{n_{ij}} = I_{n_{ij}}(\beta), \tag{5.11}$$

$$V(\theta_i, n_{ij}) = e^{+in_{ij}\theta_i}. \tag{5.12}$$

We define

$$Q_{ij} = \sqrt{I_{n_{ij}}(\beta)} e^{in_{ij}\theta_i} \tag{5.13}$$

$$\widetilde{Q}_{ij} = \sqrt{I_{n_{ij}}(\beta)} e^{-in_{ij}\theta_j}, \tag{5.14}$$

and write $W(\theta_i, \theta_j)$ as

$$W(\theta_i, \theta_j) = \sum_{n_{ij}} Q_{ij} \widetilde{Q}_{ij}. \tag{5.15}$$

We can then assign each link with a summation over product of Q and \widetilde{Q} , see Fig. 5.1 for example.

The tensor T at site i can be defined as product of all the Q and \widetilde{Q} that connect to i and integrate over the variable θ_i . Since $\frac{1}{2\pi} \int d\theta e^{i\theta(n_1 - n_2)} = \delta(n_1 - n_2)$, we will get element of tensor T as sum of delta function times modified Bessel function. In principle, we have infinite terms in the summation. In practice, we choose to keep

only the first several terms with the constraint $|n_{ij}| \leq n_{max}$. The element of T is

$$T_{xx'yy'}^i(n_{ix}, n_{iy}, n_{ix'}, n_{iy'}) = \sqrt{I_{n_{ix}}(\beta)} \sqrt{I_{n_{iy}}(\beta)} \sqrt{I_{n_{ix'}}(\beta)} \sqrt{I_{n_{iy'}}(\beta)} \quad (5.16)$$

$$* \delta(n_{ix} + n_{iy} + n_{ix'} + n_{iy'}). \quad (5.17)$$

We can apply similar trick to higher dimensions.

5.2 TRG on 2D O(3) model

The Hamiltonian for the two dimensional $O(3)$ model is defined as

$$H = - \sum_{\langle ij \rangle} \mathbf{S}_i \cdot \mathbf{S}_j, \quad (5.18)$$

where \mathbf{S}_i and \mathbf{S}_j are unit vectors staying at each site i : $(\sin\theta\cos\phi, \sin\theta\sin\phi, \cos\theta)$ and the summation is taken over all the nearest neighbor pairs of spins in the lattice.

H can then be written as

$$H = - \sum_{\langle ij \rangle} \cos\gamma_{ij}, \quad (5.19)$$

where γ_{ij} is the angle between \mathbf{S}_i and \mathbf{S}_j and $\cos\gamma_{ij}$ can be expressed in terms of θ and ϕ

$$\cos\gamma_{ij} = \cos\theta_i \cos\theta_j + \sin\theta_i \sin\theta_j \cos(\phi_i - \phi_j). \quad (5.20)$$

The partition function can be written as

$$\begin{aligned} Z_N &= \int \prod_{k=1}^N \frac{d\Omega_k}{4\pi} e^{-\beta H} \\ &= \int \prod_{k=1}^N \frac{d\Omega_k}{4\pi} e^{\beta \sum_{\langle ij \rangle} \cos\gamma_{ij}} \\ &= \int \prod_{k=1}^N \frac{d\Omega_k}{4\pi} \prod_{\langle ij \rangle} e^{\beta \cos\gamma_{ij}}. \end{aligned} \quad (5.21)$$

We can further expand $e^{\beta \cos\gamma}$ in terms of modified Bessel function of the first kind

as

$$\begin{aligned} e^{\beta \cos \gamma} &= I_0(\beta) + \sum_{n=1}^{\infty} I_n(\beta)(e^{in\gamma} + e^{-in\gamma}) \\ &= I_0(\beta) + 2 \sum_{n=1}^{\infty} I_n(\beta) \cos(n\gamma). \end{aligned} \quad (5.22)$$

In the following, three more special functions will be involved: Chebyshev polynomials of the first kind $T_n(x)$; Legendre polynomials $P_n(x)$; and spherical harmonic $Y_{lm}(\theta, \phi)$.

By definition, Chebyshev polynomials of the first kind satisfy

$$T_n(\cos \gamma) = \cos(n\gamma) \quad (5.23)$$

and the right hand side of the above equation Eq. 5.23 can be written as a sum of the Legendre polynomials $P_n(\cos \gamma)$

$$\begin{aligned} T_n(\cos \gamma) &= \cos(n\gamma) \\ &= \sum_{l=0}^n a_{nl} P_l(\cos \gamma). \end{aligned} \quad (5.24)$$

The coefficient a_{nl} can be obtained by using the orthogonality property of the Legendre polynomials

$$\int_{-1}^1 P_m(x) P_n(x) dx = \frac{2}{2m+1} \delta_{mn}. \quad (5.25)$$

Therefore,

$$a_{nl} = \frac{2l+1}{2} \int_{-1}^1 T_n(x) P_l(x) dx. \quad (5.26)$$

Combining Eq. (5.22) and Eq. (5.24), one would get

$$e^{\beta \cos \gamma_{ij}} = I_0(\beta) + 2 \sum_{n=1}^{\infty} I_n(\beta) \sum_{l=0}^n a_{nl} P_l(\cos \gamma_{ij}). \quad (5.27)$$

By using the following addition theorem for spherical harmonic

$$P_l(\cos \gamma_{ij}) = \frac{4\pi}{2l+1} \sum_{m=-l}^l Y_{lm}^*(\theta_j, \phi_j) Y_{lm}(\theta_i, \phi_i). \quad (5.28)$$

Eq. (5.27) can be written as

$$e^{\beta \cos \gamma_{ij}} = I_0(\beta) + 2 \sum_{n=1}^{\infty} I_n(\beta) \sum_{l=0}^n a_{nl} \frac{4\pi}{2l+1} \sum_{m=-l}^l Y_{lm}^*(\theta_j, \phi_j) Y_{lm}(\theta_i, \phi_i). \quad (5.29)$$

Finally, we get the expression of $e^{\beta \cos \gamma_{ij}}$ as functions of spherical harmonics

$$e^{\beta \cos \gamma_{ij}} = \sum_{l=0}^{\infty} A_l(\beta) \sum_{m=-l}^l Y_{lm}^*(\theta_j, \phi_j) Y_{lm}(\theta_i, \phi_i), \quad (5.30)$$

where

$$\begin{aligned} A_0(\beta) &= I_0(\beta) + 2 \sum_{n=1}^{\infty} I_n(\beta) a_{n0} 4\pi \\ A_1(\beta) &= 2 \sum_{n=1}^{\infty} I_n(\beta) a_{n1} \frac{4\pi}{3} \\ &\vdots \\ A_l(\beta) &= 2 \sum_{n=l}^{\infty} I_n(\beta) a_{nl} \frac{4\pi}{2l+1}. \end{aligned} \quad (5.31)$$

The element of the tensor T can then be written as

$$\begin{aligned} T_{(l_1, m_1), (l_2, m_2), (l_3, m_3), (l_4, m_4)} &= \\ \int_0^{2\pi} d\phi \int_0^\pi d\theta \sin \theta Y_{l_1 m_1}(\theta, \phi) Y_{l_2 m_2}(\theta, \phi) \\ Y_{l_3 m_3}^*(\theta, \phi) Y_{l_4 m_4}^*(\theta, \phi) \sqrt{A_{l_1} A_{l_2} A_{l_3} A_{l_4}}. \end{aligned} \quad (5.32)$$

