FINITE TEMPERATURE AND CHEMICAL
POTENTIAL IN LATTICE QCD
AND ITS CRITICAL POINT

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Abstract

We propose a method to study lattice QCD at finite temperature (T) and chemical potential (µ). We compare the method with direct results and with the Glasgow method by using \(n_f=4\) QCD at \(\text{Im}(\mu)\neq 0\). We locate the critical endpoint (E) of QCD on the \(\text{Re}(\mu)-T\) plane. We use \(n_f=2+1\) dynamical staggered quarks with semi-realistic masses on \(L_t=4\) lattices. Our results are based on \(\mathcal{O}(10^3 \ldots 10^4)\) configurations.

QCD at finite \(T\) and/or \(\mu\) is of fundamental importance, since it describes relevant features of particle physics in the early universe, in neutron stars and in heavy ion collisions. Extensive experimental work has been done with heavy ion collisions at CERN and Brookhaven to explore the \(\mu-T\) phase boundary (cf. [1]). Note, that past, present and future heavy ion experiments with always higher and higher energies produce states closer and closer to the \(T\) axis of the \(\mu-T\) diagram. It is a long-standing question, whether a critical point exists on the \(\mu-T\) plane, and particularly how to tell its location theoretically [2].

Let us discuss first the \(\mu=0\) case. Universal arguments [3] and lattice simulations [4] indicate that in a hypothetical QCD with a strange (s) quark mass \((m_s)\) as small as the up (u) and down (d) quark masses \((m_u,d)\) there would be a first order finite \(T\) phase transition. The other extreme case \((n_f=2)\) with light u/d quarks but with an infinitely large \(m_s\) there would be no phase transition only a crossover. Observables change rapidly during a crossover, but no singularities appear. Between the two extremes there is a critical strange mass \((m_s^c)\)

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at which one has a second order finite $T$ phase transition. Staggered lattice results on $L_t=4$ lattices with two light quarks and $m_s$ around the transition $T$ ($n_f=2+1$) indicated $[3]$ that $m_c^*$ is about half of the physical $m_s$. Thus, in the real world we probably have a crossover.

Returning to a non-vanishing $\mu$, one realizes that arguments based on a variety of models (see e.g. $[4,5,2]$) predict a first order finite $T$ phase transition at large $\mu$. Combining the $\mu=0$ and large $\mu$ informations an interesting picture emerges on the $\mu$-$T$ plane. For the physical $m_s$ the first order phase transitions at large $\mu$ should be connected with the crossover on the $\mu=0$ axis. This suggests that the phase diagram features a critical endpoint $E$ (with chemical potential $\mu_E$ and temperature $T_E$), at which the line of first order phase transitions ($\mu>\mu_E$ and $T<T_E$) ends $[2]$. At this point the phase transition is of second order and long wavelength fluctuations appear, which results in (see e.g. $[8]$) consequences, similar to critical opalescence. Passing close enough to $(\mu_E,T_E)$ one expects simultaneous appearance of signatures which exhibit nonmonotonic dependence on the control parameters $[9]$, since one can miss the critical point on either of two sides.

The location of $E$ is an unambiguous, non-perturbative prediction of QCD. No ab initio, lattice analysis based on QCD was done to locate the endpoint. Crude models with $m_s=\infty$ were used (e.g. $[2]$) suggesting that $\mu_E \approx 700$ MeV, which should be smaller for finite $m_s$. The goal of our exploratory work is to show how to locate the endpoint by a lattice QCD calculation. We use full QCD with dynamical $n_f=2+1$ staggered quarks.

QCD at finite $\mu$ can be given on the lattice $[10]$; however, standard Monte-Carlo techniques fail. At Re($\mu$)$\neq0$ the determinant of the Euclidean Dirac operator is complex, which spoils any importance sampling method.

Several suggestions were studied in detail to solve the problem. We list a few of them (for a recent review see Ref. $[11]$).

In the large gauge coupling limit a monomer-dimer algorithm was used $[12]$. For small gauge coupling an attractive approach is the “Glasgow method” $[13]$ in which the partition function is expanded in powers of $\exp(\mu/T)$ by using an ensemble of configurations weighted by the $\mu=0$ action. After collecting more than 20 million configurations only unphysical results were obtained: a premature onset transition. The reason is that the $\mu=0$ ensemble does not overlap sufficiently with the states of interest. Another possibility is to separate the absolute value and the phase of the fermionic determinant and use the former to generate configurations and the latter in observables $[14]$.

