O(3) model with Nienhuis action

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We study the O(3) sigma model in $D=2$ on the lattice with a Boltzmann weight linearized in $\beta$ on each link. While the spin formulation now suffers from a sign-problem the equivalent loop model remains positive and becomes particularly simple. By studying the transfer matrix and by performing Monte Carlo simulations in the loop form we study the mass gap coupling in a step scaling analysis. The question addressed is, whether or not such a simplified action still has the right universal continuum limit. If the answer is affirmative this would be helpful in widening the applicability of worm algorithm methods.
1. Introduction

In recent years the efficient worm algorithm simulation method of Prokof’ev and Svistunov [1] has been successfully generalized to the O(N) [2] and CP(N−1) [3] class of nonlinear sigma models with continuous spin manifolds. In this approach one samples the complete set of strong coupling graphs of the model instead of spin configurations. Since this expansion of the partition function in the inverse coupling \( \beta \) is absolutely convergent for finite lattices this is an equivalent representation of the model. The worm method not only deals with the vacuum graphs of the partition function itself but also with those of the two point function at an arbitrary pair of arguments. This enlarged scope of graphs simulated is essential for the efficiency of the simulation dynamics. In addition it makes the two point function a natural observable which becomes accessible with very high precision at the physically interesting large distance. Other observables may be less easily measurable in the graph representation. Further interest in the reformulated theory derives from the possibility to simulate with non-zero chemical potential without running into sign problems [4].

In the above papers standard lattice formulations have been employed. They contain exponential Boltzmann weights on each link and hence arbitrarily high powers of \( \beta \). In the graph formulation this implies that in principle arbitrarily many lines can pile up on top of each other. A significant simplification therefore arises for the O(N) model if we truncate

\[
Z = \int Ds e^{\beta \sum_{\langle xy \rangle} s(x) \cdot s(y)} \rightarrow Z = \int Ds \prod_l \left[ 1 + \tilde{\beta} s(x) \cdot s(y) \right], \quad s \in S_{N-1}. \tag{1.1}
\]

In these formulas \( s(x) \) means an \( N \)-component spin field on all sites of a \( D \) dimensional lattice and \( Ds \) implies independent integrations of the spins over the sphere. Energy bonds arise on links \( l \) of nearest neighbor pairs on a hypercubic lattice in arbitrary dimension. The truncated model trivially has the same degrees of freedom and symmetry and may therefore be naively conjectured to lie in the same universality class (same continuum limit). On the other hand, for \( \tilde{\beta} > 1 \), the weight in (1.1) oscillates and we leave the realm of standard statistical physics. In the equivalent graph form no sign problem appears and we hence use it to numerically investigate universality in this system. In the affirmative case, the permission to employ such truncations could be very useful to extend the range of models that can studied with worm-type algorithms.

The truncated O(N) model has been intensely studied before by Nienhuis and collaborators [5], [6]. They assumed universality and focused on the range \(-2 \leq N \leq 2\) on the two dimensional honeycomb lattice. Here only 3 links surround a site and the graph structure simplifies even more by not allowing any intersections. As a consequence the models in the graph representation could be solved exactly, including the famous XY model \( N = 2 \). The occurring \( \tilde{\beta} \) are smaller than one and the range above one is called unphysical in these references. We add here that \( N = 1 \) is the Ising model, and here both actions are trivially equivalent if we identify \( \tilde{\beta} = \tanh \beta \).

2. Transfer matrix

Specializing to \( D = 2 \) in this section we now investigate the truncated transfer matrix

\[
T = T_0 T_1, \quad T_0 [s', s] = \prod_z \left( 1 + \tilde{\beta} s'(z) \cdot s(z) \right), \quad T_1 [s] = \prod_{\langle xy \rangle} \left( 1 + \tilde{\beta} s(x) \cdot s(y) \right). \tag{2.1}
\]
where $x, y, z$ here are sites on a row of spins and $s, s'$ denote configurations thereon. $T$ acts on states or ‘wave functions’ $\psi[s] \rightarrow (T\psi)[s] = \int Ds' T[s, s'] \psi[s']$ and we notice that the image is a product of monomials of zeroth and first order in $s(x)$. Hence $T$ projects to a Hilbert space of finite dimension $(1 + N)^L$ if there are $L$ sites in the row which we close periodically. This allowed us to straight forwardly set up $T$ as a finite matrix and we could numerically explore its spectrum up to $L = 14$ on a PC and without any special tricks. Since as it stands $T$ is real but not symmetric we first symmetrize it by the similarity transformation

$$T \rightarrow T_0^{-1/2} TT_0^{1/2} , \quad T_0^{\gamma}[s', s] = \prod_z \left( 1 + \tilde{\beta}^{-1} N^{-1} \tau s'(z) \cdot s(z) \right). \quad (2.2)$$

Thus we know that $T$ has real but not necessarily positive eigenvalues and the two-step matrix $T^2$ is real positive.