The direction convention is shown in Figure 5.2. Equation (5.32) can be further simplified by expanding the product of two spherical harmonics in terms of spherical harmonics themselves

$$Y_{l_1 m_1}(\theta, \phi) Y_{l_2 m_2}(\theta, \phi) = \sum_{L=l_{\min}}^{l_{\max}} G_L^{(m_1, m_2, l_1, l_2)} Y_L^{m_1+m_2}(\theta, \phi), \quad (5.33)$$

where $G_L^{(m_1, m_2, l_1, l_2)}$ is Gaunt coefficients. Different methods have been developed to effectively calculate these coefficients. The summation bounds are given in the

following

$$\begin{aligned}
l_{max} &= l_1 + l_2 \\
l_{min} &= \begin{cases} \lambda_{min} & \text{if } l_{max} + \lambda_{min} \text{ is even} \\ \lambda_{min} + 1 & \text{if } l_{max} + \lambda_{min} \text{ is odd} \end{cases} \\
\lambda_{min} &= \max(|l_1 - l_2|, |m_1 + m_2|).
\end{aligned} \tag{5.34}$$

The angular integration in Eq. (5.32) can be performed in the following

$$\begin{aligned}
& \int_0^{2\pi} d\phi \int_0^\pi d\theta \sin\theta Y_{l_1 m_1}(\theta, \phi) Y_{l_2 m_2}(\theta, \phi) Y_{l_3 m_3}^*(\theta, \phi) Y_{l_4 m_4}^*(\theta, \phi) \\
&= \int_0^{2\pi} d\phi \int_0^\pi d\theta \sin\theta \sum_{L=l_{min}}^{l_{max}} G_L^{(m_1, m_2, l_1, l_2)} Y_L^{m_1+m_2}(\theta, \phi)
\end{aligned} \tag{5.35}$$

$$\sum_{L'=l'_{min}}^{l'_{max}} G_{L'}^{*(m_3, m_4, l_3, l_4)} Y_{L'}^{*m_3+m_4}(\theta, \phi). \tag{5.36}$$

By using the orthonormal property of the spherical harmonics

$$\int_0^{2\pi} d\phi \int_0^\pi d\theta \sin\theta Y_{l_1, m_1} Y_{l_2, m_2}^* = \delta_{l_1 l_2} \delta_{m_1 m_2} \tag{5.37}$$

along with Equation (5.36), the element of the tensor T (Eq. (5.32)) can then be simplified as

$$\begin{aligned}
& T(l_1, m_1, l_2, m_2, l_3, m_3, l_4, m_4) \\
&= \delta_{m_1+m_2, m_3+m_4} \sum_{L, L'} \delta_{LL'} G_L^{(m_1, m_2, l_1, l_2)} G_{L'}^{*(m_3, m_4, l_3, l_4)} \sqrt{A_{l_1} A_{l_2} A_{l_3} A_{l_4}}.
\end{aligned} \tag{5.38}$$

To summarize, the expression of the element of the tensor T is Eq. (5.38). $G_L^{(m_1, m_2, l_1, l_2)}$ and $G_{L'}^{*(m_3, m_4, l_3, l_4)}$ are Gaunt coefficients and can be calculated numerically. The summation bounds are given in Eq. (5.34). A_{l_1} , A_{l_2} , A_{l_3} , and A_{l_4} are listed in Eq. (5.31), where the coefficients a_{nl} can be obtained from Eq. (5.26).

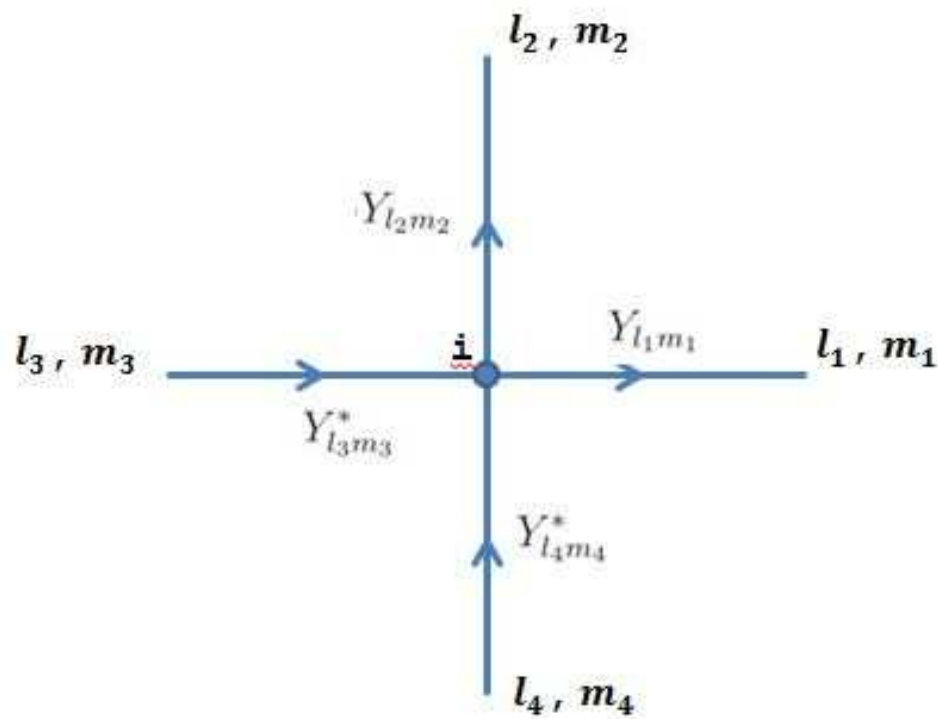


Figure 5.2: Tensor representation of the two dimensional $O(3)$ model.

CHAPTER 6 CONCLUSION

In previous chapters, we have discussed several different versions of the renormalization group (RG) methods. These methods have been first explored in spin models and some of them were then used in lattice gauge models.

In Chapter 2, we started with the conventional Ising model and then extensively discussed Dyson's hierarchical model and its variants. The partition function zeros have been calculated and finite size scaling technique has also been applied to get the critical exponents. We also applied the two lattice matching method to construct the RG flows in the complex temperature plane. More importantly, we found out that Fisher zeros, which are partition function zeros in the complex temperature plane, act as a gate controlling the flows. We further discussed the possibility of extending the RG method from discrete block-spinning procedure to the continuous one. We did not succeed here but discussed the obstacle to doing so and tried to quantitatively analyze them.

Inspired by the continuous block-spinning procedure, in Chapter 3, we studied the exact renormalization group (ERG) flow equations and the Wilson-Polchinski equation. We showed that the Wilson-Polchinski equation is equivalent to the ERG equation under the local potential approximation. We also systematically analyzed the cutoff function dependence of the critical exponents. The aim was to find the relation between the ERG equation and the hierarchical approximation and further make the connection between the discrete and continuous transformation.

In Chapter 4, we applied methods developed on spin models to lattice gauge models. We first applied the finite size scaling (FSS) technique to the pure gauge $SU(2)$ lattice gauge model and extrapolated the critical exponents ν and ω . We then analyzed Fisher zeros of the $SU(3)$ lattice gauge theory with many light flavors. This

is an on-going project and we are planning on working on it in a more systematic way.

In Chapter 5, we started to formulate the tensor renormalization group on the continuous spin models, $O(2)$ and $O(3)$ (or XY model) in two dimension. We have extended the method to higher dimensions lattice gauge models but I have not included these in the thesis. The advantages of this method are that they can be used to calculate thermodynamic quantities in arbitrary large system, and the systematical uncertainty can be improved systematically.

In summary, RG is an important tool in dealing with many degrees of freedoms, which includes lattice spin and gauge models. Combining with FSS and other numerical techniques, RG becomes an extremely powerful tool in dealing with many interesting but difficult problems in physics. An especially interesting prospect would be to study the conformal and near conformal lattice gauge theories. In those theories, there is more than one scaling variable due to the non-vanishing quark masses. Calculating the running of the coupling and the anomalous dimension associated with the running of the quark masses for those theories will be exciting projects for me in the next couple of years.

