Core Polarization Effects on the Inelastic Longitudinal C2 and C4 form Factors of $^{46,48,50}$ Ti Nuclei

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Abstract The form factors for the inelastic electron scattering to $2^+, 4^+$ states in $^{46,48,50}$Ti are studied in the framework of shell model. The calculation is performed in (0f7/2, 1p3/2, 0f5/2, 1p 1/2) model space as well as extended 6$\hbar$ω model space. The predictions of longitudinal form factors which includes core-polarization effects to first order were compared with the experimental values. It is noticed that the core polarization effects are essential in obtaining a remarkable agreement between the calculated inelastic longitudinal $F(q)$'s and those of experimental data.

Keywords Nuclear Reactions, Electron Scattering(E,E'), Inelastic Form Factors with Core-Polarization Effects

1. Introduction

The calculations of Shell model, carried out within a model space in which the nucleon are restricted to occupy a few orbits are unable to reproduce the measured static moments or transition strengths without scaling factors. Comparison between calculated and measured longitudinal electron scattering form factors has long been used as stringent tests of models for transition densities. Various microscopic and macroscopic theories have been used to study excitations in nuclei[1]. Shell model within a restricted model space is one of the models, which succeeded in describing static properties of nuclei, when effective charges are used. Calculations of form factors using the model space wave function alone is inadequate for reproducing the data of electron scattering[2]. Therefore, effects out of the model space, which are called core polarization effects, are necessary to be included in the calculations.

The intermediate one-particle one-hole states are taken up to 6$\hbar$ω excitation. These effects are found essential for obtaining a quantitative agreement with the experimental data[3,4]. A microscopic model[5,6] has been used in order to study the core polarization effect on the longitudinal form factors of fp-shell nuclei. A microscopic model which adopted the first order core polarization was considered to calculate the C2 form factors of the fp-shell nuclei. Inelastic Electron Scattering from fp Shell Nuclei had been studied by Sahu et al[7]. They calculated form factors for $^{50,52,54}$Cr, $^{54}$Fe, $^{56}$Fe, $^{46,48}$Ti, and $^{50}$Ti by the use of Hartree-Fock theory, results are in a good agreement with the experimental data. The form factors for the inelastic electron scattering to $2^+$, 4$^+$ and 6$^+$ states in $^{46,48,50}$Ti, $^{50,52,54}$Cr and $^{54,56}$Fe were studied by Sahu[8,9] in the framework of the Hartree-Fock model, also the calculation is performed in the 1f7/2, 2p3/2, 1f5/2, 2p 1/2 model space using a modified Kuo-Brown effective interaction. Magnetic dipole excitation of N = 28 isotones $^{48}$Ca, $^{50}$Ti, $^{52}$Cr, $^{54}$Fe and $^{51}$V, was studied by Muto and Horie[10], in terms of the shell model by assuming $f_{7/2}^{m=-m}(p_{3/2} f_{5/2}^{m=m} p_{1/2})$ configurations with m = 0, 1 and 2 on an inert $^{40}$Ca core. Coulomb form factors for the $0_g^{+} \rightarrow 2^+$ transitions in the even-even $^{1}$f$_{7/2}$-shell nuclei are studied in terms of the shell model within the $\pi f_{7/2}^{+}+\pi f_{7/2}^{-}\nu p_{3/2}$ configurations and with the effective interactions. It is shown that the characteristics of the C2 form factors in the higher-momentum-transfer region, not explained by the simple $\pi f_{7/2}^{+}$ model, can be interpreted by the mixing of the one-particle excitations into the $p_{3/2}$ orbit in the shell-model wave functions. E2 transition strengths and Q moments are also discussed in connection with the C2 form factors[11]. The use of modern realistic M3Y effective nucleon- nucleon interaction with two sets of fitting parameters (Ried fitting (M3Y-P1), and Paris fitting (M3Y-P0)) besides the use of MSDI has been done as a residual interactions within the calculation of core polarization effects in Inelastic longitudinal electron scattering C6 form factor in Ti-50 within the framework of first order perturbation theory (microscopic theory) with 2$\hbar$ω excitation energy coupling the core orbits to the higher configurations one across the model space at normal transition. Harmonic oscillator wave functions (H.O) has

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been adopted as a single particle wave functions in $1f^{7/2}$ and
with the aid of F7MBZ model space $1f^{7/2}$ effective interaction
to generate the model space wave functions. The present
results have been compared with the experimental data[12].
Differential cross sections for inelastic electron scattering
from $^{40}$Ca, $^{42}$Ca, $^{44}$Ca, $^{46}$Ti, $^{48}$Ti, $^{50}$Ti and $^{56}$Fe have been
measured. The cross sections for the strongly excited $2^+$, $3^+$,
$4^+$ and $5^+$ levels have been measured over a range of
momentum transfer $q$ of $0.4-2.6$ fm$^{-1}$. The data have been
fitted using two representations for the transition charge, (i)
the hydrodynamic liquid drop, and (ii) a phenomenological
model[13].

