Improving the dynamical overlap algorithm

Nigel Cundy
Universität Wuppertal
E-mail: cundy@theorie.physik.uni-wuppertal.de

Stefan Krieg∗
NIC/ZAM Forschungszentrum Jülich/Universität Wuppertal
E-mail: s.krieg@fz-juelich.de

Thomas Lippert
NIC/ZAM Forschungszentrum Jülich/Universität Wuppertal
E-mail: t.lippert@fz-juelich.de

We present algorithmic improvements to the overlap Hybrid Monte Carlo algorithm, including preconditioning techniques and improvements to the correction step, used when one of the eigenvalues of the Kernel operator changes sign, which is now $O(\Delta t^2)$ exact.
1. Improved Correction Step

The sign function in the overlap Dirac operator creates a discontinuity $-2 d$ in the pseudo-fermion contribution to the action whenever an eigenvalue of the kernel operator changes sign. To conserve energy, we integrate up to the computer time $\tau_c$ where the eigenvalue crosses, and introduce a discontinuity in the kinetic energy which exactly cancels the jump in the pseudo-fermion energy. A general area conserving and reversible update which can do this is:

$$\Pi^+ = \Pi^- + (\eta(\eta, \Pi^-)) \left( \sqrt{1 + \frac{4d_0}{(\eta, \Pi^-)^2}} - 1 \right) + \sum_{j=1}^{N} \left( \eta^j_1(\eta^j_1, \Pi^-) + \eta^j_2(\eta^j_2, \Pi^-) \right)$$

$$\times \left( \sqrt{1 + \frac{4d_j}{(\eta^j_1, \Pi^-)^2 + (\eta^j_2, \Pi^-)^2}} - 1 \right) + A,$$

where $\Pi^-$ is the original momentum, $\Pi^+$ the final momentum, $A$ is an arbitrary function of the gauge field at $\tau_c$, $\eta$ is a unit vector normal to the $\lambda = 0$ surface, and the $\eta^j_i$ are unit vectors normal to $\eta$. The original algorithm [5] set $d_1 = 4 d$ and $d_j = 0$, and had $O(\tau_c)$ errors. We can use the $d_j$ terms to cancel these errors, giving the transmission algorithm:

$$\Pi^+ = \Pi^- + \tau_c(F) - \eta \tau_c(F, F) - \frac{\tau_c}{3} \text{Tr}(F) + (\eta, \Pi^-) \left( \sqrt{1 + \frac{4d}{(\eta, \Pi^-)^2}} - 1 \right) + (\eta^2_1(\eta^2_1, \Pi^+) + \eta^2_2(\eta^2_2, \Pi^+)) \times \left( \sqrt{1 + \frac{4d_2}{(\eta^2_1, \Pi^+ - F)^2 + (\eta^2_2, \Pi^+ - F)^2}} - 1 \right)$$

$$d_2 = -2\tau_c(F^-, \eta)(\Pi^-, \eta) + 2\tau_c(F^+, \eta)(\Pi^+, \eta) + 2\tau_c(F^+, F^-).$$

$$\hat{F} = \frac{\tau_c}{2}(F^- + F^+)$$

where $F^\pm$ are the MD forces immediately before and after the crossing. We cannot use this algorithm if it would lead to complex $\Pi^+$. In this case, we have to reflect of the $\lambda = 0$ surface, and there will be no topological charge change. Figure 5 shows how the energy difference across the correction step varies as a function of $\Delta \tau$. It clearly shows that the energy has errors of at maximum $O(\Delta \tau^3)$.

2. Improved Leapfrog algorithm

In [3] an alternative leapfrog update for the molecular dynamics part of the HMC is suggested:

1. $\Pi(\tau + \lambda \Delta \tau) = \Pi(\tau) + \lambda \Delta \tau \Pi(\tau).$
2. $U(\tau + \Delta \tau/2) = e^{i(\Delta \tau/2)\Pi(\tau + \lambda \Delta \tau)}U(\tau).$
3. $\Pi(\tau + (1 - \lambda) \Delta \tau) = \Pi(\tau) + (1 - 2\lambda) \Delta \tau \Pi(\tau + \lambda \Delta \tau).$
4. $U(\tau + \Delta \tau) = e^{i(\Delta \tau/2)\Pi(\tau + (1 - \lambda) \Delta \tau)}U(\tau + \Delta \tau/2).$
5. $\Pi(\tau + \Delta \tau) = \Pi(\tau + (1 - \lambda) \Delta \tau) + \lambda \Delta \tau \Pi(\tau + (1 - \lambda) \Delta \tau).$

