The Multi-dimensional Langevin Approach to the Description of Fusion-fission Reactions

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The two stage approach to the description of fusion-fission reactions is suggested. On each stage (fusion or fission) the tree-dimensional Langevin equation for the variables describing the shape of nuclear system is solved. The results obtained on the first stage are used as the input data for description of fission dynamics. In this way it turned out possible to describe for the reaction $^{18}$O + $^{208}$Pb simultaneously both fusion and fission cross sections, the energy and mass distribution of fission fragments, the probability of the evaporation residue formation, the dependence of pre-fission neutron multiplicities on the fragment mass number. From the results of the computations it follows that the observed quantities of the fission process can be reproduced without formation of compound nucleus. At the same time the duration of the process is so large that events of quasi-fission and “true” fission of compound nucleus can not be distinguished.

1. Introduction

For many years the fusion and fission processes were studied separately. This was mainly due to the fact that the relatively light ions were used in fusion reaction and it was assumed that the ions are absorbed very fast by the target nucleus. With the increasing weight of the ions the picture changes considerably.1 Now, when the Uranium ions are used as the projectile the description of fission independently from the fusion stage is not justified.

One of the examples of such approach is the concept of dineural system (DNS).2-5 In this model it is assumed that after the touching point both target and projectile keep their “individuality” — spherical shape and shell structure. The entrance channel in this model is taken into account through by the probability of the formation of DNS which is calculated on the basis of optical model. This is the only memory about entrance channel. So the initial stage and the further evolution of the system is carried out within very different approaches.

More consistent are the approaches developed in References 6, 7. The parameters of the entrance channel (the dependence of the compound nucleus formation probability on the angular momentum)6 and the further evolution of the composite system6 are considered within the same approach. However whereas the DNS-approach starts from the contact of two nuclei in References 6,7 this moment is missing completely. Once the probability of compound nucleus is known, the evolution of the fissioning system starts in References 6,7 from the ground state. Thus the “pre-history” is ignored completely. On this way the considerable part of the information on systems evolution is lost.

In present work an attempt is undertaken to consider both the approaching phase and the fission within the same framework, namely by solving the Langevin equations for the variables which specify the shape of composite system. As an example we will consider the reaction $^{18}$O + $^{208}$Pb for which the detailed experimental information is available. Note also that both nuclei are spherical.

2. The Model

The description of the fusion-fission reactions will be carried out in two stages. Both stages are described by three-dimensional Langevin equation. On the first stage the approaching phase is considered. The information obtained on this step is used as the input data for describing the further evolution of the composite system. Unfortunately, at present there is no shape parametrization which would describe well both divided and compact shapes. Thus on both stages different parametrization will be used. On each stage the evolution of large number of collisions is considered. Each of them will be referred as ”trajectory” in the space of deformation parameters.

2.1. Equations of Motion. To describe the dynamics of the fusion-fission reaction we will use the Langevin equations. For the fusion stage we will solve the equations:8

$$\frac{dr}{dt} = \frac{p_r}{m},$$

$$\frac{dp_r}{dt} = -\frac{\partial V(r, \alpha)}{\partial r} - \frac{L^2}{mr^3} - K^p \frac{p_r}{m} - \sum_j K^j \pi_j D_j + \xi,$$

$$\frac{d\alpha}{dt} = \frac{\pi}{D},$$

$$\frac{d\pi}{dt} = -\frac{\partial V(r, \alpha)}{\partial \alpha} - K^p \frac{p_r}{m} - \sum_j K^j \frac{\pi}{D_j} - C \alpha - \frac{1}{2} \xi^2. \tag{1}$$

Here $r$ is the distance between centers of mass of colliding nuclei, $p_r$ — conjugated momentum, $m$ — the reduced mass, $V(r, \alpha)$ — the interaction energy. The latter includes the Coulomb and nuclear parts. The nuclear part $V_{Coul}$ was calculated according to Gross and Kalinowski,6,9 the $K^p$ is the tensor of friction parameters. The angular momentum $L$ is expressed in units $\hbar$. The second couple of eq 1 describes the shape evolution of colliding nuclei. The quantities $\alpha$ and $\pi$ are the collective variables describing the shape of nucleus and conjugated momentum correspondingly. The index $i$ attains the values “target” or “projectile”, the $D_i$ and $C_i$ are the friction and mass parameters with respect to $\alpha_i$. Both $D_i$ and $C_i$ were defined within liquid drop model. The quantity $\xi$ describes the random force acting upon the system and leading to the fluctuations. The magnitude of the fluctuations depends on the excitation energy of the system, so at initial moment it is equal to zero. During the approach the part of the kinetic energy turns into heat due to the friction in the system. Consequently the magnitude of fluctuations gets different from zero and increases with time.