APPENDIX A

CRITICAL VALUES FOR THE HIERARCHICAL MODEL

In order to reliably get the critical exponents, such as ν and ω , the first step is to locate the critical coupling(β_c). It has been shown that the accuracy of γ depends on the accuracy of β_c and the dependence can be approximated by $\delta\gamma \approx |\beta - \beta_c|^{\frac{1}{1+\Delta}} = 10^{-SD}$, where SD is the accuracy goal for γ and Δ is defined as $\Delta = -\text{Log}[\lambda_2]/\text{Log}[\lambda_1]$. Table A.1 listed the calculated Δ for various l^D . From Table A.1, we can see that $1 + 1/\Delta$ is between 3.3 and 3.4. That means we need to calculate β up to 10^{-34} in order to get an accurate γ up to 10^{-10} ($SD = 10$).

In our calculation, we always calculate β up to 10^{-40} or more. Theoretically, we can calculate β up to any order. There should be no computer memory problem for this part of calculation.

l^D	λ_1	λ_2	Δ	$1 + 1/\Delta$
3	1.7572064	0.7865396	0.42593834	3.3477577
4	2.0365506	0.7386475	0.42591392	3.3478923
5	2.2832894	0.7035557	0.42587323	3.3481166
6	2.50674	0.67616	0.42582453	3.3483852

Table A.1: γ and Δ for different l^D .

The latest result for β_c , γ , and λ_i with $b^D = 2, 3, 4, 5, 6, 7$, and 8 are obtained from β_c which has accuracy up to 10^{-64} . This has been checked independently by Dr. Bugra Oktay.

l^D	γ_c	β_c
2	1.29914073	1.17903017
3	1.2992289425	1.95360681158131
4	1.2993834469	2.57436660736656
5	1.2995837185	3.11114443002212
6	1.2998	3.5936743173223

Table A.2: Critical values for different l^D .

All the digits shown for β_c are exact. The digits for γ and λ are just for reference.

l_{max}	aim	t_{val}	β_c
32	40	64	1.953606811581309802176259103054353836796
32	40	70	1.953606811581309802176259103054353836796
32	40	76	1.953606811581309802176259103054353836796
32	40	82	1.953606811581309802176259103054353836796
32	40	88	1.953606811581309802176259103054353836796
32	40	94	1.953606811581309802176259103054353836796
36	40	72	1.953606811581309802176259103053851550042
36	40	79	1.953606811581309802176259103053851550042
36	40	86	1.953606811581309802176259103053851550042
36	40	93	1.953606811581309802176259103053851550042
36	40	100	1.953606811581309802176259103053851550042
36	40	107	1.953606811581309802176259103053851550042

Table A.3: β_c for $l^D = 3$. l_{max} is the initial and final truncation value; aim is the precision of β_c ; t_{val} is the intermediate truncation value.

l_{max}	aim	t_{val}	β_c
40	40	80	1.953606811581309802176259103053851542680
40	40	88	1.953606811581309802176259103053851542680
40	40	96	1.953606811581309802176259103053851542680
40	40	104	1.953606811581309802176259103053851542680
40	40	112	1.953606811581309802176259103053851542680
40	40	120	1.953606811581309802176259103053851542680
44	40	88	1.953606811581309802176259103053851542680
44	40	97	1.953606811581309802176259103053851542679
44	40	106	1.953606811581309802176259103053851542679
44	40	115	1.953606811581309802176259103053851542680
44	40	124	1.953606811581309802176259103053851542680
48	40	96	1.953606811581309802176259103053851542679
48	40	106	1.953606811581309802176259103053851542679
48	40	116	1.953606811581309802176259103053851542680
48	40	126	1.953606811581309802176259103053851542680
48	40	136	1.953606811581309802176259103053851542679
44	44	88	1.953606811581309802176259103053851542679

Table A.4: β_c for $l^D = 3$. l_{max} is the initial and final truncation value; aim is the precision of β_c ; t_{val} is the intermediate truncation value.

l_{max}	aim	t_{val}	γ
32	40	64	1.299228942539096704751713479952837548239
32	40	70	1.299228942538789997108335021215099666667
32	40	76	1.299228942538615134568811785928293767316
32	40	82	1.299228942538635863061188181038703171005
32	40	88	1.299228942539059636165227923786573772275
32	40	94	1.299228942538904499736457717354803732439
36	40	72	1.299228942538525734311222716384997381804
36	40	79	1.299228942538584986703836967414504578432
36	40	86	1.299228942539040279508918499040205580723
36	40	93	1.299228942538599666527978413932297962763
36	40	100	1.299228942538580056290327303744108375970
36	40	107	1.299228942538615900535154791106016243179

Table A.5: γ for $l^D = 3$. l_{max} is the initial and final truncation value; aim is the precision of β_c ; t_{val} is the intermediate truncation value.

l_{max}	aim	t_{val}	γ
40	40	80	1.299228942538560512974344369671636794173
40	40	88	1.299228942538997274114849595551652796182
40	40	96	1.299228942538548445718520535494144766115
40	40	104	1.299228942538537050896258421727505797021
40	40	112	1.299228942538591426805662193566184090402
40	40	120	1.299228942539051596097055241722107819011
44	40	88	1.299228942538952305920170000502989354029
44	40	97	1.299228942539069502598778321768812111667
44	40	106	1.299228942539194839714669287059491647935
44	40	115	1.299228942538473931879181756076576634668
44	40	124	1.299228942538880749149786615545974693001
48	40	96	1.299228942539095015164526158475210808315
48	40	106	1.299228942539165246170459726584161677860
48	40	116	1.299228942538602536437857185472522101024
48	40	126	1.299228942538603869472061544030207426767
48	40	136	1.299228942539043022801978917931281023475
44	44	88	1.299228942538617960143482614595703213070

Table A.6: γ for $l^D = 3$. l_{max} is the initial and final truncation value; aim is the precision of β_c ; t_{val} is the intermediate truncation value.

l_{max}	aim	t_{val}	λ_1
32	40	64	1.757206398881104776634947198492592259406
32	40	70	1.757206398881338622230979649774077358276
32	40	76	1.757206398881471944095176303096928190400
32	40	82	1.757206398881456139902619438635159199953
32	40	88	1.757206398881133039137978780437636844533
32	40	94	1.757206398881251321063569226992441832437
36	40	72	1.757206398881540106258633017023288673202
36	40	79	1.757206398881494929976377339437821323472
36	40	86	1.757206398881147797390437087766913471547
36	40	93	1.757206398881483737518999371120100046028
36	40	100	1.757206398881498689111536654311404640191
36	40	107	1.757206398881471360093245187418656220222

Table A.7: λ_1 for $l^D = 3$. l_{max} is the initial and final truncation value; aim is the precision of β_c ; t_{val} is the intermediate truncation value.

l_{max}	aim	t_{val}	λ_1
40	40	80	1.757206398881513589680443715631324328457
40	40	88	1.757206398881180586341823742384974826926
40	40	96	1.757206398881522790216170569736413107470
40	40	104	1.757206398881531478062923322335495287328
40	40	112	1.757206398881490019797311562319092120003
40	40	120	1.757206398881139169192363052790607742378
44	40	88	1.757206398881214871807198913757615008738
44	40	97	1.757206398881125516593027202997292128121
44	40	106	1.757206398881029954799236710195417258758
44	40	115	1.757206398881579602407027375276335310787
44	40	124	1.757206398881269429415989292009118082317
48	40	96	1.757206398881106064840609488858982612833
48	40	106	1.757206398881052518045160956292319459983
48	40	116	1.757206398881481549390334663903625197793
48	40	126	1.757206398881480533034254925024151311184
48	40	136	1.757206398881145705799260099596530545119
44	44	88	1.757206398881469789769363759750127218183

