SEPARATE CHEMICAL FREEZE-OUT OF STRANGE PARTICLES WITH CONSERVATION LAWS

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1. Introduction

The hadronic multiplicities measured in heavy ion collisions and in the collisions of elementary particles are traditionally described by the Hadron Resonance Gas Model (HRGM) [1–5]. It is based on the assumption that the fireballs produced in such collisions reach a full thermal equilibrium. Using this assumption, it is possible to describe the hadronic multiplicities registered in experiment with the help of two parameters: temperature $T$ and baryo-chemical potential $\mu_B$. The parameters $T$ and $\mu_B$ obtained from the fit of multiplicities for various collision energies correspond to the stage of chemical freeze-out. Its physical meaning is that the inelastic collisions at this stage cease simultaneously for all sorts of particles. However, in such a simple form, the concept of chemical freeze-out works well for the hadrons, which consists of the $u$ and $d$ (anti)quarks, while the strange hadrons demonstrate a deviation from the chemical equilibrium. At the same time, the hydrodynamic simulations (see, e.g., the review [6]) rather successfully reproduce the transverse momentum spectra of strange particles. This is an old problem of the thermal approach. In order to account for the observed deviation of strange particles from the complete chemical equilibrium, the additional parameter $\gamma_s$, the strangeness suppression factor, was suggested [7] long ago. Although the concept of strangeness suppression proved to be important both in the collisions of elementary particles [4] and in nucleus-nucleus collisions [4, 8], the problem of its justification remains unsolved. Thus, up to now, it is unclear which is the main physical reason responsible for the chemical non-equilibrium of strange hadrons.

Moreover, it is well known [2] that the fit of hadron multiplicities with the strangeness suppression factor $\gamma_s$ improves the quality of a data description, but still the fit seldom attains a good quality, especially at low collision energies. This is clearly seen from the center-of-mass energy behavior of two most prominent ratios that involve the lightest strange meson, i.e. $K^+$/\(\pi^+\), and the lightest strange baryon, i.e. $\Lambda$/\(\pi^-\), which, so far, cannot be successfully reproduced [2, 4, 8] by the traditional versions of the HRGM. In addition, the ratios involving the multistrange hyperons $\Xi$ and $\Omega$ exhibit the apparent failure of the $\gamma_s$ fit at the center-of-mass energies $\sqrt{s_{NN}} = 8.76, 12.3$, and $17.3$ GeV [2]. Since the $\gamma_s$ fit does not improve their description sizably, we conclude that there should exist a different reason for the apparent deviation of strange hadrons from the chemical equilibrium. Hence, the concept of chemical freeze-out requires a further development.
Recently, an alternative concept of chemical freeze-out of strange hadrons was suggested [9]. Instead of a simultaneous chemical freeze-out for all hadrons, two different chemical freeze-outs were suggested: one for particles containing strange charge, even hidden, (we refer to it as the strangeness freeze-out, i.e. SFO) and another one (FO) for all other hadrons, which contains only $u$ and $d$ (anti)quarks. A partial justification for the SFO hypothesis is given in [10–12], where the early chemical and kinetic FO of Ω hyperons and $J/\psi$ and $\phi$ mesons is discussed for the energies at and above the highest SPS energy. In this article, we further develop and refine the SFO concept of Ref. [9] and present a more coherent and detailed picture of two freeze-outs together with new arguments which allow us to better justify and to improve the performance of the HRGM.

The paper is organized as follows. In the next section, we discuss the concept of chemical freeze-out in some details and give the arguments that, in a meson-dominated hadronic medium, the SFO should occur earlier than the FO. Section 3 is devoted to the description of the HRGM with the multicomponent hard-core repulsion. The results are presented in Section 4, and Section 5 contains our conclusions and suggestions.

