Letter to the Editor

Negative heat capacity for hot nuclei using formulation from the microcanonical ensemble

INDRA Collaboration

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Abstract. By using freeze-out properties of multifragmenting hot nuclei produced in quasifusion central $^{129}$Xe+$^{nat}$Sn collisions at different beam energies (32, 39, 45 and 50 AMeV) which were estimated by means of a simulation based on experimental data collected by the 4π INDRA multidetector, heat capacity in the thermal excitation energy range 4 - 12.5 AMeV was calculated from total kinetic energies and multiplicities at freeze-out. The microcanonical formulation was employed. Negative heat capacity which signs a first order phase transition for finite systems is observed and confirms previous results using a different method.

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An important challenge of heavy-ion collisions at intermediate energies is the identification and characterization of the nuclear liquid-gas phase transition in hot nuclei, which has been theoretically predicted for nuclear matter [1-4,15]. At present one can say that huge progress has been made even if some points can be deeper investigated [5-7,9,10]. Statistical mechanics for finite systems appeared as a key issue to progress, revealing specific first-order phase transition signatures related to the consequences of the local convexity of the entropy [11,12,13,14,15,16,17]. By considering the microcanonical ensemble with energy as extensive variable, the convex intruder implies a backbending in the temperature (first derivative of entropy) at constant pressure [18] and correspondingly a negative branch for the heat capacity (second derivative of entropy). Experimentally, these two converging signatures have been observed in hot nuclei from different analyses [19,20,21,22,23] of homogeneous event samples. It is important to recall here that signals of phase transition in hot nuclei are only meaningful at the level of statistical ensembles constructed from the outcome of carefully selected collisions [8]. Another consequence of the entropy curvature anomaly manifests itself when systems are treated in the canonical ensemble. In this case a direct phase transition signature is the presence of a bimodal distribution of an order parameter [24] such as the charge (size) of the largest fragment ($Z_{max}$) of multifragmentation partitions [25]. As far as microcanonical heat capacity calculation is concerned, a question still subsists. It concerns one hypothesis made in the method used up to now: the same temperature was associated with both internal excitation and thermal motion of emitted fragments, which is not physically obvious if one remembers that the level density is expected to vanish at high excitation energies [26,27,28,29]. Moreover it was shown in simulations compared to data that widths of the fragment velocity spectra cannot

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be reproduced with the hypothesis of a common temperature for intrinsic and kinetic degrees of freedom [30].

In the present letter we will give heat capacity information for experimental data from microcanonical formulae and by ignoring the hypothesis of a single temperature, which requires the introduction of a limiting temperature for fragments.

We first briefly recall the method proposed for measuring heat capacity using partial energy fluctuations with a microcanonical sampling [31,32,13]. The prescription is based on the fact that, for a given total energy, the average partial energy stored in a part of the system is a good microcanonical thermometer, while the associated fluctuations can be used to construct the heat capacity. From experiments the most simple decomposition of the thermal excitation energy is in a kinetic part, $E_k$, and a potential part, $E_{pot}$ (Coulomb energy + total mass excess). These quantities have to be determined at freeze-out and consequently it is necessary to trace back this configuration on an event by event basis. The true configuration needs the knowledge of the freeze-out volume and of all the particles evaporated from primary hot fragments including the (undetected) neutrons. Consequently some working hypotheses are used, constrained by specific experimental results (see for example [33]). Then, the experimental correlation between the kinetic energy per nucleon $E_k/A$ and the thermal excitation energy per nucleon $E^*/A$ of the considered system can be obtained event by event as well as the variance of the kinetic energy $\sigma^2_k$. Note that $E_k$ is calculated by subtracting the potential part $E_{pot}$ from the thermal excitation energy $E^*$ and consequently kinetic energy fluctuations at freeze-out reflect the configurational energy fluctuations. In the microcanonical ensemble with total energy $E^*$ the total degeneracy factor is simply given by the folding product of the individual degeneracy factors $W_k = \exp(S_k(E_k))$ and $W_{pot} = \exp(S_{pot}(E_{pot}))$ where $S_k(S_{pot})$ is the entropy of the kinetic (potential) part. One can then define the micro system as well as for the two subsystems the microcanonical temperatures and the associated heat capacities $C_k$ and $C_{pot}$. If we consider now the kinetic energy distribution when the total energy is $E^*$ we get

