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Grassmannization of classical models

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Abstract

PAPER

Applying Feynman diagrammatics to non-fermionic strongly correlated models with local constraints might seem generically impossible for two separate reasons: (i) the necessity to have a Gaussian (non-interacting) limit on top of which the perturbative diagrammatic expansion is generated by Wick's theorem, and (ii) Dyson's collapse argument implying that the expansion in powers of coupling constant is divergent. We show that for arbitrary classical lattice models both problems can be solved/ circumvented by reformulating the high-temperature expansion (more generally, any discrete representation of the model) in terms of Grassmann integrals. Discrete variables residing on either links, plaquettes, or sites of the lattice are associated with the Grassmann variables in such a way that the partition function (as well as all correlation functions) of the original system and its Grassmann-field counterpart are identical. The expansion of the latter around its Gaussian point generates Feynman diagrams. Our work paves the way for studying lattice gauge theories by treating bosonic and fermionic degrees of freedom on equal footing.

1. Introduction

Feynman's diagrammatic technique is a powerful tool of statistical mechanics. Among the hallmarks of the method are the ability to deal—both analytically and numerically—with the thermodynamic limit rather than a finite-size cluster, the possibility of partial summations up to infinite order, and the fully self-consistent formulation in terms of renormalized (dressed) quantities. The latter properties allow one to go beyond the Taylor expansion in terms of the coupling constant or any other parameter.

The advantage of having a diagrammatic technique at our disposal comes however at a price. The most serious issue is the divergence of the expansion in powers of the coupling constant for systems prone to Dyson's collapse [1] (i.e., pathological system behavior when the coupling constant is rotated in the complex plane). For partial summation techniques to work, the non-perturbed part of the theory has to be Gaussian (in terms of either real, or complex, or Grassmann variables) to ensure the validity of Wick's theorem. These issues are often related; for example, the Ising and *XY* models formulated in terms of the original spin variables do not suffer from Dyson's collapse but lack the Gaussian (non-interacting) limit, while their classical (lattice) field counterparts with the well-defined Gaussian limit are subject to Dyson's collapse. It would be a mistake, however, to think that a meaningful diagrammatic series is only possible for a very limited class of Hamiltonians, namely, when the original system is that of interacting lattice fermions. As already clearly explained by Samuel in a series of papers [2–4] a broad class of classical spin and dimer models can be reformulated in terms of familiar interacting fermions and studied with field theoretical techniques. We note that similar techniques were also explored by other authors [5–9] and that also rather arbitrary quantum spin/boson lattice models can be rigorously mapped onto fermionic field theories [10–12].

As expected, grassmannian formulations of spin/link/boson models with local constraints are generically strongly-coupled theories at low temperature, and even the most advanced self-consistent treatments based on the lowest-order graphs are not supposed to provide quantitatively (and often qualitatively) accurate answers. Moreover, these theories may contain arbitrary multi-particle interaction vertexes, which further complicate the structure of the diagrammatic expansion. One of the promising numerical techniques currently under development for strongly correlated systems is diagrammatic Monte Carlo (DiagMC). It is based on the stochastic evaluation of irreducible Feynman graphs up to some high order and can be implemented in a number of ways, from perturbative expansions in powers of the coupling constant to various self-consistent skeleton schemes based on fully renormalized one- or two-body propagators. In such contexts as resonant fermions [13], frustrated magnetism [14, 15], and out-of-equilibrium impurity-like models [16, 17] the method was recently shown to be able to go significantly beyond the state of the art. Also, significant progress has been made in understanding superfluid properties of the Hubbard-type models [18-20]. Notably, the infamous signproblem preventing conventional Monte Carlo methods from simulating fermionic system with sizes large enough for reliable extrapolation to the thermodynamic limit, is absent as such in DiagMC. Instead, the computational complexity is now linked to the number of diagrams growing factorially with their order. Nevertheless, millions of diagrams can be accounted for and the approach is flexible enough to deal with an arbitrary interaction Hamiltonian/action.

The current paradigm for generic lattice gauge models, as they occur in lattice-QCD as well as in solid state and ultra-cold atomic physics, is to work with finite-size systems and to treat link variables separately from the fermionic sector. More precisely, link variables are simulated using classical Monte Carlo techniques (with local updates), and fermions (quarks) are described by determinants. This approach suffers from a severe signproblem for finite density of fermions (non-zero chemical potential) [21, 22]. If link variables are straightforwardly represented by bosonic fields, then the thermodynamic limit can be addressed within the diagrammatic approach that treats bosonic and fermionic degrees of freedom on equal footing. However, in this formulation the bosonic fields pose a fundamental problem, which manifests itself in a zero convergence radius. It is thus desirable to have a generic scheme for replacing link variables with Grassmann fields to ensure that the diagrammatic expansion has proper analytic properties around the Gaussian point.

In this paper, we introduce a general procedure of *grassmannization* for classical lattice models. It is by no means a unique one, and in certain specific cases more compact/simpler representations can be found. There is a strong connection to the anti-commuting variables approach introduced by Samuel [2–4], which can solve the 2D Ising model exactly (free fermion operators to solve the Ising model exactly were first found by Kaufman [23] and refined by Schultz, Mattis and Lieb [24]) and provides a good starting point for field theoretic studies of the 3D Ising model. For the latter system our approach amounts to an alternative but equally complicated field theory. Our goal is to build on these ideas and develop a scheme that is flexible enough to apply to a broader class of link models with arbitrary multi-bond interactions and local constraints.

The idea of grassmannization is to represent the partition function of the model as a Grassmann integral from the exponential of a Grassmann functional. The Feynman rules then emerge by Taylor-expanding the non-Gaussian part of the exponential and applying Wick's theorem to the Gaussian averages. Paradigmatic lattice systems are link and plaquette models featuring discrete degrees of freedom—integer numbers—residing on links (plaquettes) of square lattices and subject to certain local constraints in terms of the allowed values of the sum of all link (plaquette) variables adjacent to a given site (edge). It turns out that it is these constraints that require special tricks involving multiple Grassmann variables for each value of each discrete variable. Link models often emerge as high-temperature expansions of lattice systems [25] in Ising, *XY*, O(3), etc universality classes no matter whether the original degrees of freedom are discrete or continuous (e.g., classical vector-field variables). Link models may also emerge as dual (low-temperature) expansions, and specific examples are provided by the 2D Ising model [26] and the 3D $|\psi|^4$ model (the latter case leads to the so-called *J*-current model with long-range interactions). Similarly, plaquette models emerge as a high-temperature expansion of lattice gauge theories, but sometimes they represent the dual (low-temperature) expansion, as in the case of the 3D Ising model. Finally, it is worth mentioning how the models with the same general structure are generated by strong-coupling expansions in lattice-QCD [27].

The paper is structured as follows. In section 2 we explain how a partition function of a discrete link model can be written as a Grassmann integral. The equivalence between the two formulations is readily proved through term-by-term comparison. Standard properties of Grassmann variables then immediately allow one to express the Grassmann weight in the exponential form in order to define the field theory. In section 3 we discuss generalizations of the proposed grassmannization scheme. We start by describing the procedure for a broad class of plaquette models. Next we show a simple way to introduce Grassmann variables for non-local link models with pairwise interactions between the link variables. The construction is further simplified when constraints are replaced with statistical penalties for certain configurations of link (plaquette) variables. We conclude this section with defining the meaning of the term 'order of expansion' for the resulting field theory. In section 4 we

$$\overbrace{\overline{\xi}_{\alpha}\xi_{\alpha}}^{\overline{\xi}_{\alpha}\xi_{\alpha}} \qquad \overbrace{\xi_{\gamma}\xi_{\gamma}^{*}}^{\overline{\xi}_{\gamma}^{*}} = \overline{\xi}_{\gamma}'\overline{\xi}_{\gamma} \qquad \overbrace{\overline{\xi}_{\gamma}\xi_{\gamma}^{*}}^{\overline{\zeta}_{\gamma}} = \xi_{\gamma}\xi_{\gamma}'$$

Figure 1. Assignment of Grassmann fields for link (left) and site (right) factors. Upon integration, the labels of the Grassmann variables must be equal in order to connect variables from all factors (see text).

deliberately choose the most general grassmannization scheme for the 2D Ising model to illustrate how our construction works in practice. Results are shown in section 5. We conclude with prospects for future work in section 7. There are also two appendices, where we provide implementation details for the solution of the 2D Ising model in the bare and bold expansion schemes, respectively.

2. Grassmannization of local link models

2.1. Local link models

For the purposes of this article, we mean by a link model a classical statistical model with states labeled by a set of *discrete* variables $\{\alpha_b\}$ residing on links (bonds) of a certain lattice. In addition, we require that the ground state is unique. Without loss of generality, it can be chosen to be the state with $\alpha_b = 0$ on each link *b*.

We further narrow the class of link models—to which we will refer to as *local* link models—by the requirement that the statistical weight of a state factors into a product of link and site weights (to be referred to as link and site factors, respectively). A link factor, f_b , is a function of the corresponding link variable, $f_b \equiv f(\alpha_b)$. The site factor, g_j is a function that depends on all variables residing on links attached to the site *j*, denoted as $\{\alpha_b\}_j$. Then, $g_j \equiv g(\{\alpha_b\}_j)$. Solely for the purpose of avoiding heavy notations, we consider translational invariance when $f(\alpha_b) \equiv f_b$ is the same function on all links and g_j site independent, $g_j \equiv g$. Given that only the relative weights of the states matter, we set f(0) = 1 and $g(0_j) = 1$, where 0_j stands for the $\{\alpha_b = 0\}_j$ set.

