Dynamics of Chemical Equilibrium of Hadronic Matter Close to $T_c$

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Quick chemical equilibration times of hadrons (specifically, $p\bar{p}$, $K\bar{K}$, $\Lambda\bar{\Lambda}$, and $\Omega\bar{\Omega}$ pairs) within a hadron gas are explained dynamically using Hagedorn states, which drive particles into equilibrium close to the critical temperature. Within this scheme, we use master equations and derive various analytical estimates for the chemical equilibration times. We compare our model to recent lattice results and find that for both $T_c = 176$ MeV and $T_c = 196$ MeV, the hadrons can reach chemical equilibrium almost immediately, well before the chemical freeze-out temperatures found in thermal fits for a hadron gas without Hagedorn states. Furthermore the ratios $p/\pi$, $K/\pi$, $\Lambda/\pi$, and $\Omega/\pi$ match experimental values well in our dynamical scenario.

I. INTRODUCTION

(Anti-)strangeness enhancement was first observed at CERN-SPS energies by comparing anti-hyperons, multistrange baryons, and kaons to $pp$-data. It was considered a signature for quark gluon plasma (QGP) because, using binary strangeness production and exchange reactions, chemical equilibrium could not be reached within a standard hadron gas phase, i.e., the chemical equilibration time was on the order of $\tau \sim 100$ fm/$c$ whereas the lifetime of a fireball in the hadronic stages is only $\tau \approx 4 - 7$ fm/$c$ [1]. It was then proposed that there exists a strong hint for QGP at SPS because strange quarks can be produced more abundantly by gluon fusion, which would account for strangeness enhancement following hadronization and rescattering of strange quarks. Later, however, multi-mesonic reactions were used to explain secondary production of $\bar{p}$ and anti-hyperons [2, 3]. At SPS they give a chemical equilibration time $\tau_\gamma \approx 1 - 3$ fm/$c$ using an annihilation cross section of $\sigma_{\rho\bar{\rho}} \approx \sigma_{p\bar{p}} \approx 50$ mb and a baryon density of $\rho_B \approx \rho_0$ to $2\rho_0$, which is typical for evolving strongly interacting matter at SPS before chemical freeze-out. Therefore, the time scale is short enough to account for chemical equilibration within a cooling hadronic fireball at SPS.

A problem arises when the same multi-mesonic reactions were employed in the hadron gas phase at RHIC temperatures where experiments again show that the particle abundances reach chemical equilibrium close to the phase transition [4]. At RHIC at $T = 170$ MeV, where $\sigma \approx 30$ mb and $\rho_B^q \approx \rho_{\bar{q}}^q \approx 0.04$ fm$^{-3}$, the equilibrium rate for (anti-)baryon production is $\tau \approx 10$ fm/$c$. Moreover, $\tau \approx 10$ fm/$c$ was also obtained in Ref. [5] using a fluctuation-dissipation theorem. From hadron cascades a significant deviation was found from the chemically saturated strange (anti-)baryons yields in the 5% most central Au-Au collisions [6]. These discrepancies suggest that hadrons are “born” into equilibrium, i.e., the system is already in a chemically frozen out state at the end of the phase transition [7, 8]. In order to circumvent such long time scales it was suggested that near $T_c$ there exists an extra large particle density overpopulated with pions and kaons, which drive the baryons/anti-baryons into equilibrium [9]. But it is not clear how this overpopulation should appear, and how the subsequent population of (anti-)baryons would follow. Moreover, the overpopulated (anti-)baryons do not later disappear [10]. Therefore, it was conjectured that Hagedorn resonances (heavy resonances near $T_c$ with an exponential mass spectrum) could account for the extra (anti-)baryons [10, 11, 12].

Hadrons can develop according to

$$n\pi \leftrightarrow HS \leftrightarrow n'\pi + XX$$  \hspace{1cm} (1)

where $XX$ can be substituted with $p\bar{p}$, $K\bar{K}$, $\Lambda\bar{\Lambda}$, or $\Omega\bar{\Omega}$. Eq. (1) provides an efficient method for producing of $XX$ pairs because of the large decay widths of the Hagedorn states. In Eq. (1), $n$ is the number of pions for the decay $n\pi \leftrightarrow HS$ and $n'$ is the number of pions that a Hagedorn state will decay into when a $XX$ is present. Since Hagedorn resonances are highly unstable, the phase space for multi-particle decays drastically increases when the mass increases. Therefore, the resonances catalyze rapid equilibration of $XX$ near $T_c$ and die out moderately below $T_c$ [11].

Unlike in pure glue $SU(3)$ gauge theory where the Polyakov loop is the order parameter for the deconfinement transition (which is weakly first-order), the rapid crossover seen on lattice calculations involving dynamical fermions indicates that there is not a well defined order parameter that can distinguish the confined phase from the deconfined phase. Because of this it is natural to look for a hadronic mechanism for quick chemical equilibration near the phase transition. One such possibility could be the inclusion of Hagedorn states. Recently, Hagedorn states have been shown to contribute to the physical description of a hadron gas close to $T_c$. The inclusion of Hagedorn states leads to a low $\eta/s$ in the hadron gas phase [13], which nears the string theory bound $\eta/s = 1/(4\pi)$ [14]. Calculations of the trace anomaly including Hagedorn states also fits recent lattice results well and correctly describe the minimum of
the speed of sound squared, $c_s^2$, near the phase transition found on the lattice \[13\]. Estimates for the bulk viscosity including Hagedorn states in the hadron gas phase indicate that the bulk viscosity, $\zeta/s$, increases near $T_c$, which agrees with the general analysis done in \[13\]. Furthermore, it has been shown \[10\] that Hagedorn states provide a better fit within a thermal model to the hadron yield particle ratios. Additionally, Hagedorn states provide a mechanism to relate $T_c$ and $T_{\text{chem}}$, which then leads to the suggestion that a lower critical temperature could possibly be preferred, according to the thermal fits \[10\].

Previously, in Ref. \[11\] we presented analytical results, which we will derive in detail here. Moreover, we saw that both the baryons and kaons equilibrated quickly within an expanding fireball. The initial saturation of pions, Hagedorn states, baryons, and kaons played no significant role in the ratios such as $K/\pi$ and $(B + B)/\pi$.

Here we consider the effects of various initial conditions on the chemical freeze-out temperature and we find that while they play a small role on the total particle number, they still reproduce fast chemical equilibration times. Additionally, we assume lattice values of the critical temperatures ($T_c = 176$ MeV \[17\] and $T_c = 196$ MeV \[18\] \[19\]) and find that chemical equilibrium abundances are still reached close to the temperature given by thermal fits ($T \approx 160$ MeV).