$$P_{E^*}(E_k) = \exp(S_k(E_k) + S_{pot}(E^* - E_k)).$$

Then the most probable kinetic energy $E_k$ is defined by the equality of the partial microcanonical temperatures $T_k(E_k) = T_{pot}(E^* - E_k)$ and $E_k$ can be used as the microcanonical thermometer. An estimator of the microcanonical temperature of the system can be obtained by inverting the kinetic equation of state:

$$< E_k > = \langle a_i \rangle T^2 + \frac{3}{2} (M - 1) T$$

The brackets $\langle \rangle$ indicate the average on events with the same $E^*$, $a_i$ is the level density parameter and $M$ the multiplicity at freeze-out. In this expression the same temperature is associated with both internal excitation and thermal motion of fragments. An estimate of the total microcanonical heat capacity is extracted using three equations.

$$C_k = \frac{\delta < E_k / A >}{\delta T},$$

is obtained by taking the derivative of $< E_k / A >$ with respect to $T$.

Using a Gaussian approximation for $P_{E^*}(E_k)$ the kinetic energy variance can be calculated as

$$Au^2 \approx T^2 \frac{C_k C_{pot}}{C_k + C_{pot}};$$

Eq. (4) can be inverted to extract, from the observed fluctuations, an estimate of the microcanonical heat capacity:

$$\langle \frac{C}{A} \rangle_{micro} \approx C_k + C_{pot} \approx \frac{C^2 + Ap^2}{C_k + Ap^2}.$$ (5)

From Eq. (5) we see that the specific microcanonical heat capacity $(C/A)_{micro}$ becomes negative if the normalized kinetic energy fluctuations $Au^2 / T^2$ overcome $C_k$. It is interesting to note that the constraint of energy conservation leads in the phase transition region to larger fluctuations than in the canonical case where the total energy is free to fluctuate. This is because the kinetic energy part is forced to share the total available energy with the potential part: when the potential part presents a negative heat capacity the jump from “liquid” to “gas” induces strong fluctuations in the energy partitioning.

Direct formulae have been proposed in Ref. [31] to calculate heat capacity but never used to extract information from data. They are derived within the microcanonical ensemble by considering fragments interacting only by Coulomb and excluded volume, which corresponds to the freeze-out configuration. Within this ensemble, the statistical weight of a configuration $c$, defined by the mass, charge and internal excitation energy of each of the constituting $M_c$ fragments, can be written [34] as

$$W_c(A, Z, E^*, V) = \frac{1}{M_c!} \chi^{M_c} \prod_{n=1}^{M_c} \left( \frac{\rho_0(\epsilon_n)}{h^3} (m A_n)^{3/2} \right) \times \frac{2\pi}{(3/2)(M_c - d) - 1} \frac{1}{(2\pi K)^{(3/2)(M_c - d) - 1}}$$

where $A$, $Z$, $E^*$ and $V$ are respectively the mass number, the atomic number, the thermal excitation energy and the freeze-out volume of the system. $E^*$ is used up in fragment formation, fragment internal excitation, fragment-Coulomb interaction and thermal kinetic energy $K$. $I$ is the inertial tensor of the system whereas $\chi^{M_c}$ stands for the free volume or, equivalently, accounts for inter-fragment interaction in the hard-core idealization. $\rho_0(\epsilon_n)$ represents the fragment level density (see [35] for the complete expression). $d$ takes respectively values 0, 1 and 2 for energy, energy plus linear momentum and energy plus linear and angular momenta conservations. In the following $d$ will be fixed to 1 to ensure coherence.
with simulations which are used. Taking into account that 
\( S = \ln Z = \ln \sum W_c \), the microcanonical temperature is 
deduced from its statistical definition \[34\]:

\[
T = \left( \frac{\partial S}{\partial E^*} \right)^{-1} = \left( \frac{1}{\sum_c W_c} \sum_c W_c (3/2M_c - 5/2)/K \right)^{-1} \quad (7)
\]

\[
= (3/2M_c - 5/2)/K \quad \text{or alternatively} \quad (3/2M_c - 5/2)/K^2 \quad (8)
\]

