HYBRID MOLECULAR DYNAMICS FOR LATTICE SUPERSYMMETRY

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Abstract

We present the first results obtained with a Hybrid Molecular Dynamics algorithm
applied to an $N = 1$ SU(2) Super-Yang–Mills on the lattice. We derive the Hamilton
equations of motion for the system with Wilson gluinos and present preliminary results
on small lattices.
1 Introduction

The supersymmetric extension of the Standard Model is now considered the most promising solution to the gauge hierarchy problem; this problem arises whenever the SM itself is considered as the low-energy effective action of a more general theory including gravity. Moreover, it has recently been found that exact non-perturbative solutions of supersymmetric Yang–Mills theories can be found due to the peculiar properties of extended $N = 2$ supersymmetry. It is then interesting to begin the exploration of the non-perturbative aspects of supersymmetric theories by using their lattice formulation, in order to achieve a better understanding of the confinement and chiral symmetry breaking phenomena. The lattice formulation of a supersymmetric theory seems to be impossible since the discretization of the space–time explicitly breaks supersymmetry. Nevertheless, it has been shown [1] that it is not necessary to put on the lattice an explicitly supersymmetric theory: much as in the case of chiral symmetry, the main point is that the continuum limit of the theory should be supersymmetric. Thus, it is possible to simulate a non-supersymmetric theory with the same field content as the continuum one, with the requirement that in the $a \to 0$ limit the supersymmetric theory is recovered. It has been shown (for $N = 1$ Yang–Mills in [1] and for $N = 2$ in [2]) that the supersymmetric limit actually is strongly related to the chiral limit itself. In this way, in order to study $N = 1$ Super-Yang–Mills (SYM) it is only needed to introduce gluons and massive gluinos (fermion fields in the same representation as the gauge group of the gluons) and then recover the supersymmetric limit going to the $m_g = 0$ chiral one. It is evident from what was said above that in this theory it is fundamental to maintain an exact balance between bosonic and fermionic degrees of freedom. For this reason, an unquenched simulation should be performed, keeping also gluino loops. A recent paper [3] presented a first attempt to do an $N = 1$ SU(2) Yang–Mills simulation on small lattices using the Lüscher local bosonic algorithm. In order to perform a cross-check of the results, we have decided to adopt a different approach, namely the Hybrid Molecular Dynamics (HMD) algorithm of [4].

In section 2 we introduce the Wilson action for SYM theory and the Hamilton equations of motion; in section 3 we present our preliminary results, obtained on a small lattice, and finally in section 4 we draw some conclusions discussing the feasibility of these kind of simulations and giving an outline of our future projects.

2 Supersymmetric Wilson action on the lattice

In this section the basic formalism, the notation and the main properties of the $N = 1$ supersymmetric extension of the SU(2) Yang–Mills theory are introduced. The Hamilton equations for the Molecular Dynamics algorithm are also given.

We recall that the parameter $N$ refers to the number of anti-commuting generators that are introduced in order to make a theory supersymmetric. The easiest supersymmetric extension of a non-Abelian gauge theory is $N = 1$ SU(2) Yang–Mills, and so we will take this theory as our starting point for a study of the non-perturbative aspects of supersymmetric
theories with the lattice approach. The $N = 1$ Yang–Mills action contains two different pieces: the first is the usual Wilson action for the gauge bosons and the second represents the action term for the gauge fermions, the gluinos. We have adopted as fermion action the Wilson formulation in order to avoid the doubling of fermionic flavours. The full SYM action is

$$S_{SYM} = S_g + S_f,$$

where the standard Wilson action for the SU(2) gauge field is

$$S_g = -\frac{\beta}{2} \sum \mathrm{Tr}(U).$$

The fermionic action should describe Majorana spinors in the adjoint representation of the gauge group. It is possible [3] to derive an expression formally identical to the usual Wilson action for the quark in QCD, replacing the link variables $U$ that are $N_c \times N_c$ matrices (where $N_c$ is the rank of the gauge group) with new fields in the adjoint representation $V$, using $(N_c^2 - 1) \times (N_c^2 - 1)$ matrices. We introduce the Majorana fermions as

$$\lambda(x) = T^a \lambda_a(x)$$

with $T^a$ the gauge group generators, satisfying the trace condition

$$\mathrm{Tr}[T^a T^b] = \frac{1}{2} \delta^{ab}.$$ 

For SU(2) we can take $T^a = \frac{1}{2} \sigma^a$, the Pauli matrices. We recall the gauge transformation properties of links and gluinos:

$$U_\mu(x) \rightarrow \Lambda(x) U_\mu(x) \Lambda^\dagger(x + \mu),$$

$$\lambda(x) \rightarrow \Lambda(x) \lambda(x) \Lambda^\dagger(x).$$

The fermionic action, in the Wilson approach, is then given by

$$S_f = \frac{1}{2} \sum_x \left\{ \bar{\lambda}^a(x) \lambda^a(x) - K \sum_\mu \left[ \bar{\lambda}^a(x + \mu) V^{ab\dagger}(x)(1 + \gamma_\mu) \lambda^b(x) + \bar{\lambda}^a(x) V^{ab}(x)(1 - \gamma_\mu) \lambda^b(x + \mu) \right] \right\},$$

where the adjoint link $V_\mu$ is given by

$$V^{ab\dagger}_\mu(x) = \frac{1}{2} \mathrm{Tr}[U_\mu(x) \sigma^a U_\mu(x) \sigma^b].$$

It is easy to verify that $V_\mu$ belongs to O(3), and so $V_\mu^\dagger = V^T_\mu$. In the continuum limit, the fermionic action (6) gives the interaction Lagrangian

$$\mathcal{L}_I = \frac{1}{2} f^{abc} \lambda^a A^b_\mu \gamma_\mu \lambda^c.$$
The action can be rewritten in the following way:

$$S_f = -\frac{1}{4} \text{Tr} \left[ \ln(M^\dagger M) \right], \quad (9)$$

where $M$ is the fermionic matrix. The factor $\frac{1}{4}$ in eq. (9) contains a $\frac{1}{2}$ due to the square root of $\det M^\dagger M$ and another $\frac{1}{2}$ that takes into account the Majorana nature of the gluinos. A detailed description of how the gluino propagator and the $n$-point correlation functions can be obtained in the path integral formalism on the lattice can be found in [3].

In order to simulate this theory, we have adopted the Hybrid Molecular Dynamics approach introduced in [4]. Since the Super-Yang–Mills theory we want to reproduce contains just one gluino flavour, we are forced to use the so-called $R$-algorithm. For this reason we cannot adopt the popular Hybrid Monte Carlo approach, which could be used only with a multiple of 4 Majorana flavours. As usual we introduce the Hermitian field $P_\mu$ conjugate to the gauge field in order to write the Hamiltonian of the system as

$$\mathcal{H} = \frac{1}{2} \sum_{x,\mu} \text{Tr} P_\mu^2(x) + S_g + S_f. \quad (10)$$

The conjugate momenta $P_\mu$ of the link variables are generated at the beginning of every Molecular Dynamics trajectory as complex Gaussian numbers distributed as $\exp(-\frac{1}{2} \text{Tr} P_\mu^2)$.

Following [4] we introduce a noisy estimator for $(M^\dagger M)^{-1}$ in the form $\chi \chi^*$, where $\chi$ is the solution of the matrix equation

$$M^\dagger M \chi = \phi \quad (11)$$

and $\phi = M^\dagger R$. In this expression $R$ is a vector of random Gaussian complex numbers with the same indices as the original gluino field, distributed as $\exp(-R^* R)$.

The Hamilton equations for the system are

$$\dot{U}_\mu(x) = iP_\mu(x)U_\mu(x),$$

$$\dot{P}_\mu(x) = \frac{i}{\beta} \mathcal{T}[G_\mu(x)] - \frac{i}{2} K \mathcal{T}[F_\mu(x)], \quad (12)$$

where the derivative is taken with respect to the Molecular Dynamics “time” and $\mathcal{T}$ is the projection operator on traceless anti-Hermitian matrices:

$$\mathcal{T}[B] = \frac{1}{2} \left[ (B - B^\dagger) - \frac{1}{N_c} \text{Tr}(B - B^\dagger) \right]. \quad (13)$$

The evolution of the conjugate momenta $P_\mu$ fields is driven by the gauge “force”

$$G_\mu(x) = U_\mu(x) \sum_{\nu \neq \mu} \left[ U_\nu(x + \mu)U^\dagger_\mu(x + \nu)U^\dagger_\nu(x) + U^\dagger_\nu(x + \mu - \nu)U^\dagger_\mu(x - \nu)U_\nu(x - \nu) \right] \quad (14)$$

and by the fermion “force”

$$F_\mu(x) = A^{ab}_\mu(x) \text{Re}[B^{ba}_\mu(x)], \quad (15)$$
where

\[
A_{\mu}^{ab}(x) = \sigma^a U_\mu^\dagger(x) \sigma^b U_\mu(x),
\]

\[
B_{\mu}^{ba}(x) = \text{Tr} \left[ (1 - \gamma_\mu) Y^b(x) \chi^a(x + \mu) + (1 + \gamma_\mu) \chi^b(x) Y^a(x + \mu) \right]_{\text{Dirac}}
\]

(16)

and \( Y = M \chi \).