Table A.8: λ_1 for $l^D = 3$. l_{max} is the initial and final truncation value; aim is the precision of β_c ; t_{val} is the intermediate truncation value.

l_{max}	aim	t_{val}	λ_2
32	40	64	0.7865396437135632124646513820461519491398
32	40	70	0.7865396437137473349053485006524538461584
32	40	76	0.7865396437137582088903109454796719126079
32	40	82	0.7865396437137181357413199409974844743448
32	40	88	0.7865396437135465423920795604900117811111
32	40	94	0.7865396437136659571574214805271695904763
36	40	72	0.7865396437137944086620201717742327612720
36	40	79	0.7865396437137387364931654600178421334067
36	40	86	0.7865396437135860600599816895114634170213
36	40	93	0.7865396437138244031794673946506184545514
36	40	100	0.7865396437137724127159748242931991702469
36	40	107	0.7865396437137262189270360569547320945806

Table A.9: λ_2 for $l^D = 3$. l_{max} is the initial and final truncation value; aim is the precision of β_c ; t_{val} is the intermediate truncation value.

l_{max}	aim	t_{val}	λ_2
40	40	80	0.7865396437137486463462731885157631508033
40	40	88	0.7865396437136034737172745593040480607004
40	40	96	0.7865396437138297283746857789288758433631
40	40	104	0.7865396437137898263732676997543989045536
40	40	112	0.7865396437137361287801437848213971505008
40	40	120	0.7865396437135814777699735559420856758198
44	40	88	0.7865396437136465995740842830943387319001
44	40	97	0.7865396437135425472957373162902920662189
44	40	106	0.7865396437134917960439593542434177736157
44	40	115	0.7865396437138403018244167119981935273125
44	40	124	0.7865396437136951727405705670621413629338
48	40	96	0.7865396437135638966088823900835585835474
48	40	106	0.7865396437135037790020999267775010197373
48	40	116	0.7865396437137633101000482823479583475013
48	40	126	0.7865396437138348258652061309343931866066
48	40	136	0.7865396437135849492512972835102014507718
44	44	88	0.7865396437138697473782302056330013043077

Table A.10: λ_2 for $l^D = 3$. l_{max} is the initial and final truncation value; aim is the precision of β_c ; t_{val} is the intermediate truncation value.

l_{max}	aim	t_{val}	β_c
32	40	64	2.574366607366560276233393764322237051524
32	40	128	2.574366607366560276233393764322237048475
40	40	80	2.574366607366560276233393764322236128484

Table A.11: β_c for $l^D = 4$. l_{max} is the initial and final truncation value; aim is the precision of β_c ; t_{val} is the intermediate truncation value.

l_{max}	aim	t_{val}	γ
32	40	64	1.299383446968990648082420597389185027744
32	40	128	1.299383446969440744025709126999082084684
40	40	80	1.299383446969352427087654937834492881873

Table A.12: γ for $l^D = 4$. l_{max} is the initial and final truncation value; aim is the precision of β_c ; t_{val} is the intermediate truncation value.

l_{max}	aim	t_{val}	λ_1
32	40	64	2.036550640841265654441039283550142975455
32	40	128	2.036550640840763901588513926490994222893
40	40	80	2.036550640840862354543391085542627312959

Table A.13: λ_1 for $l^D = 4$. l_{max} is the initial and final truncation value; aim is the precision of β_c ; t_{val} is the intermediate truncation value.

l_{max}	aim	t_{val}	λ_2
32	40	64	0.7386474756161715818979884554824340680415
32	40	128	0.7386474756159031757269478294318271174266
40	40	80	0.7386474756159942496409988023234663687984

Table A.14: λ_2 for $l^D = 4$. l_{max} is the initial and final truncation value; aim is the precision of β_c ; t_{val} is the intermediate truncation value.

l_{max}	aim	t_{val}	β_c
16	40	32	3.111144430022622778922915294803883518779
16	40	44	3.111144430022123178568317034525175980765
16	40	56	3.111144430022123178453278705116288715924
16	40	68	3.111144430022123178453278700620123711324
16	40	80	3.111144430022123178453278700620123700812
20	40	40	3.111144430022121482846073290482461542807
20	40	55	3.111144430022121460055934935111232744852
20	40	70	3.11114443002212146005593491051657647209
20	40	85	3.11114443002212146005593491051657646914
20	40	100	3.11114443002212146005593491051657646914
24	40	48	3.111144430022121460029837242613994484801
24	40	66	3.111144430022121460029336989460676552477

Table A.15: β_c for $l^D = 5$. l_{max} is the initial and final truncation value; aim is the precision of β_c ; t_{val} is the intermediate truncation value.

l_{max}	aim	t_{val}	γ
16	40	32	1.299583719032215803258708342850073978291
16	40	44	1.299583718512967899988772557264971016965
16	40	56	1.299583718513015961760844982956631026984
16	40	68	1.299583718513000258998879319229193303839
16	40	80	1.299583718512884449786487925987916981462
20	40	40	1.299583718510391233336485024341542227967
20	40	55	1.299583718510885508385786028682654337013
20	40	70	1.299583718510860503268783721605417769867
20	40	85	1.299583718510772405623681749852427845554
20	40	100	1.299583718510866691792733336514085761198
24	40	48	1.299583718510494428821083077019940044704
24	40	66	1.299583718510317145590856700363726361095

Table A.16: γ for $l^D = 5$. l_{max} is the initial and final truncation value; aim is the precision of β_c ; t_{val} is the intermediate truncation value.

l_{max}	aim	t_{val}	λ_1
16	40	32	2.283289466193888103702553954133792956019
16	40	44	2.283289466947087857534438970978204392471
16	40	56	2.283289466947018141095169520848258742954
16	40	68	2.283289466947040918878920234229090506686
16	40	80	2.283289466947208906982029741595735355549
20	40	40	2.28328946695082546471029663501288085322
20	40	55	2.283289466950108489555796570981572785119
20	40	70	2.283289466950144760954992961127449982234
20	40	85	2.283289466950272551792902701444930541026
20	40	100	2.283289466950135784135464694263289243455
24	40	48	2.283289466950675773564409213192212199554
24	40	66	2.283289466950932933361493133742513498114

Table A.17: λ_1 for $l^D = 5$. l_{max} is the initial and final truncation value; aim is the precision of β_c ; t_{val} is the intermediate truncation value.

l_{max}	aim	t_{val}	λ_2
16	40	32	0.7035557328641656688813538819335051668311
16	40	44	0.7035557367803160850675204604737219646894
16	40	56	0.7035557367802940352394934374136841685974
16	40	68	0.7035557367803026871247587308613526214851
16	40	80	0.7035557367804147050569049496755473998989
20	40	40	0.703555736808953377405725899832537745963
20	40	55	0.7035557368091700380359497409605427778522
20	40	70	0.7035557368091838153216202672322253857599
20	40	85	0.70355573680928056477538767565833223638
20	40	100	0.7035557368091804055777899888779504020749
24	40	48	0.7035557368094688695865932830722341836117
24	40	66	0.7035557368095326614060477192309255003617

Table A.18: λ_2 for $l^D = 5$. l_{max} is the initial and final truncation value; aim is the precision of β_c ; t_{val} is the intermediate truncation value.