The notation \( \langle \rangle \) refers to the average over the ensemble 
states. The heat capacity of the system, \( C \), is related to the 
second derivative of the entropy by the equation \( \partial^2 S/\partial E^*^2 = -1/CT^2 \). 
Thus, one can evaluate the second derivative of the 
system entropy versus \( E \) (Eq. (7)) or alternatively 
the heat capacity \( C \) (Eq. (8))

\[
\frac{\partial^2 S}{\partial E^*^2} = \frac{(3/2M_c - 5/2)(3/2M_c - 7/2)}{K^2} - \frac{(3/2M_c - 5/2)}{K^2} \quad (9)
\]

\[
C = \left( 1 - T^2 \left( \frac{3/2M_c - 5/2}{3/2M_c - 7/2} \right) \right) \quad (10)
\]

These two quantities only depend on two parameters 
\( M_c \) and \( K \) which must be estimated at freeze-out.

In Refs. \[36,30\] we presented simulations which correctly 
reproduce most experimental observables and estimate 
freeze-out properties in a fully consistent way. They 
concern hot nuclei which undergo multifragmentation 
formed in central collisions and selected by event shape 
sorting (quasifused systems, QF, from \[28\]Xe+GeSn, 32-50 AMeV) \[37\]. We will use the event by event properties 
at freeze-out which were inferred to calculate the required 
quantities \( M_c \) and \( K \).

Experimental data were collected with the 4\pi multiplyector INDRA described in detail in Ref. \[38,39\]. Accurate 
particle and fragment identifications were achieved 
and the energy of the detected products was measured 
with an accuracy of 4\%. Further details can be found in 
Ref. \[40,41,42\].

The method for reconstructing freeze-out properties 
from simulations \[38,39\] requires data with a very high degree 
of completeness, (measured fraction of the available 
charge \( \geq 93\% \) of the total charge of the system), crucial 
for a good estimate of Coulomb energy. QF sources are 
reconstructed, event by event, from all the fragments 
and twice the charged particles emitted in the range \( 60-120^\circ \) 
in the relation centre of mass, in order to exclude the 
-major part of pre-equilibrium emission \[38,37\]: with such a 
prescription only particles with isotropic angular distributions 
and constant average kinetic energies are considered. 
In simulations, dressed excited fragments and particles 
at freeze-out are described by spheres at normal density. 
Then the excited fragments subsequently deexcite while 
flaying apart. All the available asymptotic experimental 
information (charged particle spectra, average and standard 
deviation of fragment velocity spectra and calorimetry) is 
used to constrain the four free parameters of simulations 
to recover the data at each incident energy: the percentage 
of measured particles which were evaporated from primary 
fragments, the collective radial energy, a minimum distance 
between the surfaces of products at freeze-out and a 
limiting temperature for excited fragments. All the details 
of simulations can be found in Ref. \[36,30\]. The limiting 
temperature, related to the vanishing of level density for 
fragments \[28\], was mandatory to reproduce the observed 
widths of fragment velocity spectra. With a single 
temperature (internal and kinetic temperatures equal) the sum 
of Coulomb repulsion, collective energy, thermal kinetic 
energy directed at random and spreading due to fragment 
decays accounts only for about 60-70\% of those widths.

By introducing a limiting temperature, which corresponds to 
intrinsic temperatures for fragments in the range 4-7 
MeV (see figure 1 of \[19\]), the thermal kinetic energy increases, 
due to energy conservation, thus producing the missing percentage for the widths of final velocity 
distributions. The agreement between experimental and 
simulated velocity spectra for fragments, for the different beam 
energies, is quite remarkable (see figure 3 of \[30\]). Relative 
velocities between fragment pairs were also compared 
through reduced relative velocity correlation functions \[43,45\] 
(see figure 4 of \[30\]). Again a good agreement is obtained 
between experimental data and simulations, which indicates that the retained method (freeze-out topology 
built up at random) and the deduced parameters are 
sufficiently relevant to correctly describe the freeze-out con-
figurations, including volumes.

For hot nuclei produced in central collisions the total 
excitation energy, \( E_{\text{tot}} \), differs from the thermal one due 
to the presence of a radial collective expansion energy, \( E_R \), 
and \( E_{\text{tot}} = E^* + E_R \). To derive event by event freeze-
out \( E_{\text{tot}} \), \( E^* \), the multiplicity, \( M_c \) (\( M^* \)) and the total kinetic energy, \( K \), of the multifragmenting hot nuclei, 
the following equations are used.

