Experimental proposal for accurate determination of the phase relaxation time and testing a formation of thermalized non-equilibrated matter in highly excited quantum many-body systems

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We estimate how accurate the phase relaxation time of quantum many-body systems can be determined from data on forward peaking of evaporating protons from a compound nucleus. The angular range and accuracy of the data needed for a reliable determination of the phase relaxation time are evaluated. The general method is applied to analyze the inelastic scattering of 18 MeV protons from Pt for which previously measured double differential cross sections for two angles in the evaporating domain of the spectra show a strong forward peaking. A new experiment for an improved determination of the phase relaxation time is proposed. The experiment is also highly desirable for an accurate test of a formation of thermalized non-equilibrated matter in quantum many-body systems.

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For some years one of us has emphasized that phase relaxation in a many-body system can be considerably longer than energy relaxation among independent particle states ¹²³. Maybe the easiest experimental access to the problem can be found in low energy (10-80 MeV) nucleon-nucleus scattering processes, where some of the oldest data have been available for half a century ². It turns out ² that the phase relaxation time, \( \tau_{\text{ph}} = \hbar/\beta \), is considerably longer than the energy relaxation time, \( \tau_{\text{erg}} = \hbar/\Gamma_{\text{spr}} \), obtained from standard estimates of the spreading width \( \Gamma_{\text{spr}} \) of independent particle states.

The significance of this problem results from the fact that an existence of long-living phase relations is of fundamental importance in the study of relaxation phenomena in nuclear, atomic, molecular and mesoscopic many-body systems, and for many-qubit quantum computation. In particular, if a phase relaxation time, which characterizes the lifetime of the “phase memory”, is longer than the energy relaxation time, this effect could extend the time for quantum computing ²³ beyond the quantum–chaos border ³.

The experimental result ⁴ showing that the intensity of the compound-nucleus evaporation for \( \theta = 60^\circ \) exceeds that for \( \theta = 150^\circ \) by a factor of \( \approx 6 \) indicates that \( \beta \sim \Gamma^{\dagger} = \hbar/\tau^{\dagger} \), where \( \Gamma^{\dagger} \) is the compound nucleus decay width and \( \tau^{\dagger} \) is the compound nucleus life-time. However, since the evaporation spectra ⁵ were measured for two angles only, these data allow only a rough estimate, \( \beta/\Gamma_{\text{spr}} \sim 10^{-5} \). The purpose of this note is to determine whether more detailed experiments of the same type could improve this estimate.

Our analysis for the improvement of the experimental determination of the phase relaxation time from nuclear evaporation data will be quite general and can readily be applied to any low energy nucleon-nucleus scattering showing forward peaking in the evaporation domain of the spectra. Yet, to be specific, we shall show numbers and graphs for inelastic scattering of 18 MeV protons from Pt, i.e. for improvement of one of the oldest experiments ⁴. The questions we shall answer are: What is the best realistic accuracy of determination of the phase relaxation time from the data on forward peaking of evaporating protons from the compound nucleus? What is the angular range and accuracy of data needed for a reliable determination of \( \tau_{\text{ph}} \)?

An advantage of the analysis is that it relies on relative values of the double differential cross sections which are usually determined experimentally with better accuracy than absolute cross sections.

The proposed experiment is also desirable for an accurate test of the formation of thermalized non-equilibrated matter in compound processes. A manifestation of such a new form of matter, introduced in Refs. ²³, would be equal slopes, i.e. nuclear “temperatures” ⁶, of the properly scaled ²³ proton evaporation spectra for forward and backward angles. Due to the insufficient statistics the data ⁴ indicate only approximate equality of the slopes for the forward and backward angles with about 20 percent uncertainty. We again point out that the proposed test only requires relative values of the proton emission intensities.

