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Reply to Comment on “alpha decay in the complex-energy shell model” by R. G. Lovas

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We reply to the Comment by Lovas, which concerns the shell model calculations of alpha-decay width of $^{212}$Po. In the Comment, the author claims that the alpha-formation amplitudes obtained in our work “look unusual.” Here we reaffirm the results and conclusions of our original article.

In his Comment, Lovas claims that the amount of clustering $S = 0.011$ obtained in our work [1] is inconsistent with the “classical result” [2, 3] of $S \approx 0.3$, and that the shapes of the formation amplitude $g(R)$ and the modified formation amplitude $G(R)$ computed in our paper [1] are defective.

The hybrid model of Ref. [2] and many cluster models, including those mentioned in the Comment, introduce alpha particle as a separate degree of freedom represented by an additional component in the wave function. While such models can often provide good fits to experimental data, they do not offer a microscopic description of the alpha decay phenomenon that is rooted in the nucleonic picture. While the calculation of the absolute width using the spectroscopic factor $S$ is a valid procedure ($\Gamma = \Gamma^p S$), the actual amount of clustering strongly depends on the model assumptions used. In particular, assuming the single-particle decay width $\Gamma^p = 0.1247 \times 10^{-12}$ MeV [1], for $S = 0.011$ [1] one obtains $\Gamma = 0.137 \times 10^{-14}$ MeV while for $S = 0.3$ [2] one gets $\Gamma = 3.74 \times 10^{-14}$ MeV.

There are significant differences between the assumptions of Ref. [2] and our work [1], which makes it difficult to compare these models directly: (i) our four-body wave function does not contain an additional alpha-cluster component whose amplitude is governed by a phenomenological Hamiltonian; (ii) our calculations are carried out in a shell model space that is vastly larger as compared to that of Ref. [2] (one major shell, including the unusual-parity intruder); (iii) the particle continuum is neglected in [2] while our extended space M4 consists of all resonant states with a width less than 1 MeV; the asymptotic behavior of the unbound resonant states is very different from that of the bound wave functions employed in [2].

In Fig. 1 we show the norm eigenvalues as in our paper [1] and according to the convention of Ref. [2]. It can be seen that this plot resembles Fig. 3 of Ref. [2] and Fig. 3 of Ref. [4]. In particular, we see a stabilization of results with respect to $\Delta R$ for $\nu \geq 12$, and this is consistent with the previous works. We confirm the point made by Lovas that the norm eigenvalues are generally ordered in the increasing node number order.

![FIG. 1. Eigenvalues of the norm kernel for $^{212}$Po for $R_{max} = 13$ fm for different values of $\Delta R$ ordered in (a) decreasing eigenvalue index as in Ref. [1] and (b) increasing eigenvalue index as in Ref. [2].](image)

We agree with Lovas that understanding of the eigenvalue problem of the norm operator is essential. As far as the small eigenvalues of the norm kernel ($n_v < 10^{-3}$) are concerned, one can notice that they are strictly zero when equal oscillator lengths are used for the alpha-particle wave functions and the shell-model basis, see Refs. [5, 6]. As stated in our paper, below Fig. 1, the omission of the small norm eigenvalues does impact the inner region. This is again consistent with Lovas's comment. However, as stated in Ref. [4], ‘States with those small eigenvalues
are partially forbidden’. In particular, they found that the allowed states must have at least 11 nodes. This observation is consistent with the finding of [1]: “To eliminate these spurious eigenvectors, we define the cutoff at the value where the eigenvalue distribution changes slope”. But – in addition to physical arguments – one also needs to consider numerical aspects of the problem. Indeed, as pointed out in our paper (cf. discussion around Figs. 5-7 and Eq. (49) therein), the spectroscopic factor (S) does depend on both ΔR and R_{max}. In particular, Fig. 5 of [1] shows the sensitivity of S as a function of R_{max} for various values of ΔR for n_{min} = 0.001. The new Fig. 2 illustrates the situation for n_{min} = 0.00016. Here, it is impossible to obtain a result that is independent on the parameters ΔR and R_{max} of the shifted Gaussian basis. So we are not talking about “slight numerical inaccuracies” here.

\[ S = \sum_{\nu} \frac{g^2_{\nu}}{n_{\nu}} \]  

(1)

While Eq. (1) is mathematically correct, it is numerically extremely unstable since the eigenvalues n_{\nu} rapidly decrease to zero. To this end, in Ref. [1] we have proposed a systematic procedure to determine the norm eigenvalue cutoff n_{min} to be sure that a local plateau in ΔR and R_{max} is achieved. We wish to emphasize that the basis of the norm kernel eigenstates is orthonormal with an accuracy of 10^{-10} for all eigenvalues so there is no question about the precision of the expansion of g(R). The lowest n_{\nu}-values obtained are around and below 10^{-11}; hence, they are practically impossible to control numerically. It is worth noting that an important consistency check is provided by the expansion of g(R) in the eigenfunctions of the norm kernel (see Fig. 4 in our paper) and by the asymptotic behavior of the overlap integral as it is determined by the Q_{o}-value of ^{212}Po. Moreover, the R-matrix result is not that far from the alpha spectroscopic factor approach.

As demonstrated in our work (see, e.g., Fig. 8 of [1]), the shape of the formation amplitudes strongly depends on the configuration space employed. We have difficulty to accept Lovas’s statement that the function g(R) is bound to have at least twelve nodes. Indeed, while it can be proven that \( \sum_{i=1}^{m} a_i f_i(r) \) has at least s nodes if f_i(r) are orthogonal polynomials [8], we are not aware of a similar theorem for a general orthogonal set, including functions that are not localized. If the formation amplitude is computed in the harmonic oscillator basis, or in a basis of localized states confined to a finite box, Lovas’s arguments would obviously hold. However, our Woods-Saxon basis includes resonant states, to which the theorem [8] does not apply. Consequently, we do not see how the absence of some nodes in the inner region of g(R) could indicate that our calculations are defective.

In his comment, Lovas has neither demonstrated flaws in the shell-model formalism employed, nor has he presented a solid evidence against our results. He has merely pointed out that some of our findings are not consistent with his previous work and schematic arguments. As discussed in the conclusions of our paper, much work still needs to be done on both the modeling and algorithmic side. But we believe that our results, and related discussion, represent a step toward microscopic understanding of alpha decay, without invoking an explicit cluster component.

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