APPENDIX B

RESCALING OF THE ERG EQUATION

In the following, we will show that the equation (B.1) is invariant under some rescaling.

$$-3u + \rho u' + \frac{1}{1 + u' + 2\rho u''} = 0 \quad (\text{B.1})$$

B.1 No A Dependence

Let us start with the following equation:

$$-3u + \rho u' + \frac{A}{1 + u' + 2\rho u''} = 0 \quad (\text{B.2})$$

We will show that the prefactor A can be eliminated after the rescaling of ρ and u .

$$-3\frac{u}{A} + \rho\frac{u'}{A} + \frac{1}{1 + u' + 2\rho u''} = 0 \quad (\text{B.3})$$

We first rescale the variable ρ

$$\begin{aligned} \alpha &\equiv \frac{\rho}{A} \\ u' &\equiv \frac{\partial u}{\partial \rho} = \frac{1}{A} \frac{\partial u}{\partial \alpha} \\ u'' &\equiv \frac{\partial^2 u}{\partial \rho^2} = \frac{1}{A^2} \frac{\partial^2 u}{\partial \alpha^2} \end{aligned} \quad (\text{B.4})$$

\Rightarrow

$$-3\frac{u}{A} + \frac{\alpha}{A} \frac{\partial u}{\partial \alpha} + \frac{1}{1 + \frac{1}{A} \frac{\partial u}{\partial \alpha} + \frac{2\alpha}{A} \frac{\partial^2 u}{\partial \alpha^2}} = 0 \quad (\text{B.5})$$

We further rescale the field u

$$v \equiv \frac{u}{A}$$

$$\frac{1}{A} \frac{\partial u}{\partial \alpha} = \frac{\partial v}{\partial \alpha} \quad (\text{B.6})$$

$$\frac{1}{A} \frac{\partial^2 u}{\partial \alpha^2} = \frac{\partial^2 v}{\partial \alpha^2} \quad (\text{B.7})$$

\Rightarrow

$$-3v + \frac{\partial v}{\partial \alpha} + \frac{1}{1 + \frac{\partial v}{\partial \alpha} + 2\alpha \frac{\partial v}{\partial \alpha}} = 0 \quad (\text{B.8})$$

$$\boxed{-3v + \alpha v' + \frac{1}{1 + v' + 2\alpha v''} = 0} \quad (\text{B.9})$$

Therefore, the coefficient A has no effect on the critical exponents ν and ω .

Note:

$$\alpha \equiv \frac{\rho}{A} \quad v \equiv \frac{u}{A} \Rightarrow \begin{cases} \rho \rightarrow A\alpha \\ u \rightarrow Av \\ u' \rightarrow v' \\ u'' \rightarrow \frac{1}{A}v'' = \frac{1}{A} \frac{\partial^2 v}{\partial \alpha^2} \end{cases} \quad (\text{B.10})$$

B.2 No B Dependence

$$-3u + \rho u' + \frac{1}{B + u' + 2\rho u''} = 0 \quad (\text{B.11})$$

By the same reasoning, we can prove that the factor B has no effect on the critical exponents. We can do the following substitution:

$$\alpha \equiv \rho B^2 \quad v \equiv uB \Rightarrow \begin{cases} \rho \rightarrow \frac{\alpha}{B^2} \\ u \rightarrow \frac{v}{B} \\ u' \rightarrow Bv' \\ u'' \rightarrow B^3v'' = B^3 \frac{\partial^2 v}{\partial \alpha^2} \end{cases} \quad (\text{B.12})$$

\Rightarrow

$$\boxed{-3v + \alpha v' + \frac{1}{1 + v' + 2\alpha v''} = 0} \quad (\text{B.13})$$

B.3 More about A and B dependence

Let us look at higher orders: $A \frac{1}{(1+u'+2\rho u'')^2}$, $\frac{1}{(B+u'+2\rho u'')^2}$

$$-3u + \rho u' + \frac{A}{(1 + u' + 2\rho u'')^2} = 0 \quad (\text{B.14})$$

$$\alpha \equiv \frac{\rho}{A}$$

$$-3\frac{u}{A} + \frac{\alpha}{A} \frac{\partial u}{\partial \alpha} + \frac{1}{(1 + \frac{1}{A} \frac{\partial u}{\partial \alpha} + \frac{2\alpha}{A} \frac{\partial^2 u}{\partial \alpha^2})^2} = 0 \quad (\text{B.15})$$

$$v \equiv \frac{u}{A}$$

$$\boxed{-3v + \alpha v' + \frac{1}{(1 + v' + 2\alpha v'')^2} = 0} \quad (\text{B.16})$$

So that the coefficient A has no effect on the critical exponents ν and ω even for higher orders.

For the coefficient B , we will have

$$-3u + \rho u' + \frac{1}{(B + u' + 2\rho u'')^2} = 0 \quad (\text{B.17})$$

\Rightarrow

$$\boxed{-3v + \rho v' + \frac{1}{B} \frac{1}{(1 + v' + 2\rho v'')^2} = 0} \quad (\text{B.18})$$

and

$$\boxed{-3v + \rho v' + \frac{1}{B^{n-1}} \frac{1}{(1 + v' + 2\rho v'')^n} = 0} \quad (\text{B.19})$$

APPENDIX C

LATTICE GAUGE THEORIES

Quantum Chromodynamics (QCD) is a non-Abelian gauge theory and is the best candidate for a theory of strong interactions. For now, only QCD is able to explain the experimental fact that the constituents of hadrons behave almost as free at short distance (asymptotic freedom). At long distance, the strong interactions are really strong! The perturbation theory breaks down. We would like not only to understand why “colored” object, such as quark, are not visible (confinement) but also to compute the real particles spectrum. Lattice QCD is a promising approach in dealing with the above problems by regularizing QCD on the space time lattice. In the Lattice QCD, fermion fields reside on the sites and gauge fields reside on the links. The fermion fields carry the color quantum number of the gauge group $SU(3)$. At each site, there are fermion fields ψ_α , where α is the color index. Gauge fields are the connections of the color coordinate.

Now consider a four-dimensional hyper-cubic lattice on the Euclidean space with lattice spacing a . $U_\nu(n)$ connects the neighboring lattice sites n and $n + \hat{\mu}$.

$$U_\nu(n) = \exp(igaA_\nu^c(n)\lambda_c) \tag{C.20}$$

where λ_c stands for 8 generators of $SU(3)$ and g is the gauge coupling constant.

$$\begin{aligned}
\lambda_1 &= \frac{1}{2} \begin{pmatrix} 0 & 1 & 0 \\ 1 & 0 & 0 \\ 0 & 0 & 0 \end{pmatrix} & \lambda_2 &= \frac{1}{2} \begin{pmatrix} 0 & -i & 0 \\ i & 0 & 0 \\ 0 & 0 & 0 \end{pmatrix} & \lambda_3 &= \frac{1}{2} \begin{pmatrix} 1 & 0 & 0 \\ 0 & -1 & 0 \\ 0 & 0 & 0 \end{pmatrix} \\
\lambda_4 &= \frac{1}{2} \begin{pmatrix} 0 & 0 & 1 \\ 0 & 0 & 0 \\ 1 & 0 & 0 \end{pmatrix} & \lambda_5 &= \frac{1}{2} \begin{pmatrix} 0 & 0 & -i \\ 0 & 0 & 0 \\ i & 0 & 0 \end{pmatrix} & \lambda_6 &= \frac{1}{2} \begin{pmatrix} 0 & 0 & 0 \\ 0 & 0 & 1 \\ 0 & 1 & 0 \end{pmatrix} \\
\lambda_7 &= \frac{1}{2} \begin{pmatrix} 0 & 0 & 0 \\ 0 & 0 & -i \\ 0 & i & 0 \end{pmatrix} & \lambda_8 &= \frac{1}{2\sqrt{3}} \begin{pmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & -2 \end{pmatrix}
\end{aligned}$$

The plaquette defined in the $\mu - \nu$ plane is :

$$U_{\mu\nu} = U_\mu(n)U_\nu(n + \hat{\mu})U_\mu^\dagger(n + \hat{\nu})U_\nu^\dagger(n) \quad (\text{C.21})$$

Recall that the continuum Yang Mills actions S_{YM} is the following

$$S_{YM} = -\frac{1}{2} \int_0^\beta d\tau \int d^3x \operatorname{tr} F^{\mu\nu} F_{\mu\nu}, \quad F_{\mu\nu}^a = F_{\mu\nu}^a T^a = \partial_\mu A_\nu^a - \partial_\nu A_\mu^a - g f^{abc} A_\mu^b A_\nu^c \quad (\text{C.22})$$

It can be written as

$$S(U) = \beta \sum_n \sum_{\mu < \nu} \left(1 - \frac{1}{3} \operatorname{Re} \operatorname{tr} U_{\mu\nu}(n) \right) \quad (\text{C.23})$$

on the lattice.