The evolution of a nuclear reaction is usually considered to proceed via a series of two-body nucleon-nucleon collisions, which successively form states of increasing complexity. On each stage of the reaction a distinction is made between continuum states and quasibound states. Emissions from the continuum states is related to multistep direct reactions ³¹¹, and decay of the quasibound states originates multistep compound processes ³¹. The compound nucleus is formed at the last stage corresponding to the most complex configuration of the chain of quasibound states. The multistep direct reactions originate from the decay of the simplest configurations of the chain resulting in forward-peaked angular
distributions. In contrast, the multistep pre-compound and compound reactions are conventionally assumed to give rise to angular distributions symmetric about 90 degrees.

We use the exciton model to evaluate the relative contributions of multistep direct, multistep pre-compound and compound nucleus processes for the p+Pt (E_p = 18 MeV) inelastic scattering for the proton outgoing energy of 7 and 6 MeV. Fitting the entire energy range for forward angles we found that, for the proton outgoing energy of 7 MeV, the compound nucleus cross section constitutes 90%, while multistep direct and multistep pre-compound are about 5% each. For the proton outgoing energy of 6 MeV, the compound nucleus cross section is about 98%. Therefore we observe that even though the low energy 6-7 MeV outgoing proton spectrum is overwhelmingly dominated by compound reactions, the angular distribution is strongly forward peaked. Clearly a description of the decay of such thermalized but non-equilibrated matter requires a major modification of conventional theory of compound nucleus (see e.g. Ref. [8]) originally formulated by Bohr, Bethe, Weisskopf, Wigner, Dyson and others. The basic assumption of the conventional theory is that thermalization of the compound nucleus guarantees a complete loss of memory of initial phase relations. A modification of this conventional picture of the compound nucleus was proposed by one of us in Refs. [2, 3]. The key element in the description of asymmetry of angular distributions around 90° c.m. for evaporating particles is total spin off-diagonal correlation between compound nucleus partial width amplitudes. Such a correlation is neglected in a conventional picture of compound nucleus. Following [2, 3] we have

\[
\frac{\gamma_{J_1 \pi_1 a_1} J_1 \pi_1 a_1}{(\gamma_{J_1 \pi_1 a_1})^2} \frac{\gamma_{J_2 \pi_2 a_2}}{(\gamma_{J_2 \pi_2 a_2})^2} \left( \frac{\beta |J_1 - J_2|}{|E_{J_1 \pi_1} - E_{J_2 \pi_2}|^2 + \beta^2 (|J_1 - J_2|^2)} \right) ^{1/2} \]

where overlines denote ensemble averaging. Here J_1 ≠ J_2 are the compound nucleus total spin values, π_1, π_2 are parity values, E_μ^J are resonance energies with μ being running indices, and D is average level spacing of the compound nucleus. The a(μ) indices specify the orbital momenta l_a(μ) , the channel spins J_a(μ) , and the microstates a(μ) of the target nucleus and residual nucleus, respectively. Accordingly, a_1 = a_2 denote the ground state of the target, and b_1 = b_2 specify the microstates of the residual nucleus. The above correlation between the partial width amplitudes leads to a correlation between fluctuating compound nucleus S-matrix elements carrying different total spin values:

\[
\langle S_{a_1 b_1}^{J_1 \pi} (E) S_{a_2 b_2}^{J_2 \pi} (E) \rangle = \frac{\langle |S_{a_1 b_1}^{J_1 \pi} (E)|^2 \rangle \langle |S_{a_2 b_2}^{J_2 \pi} (E)|^2 \rangle ^{1/2}}{1 + |J_1 - J_2| \beta \Gamma^+}.
\]

Here S_{a b}^{J}(E) are compound nucleus S-matrix elements with total spin J and the brackets ⟨...⟩ denote the energy E averaging. For finite values of β/Γ^+, nonvanishing of the spin off-diagonal correlations in Eq. (2) reflects nonvanishing of the interference between resonance levels with different total spins upon the energy averaging.