The system of eq 1 does not include the equation for angular momentum $L$ of the system. It was assumed to be a constant for each trajectory.

For the mono system the Langevin equations are of the form10

$$\frac{dq}{dt} = M_{ij} p_j,$$
\[
\frac{dp_i}{dt} = -\frac{\partial V(q_i)}{\partial q_i} - \frac{1}{2} p_i^2 \frac{\partial \mu_{i\beta}}{\partial q_i} - \gamma_i \mathbf{M}_\beta p_i + \xi_i, \tag{2}
\]
where \(q_i\) are the collective variables fixing the shape of mono system and \(p_i\) are the conjugated momenta. The \(\gamma_i\) and \(\mu_{i\beta}\) are the friction and mass parameters. For the compact shapes these parameters were computed within the linear response approach.\(^\text{16,17}\) The explicit expressions and numerical results for \(\gamma_i\) and \(\mu_{i\beta}\) can be found in Reference \(^\text{18}\).

2.2. Collective Co-ordinates. On the approach phase the projectile and target get deformed both by Coulomb and nuclear interaction in order to reduce the interaction energy. So, one should account for the deformation of the projectile and target. In the present work we will take into account only the most important quadrupole deformation. I.e., the shape of nuclear surface is parametrized by

\[ R_i(x) = \lambda_i^{-1} R_0 \left( 1 + \alpha_i^2 P_2 (x) \right). \]

Here \(R_0\) is the radius of the sphere with equal volume, index \(i\) denote “target” or “projectile”, the parameter \(\alpha_i^2\) defines the deformation of the projectile or the target, \(\lambda_i\) is the normalization factor (defined from the volume conservation condition). It is assumed that on the approaching stage the symmetry axis of projectile and target coincide (Figure 1), i.e. the rotation of ions is not taken into account. This approximation seems quite reasonable, though it may lead to some reduction of the fusion cross section.\(^\text{10}\)

On the second stage, when the projectile and target formed a bound system, its shape is parametrized in terms of Cassini ovaloids,\(^\text{11}\) i.e. by the profile function \(p(z)\) which is given parametrically as

\[
p_i(z) = \frac{1}{\sqrt{2}} \left[ \left( R^2 + 2\alpha R^2 (2z^2 - 1) + s^2 \right)^{1/2} - R^2 (2z^2 - 1) - s \right]^{1/2},
\]

\[ z(x) = \frac{\text{sign}(x)}{\sqrt{2}} \left[ \left( R^2 + 2\alpha R^2 (2z^2 - 1) + s^2 \right)^{1/2} + R^2 (2z^2 - 1) + s \right]^{1/2},
\]

\[ R(x) = \lambda_i^{-1} R_0 \left( 1 + \sum_m \alpha_m P_m (x) \right),
\]

\[ s = \epsilon R_0^2. \tag{3}\]

Here \(s\) is the square of distance from the focus of Cassini ovals to the center of co-ordinate system. The surface of the (axially symmetric) nucleus is obtained by the rotation of the profile function \(p_i(z)\) around \(z\)-axis. The shape of the surface is fixed by the deformation parameters \(\alpha\) and \(\alpha_m\). Near the scission it is convenient to use instead of \(\epsilon\) as an independent variable the parameter \(\alpha\) which relates to \((\epsilon, \alpha_m)\) as

\[
\epsilon = \frac{1}{4} (\alpha - 1) \left[ \left( 1 + \sum_{m=1,4} \alpha_m \right)^2 + \left( 1 + \sum_{m=1,4} (-1)^m \alpha_m \right)^2 \right] + \frac{1}{2} \left( \alpha + 1 \right) \left[ 1 + \sum_{m=1,4} (-1)^m \alpha_m (2m - 1)!/(2^m m!) \right]^2 \cdot \tag{4}\]

The advantage of parameter \(\alpha\) is due to the fact that for a shape with vanishing neck \(\alpha = 1\) for any value of \(\alpha_m\). So, \(\alpha = 0\) corresponds to spherical shape and \(\alpha = 1\) correspond to the shape with zero neck.

When we switch from two separated ions to a bound system, we conserve the full elongation and the mass asymmetry of the system. Namely, the ratio of the volumes of left and right part of the system (which are divided the zero neck) are the same as the one for colliding ions. And the sum of radiuses of ions along the symmetry axis is equal to the elongation of the whole system. The parameters of the compact system were defined numerically for each trajectory.