2. Framework of the Thermal Model

In 1950 in his pioneering paper [13], E. Fermi suggested to use the statistical model to find the outcome of high energy nucleon-nucleon collisions. Since there were produced from 10 to 30 hadrons in such reactions, they were named as the processes of multihadron production. According to E. Fermi, the large number of particles in the final state of these processes naturally suggests to apply the methods of statistical mechanics. The next crucial step suggested by E. Fermi was a justification of the thermal equilibrium assumption due to the strong interaction between particles. A few years later, L. D. Landau suggested to apply the relativistic hydrodynamics to the reactions of multihadron production [14], because the applicability conditions of relativistic hydrodynamics are basically the same as for the full (local) thermal equilibrium, if the strong discontinuities are absent.

Since that time, the assumption of thermal equilibrium at some stage of the multihadron production reactions was tested experimentally both in the nucleon-nucleon collisions and in the collisions of heavy ions. In other words, the outcome of such reactions was compared to the results of statistical models. The coincidence between the statistical models predictions and the experimental results appeared to be good both for the nucleon-nucleon collisions and for the heavy ion collisions at the energy range starting from the center-of-mass energy $\sqrt{s_{NN}} = 2$ GeV per nucleon in the fixed target experiments performed at the Brookhaven AGS up to the center-of-mass energy $\sqrt{s_{NN}} = 2.76$ TeV achieved at the Large Hadron Collider [2,15]. It was even suggested that, for the high energy electron-positron collisions, the statistical model can also describe the hadron multiplicities [3]. However, a more thorough analysis [16] showed later on that even within a rather sophisticated canonical ensemble consideration, the discrepancy between theory and experiment is rather large with $\chi^2/\text{dof} > 5$.

Let us now consider, in some details, a particular set of models used to describe hadron multiplicities in nucleon or heavy ion collisions, that are known as the HRGM [1–5,8,15]. A common feature of this set of models is the assumption that, at some time moment, there exists a fireball consisting of all possible hadronic states being locally in the thermal and chemical equilibria. The term “chemical equilibrium” means that the rates of forward and backward reactions are equal, i.e., for any hadron species, the rate of its production is equal to the rate of its destruction. The characteristic time of equilibration varies with collision energy, but one can safely say that it lies within the interval of 0.1–10 fm/c [17–19]. This means that one can safely ignore the weak interaction, because its characteristic time is essentially longer. Therefore, the baryon charge $B$, strange charge $S$, isospin projection $I_3$, charm charge $C$, and bottom charge are conserved in almost all hadron reactions. Some of the most frequent hadronic reactions read: $\pi\pi \to \rho \to \pi\pi$, $\pi K \to K^+ \to \pi K$, $\pi N \to \Delta \to \pi N$. They lead to the thermal equilibration, but do not change the number of particles. Another reactions, such as $\pi N \to N^* \to \Delta \pi \to N\pi\pi$, change the number of particles and lead to the chemical equilibration. Was such a system of all hadron states kept in a finite box of volume $V$, it would inevitably equilibrate both thermally and chemically at $t \rightarrow \infty$. Let us define the characteristic time of equilibration between the species $A$ and $B$ as the average time when some typical number of collisions between $A$ and
and $B$ occurred. If there are only $A$ and $B$ species in the box, then $\tau_{AB} \sim \frac{1}{N_A N_B \sigma_{AB}}$, where $\sigma_{AB}$ is the $AB$ reaction cross-section, and $n_A(n_B)$ denote the concentration of species $A$ ($B$). If one considers a gas of many species in the box out of equilibrium, then the equilibration times will be defined from the system of equations (assuming only the reactions $2 \to 1$ and $1 \to 2$):

$$
\frac{dN_i}{dt} = \sum_A \frac{N_A N_B v_{AB}^{rel}}{V} \sigma_{AB \to i} - \sum_A \frac{N_i N_A v_{Ai}^{rel}}{V} \sigma_{Ai \to B} - \sum_{CD} \Gamma_{i \to CD} N_i,
$$

