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TEST OF MODELS FOR PHOTON STRENGTH FUNCTIONS OF ELECTRIC DIPOLE PHOTOEXCITATION

Closed-form models for photon strength functions for the description of photoabsorption cross section in the energy range 5 - 30 MeV for even-even nuclei are tested. The experimental database was prepared with systematic uncertainty less than 10% from the EXFOR database. The theoretical models are compared with the experimental photoabsorption cross section data. The minimum of the least-square deviation and the root-mean-square deviation factor are used as a criteria comparison of the theoretical calculations with experimental data. It is shown that the simple modified Lorentzian model is the best approach for simulation of the photoabsorption cross section at the gamma-ray energies below ~ 30 MeV.

\textit{Keywords:} models of photon strength functions, photoabsorption, giant dipole resonance.

1. Introduction

The photonuclear data are being intensively studied in response to growing needs for these data in a variety of practical applications [1]. Photons are commonly produced as bremsstrahlung radiation by electron accelerators which are relatively simple machines present in many hospitals, industries and laboratories. These institutions deal with applications such as activation analyses, radiation shielding and radiation transport analyses, safeguards and inspection technologies (identification of materials through radiation induced by photonuclear reactions using portable bremsstrahlung devices), calculation of absorbed dose in the human body during radiotherapy etc., see [1] for the references. Photon strength functions (PSFs) are important input parameter for modelling nuclear reactions, for the calculations of gamma-ray capture cross sections, gamma-ray production spectra, competition between gamma-ray and particle emission and of various properties of the atomic nuclei (see, for example, [2, 3] for references). The PSFs are mandatory component of all modern computer codes for nuclear reaction calculations and nuclear data evaluation, such as EMPIRE and TALYS [4, 5].

The electric dipole (E1) multipolarity of gammarays is dominant in bremsstrahlung spectra and dominant in nuclear processes of photoabsorption and gamma-decay when they occur simultaneously with transitions of other multipolarities. For medium-heavy and heavy atomic nuclei, the most important contribution to the probability of these transitions in the range of gamma-ray energies \( 8 < \varepsilon_{\gamma} < 30 \) MeV is resulted from the response of Isovector Giant Dipole Resonances (GDR). A Lorentz shape is preferable for approximation of a PSF of such response (see [2, 3, 6, 7] and references therein). Here we consider the common-used analytical models with this line-shape, namely Standard Lorentzian (SLO) [2, 3], Generalized Lorentzian (GLO) [2, 3, 8], Simplified version of modified Lorentzian (SMLO) [3 - 7, 9 - 13] and Triple Lorentzian (TLO) [14, 15].

The total photoabsorption cross section \( \sigma_{\text{tot}}(\varepsilon_{\gamma}) \) is calculated as a sum of the terms corresponding to the E1 GDR excitation \( \sigma_{\text{GDR}}(\varepsilon_{\gamma}) \) and quasi-deuteron photodisintegration \( \sigma_{\text{qd}}(\varepsilon_{\gamma}) \):

\[
\sigma_{\text{tot}}(\varepsilon_{\gamma}) = \sigma_{\text{GDR}}(\varepsilon_{\gamma}) + \sigma_{\text{qd}}(\varepsilon_{\gamma}).
\]

The approach from Ref. [2, 16] is used for a quasi-deuteron contribution. The main part of experimental data on photoabsorption cross sections is obtained with the use of bremsstrahlung, which is mainly composed of gamma-rays of E1 multipolarity. Thus, the GDR component of the total photoabsorption cross section is adopted to be equal to the photoabsorption cross section of electric dipole gamma-rays \( \sigma_{\text{GDR}}(\varepsilon_{\gamma}) = \sigma_{\text{E1}}(\varepsilon_{\gamma}) \). The average photoabsorption cross section of a nucleus in the ground state of spin \( J_0 \) for excitation of levels of spin \( J \) is proportional to the photoexcitation E1 PSF and is given by [6]

\[
\sigma_{\text{E1}}' = \frac{g_J}{3} \sigma_{\text{E1}} = g_J (\pi \hbar c)^2 \varepsilon_{\gamma} \tilde{f}_\text{E1}(\varepsilon_{\gamma}),
\]

where \( g_J = (2J + 1) / (2J_0 + 1) \) is the statistical factor and \( |J_0 - 1| \leq J \leq J_0 + 1 \). The analytical model