This paper is structured in the following manner. In Section \[II\] we discuss the details of our statistical model that calculates the chemical equilibrium values of the Hagedorn states and other hadrons. Furthermore in this section, fits are shown to thermodynamical properties calculated in lattice QCD, which are used to determine the mass spectrum of the Hagedorn states and the rate equations are discussed in detail. In Section \[III\] we are able to extract the chemical equilibration time of an $XX$ pair when the pions and Hagedorn states are held constant. In Section \[IV\] we derive an analytical result of the rate equations when we consider only the decay $HS \leftrightarrow n\pi$. We then discuss the case of an expanding fireball and the results for the various $XX$ pair production in Section \[V\]. The production of $\Omega$ particles will also be considered in Section \[VI\]. We summarized and discussed our results in Section \[VII\]. In Appendix \[A\] we present some analytical and numerical results for the various equilibration stages in the hadron and Hagedorn states gas mixture.

\[ II. \; MODEL \]

Hagedorn resonances have an exponentially growing mass spectrum \[20\]. Their large masses open up the phase space for multi-particle decays. Recent analysis involving Hagedorn states is given in \[21\]. Moreover, thoughts on observing Hagedorn states in experiments are given in \[22\] and their usage as a thermostat in \[23\]. Hagedorn states can also explain the phase transition above the critical temperature and, depending on the intrinsic parameters, the order of the phase transition \[24\]. For the following discussion, the overall density of Hagedorn states in our extended Hagedorn gas model are straightforwardly described by,

$$\rho = \int_{M_0}^{M} \frac{A}{(m^2 + m_c^2)^{2\frac{B}{4}}} e^{\frac{m}{T_H}} dm.$$  \[2\]

where $M_0 = 2$ GeV and $m_c^2 = 0.5$ GeV. We note that in this work we consider only mesonic Hagedorn states with no net strangeness. The exponential in Eq. \[2\] arises from Hagedorn’s original idea that there is an exponentially growing mass spectrum. Thus, as $T_H$ is approached, Hagedorn states become increasingly more relevant and heavier resonances “appear”. The factor in front of the exponential has various forms \[21\] \[23\]. While the choice in this factor can vary, it was found in \[21\] that the present form gives lower values of $T_H$, which more closely match the predicted lattice critical temperature \[17\] \[18\] \[19\].

Returning to Eq. \[2\], its parameters ($A$, $M$, and $T_H$) are dependent on the critical temperature. We assume that $T_H = T_c$, and then we consider the two different different lattice results for $T_c$: $T_c = 196$ MeV \[18\] \[19\], which uses an almost physical pion mass, and $T_c = 176$ MeV \[17\]. Furthermore, we need to take into account the repulsive interactions and, therefore, we use the following volume corrections (as also seen in \[13\] \[25\] \[26\]):

$$T = \frac{T^*}{1 - \frac{\mu_B}{\mu_B^*}} \frac{4B}{4B}$$

$$\mu_B = \frac{\mu_B^*}{1 - \frac{\mu_B}{\mu_B^*}} \frac{4B}{4B}$$

$$p_{xx} = \frac{p_{xx}^*}{1 + \frac{\varepsilon_{xx}}{\varepsilon_{xx}^*}} \frac{4B}{4B}$$

$$\varepsilon_{xx} = \frac{\varepsilon_{xx}^*}{1 + \frac{n_{xx}}{n_{xx}^*}} \frac{4B}{4B}$$

$$n_{xx} = \frac{n_{xx}^*}{1 + \frac{s_{xx}}{s_{xx}^*}} \frac{4B}{4B}$$

$$s_{xx} = \frac{s_{xx}^*}{1 + \frac{\varepsilon_{xx}}{\varepsilon_{xx}^*}} \frac{4B}{4B},$$  \[3\]

which ensure that the our model is thermodynamically consistent. Note that $B$ is a free parameter that is based upon the idea of the MIT bag constant.

In order to find the maximum Hagedorn state mass $M$ and the “degeneracy” $A$, we fit our model to the thermodynamic properties of the lattice. In the RBC-Bielefeld collaboration the thermodynamical properties are derived from the quantity $\varepsilon - 3p$, the so called interaction measure, which is what we fit in order to obtain the parameters for the Hagedorn states. Thus, we obtain $T_H = 196$ MeV, $A = 0.5 GeV^{3/2}$, $M = 12$ GeV, and
$B = (340 GeV)^4$. The fit for the trace anomaly $\Theta/T^4$ is shown in Fig. 1. We also show the fit for the entropy density in Fig. 2. Both fits are within the error of lattice and mimic the behavior of the lattice results. As discussed in [13], a hadron resonance gas model with Hagedorn states uniquely fits the lattice data whereas a hadron resonance gas without Hagedorn states completely misses the behavior.

BMW calculates the thermodynamical properties separately and, therefore, we fit only the energy density as shown in Fig. 3. From that we obtain $T_H = 176$ MeV, $A = 0.1 GeV^{3/2}$, $M = 12$ GeV, and $B = (300 GeV)^4$. We also show a comparison to the entropy density in Fig. 4. Our results with the inclusion of Hagedorn states are able to match lattice data near the critical temperature but do not match as well at lower temperatures in Fig. 1 and Fig. 2.

Our idea is that these very massive Hagedorn states exist, as pictured in Fig. 5, and are so large that they decay almost immediately into multiple pions and $X \bar{X}$ pairs. While it can be argued that Hagedorn states are more likely to decay into a pair of particles: a lighter Hagedorn state and another particle, these reactions are so quick that we can consider the end results, which would be multiple particles (mostly pions). That being said, it would be possible to put Hagedorn states into a transport approach such as UrQMD [27] using binary reactions with possible cross-sections as described in [28]. We leave this as a challenge for the future.

Moreover, we need to consider the back reactions of multiple particles combining to form a Hagedorn state in order to preserve detailed balance. Rate equations provide us with a perfect tool for this because there is a loss and gain term that describe both the forward and back reactions. Moreover, the state of chemical equilibrium is a fixed point of the rate equations. The rate equations for the Hagedorn resonances $N_i$, pions $N_\pi$, and the $X \bar{X}$ pair $N_{X \bar{X}}$, respectively, are given by

$$\dot{N}_i = \Gamma_{i,\pi} \left[ N_i^{eq} \sum_n B_{i,n} \left( \frac{N_\pi}{N_\pi^{eq}} \right)^n - N_i \right]$$
The decay widths for the $i^{th}$ resonance are $\Gamma_{i,\pi}$ and $\Gamma_{i,XX}$, the branching ratio is $B_{i,n}$ (see below), and the average number of pions that each resonance will decay into is $\langle n_i \rangle$. The equilibrium values $N^{eq}_i$ are both temperature and chemical potential dependent. However, here we set $\mu_0 = 0$. Eq. (3) can also be rewritten in terms of fugacities ($\lambda_i$, $\lambda_\pi$, and $\lambda_{XX}$), which are found by dividing each total number by its respective equilibrium value, for example, $\lambda_i = \frac{N_i}{N^{eq}_i}$ (as seen for the baryon anti-baryon pairs in [11]). Additionally, a discrete spectrum of Hagedorn states is considered, which is separated into mass bins of 100 MeV. Each bin is described by its own rate equation.

The branching ratios, $B_{i,n}$, are the probability that the $i^{th}$ Hagedorn state will decay into $n$ pions. Since we are dealing with probabilities, then $\sum_{n} B_{i,n} = 1$ must always hold. In order to include a distribution for our branching ratios we assume that they follow a Gaussian distribution for the reaction $HS \leftrightarrow n\pi$

$$B_{i,n} \approx \frac{1}{\sigma \sqrt{2\pi}} e^{\frac{-(n-\langle n_i \rangle)^2}{2\sigma^2}},$$

which has its peak centered at $\langle n_i \rangle$ and the width of the distribution is $\sigma^2$. Assuming a statistical, micro-canonical branching for the decay of Hagedorn states, we can take a linear fit to the average number of pions in

$$\langle n_{i,x} \rangle = \frac{2.7}{1.9} (0.3 + 0.4 m_i) \approx 2 - 7.$$ (6)

where $m_i$ is in GeV. In this paper we do not consider a distribution but rather only the average number of pions when a $X\bar{X}$ pair is present. We assume that $\langle n_{i,x} \rangle = \langle n_{i,p} \rangle = \langle n_{i,k} \rangle = \langle n_{i,\Lambda} \rangle = \langle n_{i,\Omega} \rangle$ for when a kaon anti-kaon pair, $\Lambda \bar{\Lambda}$, or $\Omega \bar{\Omega}$ pair is present. Ideally, $\langle n_{i,k} \rangle$, $\langle n_{i,\Lambda} \rangle$, and $\langle n_{i,\Omega} \rangle$ should be derived separately and will be done in a future paper using a canonical model [31].