The site factors play the key role in link models. They describe interactions between (otherwise independent) link degrees of freedom. In particular, this interaction can take the extreme form of a *constraint* on the allowed physical configurations of $\{\alpha_b\}_j$ (e.g., the zero-divergency constraint in *J*-current models [28], or the evennumber constraint in the high-temperature expansion of Z_2 models), in which case $g_j(\{\alpha_b\}_j)$ is identically *zero* for each non-physical state of $\{\alpha_b\}_j$.

2.2. Grassmannization

For each label $\alpha \neq 0$ of the link *b*, introduce four Grassmann variables: $\xi_{\alpha,b}, \xi'_{\alpha,b}, \bar{\xi}_{\alpha,b}$, and $\bar{\xi}'_{\alpha,b}$. For a textbook introduction to Grassmann variables, we refer to [29]. For $\alpha = 0$ we assume that $\xi_{0,b} = \xi'_{0,b} = \bar{\xi}_{0,b} = \bar{\xi}'_{0,b} = 1$. In terms of these variables, define the Grassmann weight—a product of link, A_b , and site, B_j , factors such that tracing over all degrees of freedom yields the partition function $Z = \text{Tr} \prod A_b \prod B_j$ —by the following rules

$$A_{b} = \exp\left\{\sum_{\alpha \neq 0} \left[\frac{\bar{\xi}_{\alpha,b}' \xi_{\alpha,b}'}{\sqrt{f(\alpha)}} + \frac{\bar{\xi}_{\alpha,b} \xi_{\alpha,b}}{\sqrt{f(\alpha)}} \right] \right\}$$
$$= \prod_{\alpha \neq 0} \exp\left\{ \frac{\bar{\xi}_{\alpha,b}' \xi_{\alpha,b}'}{\sqrt{f(\alpha)}} + \frac{\bar{\xi}_{\alpha,b} \xi_{\alpha,b}}{\sqrt{f(\alpha)}} \right\},$$
(2.1)

$$B_{j} = \sum_{\{\alpha_{b}\}_{j}} g\left(\{\alpha_{b}\}_{j}\right) \prod_{b\in\{b\}_{j}} \check{\xi}_{\alpha_{b},b} \check{\xi}_{\alpha_{b},b}^{*}$$
$$= 1 + \sum_{\{\alpha_{b}\}_{j}\neq0_{j}} g\left(\{\alpha_{b}\}_{j}\right) \prod_{b\in\{b\}_{j}} \check{\xi}_{\alpha_{b},b} \check{\xi}_{\alpha_{b},b}^{*}.$$
(2.2)

Here $\{b\}_j$ stands for the set of all links incident to the site *j*, and variables $\xi_{\alpha_b,b}$ and $\xi_{\alpha_b,b}^*$ are defined differently for different links. We first introduce the notion of direction (on each link) so that one of the two link ends becomes 'incoming' and its counterpart 'outgoing' (with respect to the site adjacent to the end). Next, we assign (see figure 1 for an illustration)

$$\check{\xi}_{\alpha_b,b} = \xi_{\alpha_b,b}, \quad \check{\xi}_{\alpha_b,b}^* = \xi'_{\alpha_b,b} \text{ (for incoming end),}
\check{\xi}_{\alpha_b,b} = \bar{\xi}'_{\alpha_b,b}, \quad \check{\xi}_{\alpha_b,b}^* = \bar{\xi}_{\alpha_b,b} \text{ (for outgoing end).}$$
(2.3)

The claim is that the Grassmann integral of the weight over all variables reproduces the partition function of the original link model. For a link *b* to yield a non-zero contribution to the integral the link labels in (2.2) for the sites of the incoming (j = 1) and outgoing (j = 2) ends of the link should match each other: $\alpha_1 = \alpha_2$. Indeed, at $\alpha_1 \neq \alpha_2$, it is not possible to find an appropriate term in the expansion of the link exponential (2.1) such that —upon multiplying by the site factors $\xi_{\alpha_b,b} \ \xi_{\alpha_{a,b}}^*$ and $\xi_{\alpha_{2,b}} \ \xi_{\alpha_{2,b}}^*$ —all powers of the Grassmann variables $\xi_{\alpha_b,b}$, $\xi'_{\alpha_b,b}, \ \xi'_{\alpha_{a,b}}, \ \xi'_{\alpha_{2,b}}, \$

$$\prod_{\gamma \neq 0} \int \mathcal{D}[\bar{\xi}'\xi'\bar{\xi}\xi]_{\gamma} \exp\left\{\frac{\bar{\xi}_{\gamma}'\xi_{\gamma}'}{\sqrt{f(\gamma)}} + \frac{\bar{\xi}_{\gamma}\xi_{\gamma}}{\sqrt{f(\gamma)}}\right\}$$
$$= \prod_{\gamma \neq 0} \int \mathcal{D}[\bar{\xi}'\xi'\bar{\xi}\xi]_{\gamma} \left[1 + \frac{\bar{\xi}_{\gamma}'\xi_{\gamma}'}{\sqrt{f(\gamma)}}\right] \left[1 + \frac{\bar{\xi}_{\gamma}\xi_{\gamma}}{\sqrt{f(\gamma)}}\right]$$
$$= \prod_{\gamma \neq 0} \frac{1}{f(\gamma)} \equiv \frac{1}{f_{*}},$$
(2.4)

where we defined f_* in the last step. In the second case, the two end sites contribute the factor $\xi_{\alpha,b} \xi'_{\alpha,b} \bar{\xi}'_{\alpha,b} \bar{\xi}_{\alpha,b} = \bar{\xi}'_{\alpha,b} \xi'_{\alpha,b} \bar{\xi}_{\alpha,b} \xi_{\alpha,b}$. Now we have to consider the first term in the expansion of the link exponential for state α , while for other variables the calculation is repeated as in (2.4)

$$\prod_{\gamma \neq 0} \int \mathcal{D}[\bar{\xi}'\xi'\bar{\xi}\xi]_{\gamma} \bar{\xi}'_{\alpha} \bar{\xi}_{\alpha} \left[1 + \frac{\bar{\xi}'_{\gamma}\xi'_{\gamma}}{\sqrt{f(\gamma)}} \right] \left[1 + \frac{\bar{\xi}_{\gamma}\xi_{\gamma}}{\sqrt{f(\gamma)}} \right] \xi_{\alpha}\xi'_{\alpha}$$

$$= \prod_{\gamma \neq 0,\alpha} \frac{1}{f(\gamma)} = \frac{f(\alpha)}{f_{*}}.$$
(2.5)

We see that, apart from the irrelevant global factor $\prod_b 1/f_*$, we reproduce the configuration space and weight factors of the original link model.

2.3. Field theoretical formulation

To generate the Feynman diagrammatic expansion, we need to represent the Grassmann weight factor in the exponential form. The link factors (2.1) have the form of Gaussian exponentials already. Hence, it is only the site factors that need to be rewritten identically as

$$B_{j} = \exp\left[\sum_{\{\alpha_{b}\}_{j}} \lambda(\{\alpha_{b}\}_{j}) \prod_{b \in \{b\}_{j}} \check{\xi}_{\alpha_{b},b} \check{\xi}_{\alpha_{b},b}^{*}\right].$$
(2.6)

The constants $\lambda(\{\alpha_b\}_j)$ are readily related to the site factors $g(\{\alpha_b\}_j)$ by simple algebraic equations obtained by expanding the exponential and equating sums of similar terms to their counterparts in the rhs of equation (2.2).

By expanding the non-Gaussian part of the exponential (2.6) and applying Wick's theorem, we arrive at Feynman rules for the diagrammatic series. The reader should avoid confusion by thinking that an expansion of the exponential (2.6) takes us back to equation (2.2). Recall that connected Feynman diagrams are formulated for the free energy density, not the partition function, and summation over all lattice sites is done for a given set of interaction vertexes in the graph, as opposite to the summation over all vertex types for a given set of lattice points. Therefore, the 'coupling constants' in Feynman diagrams are λ 's, not g's.

2.4. Absorbing link factors into site factors

The separation of the weight factors into link and site ones is merely a convention. Indeed, each link factor can be ascribed to one of the two site factors at its ends. This leads to a slightly different Grassmannization protocol. This trick may prove convenient for generalization to non-local models considered below.

3. Generalizations

3.1. Plaquette models

A plaquette model can be viewed as a certain generalization of the local link model. States (configurations) of a plaquette model are indexed by a set of discrete labels residing on (oriented) plaquettes of a hyper-cubic lattice. The plaquette label α takes on either a finite or countably infinite number of values. The statistical weight of each state factors into a product of plaquette and edge weights (to be referred to as plaquette and edge factors, respectively). A plaquette factor, *f*, is a function of the corresponding plaquette variable, $f \equiv f(\alpha)$. An edge factor, *g*, is a function which depends on the labels of all plaquettes sharing this edge (this set of labels will be denoted as { α_p } i for the edge *j*); it encodes, if necessary, constraints on the allowed sets of { α_p }.

Without loss of generality (up to a global normalization factor), we identify the 'ground state' as $\alpha_p = 0$ for all plaquettes, and set f(0) = 1. The orientation of the plaquette (for some models it is merely a matter of convenience) is enforced by an ordered enumeration of sites at its boundary. For a plaquette p, the vertex label $\nu \equiv \nu_p = 0, 1, 2, 3$ enumerates four vertices in such a way that $\nu \pm 1$ modulo 4 stands for the next/previous vertex with respect to the vertex ν in the clockwise direction.

