

## Generating QCD Gauge Configurations

# The Fourth International Workshop on Numerical Analysis and Lattice QCD

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## Talk Outline

- Introduction
- Hybrid Monte Carlo
- Algorithm Improvement
- "Non-local" Algorithms
- Conclusion





## Introduction

• Lattice QCD path integral

$$\langle \Omega \rangle = \frac{1}{Z} \int [dU] e^{-S_{g}(U)} [\det \mathcal{M}(U)]^{\alpha} \Omega(U)$$

 $\alpha = \frac{N_f}{2} \left(\frac{N_f}{4}\right)$  for Wilson (staggered) fermions,  $\mathcal{M} = M^{\dagger}M$ 

- $10^8 10^9$  degrees of freedom  $\Rightarrow$  Monte Carlo integration
- Interpret  $e^{-S_{\rm g}} \det \mathcal{M}^{\alpha}$  as a Boltzmann weight, and use importance sampling

$$\langle \Omega 
angle pprox rac{1}{N} \sum_{i=1}^{N} \Omega(U_i)$$





#### Quenched Approximation

- Fermion determinant extremely non-local object
- Quenched approximation: set  $\det \mathcal{M}=1$
- Gauge action local: over relaxed heatbath algorithms very efficient
- Just plain wrong!



 $\Rightarrow$  Must include dynamical fermions to obtain QCD





#### The HMC Algorithm (Duane et al)

- De facto algorithm for including dynamical fermions
- Rewrite determinant in terms of pseudo-fermions

$$\det \mathcal{M} = \int D\phi^{\dagger} D\phi \, e^{-\phi^{\dagger} \mathcal{M}^{-1} \phi} = \int D\phi^{\dagger} D\phi \, e^{-S_{\mathsf{f}}}$$

- Need global updates since psuedofermion action is non-local
- $\bullet$  Introduce fictitious momentum field  $\pi$  and define a Hamiltonian

$$H = \frac{1}{2} \operatorname{tr} \pi^2 + S_{g} + S_{f} = T + S$$

- Integrate Hamilton's equations to propose a new configuration
- Global Accept / Reject to obtain desired probability distribution

$$P(U,\phi) = \frac{1}{Z}e^{-S_{g}-S_{f}}$$





#### The HMC Algorithm (Duane et al)

- Each update consists of
  - Hybrid Molecular Dynamics Trajectory
    - \* Momentum refreshment heatbath  $(P(\pi) \propto e^{-\pi^* \pi/2})$ .
    - \* Pseudo-fermion heatbath ( $\phi \propto M^{\dagger}\xi$ , where  $P(\xi) \propto e^{-\xi^{*}\xi}$ ).
    - \* MD trajectory with  $\tau/\delta\tau$  steps.
  - Metropolis Acceptance Test  $P_{acc} = min(1, e^{-\delta H})$





#### Molecular Dynamics

- Hamilton's equations  $\frac{dU}{d\tau} = \frac{dT}{d\pi} = \pi$  and  $\frac{d\pi}{d\tau} = -\frac{dS}{dU} = F$
- $\bullet$  Must discretize the "fictious time"  $\tau$  and integrate numerically
- $\bullet$  Define integrators in terms of evolution operators Q and P

$$Q \equiv \frac{dT}{d\pi} \frac{\partial}{\partial U} \quad \text{with} \quad e^{\delta \tau Q} : f(U, \pi) \to f(U + \delta \tau T'(\pi), \pi)$$
$$P \equiv -\frac{dS}{dU} \frac{\partial}{\partial \pi} \quad \text{with} \quad e^{\delta \tau P} : f(U, \pi) \to f(U, \pi - \delta \tau S'(U))$$

• Metropolis requires Detailed Balance

- Integration must be reversible and area preserving

- Use Symmetric Symplectic Integrators, e.g., leapfrog

$$U(\delta\tau)^{\tau/dt} = \left(e^{\delta\tau P/2}e^{\delta\tau Q}e^{\delta\tau P/2}\right)^{\tau/\delta\tau} + O(\delta\tau^2)$$