We used a linear fit for the total decay width similar to that used in Ref. [32]. The total decay width

$$\Gamma_i = 0.15 m_i - 0.0584$$ (7)

($\Gamma_i$ and $m_i$ in terms of GeV), which ranges from $\Gamma_i = 250$ MeV to 1800 MeV, is a linear fit extrapolated from the data in Ref. [33]. However, in Eq. (4) the total decay width is separated into two parts: one for the reactions $HS \leftrightarrow n\pi$, $\Gamma_{i,\pi}$, and one for the reaction in Eq. (1), $\Gamma_{i,XX}$, whereby $\Gamma_i = \Gamma_{i,\pi} + \Gamma_{i,XX}$. Then relative decay width $\Gamma_{i,XX}$ is the average number of $XX$ in the system $\langle X \rangle$ multiplied by the total decay width $\Gamma_i$. Essentially, a fraction of the decay of the $i^{th}$ Hagedorn state goes into $XX$ (set by the number of $XX$ the $i^{th}$ Hagedorn state on average decays into) and the remainder goes into pions.

We find $\langle p \rangle$ by linearly fitting the proton in Fig. 2 in Ref. [10] so that

$$p = 0.058 m_i - 0.10$$ (8)

Fig. 1 in Ref. 10 (multiplying $\pi^+$ by three to include all pions) to find $\langle n_\pi \rangle$ such that $\langle n_\pi \rangle = 0.9 + 1.2 \frac{n_\pi}{m_\pi}$ is the average pion number that each Hagedorn state decays into. Within the microcanonical model a Hagedorn state is defined by its mass and corresponding volume where the volume is taken as $V = m_i / \epsilon$. The mean energy density of a Hagedorn state is $\epsilon$ (taken as $\epsilon = 0.5 \frac{m_i}{m_\pi}$). Further discussions regarding this can be found in [10, 29].
where \( m_i \) is in GeV and \( \langle p \rangle \approx 0.01 - 0.6 \). Thus, \( \Gamma_{i,\rho\bar{\rho}} \) is between 3 and 1000 MeV. Clearly, \( \Gamma_{i,\pi} = \Gamma_i - \langle p \rangle \Gamma_{i,\pi} \). Analogously for the kaons, the decay width is \( \Gamma_{i,K\bar{K}} = \langle K \rangle \Gamma_i \) where

\[
K^+ = 0.075 \, m_i + 0.047
\]  

where \( m_i \) is in GeV, which is also taken from Fig. 2 in Ref. [10]. We find that \( \langle K \rangle = 0.2 \) to 0.95 [10] [29]. Thus, \( \Gamma_{i,K\bar{K}} \) is between 50 and 1700 MeV.

For \( \Lambda \) we use a canonical model assuming that the baryon number \( B = 0 \), the strangeness \( S = 0 \), and the electrical charge \( Q = 0 \) in order to calculate the average lambda number. The results of this are shown in Fig. 6. This corresponds to a decay width of \( \Gamma_{i,\Lambda\bar{\Lambda}} = 3 - 250 \) MeV.

Furthermore, the average number of \( \Omega \)'s is also shown in Fig. 6 from our canonical model again assuming that the baryon number \( B = 0 \), the strangeness \( S = 0 \), and the electrical charge \( Q = 0 \). In Fig. 6 we multiple \( (\Omega) \) in order to better view the results. The resulting decay width is \( \Gamma_{i,\Omega\bar{\Omega}} = 0.01 - 4 \) MeV.

The equilibrium values are found using a statistical model [34], which includes “all” the pions from resonances from the PDG [33]. Feeding is also considered for the protons, kaons, and lambdas. Additionally, throughout this paper our initial conditions are the various fugacities at \( t_0 \) (at the point of the phase transition into the hadron gas phase)

\[
\alpha \equiv \lambda_\pi(t_0), \quad \beta_i \equiv \lambda_i(t_0), \quad \text{and} \quad \phi \equiv \lambda_{XX}(t_0),
\]

which are chosen by holding the contribution to the total entropy from the Hagedorn states and pions constant i.e.

\[
s_{\text{Had}}(T_0, \alpha) V(t_0) + s_{\text{HS}}(T_0, \beta_i) V(t_0) = s_{\text{Had}+\text{HS}}(T_0) V(t_0) = \text{const}.
\]  

and the corresponding initial condition configurations we choose later can be seen in Tab. [11] \( s_{\text{Had}}(T_0, \alpha) \) is the entropy density at the initial temperature, i.e., the critical temperature multiplied by our choice in \( \alpha \). Because the hadron resonance is dominated by pions we can assume that \( \alpha \) represents the initial fraction of pions in equilibrium. \( s_{\text{HS}}(T_0, \beta_i) \) represents the entropy contribution from the Hagedorn states at \( T_c \) multiplied by the initial fraction of Hagedorn states in equilibrium. We hold \( \alpha \) constant and then find the appropriate \( \beta_i \). The volume expansion, \( V(t) \) is discussed in detail following section entitled ‘Expanding Fireball’.

III. CHEMICAL EQUILIBRATION TIME ESTIMATE

As a starting point of our analysis, we first estimate the chemical equilibration time of the \( XX \) by looking at the fugacity of the \( XX \) rate equation, i.e., Eq. (4) can be rewritten in terms of \( \lambda \) as shown for \( BB \) in Eq. (3) in [11], when both the pions and Hagedorn states are held constant. The \( XX \) rate equation then becomes

\[
\dot{\lambda}_{XX} = \sum_i \Gamma_{i,XX} \frac{N^q_{i,XX}}{N^q_{i,XX}} \left( \beta_i \right)^{\alpha(x,s)} \lambda_{XX}^2,
\]

which we can integrate

\[
\lambda_{XX} = \zeta \left[ \left( \frac{\phi + \zeta}{\phi - \zeta} \right) e^{\frac{2\theta}{\phi}} + 1 \right] e^{\frac{2\phi}{\phi - \zeta}} - 1
\]

where

\[
\tau_{XX} = \frac{N^q_{XX}}{\sqrt{\sum_i \Gamma_{i,XX} N^q_{i,XX} \beta_i \sqrt{\sum_i \Gamma_{i,XX} N^q_{i,XX} \alpha(x,s)}}},
\]

\[
\zeta = \sqrt{\sum_i \Gamma_{i,XX} \frac{N^q_{i,XX} \beta_i}{\alpha(x,s)}}, \quad \text{and} \quad \lambda_{XX}(0) \equiv \phi.
\]

Substituting in \( \alpha = 1 \) and \( \beta_i = 1 \) when the pions and Hagedorn states are in chemical equilibrium, we rederive Eq. (7) in Ref. [11]

\[
\tau_{XX} = \frac{N^q_{XX}}{\sum_i \Gamma_{i,XX} N^q_{i}},
\]

which is shown in Fig. 7. From Eq. (15) we see that the time scale has an indirect dependence on the decay width. Since the decay width has a linear dependence on the mass, the time scale decreases when more Hagedorn states are included. However, \( N^q_{i} \) also decreases.
protons, kaons, and lambdas when $\alpha = 1$ and $\beta_i = 1$ where $T_H = 176 \text{ MeV}$ (top) and $T_H = 196 \text{ MeV}$ (top). The gray band is the range of chemical equilibrium times for the Hagedorn states (see Tab. I).

