QCD Phase Diagram According to the Center Group

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We study an effective theory for QCD at finite temperature and density which contains the leading center symmetric and center symmetry breaking terms. The effective theory is studied in a flux representation where the complex phase problem is absent and the model becomes accessible to Monte Carlo techniques also at finite chemical potential. We simulate the system by using a generalized Prokof’ev-Svistunov worm algorithm and compare the results to a low temperature expansion. The phase diagram is determined as a function of temperature, chemical potential, and quark mass. The shape and quark mass dependence of the phase boundaries are as expected for QCD. The transition into the deconfined phase is smooth throughout, without any discontinuities or critical points.

Obtaining a deeper understanding of the QCD phase diagram will be one of the central goals of particle physics in the coming years. With running and upcoming experiments that drive this development, also the theoretical side is challenged to improve our understanding of the QCD phase structure. Analyzing phase transitions is clearly a nonperturbative problem, and suitable techniques have to be applied. For a vanishing chemical potential, lattice QCD is a powerful method that provides reliable quantitative information on the QCD finite temperature transition. However, for nonvanishing density the notorious complex phase problem so far limits numerical lattice QCD studies to painfully small volumes.

For quenched QCD, where the quark contributions to the path integral are neglected, the deconfinement transition is related to the center group \( \mathbb{Z}_3 \) of SU(3), which is a symmetry in the confined low temperature phase, while it is broken spontaneously above the deconfinement temperature [1]. When one couples the dynamics of the quark fields, the center symmetry is broken explicitly by the fermion determinant. This explicit breaking overlays the spontaneous breaking of the quenched theory. However, as for spin systems, one may expect that also for QCD the underlying symmetry still governs parts of the dynamics of the full theory, e.g., via the center properties of canonical determinants [2].

In order to study the role of center symmetry for the QCD phase diagram, we analyze an effective theory which contains the leading center symmetric and center symmetry breaking terms. It can be mapped exactly to a flux representation [3], without a complex phase problem. So far, the model was studied only in a very limited parameter range [3–7]. Here we apply a generalization of the worm algorithm [8] which allows us to efficiently explore the full range of temperatures and chemical potential values. This constitutes one of very few examples where a QCD-related complex phase problem can be solved. We study the phase diagram of the effective theory and analyze what role the center degrees of freedom of QCD play for the phase structure of hot and dense matter.

Effective center theory and flux representation.—The effective center theory is defined by the action [3]

\[
S[P] = -\frac{1}{C_3} \sum_x \left( \tau \sum_{\mu=1}^3 \left( P_x^\mu P_{x+\hat{\mu}}^\mu + \text{c.c.} \right) + \eta P_x + \bar{\eta} P_x^* \right),
\]

The dynamical degrees of freedom are the center variables \( P_x, \eta, \bar{\eta} \), and the action (1) is invariant under global center transformations \( P_x \to z P_x, \quad z \in \mathbb{Z}_3 \). The form of this term may be obtained from a strong coupling expansion of the effective action for the Polyakov loop which in quenched QCD is the order parameter for center symmetry and for confinement. The variables \( P_x \) take over the role of the local Polyakov loops. Although in full QCD center symmetry is broken explicitly by the quarks, the Polyakov loop is still used to monitor confinement properties and to determine the crossover temperature (see, e.g., [9,10]). The strong coupling expansion also identifies the parameter \( \tau \) as an increasing function of the temperature \( T \) of the underlying lattice QCD theory, and for brevity we refer to \( \tau \) as temperature.

The second term of (1) may be obtained from a hopping (i.e., large quark mass) expansion of the fermion determinant and contains the leading center symmetry breaking contributions. The parameters \( \eta = e^{i \mu} \) and \( \eta = e^{-i \mu} \) are related to the chemical potential \( \mu \). The expansion shows that \( \kappa = N_f \hbar(m) \), where \( N_f \) is the number of flavors and \( h(m) \) is a function of the QCD quark mass \( m \) which decreases with increasing \( m \). We refer to \( \kappa \) as the inverse mass parameter.
For $\kappa = 0$, the model reduces to the 3-state Potts model, which is known to have a first-order transition at $\tau = 0.183522(3)$ [11]. For small $\kappa$ and vanishing $\mu$, the first-order transition persists, giving rise to a short first-order line which terminates at a critical end point at $(\tau, \kappa) = [0.183127(7), 0.00026(3)]$ [11]. For a small nonzero $\mu$, the system has been analyzed with techniques based on the Swendsen-Wang cluster algorithm [4], with reweighting [5] and with imaginary $\mu$ [6]. Within the flux representation [3] local Metropolis updates were also used [3,4,7]. It has been demonstrated that turning on the chemical potential softens the transition and shifts the critical end point towards smaller values of $\kappa$. So far, no simulations were done in the parameter region where the complex phase problem of the formulation (1) becomes severe (see [4] for a discussion of that regime).