The optimal value of $\lambda$ is given in [3]. This algorithm has improved energy conservation, which more than compensates for the need to invert the overlap operator twice. We have tested it on $4^4$, $8^4$, and $12^4$ lattices, and found gains of around 30% (see section 7).
Improving the dynamical overlap algorithm

Stefan Krieg

Figure 1: Dependency of the energy on $\Delta \tau$. The red lines are from top down: $(\Delta \tau, \Delta \tau^2, \Delta \tau^3)$.

3. Stout Smearing

We use the “stout links” proposed in [3]. As mentioned in [3] this improves the condition number of the Wilson operator substantially, thus speeding up the inversions needed to construct the overlap operator. We find however that there is a “phase transition” at a critical level of the smearing parameter, leading to a sharp increase in the magnitude of the smallest eigenvalue of the Wilson operator. This reduces the effectiveness of the smearing.

4. Hasenbusch acceleration

Hasenbusch acceleration has been used to speed up dynamical simulations. We introduce an additional fermion flavour with a large mass, and by placing the two fermions on different time scales we can in principle reduce the number of low mass inversions needed during a trajectory. However, we saw little gain when using this method, partly because we were testing on large masses, and partly because our overlap operators are usually well conditioned (see section 7).

5. Overlap eigenmode preconditioning

In the case of a topological nontrivial configuration, the spectrum of the overlap matrix includes a “zero mode”. Inversions of the overlap operator become prohibitively expensive when simulating in the regime of small quark masses. Our ansatz is to calculate the smallest $m$ eigenvectors $\Psi_m$ and eigenvalues $\lambda_m$ of the overlap operator to a very low precision (e.g. $10^{-2}$) and use them as a preconditioner for our CG preconditioner in our GMRESR inverter. Our preconditioner
Figure 2: Convergence history for the preconditioning method on configuration with trivial topology

is:

\[ P = 1 + \sum_m \left( \frac{\alpha_m}{\lambda_m} - 1 \right) |\Psi_m⟩⟨\Psi_m| \]

Figures 2 and 3 show the convergence of CG with and without preconditioning using above projector. These plots were generated using a $8^4$ dynamical configuration at mass $\mu = 0.1$, with the inversions carried out at mass $\mu = 0.03$. Figure 2 shows the convergence history for the case of a configuration with trivial topology; Figure 3 shows the convergence history for a configuration with a “zero mode” induced by topology: Clearly in the latter case the preconditioning offers great possible gains, which — according to our experience — increase with the volume and decreasing of the masses.

In an HMC simulation, using the previous eigenvectors as a starting point for the next eigenvalue calculation can dramatically reduce the time needed, although it is unclear how large an effect this leads has on the reversibility of the MD.
6. Non area conserving correction step

It is possible to use a non area conserving molecular dynamics update by including the Jacobian in the Metropolis accept/reject step\(^1\). The detailed balance condition reads:

\[
P[U → U′]W_C[U] = \int d\Pi d\Pi' \exp^{-\frac{1}{2}\Pi^2} \delta([U, \Pi] - T_{MD}[U', \Pi']) \min(1, \exp^\Delta) W_C[U]
= \int d\Pi d\Pi' \exp^{-\frac{1}{2}\Pi^2} \delta([U', -\Pi'] - T_{MD}^{-1}[U, -\Pi]) \frac{\partial U', \Pi'}{\partial U, \Pi} \min(1, \exp^\Delta) W_C[U]
= \int d\Pi d\Pi' \exp^{-\frac{1}{2}\Pi^2} \delta([U', -\Pi'] - T_{MD}^{-1}[U, -\Pi]) \min(1, \exp^{-\Delta}) W_C[U']
= P[U ← U′]W_C[U′]
\]

\[
\Delta = -\ln\left[\frac{\partial U', \Pi'}{\partial U, \Pi}\right] + S_G[U] + \frac{1}{2}\Pi^2 - S_G[U'] - \frac{1}{2}\Pi'^2
\]