Even as the temperature is lowered we still see quick chemical equilibrium times. For the $p\bar{p}$ and $\Lambda\bar{\Lambda}$ pairs at $T = 0.9 T_c$ the chemical equilibrium time is still about $t < 1/\text{fm}/c$. The $K\bar{K}$ pairs do have a slower chemical equilibrium time due to their larger chemical equilibrium abundances, which is directly related to the chemical equilibrium time through Eq. (4). This again represents the main idea, which is the importance of potential Hagedorn states in understanding fast chemical equilibrium of hadrons close and below $T_c$. The Hagedorn states increase dramatically in number close to the critical temperature and, thus, by its subsequent decay and re-population they will quickly produce the various hadronic particles.

The equilibration of $XX$ pairs then shown in Fig. 8 and Fig. 9 where the analytical result in Eq. (13) matches the numerical result exactly. From Fig. 8 and Fig. 9 it can be seen that all $XX$ pairs equilibrate quickly close to the critical temperature $t < 1/\text{fm}/c$. Clearly, though, as the temperature decreases the chemical equilibrium time lengthens. However, at $T_H = 196 \text{ MeV}$ chemical equilibrium is still reached quickly, $t < 1/\text{fm}/c$.

IV. ANALYTICAL RESULTS: PIONS AND HAGEDORN RESONANCES

While the chemical equilibrium time derived in the previous section is a good estimate, it can only be strictly applied when the pions and Hagedorn states are assumed to stay in chemical equilibrium at a constant temperature (Fig. 8 and Fig. 9). Otherwise, non-linear effects that appear when the pions and Hagedorn states are allowed to equilibrate appear.

To understand the dynamics in more detail, we consider the simplified case when the Hagedorn resonances decay only into pions $HS \leftrightarrow n\pi$, which gives

$$\dot{N}_i = \Gamma_i \left[ N_i^{eq} \sum_{n=2} B_{i,n} \left( \frac{N_i}{N_{\pi}} \right)^n - N_i \right] $$

$$\dot{N}_\pi = \sum_i \Gamma_i \left[ N_i \langle n_i \rangle - N_i^{eq} \sum_{n=2} B_{i,n} \left( \frac{N_\pi}{N_{\pi}} \right)^n \right]$$

Assuming that the pions and the Hagedorn states described in Eq. (16) are then allowed to equilibrate near $T_c$ in a static system, we are able to derive analytical solutions, the derivation of which is shown in detail in Appendix X. For the analytical solutions we divide the chemical equilibration into three stages, the chemical equilibrium times of which are shown in Tab. I. The first stage (described by $\tau^{eq}_\pi$ in Tab. I) of the evolution is dominated by the chemical equilibration of the pions when the pions are still far away from their chemical equilibrium values. After the pions are close to chemical equilibrium, new dynamics take over, which are described by $\tau_\pi$ in Tab. I and Fig. 10.

In both Stage 1 and 2 the equilibrium of the Hagedorn states is set by the dynamics of the pions. Finally,
in Stage 3 the pions, which are already almost in chemical equilibrium, reach a quasi-equilibrium state with the Hagedorn states. Quasi-equilibrium is reached when at least one species of Hagedorn states has succeeded its chemical equilibrium time scale determined from the inverse of its decay width, i.e., $\tau_i = 1/\Gamma_i$. Since the heaviest Hagedorn states have the shortest $\tau_i$’s, then quasi-equilibrium is reached when $\tau_i$ of the heaviest Hagedorn state is surpassed. During this stage non-linear affects take over and, thus, a longer time scale, $\tau_{\pi^{QE}}$, is seen. While this time scale may appear long, both the pions and Hagedorn states are so close to chemical equilibrium that they are within roughly 10% (depending on the initial conditions) or less of their chemical equilibrium values before quasi-equilibrium is even reached. The detailed calculations are shown in the Appendix.

Therefore, the most important chemical equilibration time is then that from the pions in Stage 1, i.e., $\tau_0^{\pi}$. The time scale from Stage 2 is so short that it is not of much importance. Additionally, by the time that Stage 3 is
TABLE I: Chemical equilibration times from analytical estimates where QE is quasi-equilibrium at 95% of each respective \( T_H \).

| \( \lambda_\pi \approx 0 \) | \( \lambda_\pi \approx 1 \) | \( \lambda_\pi \approx 0 \) | \( \lambda_\pi \approx 1 \) |
|---|---|---|---|
| \( \tau_0 \equiv \frac{N^\pi}{\sum_i \Gamma_i N_i^\pi (\sigma_i^\pi)} \) | \( \tau_\pi \equiv \frac{N^\pi}{\sum_i \Gamma_i N_i^\pi (\sigma_i^\pi)} \) | \( \tau_{QE} \equiv \frac{N^\pi}{\sum_i \Gamma_i N_i^\pi (\sigma_i^\pi)} + \frac{\sum_{QE} N_{\pi QE} (\sigma_i^\pi)}{\sum_i \Gamma_i N_i^\pi (\sigma_i^\pi)} \) | \( \tau_{QE} \equiv \frac{N^\pi}{\sum_i \Gamma_i N_i^\pi (\sigma_i^\pi)} + \frac{\sum_{QE} N_{\pi QE} (\sigma_i^\pi)}{\sum_i \Gamma_i N_i^\pi (\sigma_i^\pi)} \) |
| 0.5 fm/c | 0.01 fm/c | 1.7 fm/c | 1.6 fm/c |

FIG. 10: Comparison of the chemical equilibration times of the pions to the total chemical equilibration time for \( T_H = 176 \) MeV (top) and \( T_H = 196 \) MeV (bottom).

In order to include the cooling of the fireball we need to find a relationship between the temperature and the time, i.e., \( T(t) \). To do this we apply a Bjorken expansion for which the total entropy is held constant

\[
\text{const.} = s(T)V(t) \sim \frac{S_\pi}{N_\pi} \int \frac{dN_\pi}{dy} dy. \quad (17)
\]

where \( s(T) \) is the entropy density of the hadron gas with volume corrections.

The total number of pions in the 5% most central collisions, \( \frac{dN_\pi}{dy} \), can be found from experimental results in [35]. There they found the phase-space yields for the pions \( \pi^+ (292.0) \) and \( \pi^- (290.9) \) using a Gaussian fit for yields as a function of the rapidity \( \frac{dN_\pi}{dy} \) where we used the rapidity range \( y = \pm 0.5 \). We then assumed that the number of \( \pi^0 \)'s were also in that same range and took the average of the two to find 291.5. Thus, our total pion number is \( \sum_i N_{\pi i} = \int_{-0.5}^{0.5} \frac{dN_\pi}{dy} dy = 874 \). While for a gas of non-interacting Bose gas of massless pions \( S_\pi/N_\pi = 3.6 \), we do have a mass for our pions, so we must adjust \( S_\pi/N_\pi \) accordingly. In [36] it was shown that when the pions have a mass the ratio changes and, therefore, the entropy per pion is close to \( S_\pi/N_\pi \approx 5.5 \). The actual \( S_\pi/N_\pi \) in our model is shown in Fig. 11 where \( S_\pi/N_\pi \approx 6 \), which is only slightly higher.