And the continuum free Euclidean fermion action is

$$S = \int d^d x \bar{\psi}(\not{\partial} + m)\psi \quad (\text{C.24})$$

The naive lattice fermion action can be obtained by replacing the derivatives by symmetric differences:

$$\partial_\mu \psi_\alpha(n) = \frac{1}{2a} (\psi_\alpha(n + \hat{\mu}) - \psi_\alpha(n - \hat{\mu})) \quad (\text{C.25})$$

$$S_{naive} = \sum_{n,\mu} \bar{\psi}(n) \frac{\gamma_\mu}{2a} (\psi(n + \hat{\mu}) - \bar{\psi}(n - \hat{\mu})) + m \sum_n \bar{\psi}(n) \psi(n) \quad (C.26)$$

The fermion propagator can be calculated as

$$G(p)_{naive} = (ma + i \sum_\mu \gamma_\mu \sin(p_\mu a))^{-1} \quad (C.27)$$

However, there are sixteen poles for the propagator $G(p)_{naive}$ in the $m = 0$ limit

$$p = (0, 0, 0, 0), \quad p = (0, 0, 0, \pi), \quad p = (0, 0, \pi, 0), \quad \cdots \quad p = (\pi, \pi, \pi, \pi) \quad (C.28)$$

This is a theory with 16 mass-less fermions instead of 1! And this is the so called fermion doubling problem: the number of mass-less fermions would be 2^d in a d space-time dimensions.

One way to deal with the doublers is to reduce the number of fermion flavors by using one component “staggered” fermion fields instead of four component Dirac spinors.

The staggered field $\chi(n)$ is defined as

$$\psi(n) = \eta_\mu(n) \chi(n), \quad \eta_\mu(n) = (-1)^{n_1 + n_2 + \cdots + n_{\mu-1}} \quad (C.29)$$

The resulting lattice action for “staggered” fermion action is:

$$S_{stag} = \frac{1}{2a} \sum_{n,\mu} \eta_\mu(n) \bar{\chi}(n) [U_\mu(n) \chi(n + \hat{\mu}) - U_\mu^\dagger(n - \hat{\mu}) \chi(n - \hat{\mu})] + m \sum_n \bar{\chi}(n) \chi(n) \quad (C.30)$$

S_{stag} is invariant under the chiral transformation as $m \rightarrow 0$.

There are also many different other fermion actions, such as Wilson fermion, Domain wall fermion, \cdots . We do not discuss them in details here.

The expectation value of an observable is:

$$\langle O \rangle = \frac{\int DU O(U) e^{-S[U]}}{\int DU e^{-S[U]}}$$

For a 10^4 lattice, there are about $8 * 4 * 10^4 = 320000$ integrations to be done.

Importance sampling methods: select a sequence of gauge configurations with a probability distribution given by the Boltzmann factor $e^{-S(U)}$.

The Monte Carlo method: Metropolis method and heat bath method.

Molecular dynamics method.

The Metropolis method:

I, Change the gauge configuration $\phi \rightarrow \phi'$

$$P(\phi' \leftarrow \phi) = P(\phi \leftarrow \phi')$$

II, Choose a random number r distributed uniformly on $[0,1]$.

III, Calculate $S(\phi)$ and $S(\phi')$.

IV, Always accept the change if $S(\phi) < S(\phi')$.

V, Accept the change if $e^{-\Delta S} > r$.

Molecular dynamics method:

I, Define a new Hamiltonian: $H(p, \phi) = \sum_n \frac{p_n^2}{2} + S(\phi)$

where p_n is the fictitious momentum conjugate to the field ϕ_n .

II, $H(p, \phi)$ governs the dynamics under the “evolution time” τ .

$$\dot{\phi}_n = p_n, \quad \dot{p}_n = -\partial S / \partial \phi_n$$

III, The above equations define a trajectory $[p(\tau), \phi(\tau)]$ through phase space.

IV, The expectation value of an observable can be computed as:

$$\langle O \rangle = \frac{1}{T} \int_{\tau_0}^{\tau_0+T} D\tau O[\phi(\tau)]$$

Molecular dynamics method for SU(N) pure gauge:

Link variable $U_\mu(n) = e^{iA_\mu(n)}$ can be treated as coordinates.

The equation of motion for $U_\mu(n)$:

$$\dot{U}_\mu(n) = iH_\mu(n)U_\mu(n), \quad H_\mu(n) = \sum_\alpha h_\mu^\alpha(n)T^\alpha$$

where T^α are generators of SU(N) Lie algebra and $h_\mu^\alpha(n)$ will be chosen from unit Gaussian random values.

We can also get

$$i\dot{H}_\mu(n) = -\frac{\beta}{N}Tr(U_\mu(n)V_\mu(n))|_{TA}$$

Hybrid method(Metropolis+Molecular dynamics):

I, Choose the coordinates $\{\phi_i\}$ in some arbitrary way.

II, Choose the momenta $\{\pi_i\}$ from the Gaussian ensemble:

$$P_G(\pi) = N_o e^{-\frac{1}{2}\sum \pi_i^2}$$

III, Calculate $\tilde{\pi}$ according to :

$$\tilde{\pi}_i = \pi_i(n) - \frac{\epsilon}{2} \frac{\partial S[\phi]}{\partial \phi_i(n)}$$

IV, Iterate the following form n times:

$$\phi_i(n+1) = \phi_i(n) + \epsilon \tilde{\pi}_i(n), \quad \tilde{\pi}_i(n+1) = \tilde{\pi}_i(n) - \epsilon \frac{\partial S[\phi]}{\partial \phi_i(n+1)}$$

V, Accept $\{\phi'_i, \pi'_i\}$ with probability: $p = \min(1, e^{-H[\phi', \pi']}/e^{-H[\phi, \pi]})$.

The Dirac fields:

I, Grassmann valued:

$$\{\eta_i, \eta_j\} = \eta_i \eta_j + \eta_j \eta_i = 0$$

II, The Dirac fields, ψ , enter quadratically into the action S .

$$S = \bar{\psi} M \psi + \dots$$

The partition function for full QCD with one fermion flavor is:

$$Z = \int [dU][d\bar{\psi}][d\psi] \exp[-S_G(U) - \bar{\psi} M(U) \psi], \quad M = D + m$$

Integrating over Grassmann variables:

$$Z = \int [dU] \exp[-S_G(U)] \det M(U)$$

Introducing a pseudo-fermion field Φ :

$$\det M(U) = \int [d\Phi^* d\Phi] \exp[-\Phi^* M^{-1} \Phi]$$

The pseudonym action can be written as:

$$Z = \int [dU][d\Phi^*][d\Phi] \exp[-S_G(U) - \Phi^* M(U) \Phi]$$

For 2 flavors, one would have:

$$Z = \int [dU] \exp[-S_G(U)] \det[M^\dagger M], \quad \det D = \det D^\dagger$$

For staggered fermion:

$$Z = \int [dU] \exp[-S_G(U)] \prod_f \det[M^\dagger M]^{N_f/4}$$

$$(M^\dagger M)^{-N_f/4} \approx r(M^\dagger M) = a_0 + \sum_n = 1^N \frac{a_n}{(M^\dagger M) + b_n}$$

where a_n and b_n are constants.