For each state $\alpha \neq 0$ of the plaquette *p*, we introduce eight Grassmann variables: ξ_{α,p,ν_p} , $\bar{\xi}_{\alpha,p,\nu_p}$, $\nu_p = 0, 1, 2, 3$. As before, for $\alpha = 0$ the variables ξ and $\bar{\xi}$ are not Grassmannian, $\xi_{0,p,\nu} = 0$, $\bar{\xi}_{0,p,\nu} = 1$. The corresponding plaquette weight in the Grassmann partition function reads

$$A_{p} = \exp\left\{\sum_{\alpha \neq 0} [-f(\alpha)]^{-1/4} \sum_{\nu_{p}=0}^{3} \bar{\xi}_{\alpha,p,\nu_{p}} \,\xi_{\alpha,p,\nu_{p}}\right\}.$$
(3.1)

Note a close analogy with equation (2.1). Site weights equation (2.2) are now replaced with edge weights B_j . Using the notation $\{p\}_j$ for the set of all plaquettes sharing the edge j, and 0_j for the state when all plaquettes in this set have $\alpha_p = 0$, we write

$$B_{j} = 1 + \sum_{\{\alpha_{p}\}_{j} \neq 0_{j}} g(\{\alpha_{p}\}_{j}) \prod_{p \in \{p\}_{j}} \xi_{\alpha, p, (\nu_{p}^{(j)}+1)} \bar{\xi}_{\alpha, p, \nu_{p}^{(j)}},$$
(3.2)

where $\nu_p^{(j)}$ is the site enumeration index within the plaquette *p*, with respect to which the edge *j* is outgoing. (Accordingly, the edge *j* is incoming with respect to site $(\nu_p^{(j)} + 1)$.) In what follows, we will associate $\nu_p^{(j)}$ not only with the site, but also with the corresponding edge.

The proof that the classical and Grassmannian partition functions are identical (up to a global factor) is similar to the one for the link model after we notice that a non-zero contribution from plaquette *p* is possible only if the same plaquette label α_p is used in all edge weights. The $\alpha = 0$ contribution comes from the term

$$-\prod_{\gamma} \frac{1}{f(\gamma)} \prod_{\nu_p=0}^{3} \bar{\xi}_{\gamma,p,\nu_p} \xi_{\gamma,p,\nu_p} \quad (\text{at } \alpha = 0)$$
(3.3)

in the expansion of the exponential (3.1). It contributes a factor $1/q_*$, where

$$q_* = \prod_{\gamma} \ (-1)f(\gamma). \tag{3.4}$$

The $\alpha \neq 0$ contribution comes from the plaquette term

$$\prod_{\gamma \neq \alpha} \frac{1}{f(\gamma)} \prod_{\nu_p=0}^{3} \bar{\xi}_{\gamma,p,\nu_p} \xi_{\gamma,p,\nu_p} \quad (\text{at } \alpha \neq 0)$$
(3.5)

multiplied by the product $\prod_{\nu_p=0}^{3} \bar{\xi}_{\alpha,p,\nu_p} \xi_{\alpha,p,\nu_p}$ originating from the boundary edge terms $\xi_{\alpha,p,(\nu_p^{(j)}+1)} \bar{\xi}_{\alpha,p,\nu_p^{(j)}}$. Because of the Grassmann anticommutation rules, this four-edge factor yields an additional minus sign, explaining the use of the negative sign in front of $f(\alpha)$ in equation (3.1). Upon Grassmann integration, the contribution to the partition function of the resulting term equals to $f(\alpha)/q_*$.

Feynman diagrammatics for the plaquette model is obtained by following the same basic steps as for the link models. The Gaussian part is given by equation (3.1) with four pairs of Grassmann fields for every non-zero plaquette state. The interaction part of the Grassmann action is contained in edge weights (3.2) after they are written in an exponential form

$$B_{j} = \exp\left[\sum_{\{\alpha_{p}\}_{j}} \lambda(\{\alpha_{p}\}_{j}) \prod_{p \in \{p\}_{j}} \xi_{\alpha,p,(\nu_{p}^{(j)}+1)} \bar{\xi}_{\alpha,p,\nu_{p}^{(j)}}\right],$$
(3.6)

with the constants $\lambda(\{\alpha_b\}_i)$ unambiguously related to the edge factors $g(\{\alpha_b\}_i)$.

3.2. Unconstrained discrete models with pair-wise interaction

The hallmark of the considered link (plaquette) models is the non-trivial interaction introduced via site (edge) factors. It is due to this type of interaction—and, in particular, its extreme form of a constraint on allowed combinations of discrete variables—that we had to introduce multiple Grassmann variables for each state of the link (plaquette). The situation simplifies dramatically if we are dealing with unconstrained discrete degrees of freedom with pair interactions between them.

Consider a link model defined by the statistical weight

$$W(\{\alpha_b\}) = \prod_{b_1, b_2} F(\alpha_{b_1}, b_1; \alpha_{b_2}, b_2),$$
(3.7)

based on products of two-link factors. Without loss of generality, these factors can be cast into the exponential form

$$W(\{\alpha_b\}) = \prod_{b_1, b_2} e^{-(1/2)\eta_{\alpha_{b_1}, b_1; \alpha_{b_2}, b_2}},$$
(3.8)

We assume that all factors in the product are bounded and properties of the η -matrix are well-conditioned. Grassmannization of this model can be done by taking advantage of properties of Gaussian integrals that allow one to express (3.8) identically (up to normalization) as

$$W(\{\alpha_b\}) = \int \mathcal{D}X \prod_b e^{iX_{\alpha_b,b}} W_G(\{X_{\alpha_b,b}\}).$$
(3.9)

Here $\{X_{\alpha_b,b}\}$ is a collection of auxiliary real continuous variables. For briefness, we do not show explicitly the Gaussian weight W_G that is uniquely defined by the values of all pairwise averages performed with this weight

$$\eta_{\alpha_{b_1}, b_1; \alpha_{b_2}, b_2} = \langle X_{\alpha_{b_1}, b_1} X_{\alpha_{b_2}, b_2} \rangle.$$
(3.10)

What we achieve for a fixed set of X variables is a link model that contains only single-link factors

$$\forall b: \qquad f_b(\alpha_b) = e^{iX_{\alpha_b,b}}. \tag{3.11}$$

For models with site constraints, link factors can be attributed to site factors at the incoming (or outgoing) ends with subsequent Grassmannization of the latter as discussed above. For unconstrained models, Grassmannization is accomplished by replacing sums over link variables with

$$\sum_{\alpha_b} f_b(\alpha_b) \to \mathcal{W}_b^{(G)} = \exp\left[\bar{\xi}_b \xi_b \left(\sum_{\alpha_b} e^{iX_{\alpha_b,b}}\right)\right].$$
(3.12)

Note that here Grassmann variables have nothing to do with the discrete index α_b , in contrast with previous considerations. The resulting formulation contains both Grassmann and real-number integrations.

Clearly, all considerations can be repeated identically (up to a trivial change in notations) for a model based on discrete variables α_s residing on lattice sites when the configuration weight is given by

$$W(\{\alpha_s\}) = \prod_{s_1, s_2} e^{-(1/2)\eta_{\alpha_{s_1}, s_1; \alpha_{s_2}, s_2}}.$$
(3.13)

3.3. Order of expansion

The notion of the order of expansion is absolutely central for practical applications when diagrammatic series are truncated. Normally, it is defined as an integer non-negative power of a certain dimensionless parameter ζ playing the role of a generalized coupling constant, such that the diagrammatic expansion corresponds to a Taylor expansion in ζ about the point $\zeta = 0$. Without loss of generality, we can always select ζ (by an appropriate rescaling) in such a way that the physical value of ζ is 1. This is especially convenient in cases when there is more than one interaction vertex, and ascribing different powers of ζ to them results in (re-)grouping of different terms in the series. A reasonable guiding principle behind such a (re-)grouping is the requirement to end up with Taylor series having finite convergence radius around $\zeta = 0$. The latter is guaranteed if the theory is analytic in ζ at the origin; the necessary condition for this to be true is the absence of Dyson's collapse when changing the sign (more generally, the phase) of ζ .

As an illustration, consider the theory (3.7) and (3.8) and its Grassmann counterpart (3.12). Introduce the ζ -dependence by the replacement

$$e^{iX_{\alpha_b,b}} \to e^{i\zeta X_{\alpha_b,b}}.$$
(3.14)

In terms of the original theory, the replacement (3.14) means $\eta \to \zeta^2 \eta$, for all η 's in equation (3.8). If amplitudes of all η values in (3.8) are bounded, we expect that such a dependence on ζ is analytic not only for a finite system, but also in the thermodynamic limit at finite temperature. In the Grassmann action (3.12), the expansion of the exponential $e^{i\zeta X_{s,o_b}}$ in powers of ζ generates an infinite series of interaction vertexes (the zeroth-order term defines the harmonic action):

$$\bar{\xi}_{b}\xi_{b}\sum_{\alpha_{b}}\left(i\zeta X_{\alpha_{b},b}-\frac{1}{2}\zeta^{2}X_{\alpha_{b},b}^{2}-\frac{i}{3!}\zeta^{3}X_{\alpha_{b},b}^{3}+\cdots\right).$$
(3.15)

Higher-order vertexes in *X* come with a higher power of ζ and this sets unambiguously the rules for defining the diagram order.

4. Illustration for the 2D Ising model

In this section we follow the most general protocol of introducing Grassmann variables and deliberately avoid any possibility is using more efficient, model specific, formulations. The 2D case is considered exclusively for reasons of presenting explicitly the final formulation in a relatively compact form. Let us mention again the crucial property we want from our theory: analyticity in the expansion parameter ζ at the origin, as seen from the absence of Dyson's collapse when changing the phase of ζ . This should become apparent in the derivation presented in this section.