The effective volume at mid-rapidity can be parametrized as a function of time. We do this by using a Bjorken expansion and including accelerating radial flow. The volume term is then

\[
V(t) = \pi ct \left( r_0 + v_0 (t - t_0) + \frac{1}{2} a_0 (t - t_0)^2 \right)^2 \quad (18)
\]

where the initial radius is \( r_0(t_0) = 7.1 \) fm for \( T_H = 196 \) and the corresponding \( t_0(196) \approx 2 \) fm/c. For \( T_H = 176 \) we allow for a longer expansion before the hadron gas phase is reached and, thus, calculate the appropriate \( t_0(176) \) from the expansion starting at \( T_H = 196 \), which is \( t_0(176) \approx 4 \) fm/c (there is a slightly variation dependent on the choice of \( v_0 \) and \( a_0 \)). The \( T(t) \) relation is shown in Fig. 12 which has almost no effect on the results as

\[
V(\tau) = \pi ct \left( r_0 + v_0 (t - t_0) + \frac{1}{2} a_0 (t - t_0)^2 \right)^2 \quad (18)
\]
seen later on in Fig. 16 and Fig. 17. Therefore, we choose $v_0 = 0.5$ and $a_0 = 0.025$ for the remainder of this paper. The relation depicted allows to translate the later shown figures labeled by the effective global temperature of the evolving system directly into the evolving system time.

Because the volume expansion depends on the entropy according to Eq. (17) and the Hagedorn resonances contribute strongly to the entropy only close to the critical temperature (see Fig. 13), the equilibrium values actually decrease with increasing temperature close to $T_c$ for the hadrons as seen in Fig. 14 and Fig. 15. One can clearly see from Fig. 13 that the Hagedorn states contribute strongly close to $T_c$ down to about 80% of $T_c$.

Therefore, one has to include the potential contribution of the Hagedorn resonances to the pions as in the case of standard hadronic resonances, e.g. a $\rho$-meson decays dominantly into two pions and, thus, accounts for them by a factor two. This is similar to what was done in Appendix A in Eq. (A7). Including the Hagedorn state contribution, we arrive at our effective number of pions

$$\tilde{N}_{n, X\bar{X}} = N_n + \sum_i N_i \left[ (1 - \langle X_i \rangle) \langle n_i \rangle + \langle X_i \rangle \langle n_{i, x} \rangle \right]$$

(19)

which are shown in Fig. 14. In Fig. 14 we see that after the inclusion of the effective pion numbers that the number of pions only decreases with decreasing temperature. Furthermore, in Fig. 14 the total number of Hagedorn states, $\sum_i N_{i, eq}$ is also shown. While there are by far fewer Hagedorn states present than pions, we see that they are important because of their large contribution to the entropy density as shown in Fig. 13. The reason that the effective number of pions increase close to $T_c$ is due to the large number of pions that the heavy Hagedorn states decay into. If $\langle n_i \rangle$ was smaller or no longer linear than it could be possible that the effective number of pions would remain constant.

Moreover, it is useful to consider the effective number of $XX$ pairs

$$\tilde{N}_{XX} = N_{XX} + \sum_i N_i \langle X_i \rangle$$

(20)

because Hagedorn states also contribute strongly to the $XX$ pairs close to $T_c$ as seen in Fig. 15. Again we see that only the effective number of $XX$ pairs have consistent decreasing behaviour with decreasing temperature whereas without the Hagedorn state contributions we see a decrease close to $T_c$.

Along with the expansion we also must solve these rate equations, Eq. (1), numerically [38]. We start with various initial conditions, as mentioned previously, that are described by $\alpha$, $\beta_i$, and $\phi$ (see table II). The initial temperature is the respective critical temperature and we end the expansion at $T = 110$ MeV, a global kinetic freezeout temperature.

For the remainder of this paper we include only results for an expanding fireball, which are solved numerically.
As an initial test we hold both the pions and Hagedorn states in chemical equilibrium and allow just $X\bar{X}$ to equilibrate as seen in Fig. 16 and Fig. 17. The black solid line in each graph is the chemical equilibrium abundances and the colored lines are the dynamical calculations for various expansions that follow the $T(t)$ shown in Fig. 12. We see that regardless of our volume expansion they all quickly approach equilibrium. In Fig. 18 and Fig. 19 we show the evolution of both the $pp$ pairs and the pions for the reaction $n\pi \leftrightarrow HS \leftrightarrow n\pi + XX$. Note that in all the following figures the effective numbers are shown so that the contribution of the Hagedorn states is included.

One can see that the chemical equilibration time does depend slightly on our choice of $\beta_i$, i.e., a larger $\beta_i$ means a quicker chemical equilibration time. For instance, if the Hagedorn states were overpopulated coming out of the QGP phase than chemical equilibrium times would be slightly shorter. However, even when the Hagedorn resonances start underpopulated the $pp$ pairs are able to reach chemical equilibrium immediately. Additionally, when the $pp$ pairs start at about half their chemical equilibrium values, it only helps the $pp$ pairs to reach equilibrium at a slightly higher temperature (on the order of a couple of MeV). Additionally, we see a greater dependence on $\beta_i$ for $T_H = 176$ MeV than for $T_H = 176$ MeV. Throughout the evolution we see from the pions that they remain roughly in chemical equilibrium. Thus, our initial analytical approximation appears reasonable.

In Fig. 20 the ratio of protons’s to $\pi$’s is shown. We also compare our results to that of experimental data. We see that for $T_H = 176$ MeV that our results enter

| $\alpha = \lambda_\pi(t_0)$ | $\beta_i = \lambda_i(t_0)$ | $\phi = \lambda_{XX}(t_0)$ |
|-------------------------|------------------------|----------------------|
| $IC_1$                  | 1                      | 1                    | 0 |
| $IC_2$                  | 1                      | 1                    | 0.5 |
| $IC_3$                  | 1.1                    | 0.5                  | 0 |
| $IC_4$                  | 0.95                   | 1.2                  | 0 |

TABLE II: Initial condition configurations, recalling Eq. (10)
FIG. 16: Results for the \( p\bar{p}, K\bar{K}, \) and \( \Lambda\bar{\Lambda} \) when the pions and Hagedorn resonances are held in equilibrium for \( T_H = 176 \) MeV. The results are slightly different. In this case, the ratios match the experimental data early on at around \( T = 190 \) MeV. However, they become briefly overpopulated around \( T = 160 - 170 \) MeV but then quickly return to the experimental values, except for the case when we have the initial conditions such that the pions are overpopulated. This could imply that there are a few too many Hagedorn states and a fit for the Hagedorn states with a lower \( A \) (degeneracy of the Hagedorn states) may produce better results.