Rational approximation for $x^{1/8}$:

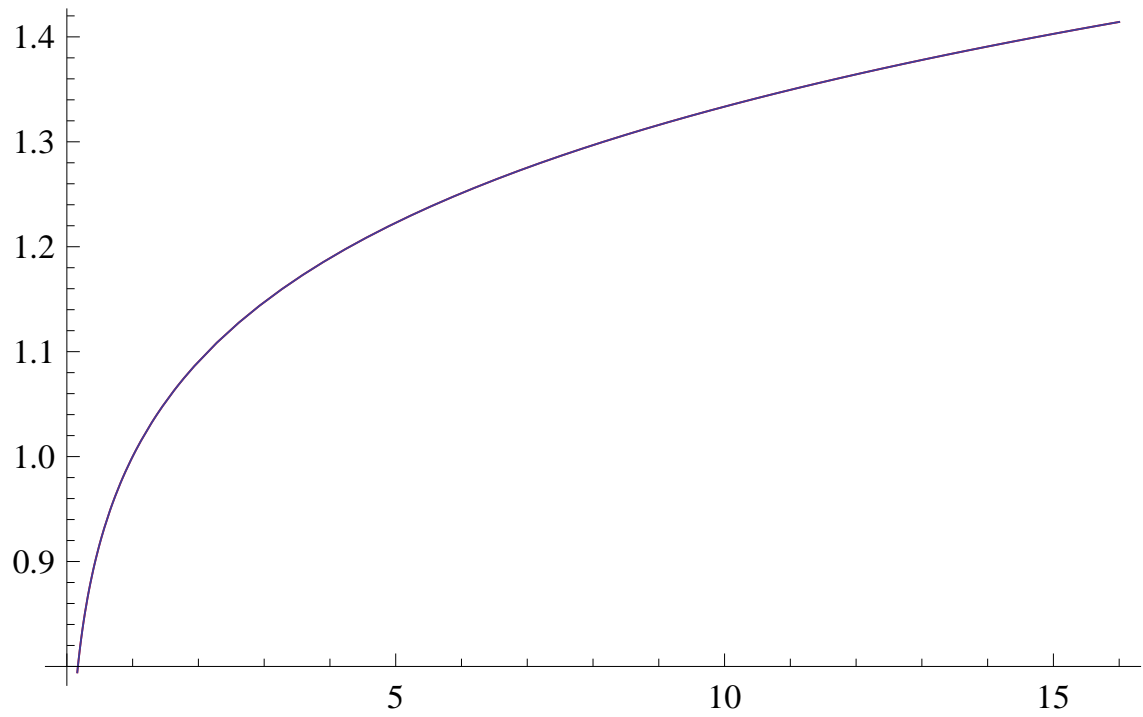


Figure C.1: Test for the rational approximation.

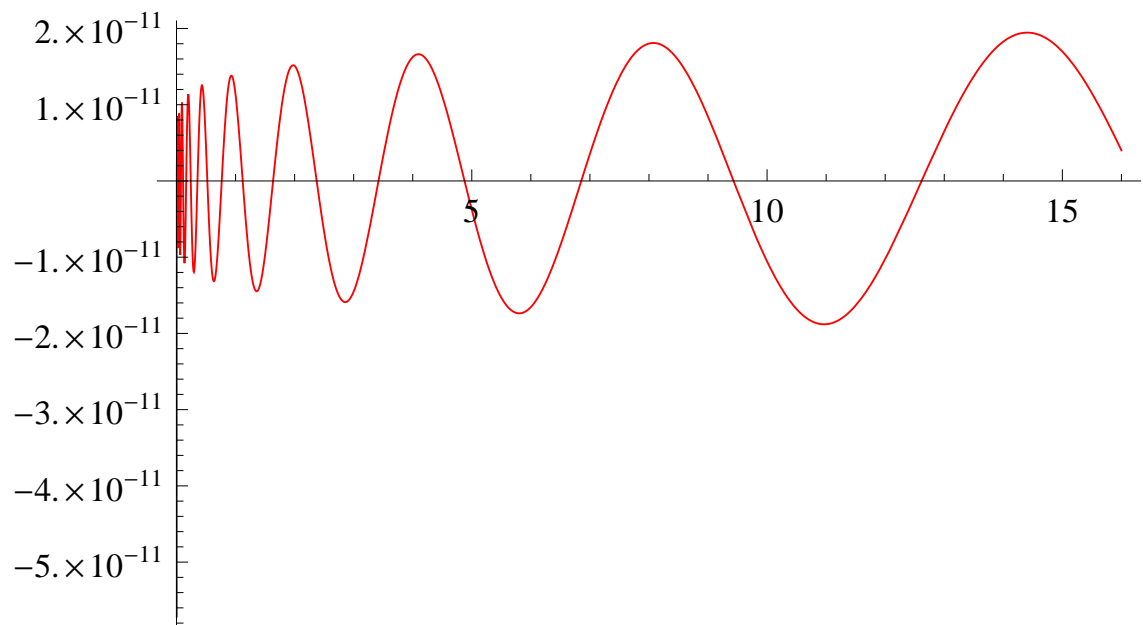


Figure C.2: Test for the rational approximation.

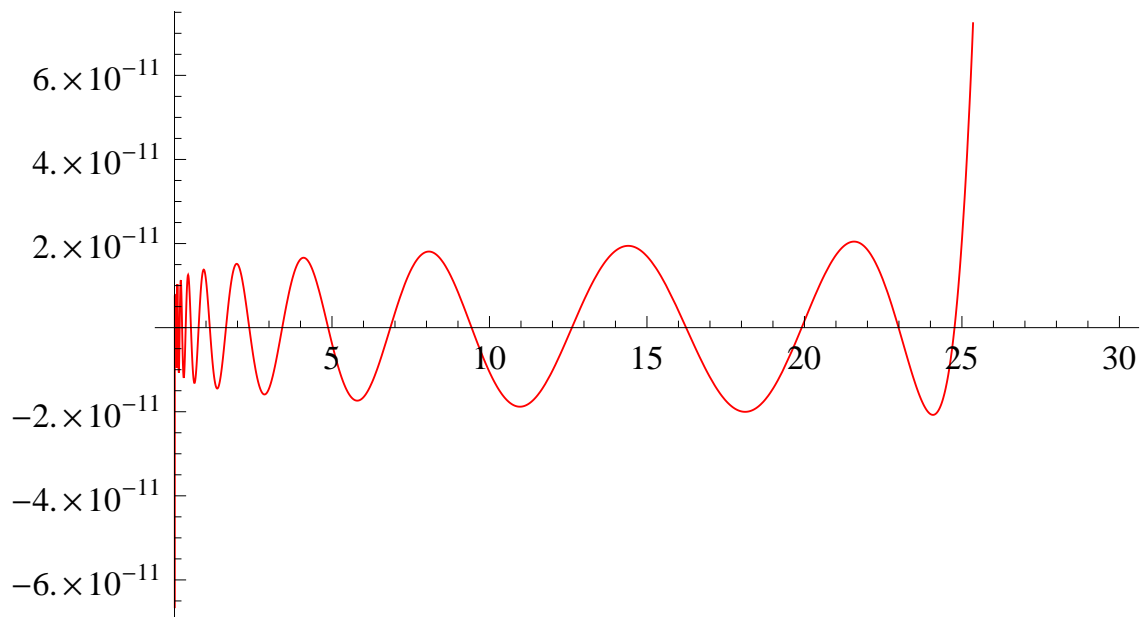


Figure C.3: Test for the rational approximation.