4.1. Model and observables

Consider the 2D Ising model on the square lattice with the Hamiltonian

$$-H/T = \beta \sum_{\langle i,j \rangle} \sigma_i \sigma_j + \sum_i h_i \sigma_i.$$
(4.1)

The Ising variables $\sigma = \pm 1$ live on the sites of the 2D square lattice and interact ferromagnetically with their nearest neighbors, as is represented by the first term in the Hamiltonian. We write the dimensionless coupling as β in units of the temperature *T*. Additionally, every spin feels a dimensionless magnetic field $h_i = h$, which can be taken $h \ge 0$ without loss of generality. The partition function of the Ising model reads

$$Z = \sum_{\{\sigma_i\}} \prod_{\langle i,j \rangle} \mathbf{e}^{\beta \sigma_i \sigma_j} \prod_i \mathbf{e}^{h_i \sigma_i}.$$
(4.2)

The most typical observable of the Ising model is the spin–spin correlation function ρ_{ii}

$$\rho_{ij} = \langle \sigma_i \sigma_j \rangle = \frac{1}{Z} \frac{\partial^2 Z}{\partial h_i \partial h_j} |_{h_i = h_j = h.}$$
(4.3)

4.2. Grassmannization of the high-temperature expansion

Using the well-known identities

$$e^{\beta\sigma_i\sigma_j} = \cosh\beta(1 + \sigma_i\sigma_j\tanh\beta)$$

$$e^{h\sigma_i} = \cosh h(1 + \sigma_i\tanh h), \qquad (4.4)$$

the partition function can be written as $Z = Z_0 Z'$ with $Z_0 = (\cosh \beta)^{2N} (\cosh h)^N$ for a lattice of N sites and 2N links. With the notation $\zeta = \tanh \beta$ and $\eta = \tanh h$ the remaining factor is given by

$$Z' = \sum_{\{\sigma_i\}} \prod_{\langle i,j \rangle} (1 + \sigma_i \sigma_j \zeta) \prod_i (1 + \sigma_i \eta).$$
(4.5)

Upon summation over spin variables we are left with a link model, where link variables take only two values, 0 or 1, to specify whether we are dealing with the first or the second term in the sum $(1 + \sigma_i \sigma_j \zeta)$. In the partition function, terms with an odd power of σ_i on any of the sites yield zero upon spin summation. The remaining terms depend on link variables in a unique way. The formalism of the previous section can be straightforwardly applied, and we obtain

$$f(0) = 1, f(1) = f_* = \zeta,$$
 (4.6)

$$g(0) = g(2) = g(4) = 1, g(1) = g(3) = \eta.$$
 (4.7)

Here we label site factors using the total sum of incident link variables, $\sum_{b \in \{b\}_j} \alpha_b$, to avoid unnecessary rank-4 tensor notations. If we further redefine $Z_0 \rightarrow Z_0 2^N f_*^{2N}$, then the Grassmann representation of the partition function Z' is given by

$$Z' = \int \mathcal{D}[\bar{\xi}'\xi'\bar{\xi}\xi]_{\{\alpha_b\}} \prod_{\{\alpha_b\}} \exp\left(\frac{1}{\sqrt{\zeta}}\bar{\xi}_{\alpha_b}\xi_{\alpha_b} + \frac{1}{\sqrt{\zeta}}\bar{\xi}_{\alpha_b}\xi_{\alpha_b}\right) \\ \times \exp\left(\sum_j \lambda_{\alpha_{\{b\}}} \prod_{b_j} \check{\xi}_{\alpha_b}\check{\xi}_{\alpha_b}^*\right).$$
(4.8)

4.3. Vertex coefficients

We now compute the factors λ . To this end, we first introduce notations (for a fixed site *j* and suppressing the site index for clarity)

$$V_{1} = \check{\xi}_{R}\check{\xi}_{R}^{*} + \check{\xi}_{U}\check{\xi}_{U}^{*} + \check{\xi}_{L}\check{\xi}_{L}^{*} + \check{\xi}_{D}\check{\xi}_{D}^{*} = n_{R} + n_{U} + n_{L} + n_{D},$$

$$V_{2} = n_{R}n_{U} + n_{R}n_{L} + n_{R}n_{D} + n_{U}n_{L} + n_{U}n_{D} + n_{L}n_{D},$$

$$V_{3} = n_{R}n_{U}n_{L} + n_{R}n_{U}n_{D} + n_{R}n_{L}n_{D} + n_{U}n_{L}n_{D},$$

$$V_{4} = n_{R}n_{U}n_{L}n_{D}$$
(4.9)

and then Taylor expand

$$\exp\left[\lambda_1 V_1 + \lambda_2 V_2 + \lambda_3 V_3 + \lambda_4 V_4\right]. \tag{4.10}$$

The only non-zero terms generated by this expansion are $V_1^2 = 2V_2$, $V_1^3 = 6V_3$, $V_1^4 = 24V_4$, $V_1V_2 = 3V_3$, $V_1V_3 = 4V_4$ and $V_2^2 = 6V_4$. All other powers and multiplications of operators yield zero. Note that operators from different sites commute and may be excluded from consideration here. The final result is

$$\exp \left[\lambda_{1}V_{1} + \lambda_{2}V_{2} + \lambda_{3}V_{3} + \lambda_{4}V_{4}\right] \\= 1 + \lambda_{1}V_{1} + \lambda_{2}V_{2} + \lambda_{3}V_{3} + \lambda_{4}V_{4} \\+ \frac{1}{2}(\lambda_{1}^{2}2V_{2} + \lambda_{2}^{2}6V_{4} + 2\lambda_{1}\lambda_{2}3V_{3} + 2\lambda_{1}\lambda_{3}4V_{4}) \\+ \frac{1}{6}(\lambda_{1}^{3}6V_{3} + 3\lambda_{1}^{2}\lambda_{2}12V_{4}) + \frac{1}{24}\lambda_{1}^{4}24V_{4}$$
(4.11)

Term-by-term matching with equation (4.7) then leads to

$$g_{\rm l} = \eta = \lambda_{\rm l},\tag{4.12}$$

$$g_2 = 1 = \lambda_2 + \lambda_1^2, \tag{4.13}$$

$$g_3 = \eta = \lambda_3 + 3\lambda_1\lambda_2 + \lambda_1^3, \tag{4.14}$$

$$g_4 = 1 = \lambda_4 + 3\lambda_2^2 + 4\lambda_1\lambda_3 + 6\lambda_2\lambda_1^4 + \lambda_1^4.$$
(4.15)

The solution is immediate

$$\lambda_{\rm l} = \eta, \tag{4.16}$$

$$\lambda_2 = 1 - \eta^2, \tag{4.17}$$

$$\lambda_3 = -2\eta + 2\eta^3, \tag{4.18}$$

$$\lambda_4 = -2 + 8\eta^2 - 6\eta^4. \tag{4.19}$$

In zero external field the only vertexes with non-zero coupling in the partition function are V_2 and V_4

1

$$\prod_{j} \exp(V_{2}^{(j)} + V_{4}^{(j)}) = \exp\left\{\sum_{j} (V_{2}^{(j)} - 2V_{4}^{(j)})\right\}$$
(4.20)

and the spin-spin correlation function is given by

$$\rho_{ij} = \frac{1}{Z'} \prod_{b} \int \mathcal{D}[...]_{b} \exp\left(\frac{1}{\sqrt{\zeta}} \bar{\xi}_{b}' \xi_{b}' + \frac{1}{\sqrt{\zeta}} \bar{\xi}_{b} \xi_{b}\right) \\
\times \exp\left\{\sum_{j} (V_{2}^{(j)} - 2V_{4}^{(j)})\right\} \\
\times (V_{1}^{(i)} - 2V_{3}^{(i)})(V_{1}^{(j)} - 2V_{3}^{(j)}).$$
(4.21)

4.4. Feynman rules

In order to derive the Feynman rules generating the diagrammatic series, we write the partition function in the form



occur at the end points of the spin correlator (indicated by the open circle), the elements in the second and fourth row are the generic basic vertexes of the theory ascribed to the sites of the underlying lattice. There are hence $4V_1$ vertexes with 1 leg (first row, U, R, D, and L), $6V_2$ vertexes with 2 legs (second row, RU, RD, LD, LU, UD and LR), $4V_3$ vertexes with 3 legs (third row, LUR, URD, LDR, and DLU), and 1 V_4 vertex with four legs (fourth row, RULD). Connected to the legs of these vertexes are pairs of bi-Grassmann fields (thick dash lines (blue and red)) that reside on the links of the underlying 2D lattice. Thin dashed lines (showing lattice links adjacent to the site of the vertex) are to guide the eye and have no other meaning than showing the underlying 2D lattice. The generalization to other dimensions is straightforward.

$$Z' = Z_{1} \left(\sum_{n=0x_{1},...,x_{n}}^{\infty} \frac{(+1)^{n}}{n!} \langle V(x_{1}) \dots V(x_{n}) \rangle_{0} \right),$$
(4.22)

where Z_{\downarrow} is the partition function of the Gaussian part (it is the product of local link contributions), $Z_{\downarrow} = \prod_b \int \mathcal{D}[...] \exp\left(\bar{\xi}'_b \xi'_b + \frac{1}{\zeta} \bar{\xi}_b \xi_b\right) = (1 + \zeta)^{(2N)}$. The Feynman rules for the correlation function of the 2D Ising model now follow from the textbook considerations:

- (i) The bare propagators $G^{(0)} = \sqrt{\zeta}$ for primed and non-primes variables are local and reside on the links of the original lattice. In the correlation function they always occur in pairs of conjugate Grassmann variables and each pair contributes a factor ζ . The propagation lines do not have arrows. The bare interaction vertexes (or pre-diagrams, see figure 2) are also local and live on the sites of the lattice. There are different types belonging to the V_2 and V_4 classes with weight 1 and -2, respectively (see equation (4.20)). On the first (and last) site of the correlator we have a vertex belonging to the class V_1 or V_3 (see figures 3–7) with weight 1 and -2, respectively (see (4.21)).
- (ii) Draw in order *n* all topologically distinct connected diagrams with *n* pairs of bi-grassmann variables living on the links of the lattice. The number of interaction vertexes, excluding the end points, is at most n 1.
- (iii) For links with multiple occupancy, a minus sign occurs when swapping 2 Grassmann variables. The minus sign can also be found by counting all closed fermionic loops.
- (iv) The total weight of the diagram in order *n* is hence $(-1)^{p}(-2)^{q}\zeta^{n}$ with *P* the signature of the exchange permutation and *q* the sum of all type-3 and type-4 vertexes.