As in [3], we assume that the sign of the Pfaffian operator (the square root of the fermion determinant) remains positive during the evolution of the system in the Molecular Dynamics “time”. In principle, we could take this phase into account by inserting it in the definition of the observables.

3 Preliminary results

In this section we present the first results we obtained with the Hybrid Molecular Dynamics algorithm for the Super-Yang–Mills SU(2) \( N = 1 \) theory.

We have performed a test of the method using the same parameters as reported in [3], namely \( \beta = 2.0 \) and \( K = 0.150 \). We have run the program on a lattice with the same size \( (4^3 \times 8) \), obtaining compatible results for the plaquette, the absolute value of the Polyakov loop and the \( 2 \times 2 \) Creutz ratio, defined in terms of \( k \times l \) Wilson loops \( W(k, l) \) as

\[
\chi^2 = \frac{W(2, 2)W(1, 1)}{W(2, 1)W(1, 2)}.
\]

(17)

In all the runs the time–length of the Molecular Dynamics trajectories between different measurements was kept fixed at 0.5. This implies a number of leap-frog steps \( N_{\text{step}} = 5, 10, 20 \) for the three simulations performed. Our statistics is of the order of \( N_m = 10000 \) measures for the full theory runs, and \( N_m = 5000 \) in the pure gauge case. In this last case, we have used method 1 of [5]. With a short heat-bath run we obtain for the SU(2) pure gauge plaquette at \( \beta = 2.0 \) the value \( \Box_{PG} = 0.5008(3) \).

It is important to stress that the algorithm used introduces an intrinsic \( O(\Delta t^2) \) error, where \( \Delta t \) is the Molecular Dynamics time step. It is then necessary to extrapolate to the \( \Delta t = 0 \) limit in order to obtain physical results. Furthermore, the approach of the plaquette to the physical value is expected to be from above. Both behaviours, namely the \( \Delta t^2 \) dependence and the approach from above can be seen in fig. 1 where the expectation value of the plaquette is presented as a function of \( \Delta t \).

We present in table 1 the results obtained for the three observables introduced above. The results are fully compatible with those reported in [3]: for example the value of the plaquette extrapolated to \( \Delta t = 0 \) is \( \Box = 0.5058(4) \). We have no data for \( |P| \) and \( \chi^2 \) at \( \Delta t = 0.025 \), but the values obtained at larger \( \Delta t \) are quite flat and already compatible with [3].

On the same observables, we have studied the autocorrelation times at different time steps, with the windowing method [8]. We build the connected autocorrelation function
Table 1: Average observables for $\beta = 2.0$, $K = 0.150$, $V = 4^3 \times 8$ at different values of $\Delta t$. We show the plaquette, the absolute value of the Polyakov loop and the $2 \times 2$ Creutz ratio. $N_m$ is the number of measurements for each run.

\begin{table}[h]
\centering
\begin{tabular}{|c|c|c|c|c|c|}
\hline
$\Delta t$ & $\Box$ & $|P|$ & $\chi_2$ & $N_m$ \\
\hline
0.100 & 0.5098(3) & 0.0505(4) & 0.5555(12) & 10000 \\
0.050 & 0.5065(3) & 0.0500(4) & 0.5559(13) & 8000 \\
0.025 & 0.5063(3) & - & - & 10000 \\
\hline
\end{tabular}
\end{table}

$C(t)$ and from this an estimate for the integrated autocorrelation time

$$\tau_{\text{int}}(T) = \frac{1}{2} + \sum_{t=1}^{T} \frac{C(t)}{C(0)}.$$  \hspace{1cm} (18)

When the ratio $T/\tau_{\text{int}}(T)$ is around 4 we should observe a plateau in the value of $\tau_{\text{int}}(T)$: this is exactly what happens, as can be seen in fig. 2 in a particular case. Results for integrated autocorrelation times are presented in table 2, together with a rough estimate of the errors.

\begin{table}[h]
\centering
\begin{tabular}{|c|c|c|c|}
\hline
$\Delta t$ & $\tau_{\Box}$ & $\tau_{|P|}$ & $\tau_{\chi_2}$ \\
\hline
0.100 & 7.5(5) & 1.0(2) & 2.8(3) \\
0.050 & 6.8(5) & 1.0(2) & 2.7(3) \\
0.025 & 6.7(5) & - & - \\
\hline
\end{tabular}
\end{table}

Table 2: Autocorrelation times for the different observables at different values of $\Delta t$.