For the correlation between S-matrix elements carrying the same total spin and parity values and the same microstates a_1 = a_2 and b_1 = b_2 but different orbital momenta and/or channel spins we have [2, 3]

\[
\langle S_{a_1 b_1}^{J_1 \pi} (E) S_{a_2 b_2}^{J_2 \pi} (E) \rangle = \frac{\langle |S_{a_1 b_1}^{J_1 \pi} (E)|^2 \rangle \langle |S_{a_2 b_2}^{J_2 \pi} (E)|^2 \rangle ^{1/2}}{1 + |J_1 - J_2| \beta \Gamma^+}.
\]

The above equation results from a strong correlation between partial width amplitudes γ_{J_π a} and γ_{J_π a} with a_1 = a_2 and b_1 = b_2 but l_1 ≠ l_2, l_1 ≠ l_2, j_1 ≠ j_2, j_1 ≠ j_2. Such a correlation is referred to [2, 3] as the continuum correlation. Note that such a strong correlation between reduced width amplitudes corresponding to the same total spin and parity values but different orbital momenta was experimentally revealed for a number of compound nuclei in the regime of isolated resonances [13].

For β >> Γ^+, the spin off-diagonal correlations in Eq. (2) result in the angular distributions symmetric around 90° c.m. recovering a conventional picture of compound nucleus. However, if β ≤ Γ^+, i.e. the phase relaxation time τ_ph is comparable or longer than the average lifetime of the compound nucleus, this allows us to describe a strong asymmetry of the angular distributions around 90° c.m. of the evaporating yield.

For the treatment presented here we follow Ref. [16]. We neglect the intrinsic spins of the scattering partners in the entrance channel and proton spin in the exit channel. Since evaporated protons carrying orbital momenta I_l > 1 are significantly sub-barrier due to both the centrifugal and Coulomb barriers, we take T_l ≥ 1 = 0, where T_l are the transmission coefficients for the inverse process of capture of the proton by the residual nucleus. Then the double-differential cross section in the evaporation domain of the spectra has the form [16]

\[
\frac{d^2 \sigma}{d\Omega d\epsilon} = \frac{1}{4\pi} \sigma(\epsilon) \sum_{L=0} A_L P_L(\cos \theta).
\]

Here, σ(ε) is the angle-integrated cross section for the evaporation of a proton with the energy ε, and P_L(\cos θ) denote the Legendre polynomials of order L. The coefficients of the angular decomposition are given by

\[
A_{L=0} = 1,
A_{L=1} = \frac{1}{1 + 3 \frac{\beta}{\Gamma^+}},
A_{L=2} = \frac{3 \frac{\beta}{\Gamma^+}}{1 + 2 \frac{\beta}{\Gamma^+}}.
\]
Indeed, in Fig. 1 any of the seven dots as well as any ambiguous determination of the phase relaxation time. In any case we have an underdetermined equation

\[ \frac{\beta}{\Gamma^*} \]

which are in accordance with the experimental data of Ref. [2]. The dots marked with numbers on the solid line correspond to:

1: (0.05, 0.72); 2: (0.1, 0.63); 3: (0.2, 0.52); 4: (0.3, 0.39); 5: (0.367, 0.231); 6: (0.3, 0.13); 7: (0, 0.05).

One can see that if this phase memory time \( \tau_{ph} = \hbar / \beta \) is about or longer than the average life-time \( \tau^1 = \hbar / \Gamma^* \) of the compound nucleus the evaporation yield is emitted asymmetrically about 90° c.m., i.e. the memory about the direction of the initial beam remains. However, if the phase memory time is much shorter than the average lifetime of the compound nucleus then the spin off-diagonal correlations vanish, the memory on the direction of the initial beam is lost, and an isotropic angular distribution around 90° c.m. is obtained.

We now turn to the experimental data available from Ref. [4] in order to estimate bounds for the possible values of \( \beta \). Since in Ref. [4] only relative yields for two scattering angles are reported we focus on the analysis of the shape of the angular distributions \( I(\theta) = K d^2 \sigma / d \Omega d \varepsilon \) without paying attention to the angle and the energy independent prefactor \( K \).