Disconnected diagrams are defined with respect to both the primed and non-primed Grassmann variables simultaneously. Thus, a link can lead to a disconnected diagram only if the primed and non-primed variables simultaneously lead to disconnected pieces (such as the upper left panel in figure 8). We check the connectivity of a diagram by the breadth-first algorithm. We have implemented the Feynman rules in two different appraoches: an algorithm that enumerates and evaluates all possible diagrams, as well as a DiagMC approach. Details about the implementation can be found in the appendix A

4.5. Example: the first element of the spin correlation function

For illustrative purposes, let us focus on the first element of the correlation function connecting the sites (0, 0) and (1, 0) (using translational invariance, any two neighboring sites $\langle \mathbf{r}_1, \mathbf{r}_2 \rangle$ can be taken). To first order, we put a V_1 vertex on the origin and target site. There is one way to combine them, thus the total contribution is ζ . By the symmetry of the lattice, even expansion orders do not contribute. In third order, we can construct a diagram by putting a V_2 (RD) vertex on the site (0, 1) and a V_2 vertex (LD) on the site (1, 1). The mirror symmetry of this diagram about the *x*-axis is also a valid diagram. Hence, the contribution is $2\zeta^3$. These diagrams contributing in first and third order are shown in figure 3.

In fifth order, there are four diagrams with a V_3 vertex on one of the endpoints, yielding a contribution $-8\zeta^5$. There are 14 diagrams consisting of only V_1 and V_2 vertexes and single pair-lines, yielding a contribution $14\zeta^5$. The contributions to fifth order are shown in figures 4, 5, 7, and 8. There are however additional diagrams with



Figure 5. Fifth-order diagrams for $\rho_{(1,0)}$ (at h = 0) based on expanding (4.3): these four counterparts of the diagrams shown in figure 4 are obtained by replacing a three-leg end vertex with a one-leg end vertex. Each diagram contributes ζ^5 .



two pairs of Grassmann variables living on the same link, as is shown in figure 8 (there are equivalent diagrams obtained by mirror symmetry around the x-axis which are not shown). They all have on the origin a V_1 and a V_2 (*RU*) vertex, and on the target site (1, 0) a V_1 and a V_2 (*UL*) vertex. On the site (1, 1) there is a V_2 (*LD*) and on site (0, 1) a V_2 (*RD*) vertex. Let us look more carefully at the link between the origin and target site:

$$\frac{1}{\zeta^2} \int \mathcal{D}[\bar{\xi}'\bar{\xi}\xi\xi']\bar{\xi}'\bar{\xi}\xi\xi'\bar{\xi}'\bar{\xi}\xi\xi'.$$
(4.23)

The origin is associated with $\bar{\xi}'\bar{\xi}$ and the target with $\xi\xi'$ by our convention. Applying Wick's theorem, there are four possible ways to pair the Grassmann variables:

(i) The pairing combination $\overline{\xi'} \, \overline{\xi\xi} \, \xi' \, \overline{\xi'} \, \overline{\xi\xi} \, \xi'$ comes with the sign +1 and leads to a connected diagram (this is the lower right panel in figure 8).





- (iii) The pairing combination $\overline{\xi'} \overline{\xi} \xi \xi' \overline{\xi'} \overline{\xi} \xi \xi'$ comes with the sign -1 and leads to a connected diagram (this is the lower left panel in figure 8).

The net contribution of these four distinct diagrams is hence -1 (also the diagrams obtained by mirror symmetry around the *x*-axis yield -1, so the total contribution to fifth order is $(-8 + 14 - 2)\zeta^5 = 4\zeta^5$. It is instructive to notice that the sum of all diagrams in which multiple Grassmann pairs live on the same link always produces zero in case all diagrams are connected, in line with the nilpotency of Grassmann variables.

Site/order	ζ	ζ^2	ζ^3	ζ^4	ζ^5	ζ^6	ζ^7	ζ^8	ζ9	ζ^{10}	ζ^{11}
(1,0)	1	0	2	0	4	0	12	0	42	0	164
(1,1)	0	2	0	4	0	10	0	32	0	118	0
(2,0)	0	1	0	6	0	16	0	46	0	158	0
(2, 1)	0	0	3	0	11	0	31	0	97	0	351
(2, 2)	0	0	0	0	6	0	24	76	0	248	0
(3,0)	0	0	1	0	12	0	48	0	152	0	506
(3,1)	0	0	0	4	0	26	0	92	0	298	0
(3, 2)	0	0	0	0	10	0	55	0	201	0	684
(3, 3)	0	0	0	0	0	20	0	120	0	480	0
(4,0)	0	0	0	1	0	20	0	118	0	7 452	0
(4, 1)	0	0	0	0	5	0	52	0	244	0	885
(4, 2)	0	0	0	0	0	15	0	118	0	521	0
(4, 3)	0	0	0	0	0	0	25	0	259	0	1176
(4, 4)	0	0	0	0	0	0	0	70	0	560	0
(5,0)	0	0	0	0	1	0	30	0	250	0	1200
(5,1)	0	0	0	0	0	6	0	92	0	574	0
(5,2)	0	0	0	0	0	0	21	0	231	0	1266
(5,3)	0	0	0	0	0	0	0	56	0	532	0
(5,4)	0	0	0	0	0	0	0	0	126	0	1176
(5,5)	0	0	0	0	0	0	0	0	0	252	0
(6,0)	0	0	0	0	0	1	0	42	0	474	0
(6,1)	0	0	0	0	0	0	7	0	149	0	1215
(6,2)	0	0	0	0	0	0	0	28	0	416	0
(6,3)	0	0	0	0	0	0	0	0	84	0	1026
(6,4)	0	0	0	0	0	0	0	0	0	210	0
(6,5)	0	0	0	0	0	0	0	0	0	0	462
(7,0)	0	0	0	0	0	0	1	0	56	0	826
(7,1)	0	0	0	0	0	0	0	8	0	226	0
(7,2)	0	0	0	0	0	0	0	0	36	0	699
(7,3)	0	0	0	0	0	0	0	0	0	120	0
(7,4)	0	0	0	0	0	0	0	0	0	0	330
(8,0)	0	0	0	0	0	0	0	1	0	72	0
(8,1)	0	0	0	0	0	0	0	0	9	0	326
(8,2)	0	0	0	0	0	0	0	0	0	45	0
(8,3)	0	0	0	0	0	0	0	0	0	0	165
(9,0)	0	0	0	0	0	0	0	0	1	0	90
(9,1)	0	0	0	0	0	0	0	0	0	10	0
(9,2)	0	0	0	0	0	0	0	0	0	0	55
(10,0)	0	0	0	0	0	0	0	0	0	1	0
(10,1)	0	0	0	0	0	0	0	0	0	0	11
(11,0)	0	0	0	0	0	0	0	0	0	0	1

Table 1. Expansion coefficients for the correlation function up to order 11.

Wick's theorem splits however these contributions in connected and disconnected diagrams, where the disconnected diagrams cancel against the denominator of the Feynman expansion. It is this non-trivial regrouping imposed by Wick's theorem that can yield non-zero contributions from terms like (4.23); and, in particular, from arbitrarily high powers of one and the same interaction vertex.

5. Results

We now proceed with the results for the spin–spin correlation function and the magnetic susceptibility obtained with the bare expansion. Let us in passing mention that in appendix B we show how a bold G^2W scheme (which conceptually distinguishes our approach from the usual high-temperature series expansion approach) can resum whole classes of diagrams and reproduce the bare series when expanded to the same order and accuracy.

5.1. Spin-spin correlation function

Our results for the spin–spin correlation function are shown in table 1. The correlation function is known recursively from [30–32]. It is also known as a Painlevé-VI nonlinear differential equation [33] but this is not so well suited to obtain the series coefficients. Along the principal axes and the diagonal it can also be expressed as a Toeplitz determinant. The first element along and the axis and the diagonal can be recast in terms of complete



Figure 9. The magnetic susceptibility versus ζ for different expansion orders from 12 to 1 (top to bottom), compared to the order 100 result—the converged answer over this plotting range—obtained from [34], which shows a divergence in good agreement with the critical exponent $\gamma = 7/4$ starting from $\beta \ge 0.38$.

elliptic integrals (see pp 200-201 in [26]), which are convenient for series expansions

$$\rho_{(1,0)} = \operatorname{coth}(2\beta) \left[\frac{1}{2} + \frac{\operatorname{cosh}^2 2\beta}{\pi} (2 \tanh^2 2\beta - 1) K(k_{>}) \right] \\
\rightarrow \zeta + 2\zeta^3 + 4\zeta^5 + 12\zeta^7 + 42\zeta^9 + \cdots, \qquad (5.1)$$

$$\rho_{(1,1)} = \frac{2}{\pi k_{>}} [K(k_{>}) + (k_{>}^{2} - 1)K(k_{>})] \rightarrow 2\zeta^{2} + 4\zeta^{4} + 10\zeta^{6} + 32\zeta^{8} + 118\zeta^{10} + \cdots.$$
(5.2)

with $k_{>} = \sinh^{2}(2\beta)$, *K*(.) and *E*(.) the complete elliptic *K* and *E* functions, respectively. The above-cited recursion relations could be initialized with these expansions and shown to yield the same results as the top two rows in table. 1.