At imaginary $\mu$ the measure remains positive and standard Monte Carlo techniques apply. The canonical partition function can be obtained by a
Fourier transform \([14, 15]\). In this technique the dominant source of errors is the Fourier transform rather than the poor overlap. One can also use the fact that the partition function away from the transition line should be an analytic function of \(\mu\), and the fit for imaginary \(\mu\) values could be analytically continued to real values of \(\mu\) \([17]\). At \(T\) sufficiently above the transition, both real and imaginary \(\mu\) can be studied by dimensionally reducing QCD \([18]\). Hamiltonian formulation may also help studying the problem \([13]\).

![Diagram](image)

**Fig. 1:** Schematic difference between the present and the Glasgow methods.

An attractive approach to alleviate the problem is the “Glasgow method” (see e.g. Ref. \([13]\)) in which the partition function \((Z)\) is expanded in powers of \(\exp(\mu/T)\) by using an ensemble of configurations weighted by the \(\mu=0\) action. After collecting more than 20 million configurations only unphysical results were obtained. The reason is that the \(\mu=0\) ensemble does not overlap enough with the finite density states of interest \([20]\).

We propose a method to reduce the overlap problem and determine the phase diagram in the \(\mu\)-\(T\) plane (for details see \([21]\)). The idea is to produce an ensemble of QCD configurations at \(\mu=0\) and at \(T_c\). Then we determine the Boltzmann weights \([23]\) of these configurations at \(\mu \neq 0\) and at \(T\) lowered to the transition temperatures at this non-vanishing \(\mu\). Since transition configurations are reweighted to transition ones a much better overlap can be observed than by reweighting pure hadronic configurations to transition ones \([13]\). Since the original ensemble is collected at \(\mu=0\) we do not expect to be able to describe the physics of the large \(\mu\) region with e.g. exotic colour superconductivity. Fortunately, the typical \(\mu\) values at present heavy ion accelerators are smaller than the region we cover (for the freeze-out condition see e.g. \([24]\)).

After illustrating the applicability of the method we locate the critical point of QCD. (Multi-dimensional reweighting was successful for determining the endpoint of the hot electroweak plasma \([22]\) e.g. on 4D lattices.)
Let us study a generic system of fermions $\psi$ and bosons $\phi$, where the fermion Lagrange density is $\bar{\psi}M(\phi)\psi$. Integrating over the Grassmann fields we get:

$$Z(\alpha) = \int D\phi \exp[-S_{\text{bos}}(\alpha, \phi)] \det M(\phi, \alpha),$$

where $\alpha$ denotes a set of parameters of the Lagrangian. In the case of staggered QCD $\alpha$ consists of $\beta$, $m_q$ and $\mu$. For some choice of the parameters $\alpha=\alpha_0$ importance sampling can be done (e.g. for $\text{Re}(\mu)=0$). Rewriting eq. (1)

$$Z(\alpha) = \int D\phi \exp[-S_{\text{bos}}(\alpha_0, \phi)] \det M(\phi, \alpha_0)$$

$$\left\{ \exp[-S_{\text{bos}}(\alpha, \phi) + S_{\text{bos}}(\alpha_0, \phi)] \frac{\det M(\phi, \alpha)}{\det M(\phi, \alpha_0)} \right\}. \quad (2)$$

We treat the curly bracket as an observable (measured on each configuration) and the rest as the measure. Changing only one parameter of the ensemble generated at $\alpha_0$ provides an accurate value for some observables only for high statistics. This is ensured by rare fluctuations as the mismatched measure occasionally sampled the regions where the integrand is large. This is the overlap problem. Having several parameters the set $\alpha_0$ can be adjusted to get a better overlap than obtained by varying only one parameter.

![Fig. 2: $\bar{\psi}\psi$ as a function of $\text{Im}(\mu)$, for direct results (squares), our technique (crosses) and Glasgow-type reweighting (dots).](image)

The basic idea of the method as applied to dynamical QCD can be summarized as follows. We study the system at $\text{Re}(\mu)=0$ around its transition point. Using a Glasgow-type technique we calculate the determinants for each configuration for a set of $\mu$, which, similarly to the Ferrenberg-Swendsen method [23], can be used for reweighting. The average plaquette values can be used to
perform an additional reweighting in $\beta$. Since transition configurations were reweighted to transition ones a much better overlap can be observed than by reweighting pure hadronic configurations to transition ones as done by the Glasgow-type techniques. The differences between the two methods are shown in Figure 1. Moving along the transition line was also suggested by Ref. [16].

Fig. 3: $\text{Im}(\beta_0^\infty)$ as a function of the chemical potential.