## Molecular Dynamics Forces

- Pure Gauge Force  $(\frac{dS_g}{dU})$  local analytic quantity - CHEAP
- Fermion force

$$\frac{dS_{f}}{dU} = \frac{d}{dU}\phi^{\dagger}\mathcal{M}^{-1}\phi = -\phi^{\dagger}\mathcal{M}^{-1}\frac{d\mathcal{M}}{dU}\mathcal{M}^{-1}\phi$$

- Each update to the momentum requires solution to  $\mathcal{M}\chi=\phi$
- Generally calculated using a Krylov solver, e.g., CG
- EXPENSIVE





#### Cost of HMC

- Condition number blows up as  $m \to 0$
- Force  $\propto 1/m$ , requires  $\delta au \to 0$  to maintain acceptance rate
- Also, as  $m \rightarrow 0$ , correlation lengths diverge
- $C \propto \left(\frac{m_{\pi}}{m_{\rho}}\right)^{-6} L^5 a^{-7}$ CP-PACS and JLQCD, 2002



 $\Rightarrow$  Require huge computers OR better algorithms





## Chronological Inverter

- Solution  $\chi(\tau) = \mathcal{M}(\tau)^{-1}\phi$  is a smooth function
- Idea: Use previous solutions to act as an initial guess
- Minimize over the space of previous solutions (Broweret al) : for

$$x_0 = \sum_i c_i \mathcal{M} \chi_i$$

solve

$$\chi_j^{\dagger}\phi = \sum_i c_i \chi_j^{\dagger} \mathcal{M} \chi_i$$

- Requires high precision solutions to maintain reversibility
- Gain around a factor of 2





#### Higher Order Integrator

• Potential for gain from using  $O(\delta \tau^4)$  integrator, e.g., Campostrini

$$U(\delta\tau)^{\tau/dt} = \left(e^{\delta\tau \,\epsilon P/2} e^{\delta\tau \,\epsilon Q} e^{\delta\tau \,(1-\sigma)P/2} e^{-\delta\tau \,\epsilon\sigma Q} e^{\delta\tau \,(1-\sigma)P/2} e^{\delta\tau \,\epsilon Q} e^{\delta\tau \,\epsilon P/2}\right)^{\tau/\delta\tau} + O(\delta\tau^4)$$

- Better volume scaling  $V^{9/8}$  vs.  $V^{5/4}$
- Constructed from sub-leapfrog steps with  $\delta\tau^{sub} > \delta\tau$
- Sub-leapfrog integrator can go unstable much sooner than  $\delta\tau$  suggests (Joó et al)
- Higher order integrators are very unstable
- $O(\delta \tau^2)$  usually found to be optimal





#### Multiple Timescale Integration (Sexton/Weingarten, 1992)

- For Hamiltonians of the form  $H = T + S_1 + S_2$
- Integrate  $S_1$  and  $S_2$  force contributions on different timescales

$$U(\delta\tau)^{\tau/\delta\tau} = \left( \left( e^{\delta\tau P_1/4m} e^{\delta\tau Q/2m} e^{\delta\tau P_1/4m} \right)^m e^{\delta\tau P_2} \right)^m \left( e^{\delta\tau P_1/4m} e^{\delta\tau Q/2m} e^{\delta\tau P_1/4m} \right)^m \right)^{\tau/\delta\tau}$$

- Two separate timescales  $\delta\tau^{S_1}=\delta\tau/m$ ,  $\delta\tau^{S_2}=\delta\tau$
- Large and cheap force =  $P_1$ , Small and expensive force =  $P_2$ ,
- Naïve partitioning:  $S_1 = S_g, S_2 = S_f$  fails as  $m \to 0$
- Can extend this recursively for N timescales  $H = T + S_1 + \ldots + S_N$
- Great idea, but dormant for 10 years...





