The absolute position of a resonance peak

Shachar Klaiman$^1$ and Nimrod Moiseyev$^{1,2}$

$^1$ Schultich Faculty of Chemistry and Minerva Center for Nonlinear Physics of Complex Systems, Technion–Israel Institute of Technology, Haifa 32000, Israel
$^2$ Department of Physics, Technion–Israel Institute of Technology, Haifa 32000, Israel

E-mail: shachark@technion.ac.il

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Abstract

It is common practice in scattering theory to correlate the position of a resonance peak in the cross section to the real part of a complex energy of a pole of the scattering amplitude. In this work, we show that the resonance peak position appears at the absolute value of the pole’s complex energy rather than its real part. We further demonstrate that a local theory of resonances can still be used even in the cases previously thought to be impossible.

There are few phenomena in physics which are as striking and ubiquitous as resonance peaks in the cross section. As such, it is of great importance to understand their origin and to be able to predict the resulting resonance profile in the cross section. Over the years, many methods have been proposed and successfully used to analyse and explain resonance peaks appearing in the cross section. Among which we mention the use of the poles of the scattering matrix (S-matrix) [1], the Fano configuration interaction [2] and the Feshbach resonance theory [3].

The main purpose of such theories is to reconstruct the resonance structure appearing in the cross section. This should be done with but a few parameters which can be used to characterize the scattering system at hand. It is in the way these parameters are chosen and calculated that the methods mentioned differ from each other. One such difference is the use of non-local versus local parameters, i.e. whether the system parameters used to describe the resonance profile are energy dependent or not.

The most famous application of a local resonance theory is the Breit–Wigner (BW) resonance profile [4], which connects an appearing resonance peak in the cross section with a pole of the S-matrix in the complex energy plane. The resulting resonance profile reads

$$
\sigma(E) \propto \frac{(\Gamma/2)^2}{(E - \varepsilon)^2 + (\Gamma/2)^2},
$$

where $\varepsilon$ is known as the resonance position and is given by the real part of the pole in the energy plane and $\Gamma$ is the width of the Lorentzian at half-maximum which is twice the imaginary part of the resonance pole in the energy plane. Although the above resonance profile has a very impressive track record, it fails whenever the pole is not isolated (by the effect of other poles), it is near the threshold, or it is far from the real axis. One possible resolution to this predicament is to move into a non-local description and define the energy-dependent width and position such that the resonance profile now reads

$$
\sigma(E) \propto \frac{(\Gamma(E)/2)^2}{(E - \varepsilon(E))^2 + (\Gamma(E)/2)^2}.
$$

Although clearly able to reproduce any resonance structure, it is neither straightforward nor simple to calculate the non-local terms in the above equation for an arbitrary scattering potential.

The use of the poles of the scattering matrix was reinvigorated by the novel method presented by Tolstikhin et al [7]. The possibility of solving the Schrödinger equation with Siegert boundary conditions [8] to obtain all the poles of the scattering matrix opened the door to describe the entire cross section using only the poles. Tracing the origin of the structures in the cross section to specific poles of the matrix has always been invaluable in order to understand the scattering mechanisms responsible for the resulting cross section. It would, therefore, be extremely beneficial if one could assign to every resonance in the cross section a specific pole. It is the failure of this possibility that necessitates the use of non-local theories and, as we will demonstrate, there is still much to be done using only local resonance profiles.

In order to demystify the above discussion, let us start with the simplest example where one cannot connect the poles of the S-matrix and the resonance appearing in the cross section using the assumptions of the BW profile. Consider a one-dimensional (1D) square well. It is well known (see for example [5]) that the transmission spectrum displays an oscillating behaviour and the transmission probability reaches...
Figure 1. The transmission probability above a square well potential of depth $V_0 = -13$ (au) and length $L = \pi/\sqrt{2}$. The full (green) and dashed (red) arrows mark the real part and absolute value of the resonance poles energy respectively. The table in the inset shows the values of the first three energies, $E_n$, of the resonance poles, the analytical position, $\lambda_n$, of the unites, and the absolute values of the resonance energies. All quantities are given in atomic units for which $\hbar = 1$ and $m = 1$.

unit value only at discrete energies satisfying $\lambda_n = -V_0 + \frac{\hbar^2\pi^2n^2}{2mL^2}$, such that $T(\lambda_n) = 1$. In the above equation $V_0$ and $L$ are the well’s depth and length respectively, $m$ is the mass of the projectile and $n$ is a positive integer. Figure 1 depicts the transmission for a specific choice of the well’s depth and length. The poles of the scattering amplitude can be calculated by solving the Schrödinger equation with outgoing boundary conditions [6]. The positions of the first three resonance poles, i.e. the real part of the complex energy of the poles appearing in the fourth quadrant, are depicted in figure 1 using full (green) arrows. As can be readily observed, one cannot directly assign a transmission unity with the real part of the resonance pole’s energy. At first this may seem obvious since the resonance poles clearly overlap, no direct connection between a single energy. At first this may seem obvious since the resonance peak has a maximum at the absolute value of the complex resonance pole energy. We have already demonstrated in the fourth quadrant, are depicted in figure 1 using full (green) arrows indicating the absolute values of the complex resonance poles’ energies.