where $N_i$, $N_A$, and $N_B$ are the number of hadrons of the corresponding kind, $\sigma$ denotes the corresponding cross-sections, and $\Gamma$ is the decay rate. The first term on the right-hand side describes the formation of particles of kind $i$, the second term stands for the particle destruction of this kind in the $2 \to 1$ reaction, and the third term stands for the decays of this kind of particles. From these equations, one can see that the larger production cross-section leads to a faster equilibration, while the larger volume leads to a slower equilibration. One can also see that, depending on the cross-sections of production and decay and on the volume, the equilibration times for different species may be different. These equations are, of course, oversimplified, because they do not include the momentum dependencies. If one introduces such dependencies, then one obtains the system of Boltzmann equations, and, hence, Eq. (1) can be regarded as the system of Boltzmann equations averaged over momenta. However, even these oversimplified equations can help to understand the way how a system approaches an equilibrium. For instance, from Eq. (1), one can see that increasing the box volume $n$ times is equivalent to decreasing all the cross-sections by $n$ times. One can also see that, for very large volumes, only the decays will occur.

If the system is expanding, i.e. $V = V(t)$, then there is no guarantee that all particle species will be at the chemical and thermal equilibria at any time. The simplest way to qualitatively characterize an expanding system is to introduce a set of characteristic times: expansion time $t_{ex}$, thermalization time $t_{th}$, and chemical equilibration $t_{ch}$ time for different species. It is known that, typically of the reactions of strongly interacting particles, there is the inequality $t_{ch} \gg t_{th}$ [17–19]. It is equivalent to the statement that the cross-sections of reactions, which lead to a chemical equilibration, are much smaller than the cross-sections of reactions, which lead to a thermalization. During the expansion process, the system volume increases, or one can say equivalently that all cross-sections effectively decrease by the same factor. Therefore, the reactions, which lead to a chemical equilibration will cease earlier, than the reactions, which lead to a thermalization, they are called, respectively, as chemical and kinetic freeze-outs. Since the cross-sections of different reactions are not the same, one can say generally about the chemical and kinetic freeze-outs for each particle species.

Typically in vacuum, the reactions involving strange particles have smaller cross-sections than the reactions involving only non-strange particles (charm and bottom are not considered here at all). Then, from our previous consideration, one can conclude that if the cross-sections and the thresholds of hadronic reactions occurring at the late stage of the expansion do not differ from their vacuum values, then the chemical equilibrium for strange particles should be lost earlier. The kinetic freeze-out for strange particles is also going to occur earlier than the kinetic freeze-out of non-strange hadrons, but later than the chemical freeze-out for any hadron species. These conclusions are based on the following hierarchy of the switching-off times of hadronic reactions:

$$
t_{K\Lambda \to \Sigma} > t_{\pi N \to \Delta \to N \pi} \gg
\gg t_{K \to K^* \to K} > t_{N \to \Delta 
\to N \pi}.
$$

It is not only cross-sections that influence the freeze-out times. As one can see from Eq. (1), the smaller the concentrations, the lower the rates of reactions are expected. The numbers of strange particles different from kaons are smaller than the number of protons, and this is one more factor that makes slower the reactions of strangeness exchange and leads to the earlier freeze-outs of strange particles. Of course, one should keep in mind that this simplified treatment is valid at low particle densities, if the approximation of binary reactions is reasonable, and if the surrounding medium does not essentially modify the reaction threshold. Therefore, the appearance of the results that contradict the conclusions above should be considered as a signal that the chemical freeze-out.
picture based on Eqs. (1) and (2) is not justified, and, hence, one has to seek for another explanation.