The flux representation [3] solves the complex phase problem. We briefly summarize it to discuss our observables and conventions: The Boltzmann factors for the nearest neighbor terms of (1) can be rewritten as

$$e^{[P,P^c]_{+ \tau + \epsilon cc.}} = C \sum_{b_{x \tau}} B^{b_{x \tau}} (P_x \tilde{P}_{x \tau})^{b_{x \tau}}. \quad (2)$$

The sum is over flux variables $b_{x \tau} \in \{-1, 0, +1\}$ attached to the links of the lattice. The constants $C$ and $B$ depend on the temperature $\tau$ via $C = (e^{2\tau} + 2e^{-\tau})/3$ and $B = (e^{2\tau} - e^{-\tau})/3C$. Similarly,

$$e^{\eta P_{x \tau} + \tilde{\eta} \tilde{P}_{x \tau}} = \sum_{s_{x \tau}} M_{s_{x \tau}} P_{x \tau}^{s_{x \tau}}, \quad (3)$$

where we sum over monomer variables $s_{x \tau} \in \{-1, 0, +1\}$ attached to the sites $x$. It is straightforward to work out the monomer weights $M_s$ for $s = -1, 0, +1$,

$$M_s = \left[ \frac{1}{3} e^{\eta + \tilde{\eta}} + 2e^{-2(\eta + \tilde{\eta})/3} \cos \left( \frac{\sqrt{3}}{2} (\eta - \tilde{\eta}) - s \frac{2\pi}{3} \right) \right]. \quad (4)$$

The weights $M_s$ turn out to be non-negative.

Inserting (2) and (3) gives rise to a complete factorization of the dependence on the dynamical variables $P_x$, and the sum over all configurations can be performed. One finds (we drop an irrelevant overall constant)

$$Z = \sum_{\{b, s\}} W(b, s) \prod_x T \left( \sum_s [b_{x,\tau} - b_{x, -\tau}] + s_x \right). \quad (5)$$

$Z$ is now a sum over configurations $\{b, s\}$ of the flux and monomer variables. Each configuration comes with a real non-negative weight factor $W(b, s) = (\prod_{x,\tau} b^{b_{x,\tau}}) \times (\prod_x M_{s_x})$. For every link with nonvanishing flux a factor $B$ is taken into account. Sites $x$ contribute with factors $M_{s_x}$ according to the $s_x$.

The flux-monomer configurations are subject to constraints: In (5), $T(n)$ is the triality function defined as $T(n) = \delta_{n \mod 3, 0}$. The constraints enforce that at every site $x$ the total flux from both, flux variables $b_{x,\tau}$ and monomers $s_x$, is a multiple of 3. In the flux form (5) the partition sum contains only real and non-negative contributions and thus the complex phase problem is solved.

In this Letter, we focus on bulk observables such as the order parameter $P$ and the corresponding susceptibility, which both are obtained as derivatives of the free energy: $\langle P \rangle = \partial \ln Z/\partial \eta$ and $\chi_P = \partial^2 \ln Z/\partial \eta^2$. In a similar way, one obtains the internal energy $U$ and the heat capacity $C$. For an efficient evaluation the identities $\partial M_{s_x}/\partial \eta = M_0$, $\partial M_0/\partial \eta = M_{-1}$, and $\partial M_{-1}/\partial \eta = M_{+1}$ are useful. In the end, all our observables are expressed in terms of the total flux and monomer numbers and their moments.

_simulation with the worm algorithm._—Having established the flux representation, we now must find a suitable algorithm for an efficient Monte Carlo update. We here use a generalized form of the Prokof’ev-Svistunov worm algorithm [8]: The worm starts at a randomly chosen site and moves along links until it returns to the starting point where it terminates. We allow for two different moves of our worm: (A) The worm randomly chooses a new direction at a site and changes the flux at the corresponding link by $\pm 1$; (B) the worm decides to change a monomer variable by $\pm 1$ and then randomly hops to another site where the monomer variable is changed by $\mp 1$. The moves are offered with equal probability, produce only configurations that are compatible with the constraint, and lead to an ergodic algorithm. The Metropolis acceptance probabilities are $p_A = B^6$ when changing a flux variable $b$ by an amount of $\delta_b$ (move A) and $p_B = M_s/M_0$ for changing a monomer variable from $s$ to $s'$ (move B). A more complete account of the algorithm and its implementation will be given elsewhere.