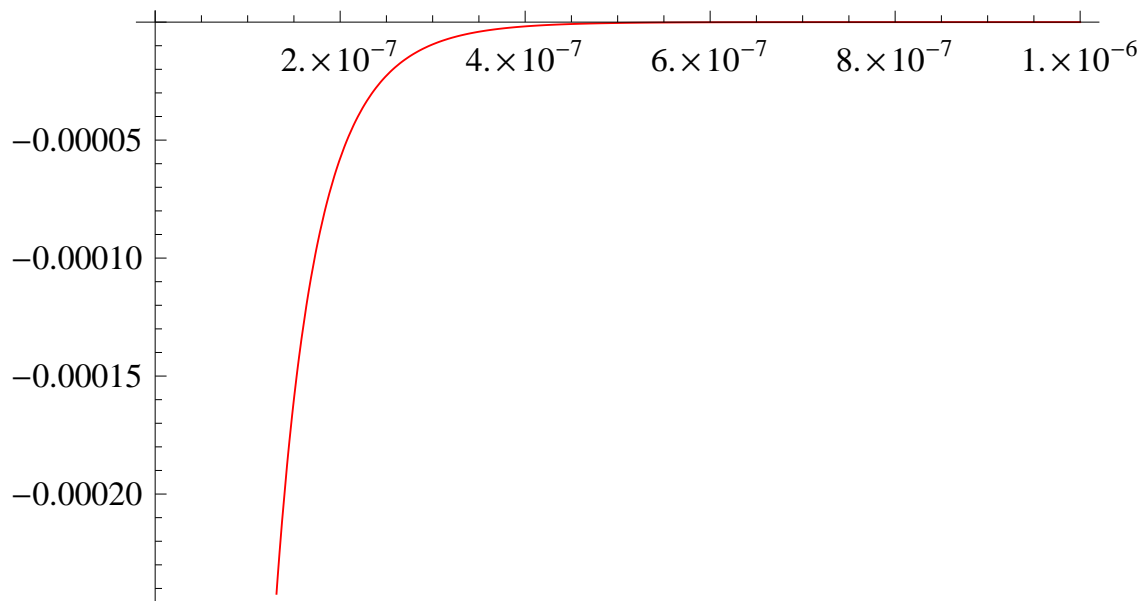


Figure C.4: Test for the rational approximation.

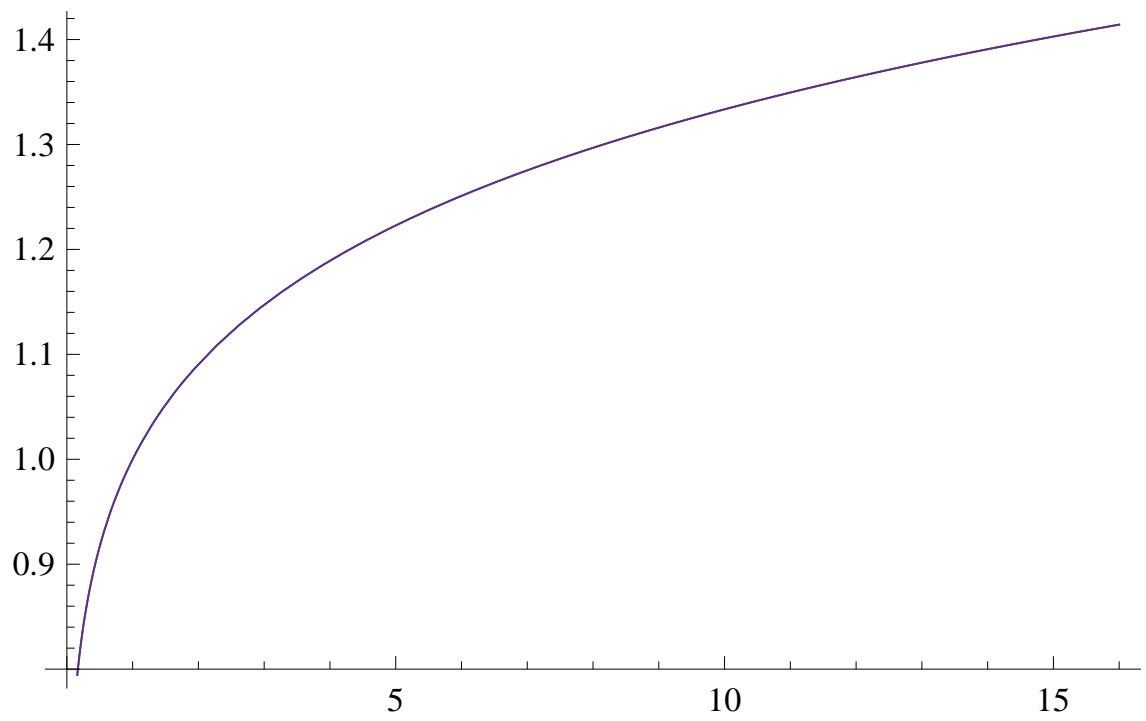


Figure C.5: Test for the rational approximation.

APPENDIX D

ERROR ANALYSIS OF THE BINDER CUMULANT

In this appendix, we presented several methods on calculating the error of the Binder Cumulant.

D.1 What We Have

$4 \times N_\sigma^3$ lattices for a pure $SU(2)$ gauge theory with $N_\sigma = 2, 4, 6, 8, 10, 12, 14, 16$. The β range decreases with increasing volume. We generated 50,000 Polyakov loops with 20 different seeds for each β .

D.2 What We Want

Calculate the Binder cumulant $g_4 = 1 - \frac{\langle P^4 \rangle}{3 \langle P^2 \rangle^2}$ and corresponding errors.

D.3 What We Have Done

- Error propagation method. First calculate the m^2 and m^4 and the corresponding errors of them. Then use error propagation to calculate g_4 . The error propagation formula is $((1/(3 \langle m^2 \rangle^2))^2 * errm^4 + ((2 * \langle m^4 \rangle) / (3 \langle m^2 \rangle^3))^2 * errm^2)^{0.5}$.
- Bootstrap method. We then tried bootstrap method. We first combine all the $5000 \times 20 = 100000$ data and then randomly pick 20000 out in order to cancel out the correlation. And then calculate g_4 from the 20000 data. We repeat the above procedure 20 times and then calculate the average and error from the calculated 20 g_4 .
- We also calculated the correlation time for one particular set of data (50000). It is around 10 for $N_\sigma = 4$ and 200 for $N_\sigma = 16$.
- We fit the 20(or less) different $g_4(\beta)$ s by following *Numerical Recipes in C++ page 667*. :

D.4 The Proper Way Of Getting The Error

- Use data from 20 different seeds to calculate 20 g_4 . We can simply ignore the correlation among data since what we want is the average value. $\langle P^4 \rangle$ is unbiased and $\langle P^2 \rangle^2$ is biased.
- Jackknife 20 g_4 s to deal with the biased $\langle P^2 \rangle^2$ and the correlation between $\langle P^4 \rangle$ and $\langle P^2 \rangle^2$. We will get the error. If we want to get the 2σ error, then we need to multiply by a factor. See *Berg's book on page 78*.
- Do the same for the rest of the β .
- Do the fitting according to *Numerical Recipes in C++ page 667*.

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