Nevertheless, the argumentation above motivates one to consider a separate chemical freeze-out of strange particles in the HRGM. This was done recently in two independent studies [9, 20] and [21]. In [21], three free parameters were taken for FO (temperature, baryon chemical potential, and volume) and three free parameters of the same kind for SFO. The electric charge chemical potential $\mu_Q$ was taken from the condition $N_B/N_Q = 2.5$ for both freeze-outs. Species subjected to the SFO were all strange particles and the $\phi$-mesons. The strange charge was treated canonically, and the particle multiplicities were fitted. The approach of [9, 20] is quite different. The parameters of FO and SFO were connected by the conservation laws, namely the baryon number conservation, $I_3$ conservation, and entropy conservation. Both freeze-outs were treated grand canonically, and the $\phi$ mesons were not subjected to the earlier freeze-out. In addition, in contrast to the oversimplified treatment of the equation of state, the HRGM of [9] includes the width of all hadron resonances and the short-range repulsion, which is taken into account via the excluded volume corrections, while these important features were neglected in [21].

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We would like to stress, although being simple and successful in describing the hadronic multiplicities, the approach suggested in [21] violates the above-mentioned conservation laws. Moreover, in such approach, it might happen that not only the entropy conservation is violated, but the entropy may decrease from an earlier freeze-out to the later one. Finally, while the number of fitted multiplicities is rarely exceeding 10 per one collision energy value, having six fitting parameters for each energy value seems to be excessive. Therefore, we outline an alternative model [9] below, which seems to be physically more relevant.

3. Model Formulation

In the simplest version, the HGRM represents the gas of hadrons being in the chemical and thermal equilibria, which is described by the grand canonical partition function. The multiplicity of particles of mass $m_i$ and degeneracy $g_i$ is given by

$$ N_i = g_i V \int \frac{d^3k}{(2\pi)^3} \frac{1}{e^{(\sqrt{m_i^2+k^2}-\mu_i)/T} \pm 1}, $$

where the sign $+(-)$ in the equation above stays for the Fermi (Bose) statistics, and $\mu_i$ denotes the full chemical potential $\mu_i = \mu_B B_i + \mu_S S_i + \mu_I I_3 i$. of particles of sort $i$, $B_i$ is their baryonic charge, $S_i$ is their strange charge, and $I_3 i$ denotes their third projection of isospin. The chemical potentials $\mu_B$, $\mu_S$, and $\mu_I$, which correspond to the conserved charges, can be found from the conservation laws

$$ \sum_i N_i B_i = B^{\text{init}}, $$

$$ \sum_i N_i S_i = S^{\text{init}}, $$

$$ \sum_i N_i I_3 i = I_3^{\text{init}}. $$

Then the temperature $T$ and the system volume $V$ will be free parameters. One can, however, take $T$ and $\mu_B$ as free parameters, and this is a conventional choice. In [22], we argued that, for mid-rapidity, the quantities $B^{\text{init}}$ and $I_3^{\text{init}}$ are anyway unknown, so one can fit the ratios and have $T$, $\mu_B$, and $\mu_I$ as the fitting parameters. Using this procedure, one gets the hadron multiplicities that correspond to the full thermal equilibrium. To get the final particle multiplicities, one has to take the decays of hadron resonances into account (see below).

An extension of the HRGM to two freeze-outs is almost obvious in the case of [21], where both non-strange and strange freeze-outs have their own parameters and are by no means connected. In such a case, one considers two separate ideal gases with their own parameters. However, if one follows the way described in [9], then some complications arise. One problem is to properly include the conservation laws, then one has to take the excluded volume into account in a consistent way. By consistency, we mean that the standard thermodynamic identities should be obeyed. One more issue is a change of the entropy between two freeze-outs due to the decays of strange resonances. However, as we argued in [9], the latter is negligible, because the time interval between two freeze-outs is short.

We would like to stress that the excluded volume for all particles remains the same after the SFO. Indeed, not all reactions between the strange and non-strange particles cease, but only those with the strangeness exchange. For instance, the reaction $\pi K \rightarrow K^* \rightarrow \pi K$ survives after the SFO. It keeps the
same excluded volume between pions and kaons, but
does not provide the chemical equilibrium for kaons.