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for E1 PSF of photoexcitation $j^\alpha(\varepsilon_r)$ defines the component of photoabsorption cross section $\sigma_{GDR}$ (1) resulted from the GDR excitation:

$$\sigma_{GDR}(\varepsilon_r) = \sigma_{GDR}^\alpha(\varepsilon_r) = 3(\pi\hbar c)^2 \varepsilon_r j^\alpha(\varepsilon_r) ,$$  \hspace{1cm} (2)

where index $\alpha$ denotes the PSF model. General analytical expression for the E1 PSF of photoexcitation of the cold nuclei governed by the GDR can be presented in the following form

$$\sigma^\alpha(\varepsilon_r) = 8.674 \cdot 10^{-8} \sum_{\alpha=1}^{\infty} \frac{\sigma_{\\text{TLO}} \alpha}{\varepsilon_r} \times$$

$$\times \frac{2}{\pi} \left( \frac{\varepsilon_r^2 \Gamma_{1j}^{\alpha}}{(\varepsilon_r^2 - (E_{1j}^{\alpha})^2)^2 + (\Gamma_{1j}^{\alpha} \varepsilon_r)^2} \right) \text{(MeV}^{-3}) .$$  \hspace{1cm} (3)

Here, the index $j$ specifies number of the normal modes of giant vibrations: for spherical nuclei $j_m = 1$, for axially symmetric $j_m = 2$ and for nuclei with triaxial shape $j_m = 3$; factor $s_\alpha^n$ is a weight of the $j$-mode; $\sigma_{\\text{TLO}}^\alpha$ is the Thomas - Reiche - Kuhn (TRK) sum rule $\sigma_{\\text{TRK}} = 15A(1 - \Gamma^2)$ (mb-MeV), with $I = (N - Z) / A$ for the neutron-proton asymmetry factor. A weight of the $j$-mode determines cross section of $j$-mode at resonance energy $E_{r,j}^\alpha$:

$$\sigma^\alpha_{r,j} = (2 / \pi) \sigma_{\\text{TLO}}^\alpha \cdot s_\alpha^n / \Gamma_{1j}^{\alpha} .$$

An approximation of axially symmetric nuclei with the effective quadrupole deformation parameter $\beta_j$ is adopted for deformed nuclei in the SLO, GLO and SMLO models. The values of effective quadrupole deformation parameters are taken from “deflib.dat” file of RIPL 2 ($\beta_j = \beta_{j,\text{eff}}$) [17]. These models for cold nuclei employing the same general expression (3) but with different determination of the shape width $\Gamma_{1j}^{\alpha}$. For the SLO model, the width is energy-independent constant which is equal to GDR width for $j$-mode: $\Gamma_{\text{SLO}}^{\alpha} = \Gamma_{1j}^{\alpha}$. The width in the GLO model is quadratic in gamma-ray energy $\Gamma_{GLO}^{\alpha}(\varepsilon_r) = \varepsilon_r^2 \Gamma_{1j}^{\alpha} / (E_{1j}^{\alpha})^2$, and the width in the SMLO model is a linear function of the gamma-ray energy $\Gamma_{\text{SMLO}}^{\alpha}(\varepsilon_r) = \varepsilon_r \cdot \Gamma_{1j}^{\alpha} / E_{1j}^{\alpha}$. For the GDR characteristics of the SLO and SMLO models, the recommended values (the energies, widths and weights) from recent database were used (see the Tables 1 and 2 from Ref. [7]). The GDR parameters of the SLO approach were taken for the GLO parameters.

Approximation of a triaxial ellipsoid is used in the TLO model for nuclear shape in deformed nuclei [14, 15], and in this approach, the GDR splits into three components and the E1 PSF is described by the expression (3) with $j_m = 3$ [18, 19]. For the input parameters of the TLO PSF, the resonance energies of the normal modes are taken from the nuclear hydrodynamics model [14, 15, 20, 21] as $E_{r,j}^{\text{TLO}} = E_{0} \cdot R_{0} / R_{j}$, where $E_{0}$ is the resonance energy of the equivalent in volume spherical nucleus with the radius $R_{0}$. The width in the TLO model is independent of gamma-ray energy and the expression with a pow er law dependence on the resonance energy was used [14, 15, 22]: $\Gamma_{j}^{\text{TLO}} = 0.045(E_{r,j}^{\text{TLO}})^{1.6}$. The approximation of equally probable excitation of the normal modes of the giant collective vibration were taken for the weights: $s_{r,j}^{\text{TLO}} = s^{\text{TLO}} / 3$ with $s^{\text{TLO}} = 0.995 \approx 1$ for the sum of the weights.