As with the protons, the total number of kaons are also slightly dependent on our chosen initial conditions, more specifically, our choice in \( \beta_i \). In Fig. 21 and Fig. 22 the temperature of the evolving system after the phase transition at which chemical equilibrium among standard hadrons is basically reached and maintained is between \( T = 160 - 170 \) for \( T_H = 176 \) MeV and they have already reached chemical equilibrium by \( T = 170 \) for...
FIG. 18: Results for the p’s and pions with various initial conditions for $T_H = 176$ MeV.

$T_H = 196$ MeV, below which the Hagedorn states basically die out. The one exception is when the Hagedorn states begin underpopulated i.e. that $\beta_i < 1$. In this case, the kaon pairs take longer to reach chemical equilibrium. However, when we look at $K/\pi$ in Fig. 23 lower $\beta_i$ actually fits the data better.

Moreover, the pions again remain roughly at chemical equilibrium throughout the expansion as seen in Fig. 21 and Fig. 22. While the pion graphs look roughly similar in Figs. 18-22 they are not. The difference is how the pions are affected in the presence of a $p\bar{p}$ pair compared to a decay that includes a kaon anti-kaon pair.

In Fig. 23 the ratio of kaons to pions is shown for $T_H = 176$ MeV and for $T_H = 196$ MeV. For $T_H = 176$ MeV our results are roughly at the upper edge of the experimental values. However, for $T_H = 196$ MeV our results are slightly higher than the experimental values. Although, the results at $T = 110$ MeV are almost exactly those of the uppermost experimental data point.

We can also observe the effects of the expansion on the $\Lambda\bar{\Lambda}$ pairs as seen in Fig. 24 and Fig. 25. We see that both reach the experimental values almost immediately ($T > 170$ for $T_H = 176$ MeV and around $T = 190$ for $T_H = 196$ MeV). The one exception is again for an underpopulation of Hagedorn states, which reaches chemical equilibrium at $T \approx 165$ for $T_H = 176$ MeV and already by $T = 170$ for $T_H = 196$ MeV.

The ratio of $\Lambda/\pi$’s is shown in Fig. 26. In both cases the $\Lambda/\pi$’s match the experimental values extremely well. For $T_H = 176$ MeV our results reach the equilibrium values at $T \approx 170$ MeV and for $T_H = 196$ MeV the experimental values are reached already by $T \approx 170$ MeV.

A summary graph of all our results is shown in Fig. 27. The gray error bars cover the range of error for the experimental data points from both STAR and PHENIX. The points show the range in values for the various initial conditions at $T = 110$ MeV. We see in our graph that our freezeout results match the experimental data well.

What the graphs in Figs. 18-26 show us is that a dynamical scenario is able to explain chemical equilibration values that appear in thermal fits by $T = 160$ MeV. In general, $T_H = 176$ MeV and $T_H = 196$ give chemical freeze-out values in the range between $T = 160 - 170$ MeV. These results agree well with the chemical freeze-out temperature found in [16].

Moreover, the initial conditions have little effect on the ratios and give a range in the chemical equilibrium temperature of about $\sim 5$ MeV, which implies that information from the QGP regarding multiplicities is washed out due to the rapid dynamics of Hagedorn states. Lower $\beta_i$ does slow the chemical equilibrium time slightly. However, as seen in Fig. 27 they still fit well within the ex-
FIG. 20: Results for the ratio of p’s with various initial conditions. Note that for STAR $p\pi^{-}/\pi^{-} = 0.11$ and $\bar{p}\pi^{-}/\pi^{-} = 0.082$. Along the top axis of each graph the corresponding time is shown in fm/c.

VI. PRODUCTION OF $\Omega\bar{\Omega}$

We can also use our model to investigate the possibility of $\Omega$'s. In [10], they discussed the possibility of $\Omega$'s being produced from the following decay channels:

$$HS \leftrightarrow \Omega + X$$

The first decay channel of a mesonic non-strange Hagedorn state we can implement straightforwardly with our model by employing the canonical branching ratio via Fig. 6. The results are shown in Fig. 28 for $T_H = 176$ MeV, in Fig. 29 for $T_H = 196$ MeV, and the $\Omega/\pi$ ratio is shown in Fig. 30. We are able to find the average number of $\Omega$'s from [31] as seen in Fig. 6. We see that, using only the first reaction, we are still impressively able to adequately populate the $\Omega\bar{\Omega}$ pairs so that they roughly match the experimental data. On the other hand, from Fig. 7 we see that for the $\Omega$ particle the equilibration time are short only very close to $T_c$. The scenario is thus more delicate. If one would take eg one half, or one fourth, respectively, of the decay width of that of eq. 7, the total production of $\Omega$ is not sufficient up to 25 %, or up to 50%, respectively, to meet the experimental yield (the other ratios are not significantly affected by such a change of the decay width).

In a future work, it would be interesting to observe the other decay channels as given in Eq. 21 and advertised in [10]. The second reaction includes a mesonic, three times strange Hagedorn state whereas the third decay channel includes a baryonic, strange Hagedorn state. Both states are much more likely to directly decay into a $\Omega$. These are, admittedly, exotic states, but should also occur in the spirit of Hagedorn states. In order to observe these decay channels a method, e.g. a microscopic quark model, must be found to find the appropriate Hagedorn spectrum for strange mesonic/baryonic Hagedorn states.

VII. CONCLUSIONS

In this paper we found that hadronic matter, at RHIC or SPS energies, can reach chemical equilibrium within a dynamical scenario using Hagedorn states close to the critical temperature. These states were able to produce quick chemical equilibration times in (anti-)proton, (anti-)kaons, and (anti-)lambdas close to the critical temperature due to their strong increase in their abundancy. The existence of such a mixture of standard hadrons and Hagedorn states just below the phase transition can explain dynamically the chemical equilibration of the hadronic species at around temperatures of 160 MeV to 170 MeV as seen within the thermal models.

From our analytical results we found that the chemical equilibration time depends on the temperature, decay widths, and branching ratios, but not the initial conditions. While this changes slightly when an expanding fireball is considered, the initial condition still only play a small role and only minimally affect the ‘freeze-out’ temperature at which chemical equilibrium is reached. This demonstrates that regardless of the population of hadrons coming out of the QGP phase, the initial conditions are washed out and everything can reach abundances which correspond to those of chemical equilibrium by the chemical freezeout temperatures found in [16].
Moreover, from our previous paper [11] we showed that particle ratios ($K/\pi^+$ and $(B + \bar{B})/\pi^+$) are not affected by the initial conditions and here we showed that $p/\pi$, $K/\pi$, $\Lambda/\pi$ and also $\Omega/\pi$ match the experimental values regardless of the initial conditions. Specially, Fig. 27 demonstrates this nicely and summarizes our findings: Regardless of the initial conditions, our dynamical scenario can match experimental data. We do find, however, that $T_H = 196$ fits within the experimental data box for $K/\pi$ whereas $T_H = 176$ is slightly above. This appears to reconfirm the findings in [16].

Our results imply that both lattice temperature can ensure that the hadrons reach their chemical equilibrium values by $T = 160 - 170$ MeV. Although the ratios for $T_H = 176$ do fit the data somewhat better, both math the experimental values reasonably well. This implies that independent of the critical temperature the hadrons are able to reach chemical freeze-out.

We see sufficiently short time scales for the chemical equilibrium of hadrons. The protons, kaons, and lambdas reach chemical equilibrium on the order of $\Delta \tau \approx 1 - 2 L_m/c$. Moreover, Hagedorn states states provide a very efficient way for incorporating multi-hadronic interactions (with parton rearrangements).