In the present work we use three deformation parameters both for bound and separated shapes. In the first case these are the distance \(r\) between the centers of mass of target and projectile and the parameters \(\alpha\) of their (quadrupole) deformation. In the second case — the deformation parameters \(\alpha, \alpha_m\), and \(\alpha\) which fix the total elongation of the system, mass asymmetry, and thickness of the neck.

2.3. The Potential Energy. The potential energy includes the Coulomb, rotational, and nuclear parts. Below only the last part is shown. The nuclear interaction of two ions in the entrance channel was described by means of Gross-Kalinowski potential \(V_{\text{GK}}\),\(^\text{12}\)

\[ V_{\text{GK}} = \frac{1}{2} (V_{12} + V_{21}), \]

\[ V_{12} = \int V(\vec{r} - \vec{r}', \alpha) \rho_2 (\vec{r}', \alpha) d\vec{r}', \]

\[ V(\vec{r}, \alpha) = \frac{V_r}{1 + \exp \left( \frac{r - R_{\text{a}}(\alpha)}{a_r} \right)}, \]

\[ \rho_2(\vec{r}, \alpha) = \frac{\rho_0}{1 + \exp \left( \frac{r - R_{\text{a}}(\alpha)}{a_r} \right)}, \]

\[ R_{\text{a}}(\alpha, x) = 1.25 A_1^{1/3} (1 + \alpha P_2(x)) \text{ [fm]}, \]

\[ R_{\text{a}}(\alpha, x) = (1 + \alpha P_2(x))(1.12 A_1^{1/3} - 0.86 A_1^{1/3}) \text{ [fm]}, \]

with \(\rho_0 = 0.17 \text{ fm}^{-3}\), \(a_r = 0.54 \text{ fm}\). The parameters of the potential \(V_r\) and \(a_r\) were chosen according to Reference 14. In the calculations the mesh \(100 \times 41 \times 41\) in the deformation space was used. The \(r\)-dependence of the potential energy for \(L = 0\) is shown in Figure 1.

The potential energy for the compact shape was computed by the shell correction method\(^\text{20}\) as the sum of liquid-drop part and the shell correction. The damping of the shell correction with growing excitation energy was taken into account according to Reference 15.

Figure 2 shows the map of the potential energy in \(\alpha, \alpha_m\) co-ordinates (the parameter \(\alpha_m\) which defines the thickness of the neck is fixed equal to \(-0.18\)). One can see that with growing temperature the potential energy gets more smooth and more close to the liquid-drop deformation energy. The two fission valley merge into one. Thus, depending on the excitation energy, the mass distribution of the fission fragments could be symmetric or not.
The potential energy of the colliding ions is shown in Figure 1 by solid line. Of course, the potential energy would change due to the deformation of ions. The mean deformations of projectile (circles) and target (squares) are shown in Figure 1 too. The value of deformation is indicated on the right y-axis. One can see that far away from the barrier parameter of deformation is zero or very slightly positive. Near the barrier deformation gets negative, i.e. both projectile and target get oblate. This lead to the increase of Coulomb interaction. The smaller is distance between centers of mass the larger is the deformation. Contrary to the case of spherical nuclei, the touching of deformed surfaces practically can not be reached. The evolution of a trajectory in entrance channel was stopped when the distance between the surfaces of ions was smaller than half of the sum of the diffuseness parameters of projectile and target. The cross section obtained in this way was identified with the touching cross section. This values are shown by open triangles in Figure 4.

By solving eq 1 we obtain the information on the entrance channel: the probability of the touching, the distribution of the excitation energy at the touching point and deformations of the fragments at the touching point. These distributions are shown in Figure 3. The top panel shows the distribution of the intrinsic excitation energy at the touching point on the initial angular momentum for the initial energy $E_{lab} = 90$ MeV. One can see that for each angular momentum the excitation energy is distributed around some mean value. With growing angular momentum, the mean value of the excitation energy goes to zero due to the redistribution of the initial energy between intrinsic excitation and rotation.

The middle panel of Figure 3 shows the probability to reach the touching point as the function of initial angular momentum. The open squares correspond to the initial energy $90$ MeV, the filled triangles — to the energy $78$ MeV. In both cases, the probability decreases from its maximal value to zero quite smoothly due to the presence of random force in eq 1. The absolute value of the probability for these two cases differs very much from each other. In the second case, when $E_{lab} = 78$ MeV the probability to reach the touching point is very small even for head-on collisions. In this case $E_{lab}$ energy is almost equal to the fusion barrier.