The triaxial nuclear shape is determined in terms of parameters $\beta$ (represents the extent of quadrupole deformation) and $\gamma$ (the degree of axial asymmetry). Two parameterizations are used here for the calculation of the semi-axis lengths $R_j$ for the TLO model. The Hill - Wheeler parameterization [22, 23] is used at experimental values of the deformation parameters ($\beta = \beta^H, \gamma = \gamma^H$), which are presented in the Table 1:

$$R_j = R_{j}^{H} = R_{0} \cdot \exp(\sqrt{5/4\pi} \cdot \beta^H \cdot \cos(\gamma^H - \frac{2}{3} \pi j)) .$$  \hspace{1cm} (4)

The Bohr parameterization is taken in [23] with the theoretical deformation parameters ($\beta = \beta^B, \gamma = \gamma^B$):

$$R_j = R_{j}^{B} = R_{0}(1 + \sqrt{5/4\pi} \cdot \beta^B \cos(\gamma^B - \frac{2}{3} \pi j)) .$$  \hspace{1cm} (5)

### Table 1. The values of deformation parameters $\beta = \beta^H, \gamma = \gamma^H$ used for PSF calculations within TLO(3) model

| Nuclei  | $^{93}\text{Mo}$ | $^{93}\text{Mo}$ | $^{140}\text{Nd}$ | $^{140}\text{Sm}$ | $^{152}\text{Gd}$ | $^{160}\text{Er}$ | $^{190}\text{Os}$ | $^{190}\text{Pt}$ | $^{209}\text{Bi}$ | $^{238}\text{U}$ |
|--------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| $\beta$ | 0.08            | 0.18            | 0.17            | 0.13            | 0.22            | 0.28            | 0.16            | 0.13            | 0.02            | 0.29            |
| $\gamma$ | 20°             | 23°             | 26°             | 25°             | 11°             | 12°             | 21°             | 29°             | 40°             | 17°             |
| Ref.   | [14]            | [14]            | [25]            | [26]            | [27]            | [26]            | [26]            | [14]            | [27]            | [25]            |

The deformation parameters in Bohr parametrization (5) were calculated in Ref.[24] for even-even nuclides within framework of the Hartree - Fock - Bogoliubov theory (HFB; $\beta^B = \beta_{B}^H, \gamma^B = \gamma_{B}^H$) and Constrained Hartree - Fock - Bogoliubov approach for five-dimensional collective Hamiltonian.
The analytical PSF expressions for mode $\alpha$ are based on photoabsorption cross sections from Ref. [7]. The absence of this contribution for first order, Bohr’s parameters $(\beta^B, \gamma^B)$ are equal to Hill - Wheeler’s parameters $(\beta^H, \gamma^H)$ [23]. The calculations within TLO(1) model were performed with three variants of the deformation parameters and denoted here as TLO(1), TLO(2) and TLO(3). The corresponding deformation parameters for each model are following: TLO(1) – $(\beta^B, \gamma^B)$; TLO(2) – $(\beta^B, \gamma^C)$; TLO(3) – $(\beta^H, \gamma^H)$. The deformation parameters within Hill - Wheeler’s parameterization for TLO(3) model were obtained from the fit of experimental data. The Table 1 shows these experimental deformation parameters and corresponding references for 10 isotopes for which the calculations within TLO(3) model were performed.

The comparison of SLO and SMLO approaches for the description of experimental data on photoabsorption cross section was done for 162 nuclei in Ref. [7]. It was shown that the low-energy tails of the photoabsorption cross sections within SLO model are higher than within the SMLO model and higher than experimental data. The comparison between SMLO model and the calculations of E1 PSF within microscopic approaches both quasiparticle random-phase approximation and shell-model was performed in Ref. [13] and demonstrated rather good agreement between them. In this contribution the comparison between SLO and SMLO models with other closed-form Lorentzian models of GLO and TLO need to be performed in order to obtain the best description of the experimental photoabsorption cross sections from EXFOR database [28]. Comparing to the Ref. [7], we additionally present the comparisons with the GLO and TLO models. For the TLO model, the atomic nuclei are considered in the triaxial shape approach. In this situation, the corresponding deformation parameters were calculated or obtained for the even-even isotopes only and due to this, we test the E1 PSF for description of the photoabsorption data for the even-even nuclei.

