We use complex Langevin simulations to explore the QCD phase diagram over a large range of chemical potentials and temperatures. For our simulations we use two flavours of dynamical Wilson fermions with a pion mass of approximately 480 MeV with a spatial volume of $24^3$. Here we report on consistency checks at zero chemical potentials and present our results for the fermion density and the Polyakov loop. We find that at the lowest temperature the fermion density remains zero until $m_N/3$, in line with the expectations from the Silver Blaze phenomenon.
1. **Introduction**

The complex Langevin method (CL), first proposed by Parisi [1] has been successfully adapted and used in various models and approximations of QCD, for instance, the heavy-dense limit of QCD [2]. Early results have been obtained for QCD with very heavy quarks [3, 4]. Here, we aim to study the QCD phase diagram with significantly lighter quarks (\(m_\pi \sim 480\) MeV). Our simulations span a large range of chemical potentials, ranging up to approximately 6500 MeV, and various temperatures, as low as 25 MeV.

2. **Complex Langevin**

The (complex) Langevin method evolves the gauge links of a lattice simulation along a new fictitious time \(\theta\) for a small step size \(\epsilon\). A first order update scheme can be written as

\[
U_{x\mu}(\theta + \epsilon) = \exp \left( i \lambda^a \left( -\epsilon D_{\mu\nu}^a S[U] + \sqrt{\epsilon} \eta_{x,\mu}^a \right) \right) U_{x\mu}(\theta),
\]

where \(\lambda^a\) are the Gell-Mann matrices, \(D_{\mu\nu}^a\) are the derivatives of the action with respect to the gauge link \(U_{x\mu}\) and \(\eta_{x,\mu}^a\) are the random white noise. The complexification is introduced by enlarging the group manifold from SU(3) to SL(3, \(\mathbb{C}\)), i.e. by allowing the coefficients of the generators to become complex numbers

\[
U_{x\mu} = \exp \left( i \lambda^a \left( A_{x\mu}^a + i B_{x\mu}^a \right) \right).
\]

The new time direction created by this procedure is analogue to the Molecular Dynamics time in standard lattice calculations. For sufficient long simulations, an observable \(O\) can be obtained by taking an average over the Langevin time \(\theta\)

\[
\langle O \rangle = \frac{1}{\theta_{\text{max}} - \theta_{\text{therm}}} \sum_{\theta = \theta_{\text{therm}}}^{\theta_{\text{max}}} O[U(\theta)],
\]

after discarding a sufficient amount \(\theta_{\text{therm}}\) to remove thermalisation effects. As we aim to take the limit of \(\epsilon \rightarrow 0\), proper treatment of autocorrelation effects is important. We use the automatic autocorrelation method presented in [5].

3. **Setup**

For our study we use two dynamical flavours of Wilson quarks without a clover term \(c_{SW} = 0\). Our simulation parameters are based on one of the setups used in [6]. The gauge coupling is fixed at \(\beta = 5.8\) throughout all simulations. This implies a fixed lattice spacing of approximately \(a \sim 0.06\) fm [6]. The hopping parameter of \(\kappa = 0.1544\) leads to a pion mass of \(a m_\pi = 0.1458(7)\), which was measured using a HMC simulation at vanishing chemical potential using \(N_t = 128\). Converting the pion mass into physical units, we find \(m_\pi \sim 480\) MeV. The spatial simulation volume is fixed to \(24^3\) and so the product \(m_\pi L = 3.5\) is sufficiently large, so that volume effects are expected to be small. The authors of [6] found a pion mass of \(a m_\pi = 0.1481(11)\) for a larger spatial volume of \(32^3\). The difference is 1.5%, which indicates small volume effects. Different
temperatures are realised by varying the temporal lattice extent. In this study, we use up to 128 points in time, resulting in temperature down to 25 MeV. The chemical potential was scaled up to \( a\mu = 2.0 \). Table 1 shows a summary of the key parameters. To improve the convergence of the complex Langevin simulation, we employ adaptive step size scaling [7], gauge cooling [8] and dynamic stabilisation [9, 10].

### Table 1: The lattice parameters used in this study.

| Parameter        | Value                                      |
|------------------|--------------------------------------------|
| \( \beta = 5.8 \) |                                            |
| \( V = 24^3 \)   |                                            |
| \( \kappa = 0.1544 \) |                                        |
| \( N_t = 4 - 128 \) |                                          |
| \( m_\Lambda L = 3.5 \) |                                    |
| \( T \sim 800 - 25 \text{MeV} \) |                                     |
| \( a\mu = 0.0 - 2.0 \) |                                          |
| \( \mu \sim 0 - 6500 \text{MeV} \) |                                      |