Let us consider a scattering process from a single 1D channel. As mentioned above, the transmission amplitude can be expressed using all the poles of the scattering matrix [7]. These are found as the solutions of the Schrödinger equation: $\left[-\frac{\hbar^2}{2m}\frac{d^2}{dx^2} + V(x)\right]\Psi_n(x) = \frac{\hbar^2}{2m}\Psi_n(x)$ along with Siegert boundary conditions imposed at $x = \pm L$ which satisfy [8] $\left(\partial_x \Psi_n(\pm L) \mp ik_n \Psi_n(\pm L)\right) = 0$. Here we use atomic units for which $\hbar = 1$. We choose to normalize the Siegert states such that $2i\kappa_0 \int_{-L}^{L} \Psi_n^*(x)dx \left(-\frac{\Psi_n^2(x) + \Psi_n^2(L)}{2} \right) = 1$. For all but piece-wise potentials, the solutions must converge with respect to $L$. One can categorize the Siegert solutions according to where they fall in the complex $k$-plane: bound (anti-bound) solutions on the positive (negative) imaginary axis and resonance (anti-resonance) solutions on the fourth (third) quadrant. Using the relation $E_n = k_n^2/2$, the solutions’ energy can be found.

Using the Siegert states solutions, the transmission amplitude for a one-dimensional potential can be written as [9] $t = 2k e^{-2kL} \sum_{n=1}^{\infty} \frac{\Psi_n(L)\Psi_n(-L)}{k_n - k^2}$, where the sum runs over all poles, i.e. bound, anti-bound, virtual and resonance poles, and $k = \sqrt{2E}$. The above sum can be somewhat simplified by transforming it into a product. Such a product formula was first given by Tolstikhin et al in [9] for symmetric potentials and by Ostrovsky and Elander in [10] for asymmetric potentials. For symmetric potentials, the product formula reads $t = \frac{1}{\sqrt{2}} e^{-2kL} \prod_{n=1}^{\infty} \frac{k_n^2 - k^2}{k_n^2 - k^2}$, where $N_s (N_v)$ correspond to symmetric (anti-symmetric) Siegert states. Since the poles $k_n$ are either purely imaginary or come in pairs $k_n$ and $-k_n^*$, the above product terms can each be written as a pure phase, the well-known Blaschke factor [11]. Considering first the pairs of resonance and virtual states, one can reformulate the product of each pair as $\frac{k_n^2 - k^2}{k_n^2 - k^2} = e^{2i\delta_n}$, where after some trigonometric manipulations, one finds that $\delta_n = \arctan(\frac{k}{\Im|k_n|}|\Gamma_{1n}/\Gamma_{2n}|)$ with the common definition of the resonance energy $E_n = k_n^2/2 = \epsilon_n - i2\Gamma$. For the bound and anti-bound poles, each term in the products reads $(k + k_n^2)/(k - k_n^2) = e^{2i\delta_n}$, where for these poles $\delta_n = \arctan(\Im|k_n|/k)$. Defining $\Delta_n = \sum_{n=1}^{N_s} \delta_n$, where the sum runs over all bound poles, anti-bound poles and resonance-virtual pole pairs, the transmission amplitude can be brought to the following form, $t = e^{-2kL} e^{i(\Delta_n + \Delta_v) \sin(\Delta_n - \Delta_v)}$, and the transmission probability reads $T = \sin^2(\Delta_n - \Delta_v)$. Using the above formula for the transmission probability, we can examine the contribution of a single resonance pole to the cross section. Without loss of generality, let us assume that the corresponding Siegert eigenstate is even. Then neglecting the contribution of all but a single resonance-virtual pair pole, the transmission probability reads

$$T \approx \frac{(\frac{\kappa}{2})^2(\Gamma/2)^2}{(E - |E_{res}|)^2 + (\frac{\kappa}{2})^2(\Gamma/2)^2}.$$  

where the resonance pole is located at $k_{res} = k_v - i\kappa_v$, $E_{res} = \frac{k_v^2}{2} = \epsilon + \frac{i}{2}\Gamma$, $k = \sqrt{2E}$, and we have used the trigonometric identity: $\sin(\arctan(x)) = \frac{x}{\sqrt{1+x^2}}$. On first sight, equation (2) is very reminiscent of the BW resonance profile, see equation (1). There are, however, several crucial differences. First and foremost, the resonance peak has a maximum at the absolute value of the complex resonance pole energy. We have already demonstrated in the above example that the transmission units above a square well potential fall on the absolute value of the resonance complex energy rather than on the expected real part (see figure 1). This, for the first time, allows the connection of each transmission unity with a single resonance pole even in the case of wide overlapping resonances. Note that for overlapping resonances one cannot hope by definition to use a single resonance profile.
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understanding the resulting transmission probability.

The resonance profile in equation (2) also differs from the

BW profile in that it is not a Lorentzian and is asymmetric

with respect to its maximum, rising sharply to the left of the

maximum and falling more slowly to the right of the maximum.