After these comments, let us formulate our ap-
proach. It is based on the multicomponent formulation
of the HRGM [5], which is currently the best at de-
scribing the observed hadronic multiplicities. There-
fore, it is natural to apply such a formulation to de-
scribe both the FO and the SFO. The present HRGM
was worked out in [5, 22–28]. The interaction be-
tween hadrons is taken into account via the hard-core
radii, with the different values for pions \( R_\pi \), kaons
\( R_K \), other mesons \( R_m \), and baryons \( R_B \). The best fit
values for such radii (\( R_\pi = 0.2 \text{ fm} \), \( R_m = 0.4 \text{ fm} \),
\( R_K = 0.1 \text{ fm} \), and \( R_K = 0.38 \text{ fm} \)) were obtained in
[5]. The main equations of the model are listed below,
but more details of the model can be found in [5, 22].

We consider the Boltzmann gas of \( N \) hadron species
in a volume \( V \) that has the temperature \( T \), baryonic
chemical potential \( \mu_B \), strange chemical potential \( \mu_S \),
and chemical potential of the isospin third component
\( \mu_{I3} \). The system pressure \( p \) and the \( K \)-th charge den-
sity \( n_i^K \ (K \in \{B, S, I3\}) \) of the \( i \)-th hadron sort are
given by the expressions

\[
p = \frac{\sum_{i=1}^{N} \xi_i}{T}, \quad \frac{n_i^K}{n_i^K} = \frac{Q_i^K \xi_i}{1 + \sum_{j=1}^{\xi_i} b_{ij}}, \quad \xi = \left( \begin{array}{c} \xi_1 \\ \xi_2 \\ \vdots \\ \xi_N \end{array} \right),
\]

(7)

where \( B \) denotes a symmetric matrix of the second
virial coefficients with the elements \( b_{ij} = \frac{2}{3} (R_i +
R_j)^3 \), and the variables \( \xi_i \) are the solutions of the following system:

\[
\xi_i = \phi_i(T) \exp \left[ \frac{\mu_i}{T} - \frac{\sum_{j=1}^{N} 2\xi_j b_{ij}}{T} \right] + \xi^T B \xi \left[ \sum_{j=1}^{N} \xi_j \right]^{-1},
\]

(8)

\[
\phi_i(T) = \frac{g_i}{(2\pi)^3} \int \exp \left( -\frac{\sqrt{k^2 + m_i^2}}{T} \right) d^3 k.
\]

(9)

Here, the full chemical potential of the \( i \)-th hadron
sort is defined as before, \( \phi_i(T) \) denotes the thermal
density of the \( i \)-th hadron sort of mass \( m_i \) and
degeneracy \( g_i \), and \( \xi^T \) denotes the row of variables \( \xi_i \).

The width correction is taken into account by aver-
eraging all expressions containing the resonance mass
by the Breit–Wigner distribution having a threshold
(see, e.g., [1] for more details). The effect of resonance
decay \( Y \to X \) with the branching ratio \( BR(Y \to X) \)
on the final hadronic multiplicity is taken into account
as \( n_i^{3m}(X) = \sum_Y BR(Y \to X)n_i^B(Y) \), where
\( BR(X \to Y) = 1 \) for the sake of convenience. The
masses, widths, and strong decay branchings of all
hadrons were taken from the particle tables used by
the thermodynamic code THERMUS [29].

The SFO is assumed to occur for all strange parti-
cles at the temperature \( T_{SFO} \), baryonic chemical po-
tential \( \mu_{B_{SFO}} \), isospin third projection chemical po-
tential \( \mu_{I3_{SFO}} \), and three-dimensional space-time ex-
tent (effective volume) of the freeze-out hypersurface
\( V_{SFO} \). The FO of hadrons, which are built of the \( u 
\)
and \( d \) (anti)quarks, is assumed to be described by its
own parameters \( T_{FO} \), \( \mu_{B_{FO}} \), \( \mu_{I3_{FO}} \), and \( V_{FO} \). The
obtained model parameters for two freeze-outs and their
\( \sqrt{S_{NN}} \) dependence are shown in Figs. 1–2. Equations
(7)–(9) for FO and SFO remain the same as for a si-
multaneous FO of all particles. In both cases, \( \mu_S \) is
found from the net zero strangeness condition. The
major difference of the SFO approach is the presence
of conservation laws and the corresponding modific-
ation of multiplicities due to resonance decays. Thus,
we assume that, between two freeze-outs, the sys-
tem is sufficiently dilute, and, hence, its evolution is
governed by the continuous hydrodynamic evolution,
which conserves the entropy. Then the equations for
the entropy, baryon charge, and isospin projection
conservation connecting two freeze-outs are as fol-
lows:

\[
\frac{s_{FO}}{n_{B_{FO}}} = s_{SFO}V_{SFO},
\]

(10)

\[
\frac{n_{B_{FO}}}{n_{SFO}}V_{SFO} = n_{SFO}V_{SFO},
\]

(11)

\[
\frac{n_{I3_{FO}}}{n_{SFO}}V_{SFO} = n_{I3_{SFO}}V_{SFO}.
\]

(12)

Getting rid of the effective volumes, we obtain

\[
\frac{s}{n_{B}} \bigg|_{FO} = \frac{s}{n_{B}} \bigg|_{SFO}, \quad \frac{n_{B}}{n_{SFO}} \bigg|_{FO} = \frac{n_{B}}{n_{SFO}} \bigg|_{SFO}.
\]

(13)

Therefore, the variables \( \mu_{B_{SFO}} \) and \( \mu_{I3_{SFO}} \) are not free
parameters, since they are found from system (13),
and only \( T_{SFO} \) should be fitted. Thus, for the SFO,
the number of independent fitting parameters is 4 for
each value of collision energy.
Fig. 1. Parameters of chemical freeze-outs in the model with two freeze-outs. Upper panel: triangles correspond to the SFO, their coordinates are \((\mu_B, T)\), while circles correspond to the FO, and their coordinates are \((\mu_B, T)\). The curves correspond to isentropic trajectories \(s/\rho_B = \text{const}\) connecting two freeze-outs. Lower panel: \(\sqrt{S_{NN}}\) dependence of the ratio of the SFO temperature to the FO temperature.

The number of resonances appeared due to decays are found from

\[
N_{\text{fin}}(X) = \sum_{Y \in \text{FO}} BR(Y \rightarrow X)n^{th}(Y) + \sum_{Y \in \text{SFO}} BR(Y \rightarrow X)n^{th}(Y)\frac{V_{\text{SFO}}}{V_{\text{FO}}}. \tag{14}
\]

Technically, this is done by multiplying all the thermal concentrations for SFO by \(n_B B_{\text{FO}}/n_B B_{\text{SFO}} = V_{\text{SFO}}/V_{\text{FO}}\) and applying the conventional resonance decays.

4. Results

4.1. Data sets and fit procedure

In our choice of the data sets, we basically followed Ref. [2]. Thus, at the AGS energy range of collisions \(\sqrt{S_{NN}} = 2.7-4.9\) GeV, the data are available for the kinetic beam energies from 2 to 10.7 AGeV. For the beam energies 2, 4, 6, and 8 AGeV, there are only a few available data points: the yields for pions [30,31], for protons [32,33], for kaons [31] (except for 2 AGeV); for \(\Lambda\) hyperons, the data integrated over 4\(\pi\) are available [34]. For a beam energy of 6 AGeV, there exist the \(\Xi^-\) hyperon data integrated over 4\(\pi\) geometry [35]. However, the data for the \(\Lambda\) and \(\Xi^-\) hyperons have to be corrected [2], and, instead of the raw experimental data, we used their corrected values of Ref. [2]. For the highest AGS center-of-mass energy \(\sqrt{S_{NN}} = 4.9\) GeV (or a beam energy of 10.7 AGeV) in addition to the mentioned data for pions, (anti)protons, and kaons, there exist the data for \(\phi\) meson [36], for \(\Lambda\) hyperon [37], and for \(\bar{\Lambda}\) hyperon [38]. Similarly to [5], we analyzed only the NA49 midrapidity data [39–44] here, since they are traditionally the most difficult to be described. Because the RHIC high energy data of different collaborations agree with
each other, we present the analysis of the STAR results for $\sqrt{S_{NN}} = 9.2$ GeV [45], $\sqrt{S_{NN}} = 62.4$ GeV [46], $\sqrt{S_{NN}} = 130$ GeV [47–50], and 200 GeV [50–52].