**Figure 1**: Complete spectrum of $T$ in the O(3) model for $L = 8$ and $\tilde{\beta} = 1.9144$. In Fig. 1 we display a complete spectrum of $T$ in the O(3) model for parameters which lead to a finite volume mass gap $m(L) L = 1.0595$ which is employed in many calculations of the step scaling function [7]. Note, that this is already in the range where (1.1) has a sign problem. The spectrum was obtained by applying the routine eig under matlab and it is ordered and plotted logarithmically against the index. The largest eigenvalue has been normalized to unity and is a single point, the non-degenerate ground state. As expected the first excitation is a triplet, presumably at zero momentum. Then many more points follow – 65536 in total – with the last 21404 of them being negative but small in modulus.

In Fig. 2 (left) we see the mass gap versus $\tilde{\beta}$ for $L = 12$ and $L = 11$. As explained in [7] the mass gap may be considered as a scale dependent renormalized coupling constant $m(L) L = \tilde{g}^2(L)$. Concentrating on the even case first, we notice that $\tilde{g}^2$ comes from large values around (and below) $\tilde{\beta} = 1$, reaches a minimum between 3 and 4 and then rises again. In other words the renormalized coupling as a function of the bare coupling $\beta^{-1}$ never gets very small. This is in contrast to the standard action where the perturbative behavior $\tilde{g}^2 = \beta^{-1} + c(L) \beta^{-2} + O(\beta^{-3})$ holds for larger $\tilde{\beta}$.

On $L = 11$ and other odd size lattices one observes a crossing of eigenvalues of the singlet ground state with the triplet, and as we have set $\exp(-m) = \lambda_1/\lambda_0$ the mass changes sign where
the triplet becomes the states with the largest eigenvalue of $T$. This symmetry breaking corresponds to a violation of the Mermin Wagner theorem which apparently does not apply with the action (1.1). The dominance of a non singlet at large $\tilde{\beta}$ can be qualitatively understood: at large $\tilde{\beta}$ it is favorable to always insert $\tilde{\beta} s' \cdot s$ in (2.1) and not the unity term. In the limit a number of $L$ spins have to be combined which precludes the formation of an over-all singlet in the case of odd $L$. This clearly is an artifact of the locally (per site) finite dimensional Hilbert space caused by our truncation.

3. Worm simulation

For each truncated bond on link $l = \langle xy \rangle$ we insert the trivial identity

$$\left[ 1 + \tilde{\beta} s(x) \cdot s(y) \right] = \sum_{k(l)=0}^{\mathbb{N}} \delta_{k(l),0} + (1 - \delta_{k(l),0}) \tilde{\beta} s_k(x) s_k(y)$$

which leads to a new field $k(l) \in \{0, 1, \ldots, \mathbb{N}\}$. For a given configuration $k$ the spin components appear in a giant monomial and can be integrated out. In this way we can write

$$\mathcal{Z} = \sum_{u, v, c} \rho^{-1}(u, v) \int \delta s \prod_{l} \left[ 1 + \tilde{\beta} s(x) \cdot s(y) \right] s_c(u) s_c(v)$$

as the exactly equivalent graph or loop gas ensemble

$$\mathcal{Z} = \sum_{u, v, c, \{k\}} \rho^{-1}(u, v) \tilde{\beta} \prod_{l} \delta_{k(l),0} \prod_{z} C[q(z)].$$

A configuration is visualized by drawing lines of $N$ different colors on those links that carry one of the $N$ values $k(l) > 0$. At sites $z$

$$q_{a}(z) = \sum_{l, l \in z} \delta_{k(l),a} + \delta_{a,c} (\delta_{a,a} + \delta_{a,v}) \quad (a = 1, 2, \ldots, \mathbb{N})$$
counts the power of the spin component $s_a(z)$ appearing in the monomial at site $z$, and upon integration over the sphere (see e.g. [2]) we get $n!! = 0$ for even $n$

$$C[q] = \int d\mu(s) \prod_{a=1}^{N} (s_a)^{q_a} = \frac{\prod_a (q_a - 1)!!}{N(N+2)\cdots(N+|q|-2)}, \quad |q| = \sum_{a=1}^{N} q_a. \quad (3.5)$$

It is now easy to establish the identity ($V =$ number of sites)

$$\langle s(x) \cdot s(y) \rangle = \rho(x,y) \frac{\langle \delta_{u,a} \delta_{v,y} \rangle}{\langle \delta_{u,v} \rangle / V}. \quad (3.6)$$

where the expectation value on the right hand side is taken in (3.3). The real positive weight function $\rho$ cancels out but is useful to optimize the signal to noise ratio [2].