### 2. Calculations and discussion

The analytical PSF expressions for models of SLO, GLO, SMLO and TLO and their input parameters were described in previous section. Quantitative comparison between these PSF models for the description of total photoabsorption cross sections is performed for the following 88 even-even isotopes: $^{24}$Mg, $^{28}$Si, $^{32}$S, $^{40}$Ar, $^{44}$Ca, $^{48}$Ti, $^{52}$Cr, $^{54}$Fe, $^{58}$Ni, $^{62}$Zn, $^{70}$Ge, $^{76}$Se, $^{88}$Sr, $^{90}$Zr, $^{92}$Mo, $^{106}$Cd, $^{112}$Sn, $^{114}$Te, $^{138}$Ba, $^{140}$Ce, $^{142}$Nd, $^{144}$Sm, $^{152}$Gd, $^{168}$Er, $^{174}$Yb, $^{176}$W, $^{186}$Os, $^{194}$Pt, $^{206}$Pb, $^{232}$Th.

The calculations within TLO(3) model are done for 10 isotopes specified in the Table 1. For comparison of the theoretical models, the experimental database on photoabsorption cross section was prepared from EXFOR database for even-even nuclei with the systematic uncertainty less than 10% (see explanations below). The same datasets were taken as used in Ref. [7] for determination of recommended GDR parameters.

The E1 PSF $\tilde{f}^a$ from equation (2) is connected with total response of the nucleus on E1 field and defines the total photoabsorption cross section which is given by the relationships:

$$\sigma(\gamma, \text{abs}) = \sigma(\gamma, \gamma) + \sigma'(\gamma, \text{abs}),$$

$$\sigma'(\gamma, \text{abs}) = \sigma(\gamma, \text{sn}) + \sigma(\gamma, \text{cp}) + \sigma(\gamma, F),$$

$$\sigma(\gamma, \text{sn}) = \sigma(\gamma, n) + \sigma(\gamma, 2n) + \sigma(\gamma, 3n) + ..., $$

$$\sigma(\gamma, \text{cp}) = \sigma(\gamma, p) + \sigma(\gamma, 2p) + ... +$$

$$+ \sigma(\gamma, d) + ... + \sigma(\gamma, dp) + ... + \sigma(\gamma, \alpha) + ....$$

Here, $\sigma(\gamma, \gamma) = \sigma(\gamma, \gamma) + \sigma(\gamma, \gamma')$ is total photon-scattering cross section with excitation of the nuclear states corresponding to total cross section of target nucleus production, i.e., a sum of the cross sections of elastic gamma-ray scattering via different type of intermediate states (without shape-elastic component like compound-nucleus elastic scattering) and non-elastic gamma-ray scattering: $\sigma'(\gamma, \text{abs})$ is photoabsorption cross section with emission of the particles together with gamma-rays; $\sigma(\gamma, \text{sn})$ is total photo-neutron reaction cross section; $\sigma(\gamma, \text{cp})$ is photo-charged-particle reaction cross section and $\sigma(\gamma, F)$ is photo-fission reaction cross section for fissionable nuclei, $\sigma(\gamma, F) = \sigma(\gamma, f) + \sigma(\gamma, nf) + \sigma(\gamma, 2nf) + ...$, with $\sigma(\gamma, f)$ (and $\sigma(\gamma, nf)$) for first (second)-chance photofission cross section.

The experimental data for total photoabsorption cross sections from EXFOR database above neutron separation energies $S_n$ do not include the contribution of the cross section $\sigma(\gamma, \gamma)$ from gamma-gamma channels. This contribution is non-negligible at gamma-ray energies $S_n + \Delta E$ below the threshold of the $(\gamma, 2n)$ reaction or, in some cases, below the thresholds of other reactions with large cross sections; typically $\Delta E < 1.5$ MeV. The absence of this contribution leads to incorrectly small values of $\sigma(\gamma, \text{abs})$ in the gamma-ray range $S_n < E_\gamma < S_n + \Delta E$.