To avoid possible biases, we fit the particle ratios rather than the multiplicities. The best fit criterion is the minimality of

$$\chi^2 = \sum_i \frac{(r_{i\text{theor}} - r_{i\text{exp}})^2}{\sigma_i^2},$$

where $r_{i\text{exp}}$ is an experimental value of the $i$-th particle ratio, $r_{i\text{theor}}$ is our prediction, and $\sigma_i$ is the total error of the experimental value.

4.2. Fit results

The FO and SFO parameters are connected by the conservation laws (13). Therefore, there is only one fitting parameter at each collision energy for the SFO, namely $T_{SFO}$, while other parameters are found from system (13). We study two things: the behavior of parameters and which ratios are improved in the SFO approach as compared to the case without SFO. First of all, we found out that, in the SFO case, $\chi^2$/dof = 58.5/55 = 1.06. At $\sqrt{S_{NN}} = 2.7, 3.3, 3.8, 4.3,$ and 4.9 GeV, the original description obtained within the multicomponent model [5] is very good. Hence, it has not improved significantly. Similar results are found at the highest RHIC energies $\sqrt{S_{NN}} > 62.4$ GeV. From Fig. 1, one can see that, within these two energy domains, the SFO temperatures demonstrate the largest deviations from the FO temperature, although they do not exceed 20%. At intermediate energies, we see a systematic improvement of the description of ratios. Three plots corresponding to the collision energies, at which an improvement after the SFO introduction is the most significant, $\sqrt{S_{NN}} = 6.3, 12,$ and 17 GeV, are shown in Fig. 3. As one can see from Fig. 3 for $\sqrt{S_{NN}} = 6.3, 12,$ and 17 GeV, the SFO approach improves the description of all ratios with more than one $\sigma$ deviation. For $\sqrt{S_{NN}} = 6.3$ GeV, the SFO greatly improves the $\Lambda/\pi^-$ and $\bar{p}/p$ ratios. For $\sqrt{S_{NN}} = 12$ GeV, four ratios of eight ones with more than one $\sigma$ deviation, namely $K^+/\pi^+$, $\Lambda/\Lambda$, $\Lambda/\pi^-$, and $\Xi^+/$ are improved. The SFO approach allows us to significantly improve the fit quality at $\sqrt{S_{NN}} = 17$ GeV. Figure 3 demonstrates that, due to the SFO fit, six of seven problematic ratios for the one-freeze-out fit moved from the region of deviations exceeding $\sigma$ to the region of deviations being smaller than $\sigma$. The most remarkable of them are $\bar{p}/\pi^-$, $\Lambda/\Lambda$, $\Xi^+/$, and $\Omega/\Omega$. Thus, the separa-

**Fig. 3.** Relative deviation of the theoretical description of ratios from the experimental value in units of experimental error $\sigma$. The symbols on the $OX$ axis demonstrate the particle ratios. The $OY$ axis shows $\frac{|r_{i\text{theor}} - r_{i\text{exp}}|}{\sigma_{i\text{exp}}}$, i.e. the relative deviation modulus for $\sqrt{S_{NN}} = 6.3, 12,$ and 17 GeV. The solid lines correspond to a model with one chemical freeze-out of all hadrons, while the dashed lines correspond to the model with the SFO.
tion of the FO and the SFO relaxes the strong connection between the non-strange and strange baryons and allows us not only to nicely describe the ratios of strange antibaryons to the same strange baryons, but also it allows us, for the first time, to successfully reproduce the antiproton to pion ratio.