The most general transmission update which is reversible and conserves \(\Delta\) is:

\[
\exp^{-((\Pi + \eta)^2/2)} = \exp^{-((\Pi - \eta)^2 - 2d)} - \exp^{-r_0^2/2 - 2d} + \exp^{-r_0^2/2}
\]

For \(r_0 = \infty\), this gives the usual area conserving transmission formula equation (1.1). One has to reflect if the transmission formula gives a complex \((\Pi^+, \eta)\). By tuning \(r_0\), we can improve the transmission rate. The results displayed in the tables of section 7 were obtained using \(r_0 = 1\), and give a 50% improvement in the transmission rate.

\(^1\)We thank A. Borici for pointing this out to us
7. Results

In this section we summarise the results referred to in the previous sections.

\[
\begin{array}{|c|c|c|c|c|c|}
\hline
\text{Type} & \text{time} & \text{Acc} & n_{\text{md}} & \text{trans./traj.} & \text{refl./traj.} & n_t \\
\hline
\text{normal} & 1897(60) & 94\% & 40 & 0.0738(240) & 1.348(100) & 325 \\
\text{has} & 1986(20) & 88\% & 40 & 0.0521(311) & 1.059(94) & 307 \\
\text{imp} & 1420(10) & 94\% & 15 & 0.0535(233) & 0.876(98) & 299 \\
\text{imphas} & 1594(40) & 75\% & 15 & 0.0772(336) & 1.093(118) & 324 \\
\text{imnnap} & 1480(10) & 95\% & 15 & 0.117(34) & 1.336(136) & 310 \\
\text{imphas} & 1611(60) & 78\% & 15 & 0.110(21) & 0.832(159) & 155 \\
\hline
\end{array}
\]

\(\mu = 0.05\)

\[
\begin{array}{|c|c|c|c|c|c|}
\hline
\text{Type} & \text{time} & \text{Acc} & n_{\text{md}} & \text{trans./traj.} & \text{refl./traj.} & n_t \\
\hline
\text{normal} & 1816(20) & 95\% & 40 & 0.447(64) & 0.938(80) & 465 \\
\text{has} & 2100(90) & 90\% & 40 & 0.569(65) & 0.880(65) & 374 \\
\text{imp} & 1479(20) & 96\% & 15 & 0.371(43) & 0.947(62) & 533 \\
\text{imphas} & 1470(60) & 90\% & 15 & 0.413(53) & 0.531(76) & 518 \\
\text{imnnap} & 1445(50) & 95\% & 15 & 0.674(89) & 1.818(147) & 209 \\
\text{imnaphas} & N/A & 94\% & 15 & 0.663(69) & 1.370(114) & 281 \\
\hline
\end{array}
\]

\(\mu = 0.2\)

In these tables “has” denotes Hasenbusch acceleration, “imp” denotes usage of the preconditioner and “nap” refers to the non area preserving update.

References

[1] N. Cundy et.al. “Numerical Methods for the QCD Overlap Operator IV: Hybrid Monte Carlo” hep-lat/0502007

[2] N. Cundy et.al. “Numerical Methods for the QCD Overlap Operator: III. Nested Iterations”, Comp. Phys. Comm. 165 (2005) 841 hep-lat/0405003

[3] C. Morningstar, M. Peardon “Analytic Smearing of SU(3) Link Variables in Lattice QCD”, Phys.Rev. D69 (2004) 054501 hep-lat/0311018

[4] T. DeGrand, S. Schaefer “Physics issues in simulations with dynamical overlap fermions” Phys.Rev. D71 (2005) 034507 hep-lat/0412005

[5] Z. Fodor, S.D. Katz, K.K. Szabo, “Dynamical overlap fermions, results with hybrid Monte-Carlo algorithm” JHEP 0408 (2004) 003 hep-lat/0311010

[6] T. Takaishi, P. de Forcrand “Testing and tuning new symplectic integrators for Hybrid Monte Carlo algorithm in lattice QCD” hep-lat/0505020