The limiting temperature for the fragments, \( T_{\text{lim}} \), 
is introduced according to the formalism of \[28\], which corre-
sponds to the following definition for the intrinsic 
temperature of the fragments, \( T_{\text{frag}} \):

\[
\frac{1}{T_{\text{frag}}} = \frac{3}{2 < K^f >} + \frac{1}{T_{\text{lim}}} \quad (11)
\]

where \( < K^f > \) is the average kinetic energy of fragments 
and particles at freeze-out. The equation used for 
calorimetry and to derive the sharing between internal ex-
citation energy and kinetic energy on the event by event 

\[
< K^f > = \sum_{k=1}^{M_f} K^f_{\text{cp}} + \Delta B_{\text{cp}} + M_f^* < K^f > + M_n^* \theta_{\text{frag}} + \Delta B_n
\]

\[
= (M^* - 1) < K^f > + \sum_{k=1}^{M_f} \theta_{\text{frag}}^2 + \Delta B_{\text{fo}} + V_{\text{Coul}} + E_R
\]

\[
= E_{\text{tot}} + \Delta B_{\text{hn}}, \quad (12)
\]

where \( \theta_{\text{frag}} \) is equal to the temperature \( T_{\text{frag}} \) in an 
ensemble average. \( \Delta B, \, \text{hn}, \, K, \, M, \, \text{cp}, \, n, \) and \( f_0 \) stand
respectively for mass excess, hot nucleus, kinetic energy, multiplicities, charged products, neutrons and freeze-out.

$$E_R = \sum_{k=1}^{M^f} \left( \frac{A_k}{r_0} \right)^2 A_k E_0$$

is the radial collective expansion energy ($R_0$ is the rms of fragment and particle distances to centre at the freeze-out volume, $E_0$ the radial expansion energy at $R_0$ and $r$ is the distance of the considered particle/fragment of mass $A$ from the centre of the fragmented source). $V_{\text{Coul}}$ is the Coulomb energy of the configuration for freeze-out previously determined. The chosen level density parameter $\alpha_k = \frac{\pi}{(3N_0)}^{3/2} A_k^{1/3}$, where $A_k$ is the mass of the $k^{th}$ fragment, well corresponds to that expected for a typical primary fragment [14]. By introducing Eq. (11) in Eq. (12) we obtain a third degree equation in $< K >$, from which we can deduce the energy sharing between the internal excitation energy of the fragments and the total thermal kinetic energy at freeze-out $K = (M^f - 1) < K >$, is shared at random between all the particles and fragments at freeze-out under constraints of conservation laws. Information on initial angular momenta of multifragmenting QF hot nuclei are unknown, which explains why $d = 1$ is used for Eqs. [9] and [10]. Systematic error on $E^*$ was estimated around 1 AMeV.

Values of heat capacity and second derivative of the entropy versus thermal excitation energy $E^*$ have been calculated respectively from Eqs. (11) and (9) for QF hot nuclei with $Z$ restricted to the range 80-100 to suppress tails of the distributions at different beam energies. The average over the ensemble states have been assimilated to an average over “event ensembles” sorted into $E^*$ bins. A binning of 0.5 AMeV was chosen to have a sufficient number of events in each bin in order to reduce statistical errors. Figure 1 shows the results. Error bars correspond to systematic plus statistical errors; systematic errors were evaluated by varying the free parameters of simulations within their limits defined by a $\chi^2$ procedure [22]. The left part of the figure shows the results for the direct calculation of $C/A$. Negative heat capacity is observed on a rather large thermal excitation energy range and the second diverging region is more visible than the first one. As the second derivative of the entropy is a very small quantity (from $\sim 2.10^{-4}$ to $\sim 3.10^{-6}$), we have kept the presentation made in [24] i.e. $A^2 \partial^2 S/\partial E^*2$ for fig. 1, right part; $A$ is replaced by the average mass, $< A >$, on the considered $E^*$ bin. This quantity, which has small error bars in regions with values close to zero, better defines the $E^*$ domain of negative heat capacity. Positive values are measured in the range 6.0 - 10.0 AMeV. The related microcanonical temperatures calculated with Eq. (8) are displayed in fig. 2. Considering the error bars, they are rather constant around 17 - 18 MeV in the $E^*$ range where negative heat capacity is observed. With the large multiplicities observed in the present study, microcanonical temperatures are close to classical kinetic temperatures (see figure 3 of [19]). Last point, the statistical nature of the carefully selected event samples, i.e. the independence of the entrance channel, was verified and fig. 3 shows the main contributions of the different bombarding energies to the signal for each $E^*$ bin.