The aim of present work is to use a realistic effective
nucleon-nucleon (NN) interaction as a residual interaction to
calculate the core polarization (CP) effects through a
microscopic theory, with a selection of model space effective
interaction which generates the model space wave functions
(shell model wave functions) and highly excited states. The
(MSDI) were used in this case as a residual interaction. The strength of the MSDI denoted by $A_T$, $B$ and $C$ are set equal to $A_T=A_1=B=6.2$ MeV and $C=0$. The single
particle wave function were those of the harmonic oscillator
potential (HO) with size parameter $b$ chosen to reproduce the
measured ground states root mean square charge radii of these
nuclei. The one-body density matrix (OBDM) elements
($\chi^T_f \Gamma_j$) are calculated using the shell model
code OXBASH[14]

2. Theory

The electron scattering form factor for a given
multipolarity $\lambda$ and momentum transfer $Q$ is expressed
as[15],

$$|F_\lambda(Q)|^2 = \frac{1}{2f_\lambda + 1} \left( \frac{4\pi}{2} \right)^2 \left| \langle \Gamma_f \mid \hat{T}^{\lambda F}_{2} \mid \Gamma_i \rangle \right|^2 \left| F_{\lambda F} \right|^2$$

Where $F_{\lambda F} = e^{-0.43b^2/4}$ is the finite nucleon-size
correction and $F_{cm} = Q^2b^2/4A$ is the center of mass
correction, $A$ is the mass number and $b$ is the harmonic
oscillator size parameter.

The effect of the core polarization on the form factors is
based on a microscopic theory, which combines shell-model
wave functions and configuration with higher energy as
particle-hole perturbation expansion. The reduced matrix
element of the electron scattering operator $\hat{T}^{\lambda F}_{2}$ is expressed
as a sum of the fp-model space (p) contribution and the
core-polarization (cp) contribution, as follows[5],

$$\langle \Gamma_f \mid \hat{T}^{\lambda F}_{2} \mid \Gamma_i \rangle = \langle \Gamma_f \mid \hat{T}^{\lambda F}_{2} \mid \Gamma_i \rangle_p + \langle \Gamma_f \mid \hat{T}^{\lambda F}_{\text{cp}} \mid \Gamma_i \rangle_p$$

selection the longitudinal (L), electric(E) and magnetic(M)
transverse form factors , The Greek symbol were used to
denote quantum numbers in coordinate space and isospace
and isospace respectively: $\lambda \equiv JT$,

$$\Gamma_j \equiv J, T_j, \Gamma_f \equiv J_j T_f$$

The fp-shell model space element can be expressed as
linear combination of the single-particle matrix element[16],

$$\langle \Gamma_f \mid \hat{T}^{\lambda F}_{2} \mid \Gamma_i \rangle_p = \sum_{\alpha, \beta} \chi^{T_f} \Gamma_j (\alpha_f, \alpha_i) \langle \alpha_f \mid \hat{T}^{\lambda F}_{2} \mid \alpha_i \rangle$$

$$\chi^{T_f} \Gamma_j (\alpha_f, \alpha_i) = \sqrt{\chi^{T_f} \Gamma_j (\alpha_f, \alpha_i)}$$

are the single-particle states $\alpha_f$ and $\alpha_i$ for the fp-shell
model space. Similarly, core-polarization matrix element as
follows:

$$\langle \Gamma_f \mid \hat{T}^{\lambda F}_{\text{cp}} \mid \Gamma_i \rangle_p = \sum_{\alpha, \beta} \chi^{T_f} \Gamma_j (\alpha_f, \alpha_i) \langle \alpha_f \mid \hat{T}^{\lambda F}_{\text{cp}} \rangle (\alpha_i)$$

Up to the first order perturbation theory, the
single-particle matrix element for the higher-energy
configuration is given by[17]

$$\langle \alpha_f \mid \hat{T}^{\lambda F}_{\text{cp}} \rangle (\alpha_i) = \langle \alpha_f \mid \frac{\hat{Q}}{E_{\alpha}-H_{\alpha}} \hat{T}^{\lambda F}_{\text{cp}} \rangle (\alpha_i)$$

The operator Q is the projection operator on the space
outside the model space. $E_{\alpha}$ and $E_{\beta}$ are the energies of
initial and final states. For the residual interaction $V_{res}$ the
MSDI and M3Y were adopted.

The two terms in right hand side of Eq.(6) can be written
as[18]

$$\sum_{\alpha} (-1)^{m_{\alpha}-m_{\alpha}} \langle \alpha_f \mid \hat{T}^{\lambda F}_{\text{cp}} \rangle (\alpha_i)$$

The electric transition strength is given by,

$$e_{\omega} = \frac{1}{2} \frac{(l+1)(\epsilon(f(r)))}{\epsilon^2}$$

With

$$B(C \lambda) = \frac{2(2l+1)!}{4\pi} \left| \frac{Z^2}{k^2} \left| F_{\lambda} (k) \right|^2 \right.$$
These parameters are taken to be $A_0=A_1=B=6.2\text{MeV}$ and $C=0$. In all of the following diagrams the dashed line give the results obtained using the fp-shell wave function and the results including cp effects are shown by solid curve.