4. Extrapolation at zero chemical potential

As a first consistency check, we compare HMC simulations with complex Langevin simulations at vanishing chemical potential. Even though this is a purely real setup and complex Langevin simulations are not necessary, we allow and start with configurations that are in the SL(3, C) manifold. As complex Langevin suffers from finite step size corrections, we simulate multiple values of the step size \( \epsilon \) and extrapolate to zero. As we employ step size scaling [7], we measure the average step size \( \langle \epsilon \rangle \) and adapt the relevant control parameter such that we achieve different values. Figure 1 shows the results for the plaquette as a function of the average step sizes for two different setups. The first setup (left panel) corresponds to large temperatures of approximately 400 MeV. The second setup (right panel) is at lower temperatures of approximately 100 MeV. The data points are augmented by three different fits to the data: a linear (lin), quadratic (quad) and cubic (cub) extrapolation. In the linear extrapolation, the four right most points have been removed for the fitting procedure. As we use a first-order integration scheme, we expect our step size correction to be of first order. This behaviour is clearly visible. Overall, we find very good agreement between
the HMC results and the extrapolated result from complex Langevin simulations. The agreement is up to the 4th significant figure or less than a permille.

5. Simulation at non-zero chemical potential

To explore the QCD phase diagram, we simulate at different chemical potential and temperatures. Our parameters are chosen such that we span a large region of the QCD phase diagram. To study the phase behaviour, we investigate in particular the fermion density and Polyakov loop. Figure 2 shows the fermion density as a function of the chemical potential. Each data point corresponds to an independent complex Langevin simulation. The left panel shows the entire simulation range, whereas the right panel focuses on smaller chemical potentials. For both, the fermion density is divided by $6V N_t$ for each flavour, i.e. the saturation density. At large $\mu$ we see aforementioned saturation, which is an intrinsic lattice artefact. This occurs when all lattice sites are filled with fermions, so that Pauli blocking does not allow additional fermions to be added to the system. The right-hand side of figure 2 also shows an enlarged view of the fermion density for small chemical potentials and tiny values of the density. Interestingly, the fermion density remains 0 up to $m_N/3$, which is a manifestation of the so-called Silver Blaze phenomenon [11]. The Polyakov loop is shown in the left panel of figure 3. Here the same lattice artefact is visible for large chemical potentials. In the right panel of figure 3 we show the Polyakov loop as function of the temperature for two chemical potentials. We find that the transition to a nonzero Polyakov loop is occurring at lower temperatures for larger chemical potentials. This allow us to quantify the transition(s). Additionally, we also look at the Binder cumulant [12], defined in the following way

$$B = 1 - \frac{\langle O^4 \rangle}{3\langle O^2 \rangle^2}.$$ 

The Binder cumulant for the fermion density is shown in the left panel of figure 4 and for the Polyakov loop in the right panel. The transition to non-zero values for the Binder cumulant occurs at smaller temperatures as the chemical potential increases, as already seen in the right panel of figure 3.
With complex Langevin towards the QCD phase diagram

Benjamin Jäger

Wilson: $24^3$, $\beta = 5.8$, $\kappa = 0.1544$

Polyakov loop

$\mu [\text{MeV}]$

$N_t = 4$

$N_t = 8$

$N_t = 16$

$N_t = 24$

$N_t = 32$

$0.2$

$0.4$

$0.6$

$0.8$

$1$

$1.2$

$1.4$

$1.6$

$1.8$

$0$ 1000 2000 3000 4000 5000 6000

Figure 3: (Left): The Polyakov loop as function of chemical potential $\mu$ for the full range of simulations.

As before the gray lines indicate $m_\pi/2$ and $m_N/3$ (Right): The Polyakov loop as function of the temperature $T$ for two choices of $\mu$.

Wilson: $24^3$, $\beta = 5.8$, $\kappa = 0.1544$

Cumulant - Fermion Density

$\mu [\text{MeV}]$

$N_t = 4$

$N_t = 8$

$N_t = 16$

$N_t = 24$

$N_t = 32$

$0$ 200 400 600 800 1000

Figure 4: The Binder cumulants for the fermion density (left) and the Polyakov loop (right) for different temperatures and chemical potentials.

We have covered a large range of temperatures and chemical potentials, however, the simulations at low temperatures become increasingly challenging, from a numerical point of view. In particular, the number of conjugate gradient steps increases sharply as the chemical potential increases, in particular for low temperature simulations. This behaviour can be seen in the left panel of figure 5. On the right panel, the average unitarity norm is shown, which stays sufficiently small for all simulations. For lower temperatures we can see an increase for larger chemical potentials.

6. Outlook & Conclusion

Here we have reported on our ongoing first-principle study of the QCD phase diagram using complex Langevin simulations. We have performed simulations at various chemical potentials and temperatures using two flavour Wilson quarks with a pion mass of approximately 480 MeV. In particular, we have found that the fermion density stays zero in the region of $m_\pi/2$ to $m_N/3$, i.e. the Silver Blaze phenomenon.

In the future, we plan to focus on improved computations and algorithms, in particular at lower temperatures. The necessary code developments are already underway. Furthermore, we plan to
Figure 5: (Left): The number of conjugate gradient iterations necessary for the computation of the drift force in the complex Langevin process. (Right): The average unitarity norm of the individual simulation, i.e. the distance to the SU(3) manifold.

repeat our study with different volumes, in order to quantify the order of the transition(s) by studying the volume scaling.

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