The bottom panel shows the distribution of the deformation of the ions at the touching point. The projectile deformation is indicated on the y-axis, and deformation of the target — on the x-axis. The negative values of deformation correspond to the oblate shapes. The mean value of deformation was shown in Figure 1 as a function of the distance between the target and projectile.

4. The Evolution of Composite System

First of all, we have to define the initial shape of the mono system. The parameter was put equal to 1, which corresponds to touching configuration with zero neck. Then for each trajectory we fix by the Neimann (“hit and miss”) method the deformations of the colliding ions, see bottom of Figure 3. These two deformations define the elongation of the whole system and the target to projectile mass ratio defines the asymmetry of the whole system. For given elongation and asymmetry we found numerically the values of the asymmetry parameter and the neck parameter. After that, we used the same procedure and distribution from the top part of Figure 3 and defined of the excitation energy and angular momentum of the mono system.

All this information was used as input data for calculation of the evolution of the mono system, eq 2. For given $\alpha$, $\alpha_1$, and...
The Coulomb, surface, and rotational energy of the mono system was calculated (the rigid body moment of inertia was used). The initial excitation energy defines the temperature of the system and, consequently, the damping of the shell effects. Besides, we assumed that the collective kinetic energy is given by the relation

\[ E_{\text{kin}} = E_{\text{tot}} - V_{\text{pot}} - E_{\text{int}}. \]  

(5)

Here \( E_{\text{tot}} \) is the total energy defined from the initial kinetic energy of ions and reaction \( Q \) value, \( V_{\text{pot}} \) — potential energy for given collective variables and temperature.

The eq 2 were solved numerically. From the solution of eq 2 it follows that the system can break apart practically at once or “travel” for some time on the surface shown in Figure 2. This “travel” can lead either to fusion or fission. The computed fusion and fission cross sections are shown in Figure 4 together with experimental values for the fusion cross section. One can see that computed and experimental values of fusion cross section and fission cross sections are shown in Figure 4 together with experimental values for the fusion cross section. One can see that computed and experimental values of fusion cross section.
barrier in the entrance channel. As it was mentioned already, the computed touching cross section in this case is smaller than the experimental value, see Figure 4. Very likely, for this energy one should account for the tunneling through the barrier. Nevertheless, the mass distribution is reproduced rather well.

In case $E_{lab} = 78$ MeV, the excitation energy is very small and shell effects lead to the "shoulders" in the mass distribution corresponding to mass asymmetric fission. One can see that the "shoulders" are reproduced by the computations. The computed neutron multiplicity is somewhat smaller than the experimental value. There is even some disagreement in the dependence of neutron multiplicity on the mass of fission fragment. Whereas experimental values show some minimum for mass symmetric fission, the computed results are rather independent of the fragment mass.

Figure 7 shows the dependence of the decay events and the number of evaporated neutrons at the smallest deformation reached by trajectory. The panel (a) shows the distribution of all decay events. This distribution has a main bump in the region of $\alpha \sim 1$ (short time events or of the deep inelastic events). The position of the second bump is in barrier region. Many trajectories can not cross the barrier. The main part of the fission trajectories returns from barrier region and minor part of the all trajectories crosses the barrier and go to ground state.

The panel (b) shows of distribution of the positions where the neutrons were evaporated. The solid line include the neutrons evaporated from all trajectories, from the trajectories with deformation smaller than $\alpha = 0.94$ (line with triangles), and from the trajectories with deformation smaller than $\alpha = 0.64$ (line with squares). Bottom — the mean number of evaporated neutrons per decay.

5. Summary

We have computed the fusion and fission cross section within the approach based on Langevin equation both for entrance and fission channels. The evaporation residue cross section is also estimated. The mass-energy distributions of the fragments and the dependence of neutron multiplicities on the fragment mass is obtained. The computed results are in good agreement with the experimental data.

The analysis of the numerical results allows us to make the conclusion that the classical compound nucleus is not formed during the collision. At the same time, the distribution of fragments is the same as for the fission of compound nucleus. Moreover, the main part of trajectories does not reach the deformations smaller than the barrier and the main part of neutrons are evaporated around the barrier. In the result of neutron evaporation, the system loses considerable part of kinetic energy, and can not overcome the barrier.

Thus, in order to reach the ground state, the system should have enough energy above the Coulomb barrier. In the synthesis of superheavy elements, these experiments were successful in which system evaporated 2–3 neutrons before it reached ground state.24

The authors realize that the results are obtained for one concrete reaction and other fusion-fission reaction should be described. Also the agreement with the experiment near the barrier energy is not very good. The investigations in this directions would be the subject of future studies.

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