#### Multiple Pseudofermions with Mass Preconditioning

• Mass-precondition the fermion determinant (Hasenbusch)

$$\det(M^{\dagger}M) = \det(\hat{M}^{\dagger}\hat{M}) \det\left(\hat{M}(M^{\dagger}M)^{-1}\hat{M}^{\dagger}\right)$$

with  $m(\hat{M}) > m(M)$ 

- Tune  $\kappa(\hat{M}^{\dagger}\hat{M}) \approx \kappa(\hat{M}(M^{\dagger}M)^{-1}M^{\dagger}) \approx \sqrt{\kappa(M^{\dagger}M)}$  (Hasenbusch-Jansen)
- Why does this work?
  - Better sampling of Gaussion integral using multiple pseudo-fermions
  - This reduces fluctuations in the fermion force
  - Fermion force  $F\propto\kappa^{\nu}$
- Factor 2 improvement through  $\delta\tau$  increase
- Use more than one dummy operator
- Gain increases as  $m_{\rm I} \rightarrow 0$





#### Multi-timescale Mass Preconditioning (QCDSF, Urbach et al)

- Tune dummy operators so that most expensive (preconditioned) force contributes the least
- Use multi-timescale integrator ( $S_1 = S_g$ ,  $S_2 = \hat{S}$ ,  $S_3 = S_f$ )
- Factor 10 improvement at light quark mass







## Domain Decomposition (Lüscher)



- Can rewrite the Dirac operator  $M = \begin{pmatrix} D_{\Omega} & D_{\delta\Omega} \\ D_{\delta\Omega'} & D_{\Omega'} \end{pmatrix}$
- $D_{\Omega}$   $(D_{\Omega'})$  is Dirac operator on black (white) blocks with Direchlet boundaries
- $D_{\delta\Omega}$  and  $D_{\delta\Omega'}$  is the Dirac operator connecting these blocks
- Rewrite determinant

$$\det M = \det \begin{pmatrix} D_{\Omega} & 0\\ 0 & D_{\Omega'} \end{pmatrix} \det \begin{pmatrix} D_{\Omega} & D_{\delta\Omega} \\ D_{\delta\Omega'} & D_{\Omega'} \end{pmatrix} \begin{pmatrix} D_{\Omega}^{-1} & 0\\ 0 & D_{\Omega'}^{-1} \end{pmatrix}$$

- Separation into large and cheap, and small but expensive ⇒ multi timescale integrator
- Large speed up over naïve algorithm





#### Determinant Preconditioning is Key

- Other methods
  - U.V. Filtering (de Forcrand)
  - Polynomial filtering (Peardon and Sexton)
  - Multistep stochastic correction (see talk by Enno)
- All improvements rely on determinant preconditioning





#### Non-local Actions

- Strange quark inclusion requires det  $\mathcal{M}^{\frac{1}{2}}$
- Finite temperature calculations typically use staggered quarks
  - Remnant chiral symmetry important here
  - Non-local action: det  $\mathcal{M}^{\alpha}$ ,  $\alpha = \frac{1}{2}, \frac{1}{4}$
- HMC cannot be applied for these cases
- Inexact algorithms traditionally used
- Is using exact algorithms more expensive?
- Are the improvements in "Local actions" applicable here?





#### The R Algorithm (Gottlieb et al)

• Rewrite fermionic determinant:

 $\det \mathcal{M}^{\alpha} = \exp\left(\alpha \operatorname{tr} \ln \mathcal{M}\right) = \exp\left(-S_{\text{eff}}\right)$ 

- Integrate Hamilton's equations as before
- Use noisy estimator  $\xi$  for trace  $\equiv$  pseudo-fermion force
- Leading order error term  $O(\delta \tau)!$
- Recover  $O(\delta \tau^2)$  with  $N_f$  dependent  $\xi$  updating
  - Non-reversible
  - Jacobian  $\neq$  1
- Cannot include Metropolis acceptance test
   ⇒ Algorithm is inexact
- Naïve cost = HMC, but requires extrapolation to zero stepsize
- Stepsize rule of thumb  $\delta au \sim rac{2}{3}m_{
  m I}$





#### Polynomial Hybrid Monte Carlo (de Forcrand-Takaishi, Frezzotti-Jansen)

• Write in pseudo-fermion notation

$$det \mathcal{M}^{\alpha} = \int \mathcal{D}\bar{\psi}\mathcal{D}\psi e^{-\bar{\psi}\mathcal{M}^{-\alpha}\psi} \\ \approx \int \mathcal{D}\bar{\psi}\mathcal{D}\psi e^{-\bar{\psi}P(\mathcal{M})\psi},$$

where  $P(\mathcal{M})$  is valid over spectrum

- Pseudo-fermion heatbath easily realised  $P(\mathcal{M}) = p^{\dagger}(\mathcal{M})p(\mathcal{M})$
- Use standard MD leapfrog  $\Rightarrow$  exact
- Generally polynomial degree  $m > N_{\text{iter}} \text{ CG}$  iterations
- Use low degree polynomial
  - Reweight acceptance test or observable
- Use high degree polynomial
  - $\alpha \neq$  1 derivative uses Leibniz rule  $\Rightarrow$  Memory, rounding