For \( \varepsilon = 7 \) MeV, we find from the experimental data a ratio \( I(\theta = 60^\circ) / I(\theta = 150^\circ) \approx 6 \). Thus, we consider the relation

\[ R_I = \frac{I(\theta = 60^\circ)}{I(\theta = 150^\circ)} = 6, \tag{6} \]

where \( I(\theta) \) is given by Eqs. (4) and (5) and depends on the two parameters \( \beta / \Gamma^* \) and \( T_1 / T_0 \). Therefore, Eq. (6) implicitly defines a curve in the parameter space of \( \beta / \Gamma^* \) and \( T_1 / T_0 \), which is plotted in Fig. 1. We can either solve Eq. (6) for \( \beta / \Gamma^* \) as a function of \( T_1 / T_0 \) or vice versa. In any case we have an underdetermined equation demonstrating that measurements of the evaporation yields for two angles only does not allow an unambiguous determination of the phase relaxation time. Indeed, in Fig. 1 any of the seven dots as well as any point on the solid line corresponds to \( R_I = 6 \). Yet, moving along the solid line, \( \beta / \Gamma^* \) changes from 0 to 0.367. Since the value of \( T_1 / T_0 \) can be obtained from model calculations this clearly demonstrates that measurement of the evaporation yields for two angles only permits to accurately determine only the upper experimental limit, \( \beta / \Gamma^* = 0.367 \). Estimating the total decay width from the systematics in Fig. 7 of Ref. [6] we obtain \( \Gamma^* = 0.02 \) keV and, therefore, \( \beta \leq 7 \) eV. Taking into account that \( \Gamma_{spr} \approx 1.5 \) MeV [2] we observe that the phase relaxation time is at least five orders of magnitude longer than the energy relaxation time. Yet, it is still about fifteen orders of magnitude shorter than the Heisenberg time, \( \hbar / D \), where the average level spacing of the compound nucleus, \( D \sim 10^{-20} \) MeV, has been estimated using the Fermi-gas model [18].

Due to the uncertainties of the data reported in Ref. [4], we also included curves of \( T_1 / T_0 \) and \( \beta / \Gamma^* \) in Fig. 1 resulting from Eq. (6) but with its right hand side being 5 and 7.

For a more accurate determination of the \( \beta \), rather than just an estimation of its upper limit, one has to measure the evaporation yields for more than two angles. Therefore, we analyze the sensitivity of the shapes of the angular distributions for different sets of \( \beta / \Gamma^* \) and \( T_1 / T_0 \) belonging to the manifold obtained from the solution of Eq. (6).

In Fig. 2 we present the angular distributions for the seven sets of \( \beta / \Gamma^* \) and \( T_1 / T_0 \) marked with dots and corresponding numbers in Fig. 1. The angular distributions are normalized in such a way that \( I(\theta = 60^\circ) = 6 \) and \( I(\theta = 150^\circ) = 1 \) for each of these curves. One does observe that the angular distributions change appreciably with the change of \( \beta / \Gamma^* \) and \( T_1 / T_0 \) values. To quantify this sensitivity we determine \( \theta_{opt} \) for which the ratio \( I(\theta_{opt}) / I(\theta = 170^\circ) \) is most sensitive to different \( \beta / \Gamma^* \) and \( T_1 / T_0 \) values. We find that, for \( R_I = 6, 5, 7 \), \( \theta_{opt} = 118^\circ, 116.3^\circ, 119.4^\circ \), respectively.

The dependence of \( I(\theta_{opt}) / I(\theta = 170^\circ) \) on \( \beta / \Gamma^* \) is pre-
FIG. 3: The dependencies of $I(\theta_{\text{opt}})/I(\theta = 170^\circ)$ on $\beta/\Gamma^\uparrow$. The solid line is obtained for $R_t = 6$ (see Eq. (6)) the dashed line for $R_t = 5$, and dotted line $R_t = 7$. Dots marked with numbers on the the solid line correspond to the $\beta/\Gamma^\uparrow$ and $T_1/T_0$ values specified in Fig. 1.

sented in Fig. 3 with the solid line corresponding to the ratio $R_t = 6$. We find that for values of $I(\theta_{\text{opt}})/I(\theta = 170^\circ) \leq 2.7 - 2.8$ the ratio $I(\theta_{\text{opt}})/I(\theta = 170^\circ)$ is rather sensitive to the $\beta/\Gamma^\uparrow$, as can be seen in the lower part of Fig. 3. If the experimental value lies in this range, an accuracy of $I(\theta_{\text{opt}})/I(\theta = 170^\circ)$ of about 5% allows a determination of $\beta/\Gamma^\uparrow$ with a minimal uncertainty of about 10%. For a too low experimental value of $I(\theta_{\text{opt}})/I(\theta = 170^\circ)$, say less than 0.84, one can determine only the upper limit of $\beta/\Gamma^\uparrow \leq 0.02$.