To simulate the ensemble (3.3) we perform iterations constructed from the following blend of standard worm steps:

i. A move of $u$ to one of its nearest neighbors is proposed, which defines a link $l$; the proposal is immediately rejected unless $k(l) = 0$ or $k(l) = c$; in this case the proposal includes swapping $k(l)$ between these two values and the proposal is metropolis accepted according to the ratio of (finite) weights in (3.3).

ii. If after a move i. the situation $u = v$ is encountered, a joint move to another randomly chosen site is proposed together with a random change of $c$ to a new value. Again an accept test follows.

iii. An iteration consists of $V$ steps of i./ii. followed by a change of the ‘active’ color $c \rightarrow c' \neq c$; this requires us to construct a path from $u$ to $v$ travelling through links with $k(l) = c$ which upon traversal are immediately changed to $c'$; if at a site there are several possibilities to continue the path, we choose randomly between them with equal probabilities. One can prove that the walk necessarily arrives at $v$ where it completes. In addition these moves preserve the weight and no conditional accept is required. The key to the proof is the observation that $(q_a - 1)!!$ is precisely the number of ways to pair objects of color $a$ at a site and that $|q|$ does not change under re-coloring.

Steps of type i. may also be applied to $v$, but this is not required. The combination of i. and ii. alone would be ergodic. However steps of type iii., which were discovered by Holger Stephan [8], are essential to avoid critical slowing down.

4. Numerical results

Based on (3.6) we determine the time-slice (zero momentum) correlation and extract the finite volume mass gap by standard techniques [2], [7]. In Fig. 2 (right) we thus confirm for larger lattices the behavior found with the transfer matrix. To check universal behavior a convenient tool is the step scaling function

$$\sigma(u) = \lim_{L \to \infty} \Sigma(u,L^{-2}), \quad \Sigma(u,L^{-2}) = \bar{g}^2(2L) \bar{g}(L)^2(u) \quad (4.1)$$
where $\sigma$ is known exactly [9] in the asymptotically free $O(N \geq 3)$ models. In this case $\sigma(u) - u > 0$ is the characteristic sign which is true for $\Sigma$ extracted from the pair of lattices in Fig. 2 (right) only in the left part of the figure. We thus are motivated to tune a number of even sized lattices with $L = 8, \ldots, 48$ to the ‘traditional’ value $\bar{g}^2(L) = 1.0595$, which occur at $\tilde{\beta} = 1.9128, \ldots , 3.0800$, and to then measure $\bar{g}^2(2L)$ on the doubled lattices. The result is Fig. 3. We notice that the data points obviously tend toward the continuum value with great precision. The fit in the right panel is given by

$$\Sigma - \sigma = \frac{1}{L^2} \left[ 0.97L^3 - 6.8L^2 + 13L - 7.3 \right]$$

which is the form derived in [10] for a large class of discretizations which however does not include the Nienhuis action employed here. Nevertheless it can accommodate our data with competing powers of logarithms as it has been observed before.

As witnessed by Fig. 2 the range of values $\bar{g}^2$ that can be realized is bounded toward small couplings. We thus decided to repeat an extrapolation for $u = 0.9$ which is about the smallest value where reasonably large $L$ are still possible. This is shown in Fig. 4 with fit coefficients 0, 0.36, $-1.6, 1.7$. While the bound obstructs the use of much larger $L/a$, we notice that on the other hand the lattice artifacts for the available lattices are particularly tiny here. As a final remark we mention that in a brief test we verified that odd size $L$ are of no help to obtain asymptotically free physics for smaller $mL$. Small values are available at $L$, but $2L$ is even again, and even if we the average $2L + 1$ and $2L - 1$ results for the step scaling function are far away from the universal values.

5. Conclusions

We have found in our investigation that the answer to the question whether the Nienhuis discretization of the 2D $O(3)$ model reproduces the known universal physics is not a simple yes or no. Using the well known finite size renormalized coupling based on the mass gap, we found that for each $\bar{g}^2$ there is a limit $L/a \leq f(\bar{g}^2)$ on the available lattice resolutions. In other words, the continuum limit can not be taken ‘all the way’. On the other hand, switching to an effective field
theory point of view familiar from many (all?) areas of physics, we have demonstrated a very precise approximation of the continuum limit by rather unambiguously extrapolating the realizable lattices. As the function $f$ seems to grow with $g^2$, the ‘difficult’ end is the perturbative regime.

Acknowledgements: We thank Erhard Seiler and Peter Weisz for discussions.

References