For every nucleus the specific interval $\Delta E$ was found from the condition of ten percent contribution of the cross section $\sigma(\gamma, \gamma)$ from gamma-gamma...
transitions to the total photoabsorption cross section \( \sigma(\gamma, \text{abs}) \), i.e. ten percent deviation of \( \sigma(\gamma, \text{abs}) \) from \( \sigma'(\gamma, \text{abs}) \) or ten percent systematic uncertainties of the PSF:

\[
\delta \sigma(\gamma, \psi) = \frac{\sigma(\gamma, \text{abs}) - \sigma'(\gamma, \text{abs})}{\sigma(\gamma, \text{abs})} = 0.1. \quad (6)
\]

![Cross section, mb](image)

Fig. 1. Theoretically calculated photo cross sections for isotope \(^{94}\text{Mo}\) in comparison with experimental data from EXFOR database (circles [29], squares [30]). The vertical lines correspond to \( S_n \) (black dashed line) and the gamma-ray energy \( S_n + \Delta \varepsilon \) (black solid line) for which evaluated systematic uncertainties of the PSF are equal to 10 \%. (See color Figure on the journal website.)

Based on the above, for test of the PSF models for photoabsorption data description, the new database on experimental E1 PSF was prepared using photoabsorption/photo-neutron cross sections from EXFOR database for even-even isotopes with systematic uncertainty less than 10 \%. In this database the gamma-ray energy interval for each isotope is starting from \( \varepsilon_{\text{min}} = S_n + \Delta \varepsilon \) and ends in the maximal value of energy \( \varepsilon_{\text{max}} \leq 30 \text{ MeV} \) presented in the data file. In these intervals above neutron separation energies the gamma-gamma contributions to photo cross sections do not exceed 10 \%. The specific low boundaries of gamma-ray energy intervals were calculated using simulations of the photo cross sections by the nuclear reaction code TALYS 1.6 [5] with gamma-strength within SLO and others input parameters by default (for example, with Gilbert - Cameron model for nuclear level density). It should be noted that in an accordance with TALYS specifications, the cross section \( \sigma(\gamma, \psi) \) corresponds to the reaction cross section for incident photons \( \sigma_{\text{reac}}(\gamma) \), and \( \sigma(\gamma, \psi) \) is a sum of production cross section \( \sigma_{\text{prod}}(Z_i, N_i) \) of the target nucleus \( (Z_i, N_i) \) and the compound elastic cross section \( \sigma_{\text{comel}}(\gamma, \psi) \): 

\[
\sigma(\gamma, \psi) = \sigma_{\text{prod}}(Z_i, N_i) + \sigma_{\text{comel}}(\gamma, \psi). 
\]

Fig. 1 demonstrates calculated photo-cross sections in comparison with the experimental data for the isotope \(^{94}\text{Mo}\).

Two criteria were taken for comparison of the quality of the description of the experimental photo-nuclear data using different E1 PSF models: 1) minimum of the least-square deviation \( \chi_n \), 2) minimum of the root-mean-square (rms) deviation factor \( f_a \) [31]:

\[
\chi_n^2 = \frac{1}{n} \sum_{i=1}^{n} (\sigma_{\text{exp}}(\varepsilon_i) - \sigma_{\text{the}}(\varepsilon_i))^2 / (\Delta \sigma(\varepsilon_i))^2, \quad (7)
\]

\[
f_a = \exp{\chi_{\text{in,a}}} \Rightarrow (\chi_{\text{in,a}})^2 = \frac{1}{n} \sum_{i=1}^{n} \left( \ln \sigma_{\text{the}}(\varepsilon_i) - \ln \sigma_{\text{exp}}(\varepsilon_i) \right)^2 = \frac{1}{n} \sum_{i=1}^{n} \ln^2 \left( \frac{\sigma_{\text{the}}(\varepsilon_i)}{\sigma_{\text{exp}}(\varepsilon_i)} \right). \quad (8)
\]