5.2. Magnetic susceptibility

The spin susceptibility is related to the zero momentum value of the Green function by $\beta^{-1}\chi = 1 + \rho(\mathbf{p} = 0)$. We can hence sum over the entire lattice to obtain

$$\beta^{-1}\chi = 1 + 4\zeta + 12\zeta^{2} + 36\zeta^{3} + 100\zeta^{4} + 276\zeta^{5} + 740\zeta^{6} + 1972\zeta^{7} + 5172\zeta^{8} + 13492\zeta^{9} + 34876\zeta^{10} + 89764\zeta^{11} + 229628\zeta^{12} + 585508\zeta^{13} + 1486308\zeta^{14} + \cdots.$$
(5.3)

To this order the series expansion agrees with the ones from [34, 35]. For a library of high-temperature series expansions, see [36]. Currently, the series is known (at least) up to order 2000 and still topic of active research [32, 34]. The series is convergent for any finite expansion order, i.e., in the thermodynamic limit the infinite series will diverge first at the phase transition point. It is hence possible to study the critical behavior of the susceptibility, which is governed by the critical exponent $\gamma = 7/4$. We plot in figure 9 the susceptibility versus β for different expansion orders, and also plot the asymptotic behavior for comparison.

The critical temperature and the exponent γ can be found from a study of the convergence radius of the series. Since

$$\beta^{-1}\chi = \sum_{n} \chi_n \zeta^n \propto (1 - \zeta/\zeta_c)^{-\gamma}$$
$$= 1 + \sum_{n=1}^{\infty} \frac{\gamma(\gamma - 1) \cdots (\gamma + n - 1)}{n!} \left(\frac{\zeta}{\zeta_c}\right)^n$$
(5.4)

the ratio of coefficients asympotically behaves as

$$\frac{\chi_n}{\chi_{n-1}} = \frac{1}{\zeta_c} + \frac{\gamma - 1}{\zeta_c} \frac{1}{n}.$$
(5.5)



Figure 10. Ratio of consecutive coefficients $\chi[n - 1]$ and $\chi[n]$ in the expansion of the susceptibility as a function of the inverse of the expansion order 1/n. Linear regression according to equation (5.4) allows to determine the critical temperature with an accuracy of 0.5% and the critical exponent γ with an accuracy of 5%. The fitting regime included orders nine through 14.

In figure 10 we extract the critical point ζ_c from the intercept and the critical exponent γ from a linear fit through the ratio of the coefficients. The critical point could be determined with an accuracy of 0.5%, whereas the error on γ is of the order of 5%. However, according to more advanced extrapolation techniques discussed in [37], γ can be determined independently from ζ_c as $\gamma \approx 1.751949$ on the square lattice when the series is known up to 14th order, i.e., an accuracy of 0.5%.

6. The Ising model in one-dimension

Let us show that the proposed approach solves the 1D Ising model exactly, both in the bare formulation as well as in the G^2W skeleton formulation (see section appendix B).

6.1. Bare series

In 1D, the only allowed vertex is *RL* (the last one in the second line of figure 2). It has weight +1. The only allowed endpoints are *L* and *R* (the second and fourth vertexes shown in the first line of figure 2). As expected, this means that there are no loops, no fermionic exchanges, and no minus signs in 1D. At order *n* of the expansion for the spin correlator there is only one contributing diagram with weight ζ^n (up to the lattice symmetry). The susceptibility is hence

$$T\chi = 1 + 2(\zeta + \zeta^2 + \dots) = 1 + 2\frac{\zeta}{1 - \zeta},$$
(6.1)

reproducing the exact solution with asymptotic behavior $\chi \propto \beta \exp(2\beta)$ as $T \to 0$.

6.2. G^2W formulation

The G^2W skeleton expansion becomes exact already in 0th order

$$\Pi = \Pi^0 = \zeta \tag{6.2}$$

$$\Sigma = 0 \tag{6.3}$$

which yields $G = G_0 = \sqrt{\zeta}$, $W = V/(1 - V\Pi) = 1/(1 - \zeta)$, and also $\Pi = \zeta/(1 - \zeta)$. This immediately leads to the same result as in equation (6.1) when adding the end-point vertexes *L* and *R* to Π .

7. Conclusion

We have developed a general scheme for mapping a broad class of classical statistical link (plaquette) models onto interacting Grassmann-field theories that can be studied by taking full advantage of the diagrammatic technique. This mapping, in particular, would allow to formulate an all-diagrammatic approach to (d + 1)-dimensional lattice gauge theories with finite density of fermions. The resulting field theory looks very complex because it contains a large number of Grassmann variables with numerous multi-point interaction vertexes. Moreover, it is generically strongly coupled at low temperature meaning that an accurate solution using diagrammatic methods is only possible when calculations are performed to high order and extrapolated to the infinite-order limit.

The complexity of the problem should not be taken as an indication that the entire idea is hopeless. Monte Carlo methods were designed to deal with configuration spaces of overwhelming size and complexity and arbitrary weights. In this sense, DiagMC methods simulating the configuration space of irreducible connected Feynman graphs are based on the same general principles and one should not be surprised that they can evaluate the sum of millions of bare (or skeleton) graphs, enough to attempt an extrapolation to the infinite-order limit. What makes DiagMC distinctly unique (apart from working with ever-changing number of continuous variables without systematic errors) is the radical transformation of the sign problem. It is completely eliminated in conventional sense because the thermodynamic limit is taken first. Given that the number of diagrams increases factorially with their order, finite convergence radius in ζ is only possible if same-order diagrams cancel each other to such a degree that at high order their combined contribution is not increasing factorially. In other words, non-positive weights are *required* for the entire approach to work and we call it the 'sign-blessing' phenomenon. Diagram weights for Grassmann/fermion fields alternate in sign depending on the diagram topology; this leads to the sign-blessing phenomenon for lattice models.

We illustrated the proposed approach by considering the 2D Ising model as a prototypical example. We have deliberately chosen to work with the generic formulation to avoid model specific simplifications because our goal was to demonstrate how one would proceed in the general case. The ultimate goal is to explore how this field theoretical approach can help with understanding properties of lattice gauge models.

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The appendix contains two technical developments: in appendix A we provide details on how we generated the bare diagrammatic series for the 2d Ising model using a DiagMC approach and a direct enumeration scheme. In appendix B we discuss the lowest order diagrams of a bold expansion in the G^2W scheme, and show how it reproduces the bare series when expanded to the same order and accuracy. This illustration constitutes a major difference between our field theoretic approach and high-temperature series expansions, where such a bold ification is not possible.

Appendix A. Implementation

We explored two ways of evaluating the (bare) series for the spin-correlator: a stochastic Monte Carlo approach and a deterministic full evaluation of all diagrams.

A.1. Monte Carlo sampling

In order to perform a Monte Carlo sampling over all Feynman diagrams, we introduce a head and a tail that represent the endpoints of the correlation function. By moving them around the lattice and changing the diagrammatic elements in between the head and tail, we are able to reach an ergodic sampling. The algorithm can be formulated as follows: the tail remains stationary at the origin whereas the head can move around the lattice. When the head and tail are on the same site and the expansion order is 0, the value of the correlation function is 1 which can be used for normalization of the Monte Carlo process. A Monte Carlo measurement contributes +1 or -1 depending on the sign of the diagram weight. The simplest Monte Carlo procedure samples according to the absolute weights of the diagrams and consists of the following pairs of reciprocal updates:

(i) MOVE–RETRACT. We choose one of the four directions randomly, and attempt to place the head on the site adjacent to the current head site according to this direction. In case this direction does not correspond to backtracking, the current V_1 type of the tail turns into a V_2 , otherwise the head goes back and changes the previous V_2 into a V_1 type (unless the diagram order is 0 or 1, when only V_1 types are possible). When

moving forward, the way of pairing primed and non-primed variables is always unique which in turns implies that we can only retract when the head is connected via a 'straight pair connection' to the previous vertex (both primed and non-primed Grassmann variables of the head are connected to the same vertex on the previous site). We only allow the MOVE–RETRACT updates if the end vertex types are V_1 .

- (ii) SWAP VERTEX. Swaps between the vertexes $V_1 + V_2 \leftrightarrow V_3$ (for head and/or tail) and $V_2 + V_2 \leftrightarrow V_4$ (anywhere in the diagram). This update is its own reciprocal.
- (iii) RELINK. On a given link, relink primed and non-primed Grassmann variables. This can change the sign of the weight only. This update is its own reciprocal.

The second and third type of updates may lead to disconnected diagrams. In such cases, the configuration is unphysical. We opt to allow such configurations, but a Monte Carlo measurement is forbidden and type-1 updates remain impossible until the diagram is connected again. For small values of ζ the sign problem is nearly absent, but only low expansion orders can be reached. For higher values of ζ (close to and above the critical one) an increasing number of orders contributes significantly, consequently more time is spent in higher orders and the sign problem significantly worsens.