We have directly tested these ideas in $n_f=4$ QCD with $m_q=0.05$ dynamical staggered quarks. We first collected 1200 independent $V=4\cdot 6^3$ configurations at $\text{Re}(\mu)=\text{Im}(\mu)=0$ and some $\beta$ values and used the Glasgow-reweighting and also our technique to study $\text{Re}(\mu)=0$, $\text{Im}(\mu)\neq 0$. At $\text{Re}(\mu)=0$, $\text{Im}(\mu)\neq 0$ direct simulations are possible. After performing these direct simulations as well, a clear comparison can be done. Figure 2 shows the predictions of the three methods for the average quark condensates at $\beta=5.085$ as a function of $\text{Im}(\mu)$. The predictions of our method agree with the direct results, whereas for larger $\text{Im}(\mu)$ the predictions of the Glasgow method are by several standard deviations off. We expect that our method can be applied at $\text{Re}(\mu)\neq 0$.

In QCD with $n_f$ staggered quarks one should change the determinants to their $n_f/4$ power in our two equations. Importance sampling works also in this case at some $\beta$ and at $\text{Re}(\mu)=0$. Since $\det M$ is complex an additional problem arises, one should choose among the possible Riemann-sheets of the fractional power in eq. (4). This can be done by using [21] the fact that at $\mu=\mu_w$ the ratio of the determinants is 1 and it should be a continuous function of $\mu$.

In the following we keep $\mu$ real and look for the zeros of $Z$ for complex $\beta$. At a first order phase transition the free energy $\propto \log Z(\beta)$ is non-analytic. A phase transition appears only in the $V\to \infty$ limit, but not in a finite $V$. Nevertheless, $Z$ has zeros at finite $V$, generating the non-analyticity of the free energy, the Lee-Yang zeros [25]. These are at complex parameters (e.g. $\beta$).
For a system with first order transition these zeros approach the real axis as $V \to \infty$ by a $1/V$ scaling. This $V \to \infty$ limit generates the non-analyticity of the free energy. For a system with crossover $Z$ is analytic, and the zeros do not approach the real axis as $V \to \infty$.

Fig. 4: The $T$-$\mu$ diagram. Direct results are given with errorbars. Dotted line shows the crossover, solid line the first order transition. The box gives the uncertainties of the endpoint.

At $T \neq 0$ we used $L_t=4, L_s=4, 6, 8$ lattices. $T=0$ runs were done on $10^3, 16$ lattices. $m_{u,d}=0.025$ and $m_s=0.2$ were our bare quark masses. At $T \neq 0$ we determined the complex valued Lee-Yang zeros, $\beta_0$, for different $V$-s as a function of $\mu$. Their $V \to \infty$ limit was given by a $\beta_0(V) = \beta_0^\infty + \zeta/V$ extrapolation. We used 14000, 3600 and 840 configurations on $L_s=4, 6$ and 8 lattices, respectively. $\text{Im}(\beta_0^\infty)$ is shown on Figure 3 as a function of $\mu$. For small $\mu$-s the extrapolated $\text{Im}(\beta_0^\infty)$ is inconsistent with a vanishing value, and predicts a crossover. Increasing $\mu$ the value of $\text{Im}(\beta_0^\infty)$ decreases, thus the transition becomes consistent with a first order phase transition (overshooting is a finite $V$ effect). Our primary result is $\mu_{end} = 0.375(20)$.

To set the physical scale we used a weighted average of $R_0$, $m_\rho$ and $\sqrt{\sigma}$. Note, that (including systematics due to finite $V$) we have $(R_0 \cdot m_\pi) = 0.73(6)$, which is at least twice, $m_{u,d}$ is at least four times as large as the physical values.

Figure 4 shows the phase diagram in physical units, thus $T$ as a function of $\mu_B$, the baryonic chemical potential (which is three times larger then the quark chemical potential). The endpoint is at $T_E = 160 \pm 3.5$ MeV, $\mu_E = 725 \pm 35$ MeV. At $\mu_B=0$ we obtained $T_c = 172 \pm 3$ MeV.

We proposed a method– an overlap improving multi-parameter reweighting technique– to numerically study non-zero $\mu$ and determine the phase diagram in the $T$-$\mu$ plane. Our method is applicable to any number of Wilson or staggered quarks. As a direct test we showed that for $\text{Im}(\mu) \neq 0$ the predictions
of our method are in complete agreement with the direct simulations, whereas
the Glasgow method suffers from the well-known overlap problem. We studied
the $\mu$-$T$ phase diagram of QCD with dynamical $n_f=2+1$ quarks. Using our
method we obtained $T_E=160\pm3.5$ MeV and $\mu_E=725\pm35$ MeV for the endpoint.
Though $\mu_E$ is too large to be studied at RHIC or LHC, the endpoint would
probably move closer to the $\mu=0$ axis when the quark masses get reduced. At
$\mu=0$ we obtained $T_c=172\pm3$ MeV. More work is needed to get the final values
by extrapolating in the R-algorithm and to the thermodynamic, chiral and
continuum limits. The details of the presented results can be found in [21].

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