Here \( \sigma_{\text{the}}(\varepsilon_i) = \sigma_{\text{GDR}}(\varepsilon_i) + \sigma_{\text{sd}}(\varepsilon_i) \) is the theoretical cross section corresponding to Eqs. (1), (2), at gamma-ray energy \( \varepsilon_i \); \( \sigma_{\text{exp}}(\varepsilon_i) \) is the experimental photoabsorption cross section from the EXFOR database; \( \Delta \sigma(\varepsilon_i) \) is the data uncertainty; \( n \) is the number of experimental data points. The rms deviation factor \( f_a \) corresponds to criterion of a minimum at given gamma-ray energy of the weighted sum of squared deviations of the theoretical cross section from their experimental data in natural logarithmic scale. Because of different estimations of uncertainties in different datasets and, as a rule, for lack of reliable estimations of the systematic errors, a weight of every point is taken as empirical probability 1/n. A logarithmic scale is used due to large-range changes of the photoabsorption cross sections.

The \( \chi_n^2 \) and \( f_a \) values were calculated for two gamma-ray energy intervals. The gamma-ray energy intervals from \( S_n + \Delta \varepsilon \) till 30 MeV (or \( \varepsilon_{\text{max}} \) with the energy \( \varepsilon_{\text{max}} \) for the last data point) were taken for an estimation of the quality of fitting in the wide energy range and marked below in the tables as “broad range”. The short energy ranges near the GDR peaks were taken from the Table 1 of Ref. [7].

For \( \chi_n^2 \) calculation, the cross section uncertainties \( \Delta \sigma(\varepsilon) \) were taken in two versions: 1) the energy-dependent trapezoidal relative uncertainties \( \delta \sigma_{\text{exp}}(\varepsilon_i) \) (see below) were applied for all nuclei and marked below as “trap”; 2) the experimental uncertainties \( \Delta \sigma_{\text{exp}}(\varepsilon_i) \) given in EXFOR datafiles were used and trapezoidal uncertainty were taken only in case of absence of \( \Delta \sigma_{\text{exp}}(\varepsilon_i) \) and marked below as “exp + trap”. The energy-dependent trapezoidal rela-
tive uncertainties \( \delta \sigma_{\text{exp}}(\epsilon_i) = \Delta \sigma_{\text{exp}}(\epsilon_i) / \sigma_{\text{exp}}(\epsilon_i) \) were taken in accordance with Ref. [7, 12]. These energy dependent errors were chosen to simulate the statistical error that is inversely proportional to the square root of the counting rate, which has maximum near the GDR. Therefore, the energy-dependent relative uncertainties were assumed to take minimum values (10%) near the GDR peaks and maximum values (50%) on the GDR tails. These values are based on the typical uncertainties of the experimental data. For deformed nuclei, a trapezoidal dependence with the GDR peaks as the top corners of the trapezium was assumed

\[
\delta \sigma(\epsilon_i) = \frac{\Delta \sigma(\epsilon_i)}{\sigma_{\text{exp}}(\epsilon_i)} = \begin{cases} 
\delta_{\text{min}} + b |E_{\epsilon,i} - \epsilon_i|, & \epsilon_i < E_{\epsilon,1} \\
\delta_{\text{min}}, & E_{\epsilon,2} \leq \epsilon_i \leq E_{\epsilon,1} \\
\delta_{\text{min}} + b |E_{\epsilon,2} - \epsilon_i|, & \epsilon_i > E_{\epsilon,2}
\end{cases}
\] (9)

with \( b = (\delta_{\text{max}} - \delta_{\text{min}}) / (E_{\epsilon,1} - \epsilon_i) \), where \( \delta_{\text{min}} = 0.1 \) and \( \delta_{\text{max}} = 0.5 \) are the minimal and maximal values of uncertainty. For spherical nuclei, a triangular dependence on gamma-energy was used

\[
\delta \sigma(\epsilon_i) = \delta_{\text{min}} + b |E_{\epsilon,i} - \epsilon_i|.
\]

Figs. 2 and 3 present the theoretical photoabsorption cross sections calculated within different PSF models in comparison with experimental data for the isotopes of \(^{124}\text{Sn}\) and \(^{238}\text{U}\). For these isotopes, the relative values of \( \chi^2 / \chi^2_{SL0} \) and \( f_a / f_{SL0} \) are given in the Table 2. For the nucleus \(^{124}\text{Sn}\), lowest values of \( \chi^2 / \chi^2_{SL0} \) and \( f_a \) were obtained by the use of the SMLO and GLO models. For \(^{238}\text{U}\), the minimal values of \( \chi^2 / \chi^2_{SL0} \) and \( f_a \) correspond to the SMLO and TLO(3).