Finally, we present a rough quantitative comparison of the relative speed of the two algorithms used for these simulations (namely, HMD and the Lüscher local bosonic one adopted in [3]); an extensive study of the scaling of the autocorrelation times with the different parameters relevant to the simulations has yet to be performed and, moreover, it is likely that further improvements can be implemented. Since the most time-consuming part of both algorithms is the application of the fermion matrix to a vector, we consider the “effort” $E$ needed to obtain an independent configuration from a previous one as the number of times this application is performed, times the autocorrelation time of a particular observable (for definiteness, the plaquette). For each simulation, introducing $N_{\text{app}}$ as the number of times the fermion matrix is applied per step, we obtain

$$E_{\text{HMD}} = N_{\text{step}} \times N_{\text{app}} \times \tau.$$  \hspace{1cm} (19)

For this preliminary study we have not yet tried to optimize the algorithm: in particular, so as to solve the matrix equation (11), we used the Stabilized Bi-Conjugate Gradient without preconditioning. With this particular inverter and with the \textit{leap-frog} scheme used to integrate the Hamilton equation,

$$N_{\text{app}} = 2 + 4N_{\text{iter}},$$  \hspace{1cm} (20)
Figure 1: Average plaquette versus $\Delta t$ in the pure gauge case (squares) and within the full theory (circles). The upper cross at $\Delta t = 0$ is reported by Montvay, while the lower one is the heat bath pure gauge result. The dotted line is the extrapolation to $\Delta t = 0$.

Figure 2: Integrated autocorrelation time versus $T/\tau_{\text{int}}$ for the plaquette at $\Delta t = 0.025$. 
where $N_{\text{iter}}$ is the average number of iterations required to invert the fermion matrix. The convergence criterion we used for all our runs was $|M^\dagger M_{\chi} - \phi|^2 < 10^{-8}|\chi|^2$, obtaining $N_{\text{iter}} \simeq 25$. For the most time-consuming simulation ($\Delta t = 0.025, N_{\text{step}} = 20$), $E_{\text{HMD}} \simeq 14000$; for the whole set of three simulations at different $\Delta t$, we obtain $E_{\text{HMD}} \simeq 25000$.

For the local bosonic algorithm used in [3], the “effort” is computed as

$$E_{\text{LB}} = N_{\text{field}} \times N_{\text{app}} \times \tau,$$

where $N_{\text{field}}$ is the number of bosonic field used in the simulation. Using the results reported in [3], we estimate (for the two most time-consuming simulations) $E_{\text{LB}} \simeq 90000$ for the one-step algorithm with 24 bosonic fields, and $E_{\text{LB}} \simeq 25000$ for the two-steps algorithm with 8 fields.\footnote{These numbers can be decreased to 44000 and 15000 respectively, by changing the mixture of heat-bath and over-relaxation sweeps in the update of the bosonic fields [6].}

For a realistic study of the theory with the HMD algorithm, it is obvious that the inversion algorithm will have to be accelerated, for example by means of the standard red–black preconditioning, which we are currently implementing. Another important improvement of the algorithm can be achieved by using a higher-order Hamilton equations integration scheme, of the kind proposed in [7], in order to accelerate the convergence towards the $\Delta t = 0$ limit.

4 Conclusions

In this paper, following the outline presented in recent papers [3, 2], we have begun an exploration of the feasibility of supersymmetric gauge theories simulations on the lattice. From a theoretical point of view, it is possible to study these kinds of theories on the lattice using the same approach adopted for chiral theories as usual QCD, namely the implementation of a non-chiral non-supersymmetric theory requiring the recovery of these symmetries in the continuum limit. We have obtained results compatible with those presented by Montvay in [3] using a completely different algorithm, the Hybrid Molecular Dynamics one. From these first simulations it seems possible, with the present computer power and the present knowledge of fermionic algorithms (such as the local bosonic, the hybrid family or even with the “negative flavour number extrapolation” approach [9]), to afford the study of non-perturbative aspects of supersymmetric extensions of gauge theories.

The next step is to begin the investigation of the supersymmetric limit of the theory, which is expected to be reached jointly with the chiral limit [1]. In order to do that, we are planning to study the asymptotic behaviour of the scalar and the pseudoscalar gluino “meson”. In the OZI approximation it is expected that the pseudoscalar mass approaches zero when going to the chiral limit, while the scalar should go to a non-zero value.

Finally, we plan to investigate the behaviour of the gluino condensate. This quantity is expected to be a relevant parameter in the study of non-perturbative supersymmetry
It is worth while to mention that a clear comprehension of the supersymmetry-breaking phenomena is of the uttermost importance if the supersymmetric extension of the Standard Model has to be a viable solution to the gauge hierarchy problem. In order to study this observable, it is necessary to understand its renormalization properties. It is perhaps important to introduce a non-perturbative renormalization scheme both on the action and on the operators [10], which seems to be relevant in the study of the chiral (supersymmetric) limit of the theory with Wilson fermions.

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References