On the other hand, in the upper part of the curve, for $I(\theta_{\text{opt}})/I(\theta = 170^\circ) \geq 2.7 - 2.8$, the ratio $I(\theta_{\text{opt}})/I(\theta = 170^\circ)$ shows a rather weak dependence on $\beta/\Gamma^\uparrow$. In particular, for an assumed accuracy of the data of about 5%, the value of $\beta/\Gamma^\uparrow$ can be determined with an uncertainty of only $\sim 50 - 100\%$. If the experimental value of $I(\theta_{\text{opt}})/I(\theta = 170^\circ)$ is larger than 2.85 one can determine only the upper limit, $\beta/\Gamma^\uparrow \leq 0.2$.

Similar conclusions about the sensitivity of $I(\theta_{\text{opt}})/I(\theta = 170^\circ)$ to $\beta/\Gamma^\uparrow$ can be drawn from Fig. 3 for $R_t = 5$ and 7. According to Fig. 2 an improvement of the accuracy might be found if the data for forward scattering ($\theta < 30^\circ$) would also be available.

It should be noted that a manifestation of a formation of the thermalized non-equilibrated matter has been also identified from a strong asymmetry around 90° c.m. of evaporating protons in the Bi$(\gamma,p)$ photonuclear reaction, see [8, 9] and references therein. Other examples of a strong forward peaking of evaporating protons are found, e.g., in Bi$(p,p')$ and Bi$(n,p)$ processes with the 62 MeV energy of initial beam, see [9] and references therein. Even though the later examples do demonstrate a formation of the thermalized non-equilibrated matter in compound processes, a determination of the phase relaxation time from these data is not unambiguous since, for such high energy of the initial beam, there is a high probability for a second and third chance proton evaporation.

The possibility that in highly excited many-body systems the phase relaxation can be much longer than the energy relaxation may have significant implications for quantum computing [10, 11] as well as, e.g., time-delayed “statistical” ionization of many-electron quantum dots and atomic clusters (see, e.g., [12] and references therein). A possible presence of the effect of anomalously slow phase relaxation [20] in chemical reactions (see [20] and references therein) would require a modification of the statistical theories - phase space and transition state theories (see, e.g., [21] and references therein). Yet, the nuclear data indicating an existence of anomalously slow phase relaxation, which is much slower than the energy relaxation, are largely unrecognized by nuclear physicists and unknown outside the nuclear physics community. In many fields, including statistical physics, the notion “thermalization” or “energy equilibration” is considered to be equivalent to the notion “statistical equilibrium”. This note is a step towards changing this undesirable situation.

In conclusion we have proposed a general method to estimate the accuracy of the determination of the phase relaxation time from data on forward peaking of evaporating protons from compound nucleus. The angular range and accuracy of the data needed for a reliable determination of the phase relaxation time have been evaluated. The general method has been applied to the analysis of inelastic scattering of 18 MeV protons from Pt for which previously measured double differential cross sections for two angles in the evaporating domain of the spectra show a strong forward peaking. We found that a new measurement of the angular distributions of evaporating protons in the Pt$(p,p')$ inelastic scattering for a wider angular range should permit an accurate determination of the phase relaxation time. The experiment is also highly desirable for an accurate test of the formation of thermalized non-equilibrated matter in compound processes. Our analysis for the improvement of the experimental determination of the phase relaxation time from nuclear evaporation data can readily be applied to any low energy nucleon-nucleus scattering showing forward peaking in the evaporation domain of the spectra.

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