### 3.1. Form factors for the $0^+\rightarrow 2^+$ transition

A comparison between the experimental and theoretical form factors for the C2 transition for $^{46,48,50}$Ti is given in figures(1 - 3) respectively. For each of these three nuclei, the form factors shows three peaks. Figure (1) shows longitudinal C2 electron scattering form factors as a function of momentum transfer for $^{46}$Ti. We observed that the first peak occur at $q=0.7\text{fm}^{-1}$,the second at $1.5\text{ fm}^{-1}$ and the third peak at $2.5\text{fm}^{-1}$. It is noticed from figure.(1) that the inclusion of cp enhance the C2 form factor. This enhancement brings the total form factors very close to the experimental data for the first peak $q<1\text{fm}^{-1}$, while the second and third peak are overestimated. We notice that from figure(1) the model space fail to describe the form factors in all moment transfer.

**Figure (1).** Inelastic longitudinal form factors for the transition to the $2^+$ in the $^{46}$Ti the experimental data are taken from ref.[8]

Figure. (2) displays the calculation of the C2 form factors $J^P=2^+,T=2$ at $E_x=0.98\text{MeV}$ for $^{48}$Ti. The first peak occur at $0.7\text{fm}^{-1}$ the second at $1.7\text{ fm}^{-1}$ and the third peak at $3\text{fm}^{-1}$. The model space calculation fail to describe the form factors and the inclusion of the core polarization enhances the calculations and bring the form factors to the experimental values up to momentum transfer $q=1\text{fm}^{-1}$,while the second peak is overestimated and the third peak is underestimated. Sahu et. al[8] have quite successfully reproduced the first two peaks where we get similar results for the first and third peaks.

**Figure (2).** Inelastic longitudinal form factors for the transition to the $2^+$ in the $^{48}$Ti the experimental data are taken from ref.[8]

Figure(3) displays the calculation of the C2 form factors to the $J^P=2^+,T=3$ at $E_x=1.55\text{MeV}$ for $^{50}$Ti. We observe three peaks, the first peak occur at $q=0.7\text{fm}^{-1}$ the second at $q=1.7\text{fm}^{-1}$ and the third peak at $q=3\text{fm}^{-1}$. We notice that the model space calculation fail to describe the form factors in all momentum transfer region. The (cp) effects enhance the C2 form factors in all momentum transfer, where we notice that the results of cp effects give good agreement with exp. data especially for the first peak up to $q=1.3\text{fm}^{-1}$, while the second peak is overestimated and the third peak is underestimated. Sahu et. al[8] observe two peaks and have quite successfully reproduced the first peak, but the second peak is underestimated and they fail to reproduced the third peak. However the results for the second and third peaks in $^{50}$Ti are better in our calculations. The three figures show that the first region of angular momentum gives good agreement for $0.5\leq q\leq 1.2$ by using the core polarization effects,where the results at $q\geq 1.2$ need more modification to get good agreement with the experimental data.

**Figure (3).** Inelastic longitudinal form factors for the transition to the $2^+$ in the $^{50}$Ti the experimental data are taken from ref.[8]
3.2. Form factors for the $0^+$ to $4^+$ transition

Figure (4) compares the calculated and experimental longitudinal C4 form factors for $^{48}\text{Ti}$ at $E_x=1.967\text{MeV}$. The model space calculation underestimates the exp. and the inclusion of cp enhances the calculations. It is seen that the present calculation is quite successful in reproducing the magnitude of the form factors at the first maximum. However, one observes discrepancies in the momentum transfer range $1.3 < q<1.8$.

Raina et al[19] were unable to reproduce the first maximum within their projected Hartree-Fock-Bogoliubov formalism even though they have changed the effective charges from one nucleus to another, where we get a better result.

Figure (5) displays the calculation of the C4 form factors to the $J=F=3$ at $E_x=2.675\text{MeV}$. The cp effects enhance the C4 form factors and the model space fail to describe the data in the form factors as shown by dashed curve. The results of the fp-shell model space with cp effects give good agreement with the exp. data. Raina et al[19] have quite successfully reproduced the first peak in term of shell model within their HFB formalism. However, the results of the first peak are better in our calculation. In these two transitions for $^{48}\text{Ti}$ and $^{50}\text{Ti}$ the effects of core-polarization shows is clear in figure(4) than figure(5) which need more corrections to get good results.

4. Conclusions

The fp-shell models, which can describe static properties and energy level are less successful in describing dynamic properties such as C2 and C4 transition rates and electron scattering form factors. The inclusion of higher-excited configurations by means of (cp) enhances the form factors and brings the theoretical results closer to the experimental data.

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