We generated ensembles for lattice sizes $36^3$, $48^3$, $64^3$, and $72^3$. For the inverse mass parameter we used $\kappa = 0.1$, $\kappa = 0.01$, $\kappa = 0.005$, and $\kappa = 0.001$. The evaluation of our observables $\langle P \rangle$, $\chi_P$, $U$, and $C$ is based on up to $10^6$ configurations, separated by 10 worms for decorrelation. Autocorrelation times were determined and used in the estimate for the statistical errors. Finite volume effects were analyzed by comparing the different system sizes and are negligible for our final results.

The results from the new worm algorithm were checked by using several strategies. For vanishing $\mu$ the known results [11] for the 3-state Potts model with external magnetic field were reproduced. For small values of $\tau$ and arbitrary $\kappa$ and $\mu$, we used low temperature expansion techniques up to $\tau^3$. For small $\tau$ we found excellent agreement between the Monte Carlo results and the perturbative series (see Fig. 2). Finally, for all our production and analysis codes two independent programs were written for cross-checks.

_results from the Monte Carlo calculation._—We begin the discussion of our results with the order parameter $\langle P \rangle$, which—as discussed above—is identified with the Polyakov loop of QCD. In Fig. 1, we show the results for
the order parameter approaches 0. The order parameter $\langle P \rangle / V$ as a function of temperature $\tau$ and chemical potential $\mu$ from our 36$^3$, $\kappa = 0.01$ ensembles. Near the rear left corner, no data were computed. $\langle P \rangle / V$ has values close to 1 there. For small $\tau$ and $\mu$ the order parameter approaches 0.

$\langle P \rangle / V$ as a function of $\tau$ and $\mu$ for our 36$^3$ ensembles at $\kappa = 0.01$. For roughly 450 points in the $\tau$-$\mu$ plane, the values of $\langle P \rangle / V$ were evaluated and used for the 3D plot. In the rear left corner and for small $\tau$ and $\mu$, no data were computed. $\langle P \rangle / V$ is expected to be close to 1 in the rear left corner. For small $\tau$ and $\mu$ there is a sizable region where the expectation value $\langle P \rangle / V$ is small and center symmetry is broken only very mildly. Transferring this finding from the effective center theory to QCD implies that for small temperature and density, matter is confined. When $\tau$ or $\mu$ are increased, the system undergoes a change and $\langle P \rangle / V$ reaches values close to 1. For QCD this implies that both temperature and $\mu$ may be used to drive the system into the deconfined phase characterized by a large Polyakov loop.

Next we identify the phase boundary. For that purpose we studied the susceptibility $\chi_P$ and the heat capacity $C$ as a function of $\mu$ at fixed $\tau$ (symbols with horizontal error bars in Figs. 2 and 3) or as a function of $\tau$ at fixed $\mu$ (vertical error bars) and determined the position of the maximum: We fitted $\chi_P$ and $C$ near the maxima with a parabola and obtained the position of the maximum as one of the fit parameters. The corresponding statistical error was computed with the jackknife method. In Fig. 2, we show the positions of the maxima of $\chi_P$ in the $\tau$-$\mu$ plane. We compare the results for 4 values of the inverse mass parameter $\kappa$ and connect data at the same $\kappa$ with a dotted line to guide the eye. The dashed horizontal line at the top marks the value of the critical $\tau$ for the 3-state Potts model, i.e., the situation at $\kappa = 0$. The dashed curves near the bottom of the plot are the results from the perturbative series for small $\tau$ discussed above. The Monte Carlo data nicely approach these curves for $\tau \to 0$.

The curves in Fig. 2 separate the phases with a small order parameter and with $\langle P \rangle / V \sim 1$, i.e., the confined and the deconfined phases. The phase boundaries depend on the inverse mass parameter $\kappa$, and their behavior is as expected for QCD: The intercept of the phase lines with the $\mu$ axis shifts to the left with decreasing quark mass (increasing $\kappa$) because a smaller $\mu$ is sufficient to excite lighter states. Also the intercept with the $\tau$ axis drops with increasing $\kappa$, corresponding to the fact that for quenched QCD (infinite quark mass) the transition temperature is considerably higher than the crossover temperature of QCD with physical quark masses. The mass dependence of the phase boundaries thus is as expected for QCD.