Fig. 2. The experimental cross section data for \(^{124}\text{Sn}\) in comparison with calculations within different PSF models. Experimental data are taken from Ref. [32]. Left-hand Figure: —— SMLO (red solid line); —— SLO (black dashed line); ——— GLO (blue dotted line). Right-hand Figure: —— SMLO (red solid line); ——— TLO(1) (light green dashed followed by one dot line); ——— TLO(2) (dark green dashed followed by two dots line). (See color Figure on the journal website.)

Fig. 3. The experimental cross section data for \(^{238}\text{U}\) in comparison with calculations within different PSF models. Experimental data are taken from Ref. [33]. Left-hand Figure: —— SMLO (red solid line); —— SLO (black dashed line); ——— GLO (blue dotted line). Right-hand Figure: —— SMLO (red solid line), ——— TLO(1) (light green dashed followed by one dot line); ——— TLO(2) (dark green dashed followed by two dots line); ——— TLO(3) (black dashed followed by three dots line). (See color Figure on the journal website.)
### Table 2. The values of $\chi^2_{a}/\chi^2_{SLO}$ and $f_a/f_{SLO}$ for $^{124}$Sn and $^{238}$U

| Isotope | Criteria | Gamma-ray energy intervals | Uncertainty | SMLO | GLO | TLO(1) | TLO(2) | TLO(3) |
|---------|----------|---------------------------|-------------|------|-----|--------|--------|--------|
| $^{124}$Sn | $\chi^2_{a}/\chi^2_{SLO}$ | broad range | trap | 0.17 | 0.11 | 0.42 | 0.38 | – |
|          |          | near GDR | trap | 0.30 | 5.15 | 21.98 | 21.15 | – |
| $f_a/f_{SLO}$ | broad range | – | 0.85 | 0.82 | 0.91 | 0.89 | – |
|          | near GDR | – | 0.99 | 1.04 | 1.13 | 1.09 | – |
| $^{238}$U | $\chi^2_{a}/\chi^2_{SLO}$ | broad range | exp | 0.81 | 0.97 | 1.83 | 1.90 | 1.55 |
|          |          | trap | 1.04 | 1.51 | 1.28 | 1.50 | 1.02 |
| $f_a/f_{SLO}$ | near GDR | exp | 1.02 | 1.90 | 3.06 | 3.35 | 2.17 |
|          |          | trap | 1.15 | 1.63 | 1.34 | 1.64 | 1.03 |

### Table 3. The ratio $K_a=<\chi^2_{a}/\chi^2_{SLO}>$ for different PSF models

| Mass numbers of the isotopes | Gamma-ray energy intervals | Uncertainty | SMLO | GLO | TLO(1) | TLO(2) | TLO(3) |
|-----------------------------|---------------------------|-------------|------|-----|--------|--------|--------|
| $24 \leq A \leq 238$ | broad range | exp + trap | 1.13 | 3.05 | 10.70 | 10.71 | 4.58 |
|          | trap | 1.03 | 2.25 | 8.61 | 9.73 | 2.68 |
|          | near GDR | exp + trap | 1.19 | 5.43 | 42.98 | 47.88 | 17.28 |
|          | trap | 1.22 | 4.86 | 43.07 | 50.08 | 18.17 |
| $80 \leq A \leq 238$ | broad range | exp + trap | 1.22 | 3.58 | 10.94 | 11.33 | 4.58 |
|          | trap | 1.04 | 2.54 | 8.28 | 9.58 | 2.68 |
|          | near GDR | exp + trap | 1.28 | 6.52 | 49.39 | 57.20 | 17.28 |
|          | trap | 1.29 | 5.82 | 49.82 | 60.15 | 18.17 |

### Table 4. The ratio $F_a=<f_a/f_{SLO}>$ for different PSF models

| Mass numbers of the isotopes | Gamma-ray energy intervals | SMLO | GLO | TLO(1) | TLO(2) | TLO(3) |
|-----------------------------|---------------------------|------|-----|--------|--------|--------|
| $24 \leq A \leq 238$ | broad range | 0.97 | 1.05 | 1.18 | 1.14 | 1.10 |
|          | near GDR | 1.00 | 1.06 | 1.24 | 1.22 | 1.10 |
| $80 \leq A \leq 238$ | broad range | 0.97 | 1.06 | 1.16 | 1.14 | 1.12 |
|          | near GDR | 1.00 | 1.05 | 1.20 | 1.18 | 1.10 |