- Generated using Remez algorithm
- Real non-degenerate roots (poles are always +ve)
- Partial fractions  $r(x) = \sum_{k=1}^{m} \frac{\alpha_k}{x + \beta_k}$
- Evaluate using multi-shift solver
- Numerically stable ( $\alpha_k$  have same sign)





#### Rational Hybrid Monte Carlo (Clark-Kennedy)

• Rewrite fermionic determinant

$$\det \mathcal{M}^{\alpha} = \int \mathcal{D}\bar{\phi}\mathcal{D}\phi e^{-\bar{\phi}\mathcal{M}^{-\alpha}\phi}$$
$$\approx \int \mathcal{D}\bar{\phi}\mathcal{D}\phi e^{-\bar{\phi}r^{2}(\mathcal{M})\phi},$$

with  $r(x) = x^{-\alpha/2}$ 

- Precision is cheap: Conventional Metropolis
- RHMC:
  - Hybrid Molecular Dynamics Trajectory
    - \* Momentum refreshment heatbath  $(P(\pi) \propto e^{-\pi^* \pi/2})$ .
    - \* Pseudo-fermion heatbath ( $\phi \propto r(\mathcal{M})^{-1}\xi$ , where  $P(\xi) \propto e^{-\xi^*\xi}$ ).
    - \* MD trajectory with  $\tau/\delta\tau$  steps.
  - Metropolis Acceptance Test  $P_{acc} = min(1, e^{-\delta H})$





#### Rational Hybrid Monte Carlo (Clark-Kennedy)

- MD trajectory
  - Double inversion from  $r^2(\mathcal{M})$
  - Use low degree approx  $\bar{r}\approx \mathcal{M}^{-\alpha}\approx r^2$
  - Pseudo-fermion force

$$S'_{\text{pf}} = -\sum_{i=1}^{\bar{m}} \bar{\alpha}_i \phi^{\dagger} (\mathcal{M} + \bar{\beta}_i)^{-1} \mathcal{M}' (\mathcal{M} + \bar{\beta}_i)^{-1} \phi.$$

- CG cost per trajectory  $\approx$  HMC
  - One extra inversion required for heatbath





Exact vs. Inexact (RBC-Bielefeld)



- P4 staggered fermions
- RHMC allows an O(10) increase in stepsize
- Speedup greater as  $m_{\rm I} \rightarrow 0$







#### Exact vs. Inexact (de Forcrand-Philipsen)

(Naïve Staggered Fermions,  $N_f = 3$ ,  $V = 8^34$ )

- Results:
  - Binder cumulant increases
  - Stepsize extrapolation is vital for R algorithm
  - 25% reduction in critical quark mass at  $\delta \tau^R = \frac{1}{2}m_{\rm I}$
  - 20% change in renormalised quark mass
- Conclusion: "an exact algorithm is mandatory"





## Multiple Pseudofermions with RHMC (Clark-Kennedy)

• Rewrite determinant

$$\det \mathcal{M} = [\det \mathcal{M}^{1/n}]^n \\ \propto \prod_{j=1}^n d\phi_j \, d\phi_j^{\dagger} \exp\left(-\phi_j^{\dagger} \mathcal{M}^{-1/n} \phi_j\right),$$

- So called  $n^{th}$  root trick
- Speedup through  $\delta \tau$  increase
- No dummy mass parameters to tune  $\Rightarrow$  easy to increase n
- Single fermion timescale





## Integrator Instability (Clark-Kennedy)



(Staggered fermions,  $V = 16^4$ ,  $\beta = 5.6$ ,  $N_{f} = 2$ , m = 0.005)

- With n = 1 integrator breaks down as  $\delta \tau$  is increased
- Instability "tickled" by low fermion modes  $\sim O(\frac{1}{m_1})$  (Joó et al)
- Higher order integrators are more ticklish
- What happens with multiple pseudofermions?