A.2. Deterministic full evaluation

For the case of the 2D Ising model, a Monte Carlo approach offers no advantages over a full series expansion approach. With this we mean the explicit listing and evaluation of all possible diagrams as opposed to the stochastic sampling over all topologies. This is because all diagrams in a given expansion order contribute a number of order unity (times the same power of ζ), often with alternating sign, leading to huge cancellations. Only the exact cancellation has physical information, and this requires that every diagram is evaluated multiple times before the correct convergence can be seen. A Monte Carlo approach makes much more sense if the dominant contributions to the total weight are coming from a narrow parameter region, which is usually the case if there are additional integrals over internal momenta.

We therefore wrote a code that evaluates all diagrams for the correlation function up to a maximum order. The construction is based on the fact that there is an easy way to construct all the 'easy' diagrams (the ones that formally look like originating from a high-temperature series expansion). These can serve as parent diagrams, from which further offspring diagrams can be constructed which have one or multiple V_3 and V_4 vertexes as well as possible fermionic exchanges. All diagrams in order *n* can be found as follows:

- (i) Write down all possible words of the form $X_1X_2 \dots X_n$ with the alphabet $X_j \in \{0, 3\}$ corresponding to the four directions on the square lattice. Make sure that subsequent directions are not backtracking. For example, if X_4 is in the positive $+\hat{x}$ direction, then X_5 cannot be in the negative $-\hat{x}$ direction. From this word we also know all sites and links that are visited, as well as all type-1 and type-2 vertexes that are used to make this diagram.
- (ii) Such a parent diagram is added to a list of different topologies only if it has a unique topology. To store the topological information of a bare vertex, we need to store a pair consisting of a site index and a vertex type. The diagram is then stored as an ordered map where the 'key' values are given first by the lattice site index and second by the vertex type (in binary format). The ordered map may have multiple entries with the same key if multiple vertexes reside on the same site and if they are of the same type (e.g., two *RL* vertexes on the same site).
- (iii) We iterate over this configuration list and check if the tail and head sites can be merged into a type-3 vertex by combining them with type-2 vertexes that reside on the same lattice site. If so, and if the resulting topology is unique, the diagram is added to the list. This step is performed in three parts: first for the head and tail together (in order to find all diagrams with 2 V_3 ends), then for the head alone, and finally for the tail alone.
- (iv) We iterate again over the full configuration list and check if 2 type-2 vertexes that live on the same site can be
 merged into a type-4 vertex. This last step has to be repeated until no further merges are possible (since it
 may happen that a diagram has multiple type-4 vertexes or even multiple type-4 vertexes on the same site).
 Diagrams thus created are also added the configuration list if their topology is unique. After completion of
 this step, all possible topologies have been generated.
- (v) We compute the product of all the vertex weights, according to the Feynman rules.
- (vi) From this list of parent diagrams we need to generate all offspring diagrams which feature all possible fermionic permutations for multiply occupied links. This first requires that we know how the vertexes are



Figure B1. Two low-order contributions to the generalized Luttinger–Ward functional Ψ . Dashed lines denote bold Green functions for primed and non-primed Grassmann variables, and wavy solid lines are effective potential lines.

connected in the parent diagram, which is stored in the configuration list. The parent diagram always has permutation sign +1 (because the connections of the primed and non-primed Grassmann variables are always the same). Next we generate all possible permutations by relinking the primed and/or the non-primed Grassmann variables using Heap's algorithm. If a link has occupation number *m*, then there are $(m!)^2$ combinations to be generated (and there may be more than one multiply occupied link). The permutation signature is also stored.

- (vii) We check the connectivity of the diagram using the breadth-first algorithm. Disconnected diagrams contribute 0.
- (viii) Finally, we compute the isomorphism factor: if *m* identical vertexes on the same site are found, a factor 1/m! must be taken into account. This is a consequence of how we construct the diagrams: topology checks were only performed on the parent diagrams (and based on vertexes only), not on offsprings obtained by fermionic exchange. (It would be prohibitively expensive to add the offspring diagrams to the list of all possible diagrams.) Hence, just as we generate illegal disconnected diagrams, we also have a double counting problem when identical vertexes occur in the list.

In order 14, there were about 140 000 parent diagrams contributing to the first entry on the diagonal of the correlator. The hugest number of permutations was $(4!)^4(3!)^4 \approx 10^8$. Since the sum of these permutations has a net contribution of order 1, Monte Carlo has roughly a sign problem of the order of 10^{-8} for these diagrams. The first time a nontrivial isomorphism factor is seen is in order 6 for the first element on the diagonal of the spin correlator: there are diagrams in which two links are doubly occupied, and those links are connected by an identical V_2 vertex, hence the isomorphism factor 1/2. More efficient ways of evaluating and storing the diagrams can probably be devised and implemented, but the above scheme is sufficient to check the validity of the technique and study the transition.

Appendix B. The G²W skeleton scheme

The expansion of susceptibility in terms of ζ is, of course, identical to the one found by the high-temperature series expansion method. To make the distinction between the high-temperature series formalism and Grassmannization approach clear, we discuss the skeleton formulation of the interacting fermionic field theory based on dressed (or 'bold') one-body propagators (*G*) and bold interaction lines (*W*). This leads to the so-called G^2W skeleton scheme (see for instance [38, 39] for the terminology): all lines in all diagrams are assumed to be fully renormalized propagators and effective potentials, but vertex functions remain bare. In section 6 we show that the G^2W -expansion scheme offers a very simple way to solve the 1D Ising model exactly.

B.1. Objects and notation

The key objects in the standard skeleton scheme are the selfenergy (Σ) and the polarization function (Π) . They are related to the Green function (G) and the effective potential (W) by their respective Dyson equations. The diagrams for Π and Σ are obtained by removing one W- or G-line, respectively, from connected graphs for the generalized Luttinger–Ward functional Ψ , shown to second order in figure B1 . In this setup, the expansion order is defined by the number of W-lines (obviously, the discussion of section 3.3 does not apply to the selfconsistent skeleton sequence). All objects of interest are tensors; they have a coordinate (or momentum) dependence, as well as the legs orientation dependence for the incoming and outgoing parts. This conventional scheme has to be supplemented with Ψ -graphs involving V_4 vertexes to account for all contributions. We start with neglecting V_4 vertexes, and discuss their role later.

In more detail, the formalism of the G^2W expansion in the absence of V_4 vertexes is as follows:

(i) There are six bare two-body interaction vertexes V_2 (*RU*, *RL*, *RD*, *LU*, *UD*, *LD*), see the second line in figure 2. They reside on the sites of the original square lattice and all have weight 1. Symbolically, we encode the tensor structure of V_2 using a convenient short hand notation $V_2 = \sum_{\alpha,\gamma=1}^4 V(\alpha, \gamma) n_\alpha n_\gamma$, where

$$V(\alpha, \gamma) = \begin{bmatrix} 0 & 1 & 1 & 1 \\ 1 & 0 & 1 & 1 \\ 1 & 1 & 0 & 1 \\ 1 & 1 & 1 & 0 \end{bmatrix}.$$
 (B.1)

The row index represents the first leg enumerated according to the convention $(R, U, L, D) \rightarrow (0, 1, 2, 3)$, and the column index represents the second leg. By doing so, we artificially double the number of vertexes from 6 to 12. For example, the element (0, 2) corresponds to $n_L n_R$ whereas (2, 0) corresponds to $n_R n_L$, which is exactly the same term.

(ii) The selfenergies Σ for the primed and non-primed Grassmann variables take the same value. Thus, we have to compute only one of them and we can suppress the index that distinguishes between the two Grassmann fields. The selfenergy defines the Green function through the Dyson equation

$$G(\alpha, \gamma) = G^{(0)}(\alpha, \gamma) + \sum_{\mu,\nu} G^{(0)}(\alpha, \mu) \Sigma(\mu, \nu) G(\nu, \gamma).$$
(B.2)

For a link going from site *i* to site *j*, the first index α refers to site *i* (in the above-defined sense), and the second index γ refers to site *j*. Note the absence of the momentum dependence in equation (B.2): the bold Green function remains local on the links in any order of renormalization. It means, in particular, that the only non-zero element for a link between sites (0, 0) and (1, 0) is G_{02} ; it can be alternatively denoted as G_x and, by 90° rotation symmetry of the square lattice, is the same for all links.

(iii) The matrix structure of polarization Π is similar to that of V. The 0th order expression based on bare Green functions is given by

$$\Pi_{(x,y)}^{(0)}(\alpha, \gamma) = \zeta \begin{bmatrix} 0 & 0 & \delta_{x,1}\delta_{y,0} & 0 \\ 0 & 0 & 0 & \delta_{x,0}\delta_{y,1} \\ \delta_{x,-1}\delta_{y,0} & 0 & 0 & 0 \\ 0 & \delta_{x,0}\delta_{y,-1} & 0 & 0 \end{bmatrix}.$$
(B.3)

(iv) The effective potential W is defined through the Dyson equation in momentum representation

$$W_{\boldsymbol{q}}(\alpha,\gamma) = V(\alpha,\gamma) + \sum_{\mu,\nu} V(\alpha,\mu) \Pi_{\boldsymbol{q}}(\mu,\nu) W_{\boldsymbol{q}}(\nu,\gamma).$$
(B.4)

We expect to see signatures of the ferromagnetic transition in matrix elements of $W_{q=0}$ because they directly relate to the divergent uniform susceptibility χ .