As we discussed above, it is expected that the SFO occurs earlier, when the system is smaller, and, hence, \( V_{\text{SFO}} < V_{\text{FO}} \) or \( \frac{1}{V_{\text{SFO}}} > 1 \). In the Fig. 2, one can see that this is, indeed, the case for most values of collision energy. But at low energies, our expectation does not come true. One possible formal reason is the same as for the unexpected behavior of \( \chi^2/\text{dof} \). In this energy range, the number of data points is just slightly larger than the number of fitting parameters, and, because of that, the fit quality at low energies of collisions is very good without assumption of two freeze-outs. There might be also a physical reason for such a behavior. Namely, the freeze-out at low collision energies occurs at large baryonic densities, which may essentially affect the in-medium cross-sections of the reactions with strangeness exchange due to the additional attraction. Therefore, such reactions do not freeze-out earlier than other reactions.

Finally, we would like to suggest a generalization of the double freeze-out HRGM that will be able to ultimately improve the description of multiplicities. The first step is to identify the hadronic reactions that freeze-out noticeably earlier or later than most of the others. This should be done separately for each collision energy, since the reaction cross-sections, particle concentrations, and fireball expansion rate are different at different energies. Such reactions may be identified, by using system (1) or by running the transport model code and by counting for the reaction rates versus the time. If such reactions exist, then their separate freeze-out should be considered. It is clear that the conservation laws between the freeze-outs may be different, by depending on which reactions are switched-off. For instance, if all reactions with the \( \Omega \) hyperon are frozen, then the conservation law of the number of \( \Omega \) hyperons should be introduced. Probably, the charmed particles are good candidates for the separate freeze-out.

5. Conclusions

We have thoroughly discussed the assumption that, in heavy ion collisions, the strangeness exchange reactions may freeze-out earlier. Using such assumption, we have constructed a modification of the HRGM with two freeze-outs, which is connected with the conservation laws. One freeze-out corresponds to all strange particles, and another freeze-out is for all non-strange ones. The conservation laws allow us to get just one additional fitting parameter for each collision energy, as compared to the HRGM with a simultaneous chemical freeze-out of all hadrons. We have shown that such a model describes 111 independent hadron ratios measured at \( \sqrt{s_{NN}} = 2.7-200 \) GeV even better than the most elaborate version of the HRGM with a single freeze-out (\( \chi^2/\text{dof} = 1.06 \) for the model with two freeze-outs versus 1.16 for one freeze-out).

We suggest to go even further: for each collision energy, to separately identify the processes, which freeze-out at considerably different times than all the others and to construct a corresponding HRGM with two freeze-outs. The identification of such reactions can be done, by using the transport models.

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Separate Strangeness Freeze-Out


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ВІДОКРЕМЛЕНІЙ ХІМІЧНИЙ ФРІЗАУТ
ДИВНИХ ЧАСТИНОК ІЗ ЗАКОНАМИ ЗБЕРЕЖЕННЯ

Р е з ю м е

Розглянуто модель адронного резонансного газу із двома фрізаутами, що пов’язані між собою законами збереження. Наведено аргументи на користь того, що хімічний фрізаут дивних адронів має відбуватися раніше, аніж хімічний фрізаут недивних адронів. За допомогою представленої моделі виконано високоякісний фіт адронних множинностей, виміряних у зіткненнях важких іонів при енергіях зіткнень в системі центра мас від 2,7 до 200 ГеВ. Грунтуючись на успіху даного підходу, нами запропоновано спосіб радикально-го вдосконалення моделі адронного резонансного газу. Для цього ми пропонуємо визначати адронні реакції, які приписують раніше або пізніше, аніж більшість інших реакцій (для різних енергій зіткнень вони можуть бути різними), та розглядати для цих реакцій відокремлений фрізаути.