In an upcoming paper we will use a canonical model to derive all the branching ratios included in our calculations. We can then look at reactions that include a mixture of strange and non-strange baryons (for instance, $\bar{p} + \Lambda$) and multi-strange baryons. However, considering that our initial results produce quick chemical equilibration times for the baryons, kaons, and lambdas, it is reasonable to believe that this will occur for mixed reactions and multi-strange baryons as well. In addition, the machinery of standard hadronic reactions, i.e. binary scattering processes and resonance production processes, help also to equilibrate the various hadronic degrees of freedom. Still, our work indicates that the population and repopulation of potential Hagedorn states close to phase boundary can be the key source for a dynamical understanding of generating and chemically equilibrating the standard and measured hadrons.

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APPENDIX A: ANALYTICAL SOLUTIONS OF VARIOUS EQUILIBRATION PROCESSES

If our initial conditions are such that both the pions and Hagedorn states begin far out of chemical equilibrium, we can find an analytical solution by subdividing the analysis into three distinct stages. Initially, during stage 1 the pions are underpopulated such that we can make the approximation that the Hagedorn states are held at their initial value of $\beta_1$. One can see this from the difference in the time scales from Tab. II where $\tau_0 > \tau_0^p$ and $\tau_1 > \tau_\pi$. Since $\alpha \approx 0$ we let $\lambda_\pi^0 \approx 0$, then substituting this into Eq. (16) we obtain

$$\dot{\lambda}_\pi = \sum \Gamma_i \frac{N_i^{eq}}{N_{i^p}^{eq}} \beta_i \langle n_i \rangle,$$

$$\lambda_\pi = \left(\frac{t}{\tau_\pi^0} + \alpha\right), \quad (A1)$$

which is the fugacity of the pions in stage 1 and gives $\tau_\pi^0 = \sum \Gamma_i \frac{N_i^{eq}}{N_{i^p}^{eq}} \beta_i$. Again using the approximation $\alpha \approx 0$ and substituting Eq. (A1) into the Hagedorn state rate equation in Eq. (16), with the solution

$$\lambda_i = \Gamma_i \left(\frac{t}{\tau_i^0}\right)^{\langle n_i \rangle} - \lambda_i$$

$$\lambda_i = \left[1 - \langle n_i \rangle \left(\frac{t}{\tau_i^0}\right)^{-\langle n_i \rangle} \right] e^{-\left(\frac{t}{\tau_i}\right) \sum_{j=0}^{\infty} \frac{\langle n_i \rangle}{j! \langle n_i + j \rangle} (\frac{t}{\tau_i})^j}$$

$$\alpha e^{-\left(\frac{t}{\tau_i}\right)} + \beta_1 e^{-\left(\frac{t}{\tau_1}\right)}. \quad (A2)$$

Substituting $x = \frac{t}{\tau_1}$ into the integral in Eq. (A2), expanding the exponential inside the integral so $e^{-x} = \sum_{j=0}^{\infty} \frac{x^j}{j!}$, and integrating over $\xi$, provides us with the fugacity of the Hagedorn states in stage 1

$$\lambda_i = \left(\frac{t}{\tau_i^0}\right)^{\langle n_i \rangle} \left[1 - e^{-\left(\frac{t}{\tau_i}\right)} \sum_{j=0}^{\infty} \frac{\langle n_i \rangle}{j! \langle n_i + j \rangle} (\frac{t}{\tau_i})^j\right]$$

$$\alpha e^{-\left(\frac{t}{\tau_i}\right)} + \beta_1 e^{-\left(\frac{t}{\tau_1}\right)}. \quad (A3)$$

Therefore, Eq. (A1) and Eq. (A3) describe the behaviour of the pions and Hagedorn states during the initial stage of the evolution towards chemical equilibrium. They are then compared to the numerical results in Fig. 21.

As the pions near equilibrium our approximation of $\lambda_\pi \approx 0$ no longer holds and we switch to stage 2 where we assume $\lambda_\pi \approx 1$ at time $t_1$. Here $t_1$ is a time when the pions are almost in chemical equilibrium, which is normally taken when the pions reach about $\sim 95\%$ of their chemical equilibrium value. Returning to the pion equation in Eq. (16), we can substitute in $\lambda_\pi = 1 - \epsilon$ and use the approximation $(1 - \epsilon) \approx 1 - \epsilon\epsilon$

$$\dot{\epsilon} = -\sum \Gamma_i \frac{N_i^{eq}}{N_{i^p}^{eq}} \left(\beta_i - 1\right) \langle n_i \rangle + \langle n_i^2 \rangle \epsilon. \quad (A4)$$

Additionally, we substituted in $\beta_i$ for $\lambda_i$ as an approximation since the Hagedorn states do not change significantly in Stage 1 (the majority of the evolution is done by the pions). Recall that $\beta_i = \lambda_i(t = 0)$ and it is a constant. In its present form, Eq. (A4) can be integrated. We also define $\epsilon(t_1) = 1 - \eta$ where $\eta$ is close to 1 ($\eta$ is the measurement of how close the pions are to their equilibrium value when we switch from Stage 1 to Stage 2). Then, after integration

$$\lambda_\pi = \left(1 + \gamma - (1 + \gamma - \eta) e^{-\frac{t-t_1}{\tau_\pi}}\right) \quad (A5)$$
where \( \gamma = \frac{\sum_i \Gamma_i \langle n_i \rangle}{\sum_i \Gamma_i \langle n_i \rangle} \) and \( \tau_\pi = \frac{N_{\pi}^{eq}}{\sum_i \Gamma_i N_{eq}^i \langle n_i^2 \rangle} \). Analogously to stage 1, we substitute the pion equation, i.e., Eq. (A5) into the Hagedorn resonance equation in Eq. (16) and integrate

\[
\lambda_i = \left[ de^{-\frac{1-e}{\gamma}} + 1 + \langle n_i \rangle \gamma \right] - \left( \frac{\tau_\pi}{\tau_\pi - \tau_i} \right) \langle n_i \rangle (1 + \gamma - \eta) e^{-\frac{1-e}{\gamma}}
\]

where \( d = \omega_i - 1 + \langle n_i \rangle \gamma + \left( \frac{\tau_\pi}{\tau_\pi - \tau_i} \right) \langle n_i \rangle (1 + \gamma - \eta) \) and \( \lambda_i(t_1) = \omega_i \). Thus, our equations for the evolution of the pions and Hagedorn states are Eq. (A5) and Eq. (A6), respectively. As with stage 1, the evolution equation for the Hagedorn states is dictated by that of the pions.