A key problem of the QCD phase diagram is the question about the nature of the various transitions and phases. Unless one goes to very high densities where more exotic phases exist, two principal phases are expected. A phase with conventional matter (confined with broken chiral symmetry) and a plasma phase (deconfined and chirally symmetric). In some parameter regions also a quarkyonic phase with confinement but restored chiral symmetry has been discussed. For the transition lines a standard scenario

FIG. 1 (color online). The order parameter $\langle P \rangle / V$ as a function of temperature $\tau$ and chemical potential $\mu$ from our 36$^3$, $\kappa = 0.01$ ensembles. Near the rear left corner, no data were computed. $\langle P \rangle / V$ has values close to 1 there. For small $\tau$ and $\mu$ the order parameter approaches 0.

FIG. 2 (color online). Phase diagram as obtained from the maxima of the Polyakov loop susceptibility $\chi_P$. We show results at 4 values of the inverse mass parameter $\kappa$. The dashed curves at the bottom are the results of the $\tau$ expansion, and the horizontal line marks the critical value of $\tau$ for the $\kappa = 0$ case.

FIG. 3 (color online). Comparison of phase boundaries obtained from the maxima of susceptibility $\chi_P$ and heat capacity $C$. 

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is that at \( \mu = 0 \) for physical quark masses the finite temperature transition is merely a crossover [12], where different second derivatives of the free energy peak at different temperature values [9,10]. With increasing \( \mu \) the crossover region narrows and terminates at a critical end point. From the end point on a first-order transition line continues (which at some point might hit other transitions to the exotic phases mentioned). Alternative scenarios suggest that either no critical end point appears and crossover type of behavior persists also for large \( \mu \) and low temperature or that even more than one end point might exist. For a glimpse of the current debate, see, e.g., [13].

While we cannot address questions concerning chiral symmetry in the effective center theory, we can analyze the type of transitions that take place at the phase boundaries. To determine the nature of the transitions, we used two techniques. We analyzed histograms for the distribution of the order parameter and the action near the phase boundaries. For a first-order transition they would display a double peak structure near the critical line. We found only single peaks and thus rule out first-order behavior. The second approach is a comparison of the normalized susceptibilities and heat capacities \( \chi_p/V \) and \( C/V \) from lattices with different volumes. For a first-order transition the height of the maxima diverges proportional to \( V \), while for a continuous transition the divergence is modified by a critical exponent. A height which is independent of \( V \) indicates a smooth crossover. Our analysis shows that for the phase boundaries of Figs. 2 and 3 the height is independent of \( V \) for all volumes studied, and that the transitions in the effective center theory are smooth crossover lines.

Once the crossover nature is established, one may ask how wide the transition region is—similar to the finite temperature crossover of QCD at zero density, which is 20–30 MeV wide. In order to get an estimate for the width of the crossover region, in Fig. 3 we compare the positions of the maxima of \( \chi_p \) and of \( C \) for two values of \( \kappa \). The fact that the corresponding lines do not coincide stresses the crossover nature of the transition, and the plot demonstrates that the crossover region is rather wide for most of the parameter values. Only for small \( \kappa \) and \( \mu \) do the lines approach each other, anticipating the first-order behavior known for very small \( \kappa \) and \( \mu \) [4,5].

**Concluding remarks.**—In this Letter, we report on our results for the phase diagram of an effective theory for the center degrees of freedom of QCD. The flux representation solves the complex phase problem, and we develop a worm algorithm for a Monte Carlo calculation in a wide range of temperatures and chemical potential.

The outcome of our analysis is a version of the QCD phase diagram when only the center degrees of freedom are considered. The phase diagram shares many features with the conjectured full QCD phase diagram: The transition to the deconfined phase can be driven by both, temperature or \( \mu \), and the quark mass dependence is as expected for QCD. The phase boundaries between a phase with only very small center symmetry breaking (\( (P)/V \sim 0 \)) and a phase with (\( (P)/V \sim 1 \)) has a shape which is similar to the one conjectured for QCD. For all parameter values studied, the transition is of smooth crossover type. We conclude that center symmetry alone does not provide a mechanism for first-order behavior in the QCD phase diagram.

Various future research directions may be followed: The effective theory can be made more realistic by replacing the \( Z_3 \) spins by continuous SU(3) valued variables (here some work is in progress, and also for this theory the complex phase problem can be solved by a suitable flux representation). Furthermore, it would be desirable to take into account also the fermion nature of the problem—an aspect which is absent in the current effective action. Another interesting direction is of a more technical nature: With our effective theory we have a reference example of a QCD related system where the complex action problem is solved. This reference theory can and should be used to test the reliability and limitations of various techniques for QCD with chemical potential, such as reweighting, series expansions, or complex Langevin methods.

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