#### Integrator Instability (Clark-Kennedy)



(Staggered fermions,  $V = 16^4$ ,  $\beta = 5.6$ ,  $N_f = 2$ , m = 0.005)

- Removes instability in the integrator!
- Why does this work? Lowest modes now  $O(\frac{1}{m_{I}})^{1/n}$
- Force now bulk dominated
- Higher order integrators now beneficial





Who's the fastest of them all? ( $N_f = 2$  Wilson)

- Compare multi-timescale mass preconditioning and high order RHMC
- Use popular testing parameters ( $V = 24^3.32$ ,  $\beta = 5.6$ , Plaquette + Wilson fermions)
- Use measure  $C = \tau_{\text{int}}^{\text{plag}} N_{\text{mv}} . 10^{-4}$

	C		
$\kappa$	RHMC	Urbach et al	Orth et al
0.15750	9.6	9.0	19.1
0.15800	29.9 <mark>*</mark>	17.4	128
0.15825	52.5 <mark>*</mark>	56.5	_

\*Using 4MN5 fourth order integrator (de Forcrand and Takaishi)

- RHMC similar in cost to mass preconditioning
- Look at integrator stability with mass preconditioning
  - Gain from higher order integrators also?





## 2+1 QCD

• e.g. Domain Wall 2+1 flavour determinant

$$\left( \frac{\det M_{\mathsf{I}}^{\dagger} M_{\mathsf{I}}}{\det M_{\mathsf{pv}}^{\dagger} M_{\mathsf{pv}}} \right) \left( \frac{\det M_{\mathsf{s}}^{\dagger} M_{\mathsf{s}}}{\det M_{\mathsf{pv}}^{\dagger} M_{\mathsf{pv}}} \right)^{1/2} = \left( \frac{\det M_{\mathsf{I}}^{\dagger} M_{\mathsf{I}}}{\det M_{\mathsf{s}}^{\dagger} M_{\mathsf{s}}} \right) \left( \frac{\det M_{\mathsf{s}}^{\dagger} M_{\mathsf{s}}}{\det M_{\mathsf{pv}}^{\dagger} M_{\mathsf{pv}}} \right)^{3/2}$$

- Mass Precondition using the strange quark
- Use  $n^{th}$  root trick for triple strange
- NOT mutually exclusive improvements
- Use multi-timescale integrator (gauge, triple strange, light)
- Light quark mass constitutes around 10% CG cost
- Cost dependence on mass comes mostly from autocorrelation





## Berlin Wall Plot

- Compare the cost of fermion formulations and/or algorithms
  - $N_f = 2 + 1$  DWF RHMC (RBC-UKQCD)
  - $N_f = 2$  Mass preconditioned Wilson (Urbach *et al*)
  - $N_{\rm f} = 2$  Mass preconditioned Clover (QCDSF)
  - $N_f = 2 + 1$  Mass preconditioned Clover + RHMC

(Wuppertal-Jülich)

- $N_{\rm f} = 2$  Mass preconditioned Twisted Mass (ETM)
- $N_{\rm f} = 2 + 1$  AsqTad R (MILC)
- $N_f = 2 + 1$  AsqTad RHMC (Clark-Kennedy)
- All data scaled to  $V = 24^3 \times 40$ , a = 0.08
- Box is too small and too coarse









-31- Yale University





-32- Yale University





-33- Yale University



#### Conclusions

- After 20 years HMC is still the best dynamical algorithm
- The last 5 years has seen an explosion in HMC improvement
- Determinant preconditioning is the key behind all improvement
- Non-local actions are no problem
- Pick and mix the most appropriate algorithm
  - Multiple time scale mass preconditioning
  - Domain decomposition
  - $n^{th}$  rootary with RHMC
- HMC cost is now  $C \propto \left(\frac{m_{\pi}}{m_{\rho}}\right)^{-2} L^5 a^{-6}$  (Lüscher)
- Physical point no longer a pipedream
- Further improvement must come from autocorrelation?