B.2. Zeroth order result

To obtain the 0th order result, we replace Π with $\Pi^{(0)}$ in equation (B.4). For any ζ we compute $W_{q=0}$ from equation (B.4) by matrix inversion. We find a divergence at $\zeta_c = 1/3$ (shown in figure B2) that can be also established analytically. We see that W diverges as $(\zeta_c - \zeta)^{-1}$. We get the same power law behavior for the (0, 1) matrix element as well as for the Frobenius norm—they just differ by a constant factor. It is not surprising that our ζ_c is below the exact value for this model; the skeleton approach at 0th order is based exclusively on simple 'bubble' diagrams in terms of bare Green functions that are all positive, leaving to an overestimate of the critical temperature. Fermionic exchange cycles and vertexes with negative weights do not contribute at this level of approximation.

B.3. First order result

We now include the diagrams with one W line for the selfenergy and the polarization. In real space we find

$$\Sigma_x^{(1)} = \Sigma^{(1)} = -G_x W_{(1,0)}(2, 0) = -GW_{(1,0)}(2, 0)$$

$$\Pi^{(1)} = G^4 W_{(0,0)}(0, 0) + \text{cycl.}$$
(B.5)

The matrix structure of $\Pi^{(1)}$ is identical to that of $\Pi^{(0)}$ and is not shown here explicitly. Coupled self-consistent equations (B.2), (B.4) and (B.5), are solved by iterations.







B.4. Second order result

As mentioned previously, to account for second-order terms in Σ , one goes to the second order graphs for Ψ and removes a *G* line, whereas the second-order terms for Π are obtained by removing one *W*-line from the third-order graphs for Ψ . The corresponding expressions in real space are

$$\Sigma^{(2)} = -W_{(0,0)}(0, 0) W_{(0,0)}(2, 2) G^{3}$$

$$\Pi^{(2)}_{(0,0)}(0, 0) = G^{6} W_{(0,0)}(2, 2) W_{(1,0)}(0, 2)$$

$$\Pi^{(2)}_{(1,0)}(0, 2) = G^{6} W^{2}_{(1,0)}(0, 2)$$

$$+ G^{6} W_{(0,0)}(0, 0) W_{(0,0)}(2, 2).$$
(B.6)

The remaining non-zero contributions are obtained by invoking discrete lattice symmetries. Note that to this order the polarization function is extremely local and contains only same site and n.n. terms Again, coupled self-consistent GW-equations are solved by fixed-point iterations. The resulting behavior for *W* is analyzed in figure B3. The transition point has slightly shifted to larger values of ζ compared to the zeroth-order result, and the exponent has also slightly increased.

B.5. Relating Π to the spin correlation function

The G^2W -expansion scheme treats different bare vertexes (see figure 2) on unequal footing: the V_2 vertexes are fully dressed, but the V_4 vertexes are included perturbatively (we neglected them so far). These higher-rank

vertexes have a weight of comparable magnitude to the V_2 vertexes (-2 for V_4 versus +1 for V_2). In addition, the difference in sign between the weights is expected to result in important cancellations between the diagrams and better convergent series for the spin correlation function (this is how ζ_c increases towards its exact value).

Formally, there is no valid reason for neglecting the V_4 vertexes altogether. Let us show how they can be taken care of in the spirit of the shifted action approach [40]. This discussion also gives us the opportunity to explain how the spin correlator is related to the G^2W skeleton expansion, which is most easily understood in the limit $\zeta \ll 1$. By assuming that the skeleton sequence (without V_4) is solved, we introduce the full polarization function $\overline{\Pi}(\hat{\alpha}, \hat{\beta})$ through the Dyson equation

$$\bar{\Pi}_{\boldsymbol{q}}(\alpha,\,\gamma) = \Pi_{\boldsymbol{q}}(\alpha,\,\gamma) + \sum_{\mu,\nu} \Pi_{\boldsymbol{q}}(\alpha,\,\mu) V(\mu,\,\nu) \bar{\Pi}_{\boldsymbol{q}}(\nu,\,\gamma). \tag{B.7}$$

To be specific, we focus on the n.n. element $\rho_{(1,0)}$; similar manipulations hold for any other distance. Now consider all diagrams for this correlator without the V_4 vertexes within the G^2W formulation (see [40]):

- Put one V₁ vertex on the origin site (0, 0) and the other V₁ vertex on the target site **r** = (1, 0), see equation (4.21). There are 4 × 4 = 16 different ways of doing that depending on the directions of legs. Connect the legs with Π_r(α, γ). For example, in the limit of ζ ≪ 1, choosing the (α = 0)-leg on site (0, 0) and the γ = 2-leg on site (1, 0) results in the contributions ζ 4ζ⁵ + …. Similarly, the choice of α = 1 and γ = 1 leads to the contribution ζ³.
- Put V_3 on (0, 0) and V_1 on (1, 0), and connect all legs with $\overline{\Pi}$ lines. There are four ways to orient the V_3 vertex and for each one there are two choices for connecting legs with $\overline{\Pi}$ propagators. The leading contribution to $\rho_{(1,0)}$ goes hence as $-8\zeta^5$.
- Putting V_1 on (0, 0) and V_3 on (1, 0) gives the same contribution by symmetry.
- Put one V_3 vertex on (0, 0) and the other V_3 vertex on (1, 0). Now there are 16 ways of orienting both V_3 vertexes, and for each orientation there are 15 choices for connecting the legs. These contributions start at order $\propto \zeta^9$.

Next, we repeat the above procedure of connecting legs by adding one V_4 vertex, which can be put on any site, after that we can add two V_4 vertexes etc to generate a perturbative expansion in the number of V_4 terms Compared to the original bare series in powers of ζ , we have reordered the series: the effective potential is summing up all V_2 vertexes, whereas we expand (and sample in a Monte Carlo framework) in powers of λ_4 .

To illustrate this framework, let us take $\zeta = 0.01$ and recall that in the bare series $\rho_{(1,0)} = \zeta + 2\zeta^3 + 4\zeta^5 + 12\zeta^7 + \cdots$. The first three terms can be reproduced without V_4 vertexes and with only 1 V_3 on either the origin or the target site, see figures 3–8. The fifth order coefficient originates from 16 'simple' diagrams containing just V_1 and V_2 vertexes without any exchange. The diagrams containing a V_3 vertex yield a coefficient –8, and the exchange diagrams yield a coefficient –4. On a 16 × 16 lattice, the propagators obtained in section B.2 (i.e., to zeroth order) are

$$\bar{\Pi}_{(x,y)=(1,0)}^{(0)}(0,\,2) = 1.00000002 \times 10^{-02},\tag{B.8}$$

$$\bar{\Pi}_{(x,y)=(1,0)}^{(0)}(1,1) = 1.00010011 \times 10^{-06}, \tag{B.9}$$

$$\bar{\Pi}_{(x,y)=(1,0)}^{(0)}(1,2) = 1.00080057 \times 10^{-10}, \tag{B.10}$$

$$\bar{\Pi}_{(x,y)=(0,0)}^{(0)}(1,2) = 1.00020021 \times 10^{-08}.$$
(B.11)

We do not mention explicitly other symmetry-related elements. The sum of all matrix elements for $\bar{\Pi}_{(x,y)=(1,0)}^{(0)}$ is 0.01 000 200 160. One clearly recognizes the coefficients 1, 2 and 16 for the first-, third- and fifth-order contributions to the bare series. Contributions from the V_3 vertexes can be estimated from multiplying $\bar{\Pi}_{(x,y)=(0,0)}^{(0)}(1, 2) \times \bar{\Pi}_{(x,y)=(1,0)}^{(0)}(0, 2)$ which yields $\approx 10^{-10}$. There are four different diagrams, each with weight -2, resulting in the above-mentioned coefficient -8.

On a 16 \times 16 lattice, the propagators obtained in section B.4 (i.e., to second order) are

 $\bar{\Pi}_{(x,y)=(1,0)}(0, 2) = 9.99999980 \times 10^{-03},$ (B.12)

 $\bar{\Pi}_{(x,y)=(1,0)}(1,1) = 1.00009999 \times 10^{-06},$ (B.13)

$$\bar{\Pi}_{(x,y)=(1,0)}(1,2) = 1.00120089 \times 10^{-10},$$
(B.14)

$$\bar{\Pi}_{(x,y)=(0,0)}(1,2) = 1.00020005 \times 10^{-08}.$$
 (B.15)

The sum of all matrix elements for $\overline{\Pi}_{(x,y)=(1,0)}^{(0)}$ is 0.01 000 200 120. One clearly recognizes the coefficients 1, 2 and 12 for first, third and fifth order contributions to the bare series. For the fifth order contribution, we now

obtain 12 instead of 16 thanks to the Grassmann exchange contribution that is accounted for properly at this level of approximation. By adding the V_3 diagrams in the way described above we recover the correct result to this order in ζ (which is +4).

The first instance of a V_4 vertex occurs in order ζ^6 in the bare series. The relevant bare diagrams are the ones for $\rho_{(1,1)}$ with a V_4 vertex on site (1, 0) (and all cases related by the lattice symmetry). Our bold expansion can correctly account for this contribution if we put a V_4 vertex on this site and connect all unpaired legs with $\overline{\Pi}$ propagators. However, with the propagators obtained in section B.4 we are not supposed to account for all possible diagrams in the bare series to order 6 because our bold expansion in section B.4 is only accurate up to order ζ^3 : consider again $\rho_{(1,1)}$ and the bare diagrams where exchanges are possible on the links between the sites (0, 0)–(1, 0) and (1, 0)–(1, 1). Then there are irreducible non-local contributions that are not accounted for in section B.4 with a positive weight that involves exchanges on both links in a correlated fashion. These contributions would obviously be accounted for in higher order corrections to Ψ , when Π becomes non-local. This is also seen in the numerics: the G^2W approach to second order yields a coefficient of 6 for ζ^6 contribution to $\rho_{(1,1)}$, which is below the correct value of 10.

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