Fig. 4. The relative least square values $\chi^2_{a}/\chi^2_{SLO}$ (left-hand Figure) and rms deviation factors $f_a/f_{SLO}$ (right-hand Figure) for even-even isotopes calculated within different PSF models. Presented results correspond to 5 - 30 MeV gamma-ray energy interval and trapezoidal relative uncertainties. On both Figures: SMLO – red circles (●); GLO – blue empty triangles (▲); TLO(1) – green crosses (×); TLO(2) – dark green pluses (+); TLO(3) – black empty squares (□). (See color Figure on the journal website.)
The arithmetic mean values $K_a = \langle \chi_a^2 / \chi_{SLO}^2 \rangle$ and $F_a = \langle f_a / f_{SLO} \rangle$ are presented in the Tables 3 and 4. The relative values of $\chi_a^2 / \chi_{SLO}^2$ and $f_a / f_{SLO}$ are presented in Fig. 4. On average, the $K_a$ and $F_a$ have lower values for the SMLO model. The description of the experimental data within the SMLO and GLO models with energy-dependent width is better than within TLO models with standard values of deformation parameters (4), (5).

3. Conclusions

Quantitative comparison between the E1 PSF models for the description of total photoabsorption cross sections is performed for 88 even-even isotopes. The database of the photoabsorption cross section was prepared from EXFOR data with systematic uncertainty less than 10 %. The uncertainty calculations were performed using TALYS code. The experimental values are compared with theoretical predictions within framework of the models SLO, GLO, SMLO and TLO. The criteria of minimum of least-square factor and root-mean-square deviation factor were used for comparison of the theoretical calculations with experimental data. It was shown that the SMLO model gives better description of the photoabsorption data in the range of gamma-ray energies till ~30 MeV. In our opinion, this model can be recommended in the nuclear reaction codes for adequate modelling of the E1 photoexcitation PSF using simple closed-form expressions.

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ТЕСТУВАННЯ МОДЕЛЕЙ ФОТОННИХ СИЛОВИХ ФУНКЦІЙ ЕЛЕКТРИЧНОГО ДИПОЛЬНОГО ФОТОЗБУЖДЕНИЯ

Тестуються аналітичні моделі E1 фотонних силових функцій з опису перерізів фотопоглинання в області енергій гамма-випромінювання 5 - 30 MeV для парно-парних ядер. Було побудовано експериментальну базу перерізів фотопоглинання із систематичною похибкою менше 10 % з даних EXFOR. Похибки оцінюються за допомогою теоретичних розрахунків перерізів з використанням коду TALYS 1.6. Теоретичні моделі порівнюються з експериментальними даними перерізів фотопоглинання. В якості критерій використовуються метод найменших квадратів та середньоквадратичний фактор відхилення. Показано, що проста модифікована модель Лоренца краще описує перерізи при енергіях гамма-випромінювання нижче від ~30 MeV.

Ключові слова: моделі фотонних силових функцій, фотопоглинання, гіганський дипольний резонанс.

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ТЕСТИРОВАНИЕ МОДЕЛЕЙ ФОТОННЫХ СИЛОВЫХ ФУНКЦИЙ ЭЛЕКТРИЧЕСКОГО ДИПОЛЬНОГО ФОТОЗАБУЖДЕНИЯ

Тестируются аналитические модели E1 фотонных силовых функций для описания сечений фотопоглощения в области энергий гамма-излучения 5 - 30 МэВ для четно-четных ядер. Подготовлена экспериментальная база сечений фотопоглощения с систематической погрешностью менее 10 % из данных EXFOR. Погрешности оцениваются при помощи теоретических расчетов сечений с использованием кода TALYS 1.6. Теоретические модели сравниваются с экспериментальными данными сечений фотопоглощения. В качестве критерий используются метод наименьших квадратов и среднеквадратичный фактор отклонения. Показано, что простая модифицированная модель Лоренца лучше описывает сечения фотопоглощения при энергиях гамма-излучения ниже ~30 МэВ.

Ключевые слова: модели фотонных силовых функций, фотопоглощение, гигантский дипольный резонанс.

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