Negative heat capacity is thus confirmed for systems with $A$ around 200 without any constraint, which demonstrates once again the robustness of this signal and supports the predictions made in [18] i.e. observation at constant pressure or at constant average volume. It is important to stress here that negative heat capacity cannot be observed at constant volume. For the liquid-gas transition in hot nuclei the volume is not fixed but multiplicity and partition-dependent, which means that on the theoretical side one is forced to consider a statistical ensemble for which the volume can fluctuate from event to event around an average value [8].

As compared to previous estimates of heat capacity [21][22], there is a significant difference in the $E^*$ range for negative values: 6.0 ± 1.0 - 10.0 ± 1.0 AMeV in the present study and < 4.0 ± 0.1 - 6.0 ± 1.0 AMeV in the previous one. For fusionsion hot nuclei selection the same shape event sorting was used. The degree of completeness was different (93% here to be compared to 80% before) but it does not affect significantly the thermal excitation energy per nucleon. The method to reconstruct freeze-out properties was also different. But the main difference seems to be related to the average freeze-out volume. In previous estimates the average freeze-out volume was kept constant at 3 times the volume at normal density ($V_0$) over the whole thermal excitation energy range whereas in the present study it varies from 3.9 to 5.9 $V_0$ (see fig. 1). A direct consequence in the approximate method is an increase of Coulomb energy and consequently a decrease and a distortion with excitation energy of $< E_k >$ obtained by subtracting the potential part (Coulomb energy + total mass excess) to the thermal energy. To verify this reasoning the approximate method has been used. From our simulation, for each $E^*$ bin, we have calculated $< E_k >$ (see Eq. (2)) and derived an apparent single temperature, $T_S$, needed to build the normalized kinetic energy fluctuations, $\sigma_{E^2}/T_S^2$, to be compared to $C_k$ (see Eqs. (3) and (5)). Figure 5 shows that heat capacity becomes negative in the $E^*$ range 5.5 ± 1.0 - 9.0 ± 1.0 AMeV, i.e. when $\sigma_{E^2}/T_S^2$ overcomes $C_k$. This clearly confirms that the main difference, as compared to previous estimates, comes from different average freeze-out volumes. We also note a small decrease of the $E^*$ domain as compared to the present study, which possibly comes from the previous method. Last information, the apparent single temperature, $T_S$, exhibits a monotonic behaviour, varying from 5.2 to 9.7 MeV over the whole $E^*$ range and values increasing from 6.5 to 8 MeV on the domain of negative heat capacity.

In conclusion, heat capacity measurements have been revisited without approximation and by ignoring the hypothesis of a single temperature associated with both internal excitation and thermal motion. For those measurements microcanonical formulæ and data reconstructed at freeze-out with the help of a simulation have been used. Negative heat capacity has been confirmed for hot nuclei with $A$ around 200 in the coexistence region of the phase transition.
Fig. 1. Heat capacity (left) and second derivative of the entropy (right) versus thermal excitation energy $E^*$. Error bars correspond to systematic plus statistical errors (see text).

Fig. 2. Microcanonical temperatures calculated with Eq. (8). Error bars include systematic plus statistical errors.

Fig. 3. Similar to the right part of fig. 1. The main contributions of the different bombarding energies are shown.

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\begin{figure}
\centering
\includegraphics[width=0.8\textwidth]{fig4}
\caption{Evolution of the average freeze-out volume of hot nuclei, $V_{fo}$, with thermal excitation energy. Freeze-out volumes are normalized to volumes at saturation density, $V_0$, and error bars correspond to standard deviations of $V_{fo}$ distributions.}
\end{figure}

\begin{figure}
\centering
\includegraphics[width=0.8\textwidth]{fig5}
\caption{Normalized kinetic energy fluctuations (filled circles) and estimated $C_k$ values (open squares) related to the approximate method (see text). Error bars correspond to systematic plus statistical errors.}
\end{figure}

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