Stage 3 i.e. quasi-equilibrium begins once the pions and at least one species of Hagedorn resonances (\( \tau_{\pi GeV} \) is the shortest chemical equilibration time) has surpassed its equilibration time (\( \tau_\pi \) and \( \tau_i \), respectively). To understand quasi-equilibrium we must use the effective pion number

\[
\tilde{N}_\pi = N_\pi + \sum_i N_i \langle n_i \rangle , \quad (A7)
\]

because we need a variable that can observe the effects of both the pions and resonances. The effective pion number essentially includes the number of effective pions that each Hagedorn state could decay into. Thus, we start by taking the derivative of Eq. (A7) in terms of its fugacity

\[
\dot{\lambda}_\pi = \frac{1}{N_{\pi}^{eq}} \left[ N_{\pi}^{eq} \dot{\lambda}_\pi + \sum_i N_{eq}^i \dot{\lambda}_i \langle n_i \rangle \right]
\]

\[
= \sum_i \Gamma_i N_{eq}^i \frac{1}{N_{\pi}^{eq}} \left[ \langle n_i \rangle \sum_n B_{i,n} \lambda_n^a - \sum_n B_{i,n} n \lambda_n^0 \right] \quad (A8)
\]

Once again we make the substitution \( \lambda_\pi = 1 - \epsilon \) so that

\[
\dot{\epsilon} = -\frac{1}{N_{\pi}^{eq}} \sum_i \Gamma_i N_{eq}^i \sigma_i^2 \epsilon . \quad (A9)
\]

where \( \sigma_i^2 = \langle n_i^2 \rangle - \langle n_i \rangle^2 \) in the Gaussian distribution of our branching ratios. To relate \( \epsilon \) and \( \dot{\epsilon} \) we return to Eq. (A7) and separate \( \lambda_i \) into a sum over the resonances in quasi-equilibrium and one over the “freely” equilibrating
resonances
\[
\tilde{\lambda}_\pi = \frac{1}{N^\text{eq}_\pi} \left[ N^\text{eq}_\pi \lambda_\pi + \sum_{i=1}^{Q} N^\text{eq}_i (n_i) \lambda_i + \sum_{i=1}^{\text{free}} N^\text{eq}_i (n_i) \right].
\]  
(A10)

Since the pions reach quasi-equilibrium first, i.e., \( \tau_\pi < \tau_i \) near \( T_i \), we set the \( \pi \) rate equation in Eq. (16) equal to zero, which gives \( \lambda_i \approx \frac{1}{(n_i)} B_i, n_i \lambda_i, \) so
\[
\tilde{\lambda}_\pi \approx 1 - \frac{\left( N^\text{eq}_\pi + \sum_{Q} N^\text{eq}_i (n_i) \right) \epsilon}{\sum_{\text{free}} (n_i) (N_i - N^\text{eq}_i \lambda_i)}. \quad \text{(A11)}
\]

Eq. (A11) then has the form \( \tilde{\lambda}_\pi \approx 1 - \tilde{\epsilon} \) where
\[
\tilde{\epsilon} = \frac{\left( N^\text{eq}_\pi + \sum_{Q} N^\text{eq}_i (n_i) \right) \epsilon}{\sum_{\text{free}} (n_i) (N_i - N^\text{eq}_i \lambda_i)}. \quad \text{(A12)}
\]

We can then solve for \( \epsilon \) in Eq. (A12) and substitute \( \epsilon \) into Eq. (A10), which in turn can be integrated. This leads us to the solution
\[
\tilde{\epsilon} = \epsilon \left[ e^{\frac{\tau_j - \tau_i}{\tau_\pi}} + \sum_{\text{free}} \langle n_i \rangle N_i^\text{eq} - \frac{\sum_i \Gamma_i N^\text{eq}_i \sigma_i^2}{N^\text{eq}_\pi \sum_{Q} N_i (n_i^2)} \right]
\cdot \sum_{\text{free}} N_i^\text{eq} \langle n_i \rangle e^{\frac{\epsilon - \tau_j}{\tau_\pi}} \epsilon \cdot \frac{\tau_j - \tau_i}{\tau_\pi} \lambda_i(x) \, dx. \quad \text{(A13)}
\]

where \( j \) stands for the latest resonance to reach chemical equilibrium at that point in time and
\[
\tau^\text{QE} = \frac{\sum_i \Gamma_i N_i^\text{eq} \sigma_i^2}{\sum_i \Gamma_i N_i^\text{eq} \sigma_i^2} + \sum_{Q} N_i^\text{eq} (n_i^2) \quad \text{(A14)}
\]

is the quasi-equilibrium time. Clearly, once all the Hagedorn states have reached chemical equilibrium than \( j \) symbolizes the resonance of \( M = 2 \) GeV, since it is the slowest Hagedorn state to equilibrate. The sums over “free” is the sum over the Hagedorn states that have not yet surpassed their respective chemical equilibrium time, \( \tau_i \). Once \( \tau_2^{2 \text{GeV}} \) is reached those sums equal zero. Therefore, after \( \tau_2^{2 \text{GeV}} \) all that remains is
\[
\tilde{\epsilon} = \epsilon e^{\frac{-\tau_2^{2 \text{GeV}}}{\tau_\pi}} \quad \text{(A15)}
\]

where \( \tau^\text{QE}_\pi \) is shown in Tab. I. Finally, we rewrite Eq. (A15) in terms of the pion evolution equation
\[
\lambda_\pi = 1 - (1 - \kappa) e^{\frac{-\tau_2^{2 \text{GeV}}}{\tau_\pi}} \quad \text{(A16)}
\]

where \( \kappa = \lambda_\pi (\tau_2^{2 \text{GeV}}) \).

Because the resonance equation depends on the population of the pions we substitute Eq. (A16) into the Hagedorn resonance rate equation in Eq. (10), assuming
the pions are near equilibrium (i.e., we use the approximation \( \lambda = 1 - \epsilon \) and \((1 - \epsilon)^n \approx 1 - n\epsilon)\)

\[
\lambda_i = \left( \frac{1}{\tau_\pi^E} - \frac{1}{\tau_i} \right) (1 - \kappa_i) e^{-\frac{\tau_\pi}{\tau_\pi^E}} \\
+ 1 - \frac{\tau_\pi^E}{\tau_\pi} (1 - \kappa_i) e^{-\frac{\tau_\pi}{\tau_\pi^E}}. \tag{A17}
\]

where \( \theta_i = \lambda_i (T_{2\text{GeV}}) \). Thus, for stage 3 the population equations for the pions and the Hagedorn states are Eq. [A16] and Eq. [A17] so long as \( t \geq \tau_{2\text{GeV}} \).

Fig. 31 reveals a remarkable close fit with our numerical results for \( T = 175 \text{ MeV} \) i.e. \( T < T_H \). Thus, the quasi-chemical equilibrium time, \( \tau^{QE} \), depends only on \( \Gamma_i, \langle n_i \rangle, \sigma_i^2, \) and \( N^{eq} \), which is temperature dependent, but not on our initial conditions. As mentioned in the text, though, \( \tau^{QE} \) includes many non-linear affects that only occur close to the chemical equilibrium. Thus, the more appropriate time scale is \( \tau_\pi^0 \) in order to describe the dynamics.

We also see from Fig. 31 that when \( K\bar{K} \) pairs are included that the pions and Hagedorn resonances equilibrate in roughly the same amount of time, which implies that our analytical solution can still be approximately applied when \( K\bar{K} \) pairs are present.

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FIG. 30: Results for the ratio of $\Omega/\pi$'s with various initial conditions. Note that for STAR $\Omega/\pi = 9.5 \times 10^{-4}$ and $\overline{\Omega}/\pi = 9.6 \times 10^{-4}$. Along the top axis of each graph the corresponding time is shown in $fm/c$.

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FIG. 31: Numerical and analytical results for the pions and Hagedorn states where $\eta = 0.9$ at $T = 175$ MeV for $T_H = 176$ MeV when $\beta_i = 1.1$ and $\alpha = 0.9$ (top) and the numerical results for the same initial conditions including $KK$ pairs with $\phi = 0$.

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[38] We solve our coupled non-linear differential equations using \texttt{NDSolve} in Mathematica.