Even though extension of the BW profile into an asymmetric

one is known [1], the current profile formula characterizes the

asymmetry only with the resonance complex energy, i.e. with

the same information needed to construct the BW profile and
do not require the calculation of additional parameters.

Wide overlapping resonances are not the only case where

the local description of resonance is abandoned for a non-local

one. For resonances near the threshold, a similar belief that

one cannot correlate between a resonance peak and a single

pole exists. However, the resonance profile developed here

shows that even very close to the threshold the resonance peak

is due to a single resonance pole and one can reconstruct the

resonance peak in the cross section using only the resonance

pole energy. Figure 2 depicts the transmission probability

near an isolated resonance in a double barrier potential of the

form \( V(x) = (\beta x^2 - \gamma) e^{-\alpha x} \) (see the inset). Clearly, the

profile in equation (2) reproduces the peak in the transmission probability

very well using the same information used in the BW resonance profile which as seen in the figure fails to

reproduce the BW resonance peak quantitatively. Thus one does

not need to resort to non-local theories to describe resonance

peaks near the threshold even when, as is in the case portrayed, the imaginary part and real part of the resonance energy are comparable in magnitude.

We turn now to discuss yet another important case where

the local approximation has be erroneously discarded. In his

seminal paper, Nussenzveig [12] studied the effect of varying

the potential on the poles of the scattering matrix. One of the most studied aspects has been the behaviour of the poles

as the potential is varied to support an additional bound state. It is well established that for a single \( s \)-wave-like channel, looking in the \( k \)-plane, the resonance poles located in the fourth quadrant and the virtual state poles located in the third quadrant move towards the negative imaginary \( k \)-axis as the potential well is, for example, deepened. Eventually, two poles approaching from either side of the imaginary axis coalesce, thereafter, forming two anti-bound state poles which lie on the negative imaginary axis. Following the creation of two anti-bound poles, one of these poles moves up towards the origin and eventually leads to the formation of a new bound state, while the other moves down on the imaginary axis.

The connection between the motion of the resonance pole described above and the corresponding motion of the resonance peak in the cross section remains somewhat elusive, even today. The main problem that arises in the analysis is that at some point, while approaching the negative imaginary axis, the resonance pole crosses the \( 45^\circ \) bisector of the fourth quadrant of the \( k \)-plane making the resonance position

negative. The resonance peak in the cross section, however, is still observable and is clearly situated above the threshold. Nussenzveig deemed these resonance poles with the negative position as the only ‘meaningless’ [12] poles. As we shall show, they are just as meaningful as any other pole of the scattering matrix.

Consider the effect of deepening the well between the two barriers (see the potential depicted in figure 2) on the transmission spectrum. At some critical well depth, a new bound state appears. We study the behaviour of the resonance peak in the transmission spectrum as the potential depth is increased towards this critical value. Previous studies noted that as the well deepens, the resonance peak approaches the origin. Figure 3 portrays the first resonance peak in the transmission spectra for three different depths of the potential well. As can be seen in the figure, the resonance peak moves towards the origin as we approach the critical value for which a new bound state is formed. Aside from the obvious shift in position, the different resonance peaks look almost the same. This is rather surprising since, as we explain in the following, the potential responsible for the leftmost peak (marked (c)) does not support a resonance pole in the vicinity of the position of the peak.

Let us examine first the two peaks in figure 3 marked (a) and (b). The potential responsible for peak (a) supports a resonance pole above the threshold similar to that depicted in figure 2. As can be seen, the resonance profile given in equation (2) (dashed (red) line) reproduces the numerically exact resonance peak in the spectrum (solid (black) line). For the peak marked (b), however, the potential is such that the resonance pole moved passed the \( 45^\circ \) bisector of the fourth quadrant of the \( k \)-plane (see the inset in figure 3). Therefore, the position of the resonances pole is now negative. As the figure shows, the profile in equation (2) can be used very successfully to reproduce the resonance peak using only the resonance pole with the negative position. This puts the resonance poles with negative positions on the same footing as other resonance poles of the scattering matrix. The resonance pole’s trajectory in the complex \( k \)-plane brings about
yet another difficulty. As the pole approaches the negative imaginary axis, the resonance width defined as \( \Gamma = 2k_r k_i \) goes to zero, since \( k_r \to 0 \). One might therefore conclude that such narrow resonances will be very hard to observe. Such a misconception stems from the expectation of a BW profile whose full-width half-maximum (FWHM) is \( \Gamma \). The profile presented in equation (2), however, is asymmetric and has a FWHM of \( \sqrt{\Gamma^2 + 2(k_r^2 + k_i^2)} \). For, over the threshold, narrow resonances this is approximately \( \Gamma \). However, for negative position resonances close to the negative imaginary axis, one can no longer neglect the change in the FWHM. For the peak marked (b) in figure 3, the resonance pole width is of the order of \( 10^{-3} \) whereas the peak appearing in the figure has a FWHM of the order of \( 10^{-2} \). We re-emphasize that the resonance peak, position and FWHM are characterized using only the resonance pole complex energy, i.e. the same information used in the BW profile.

Further increase in the depth of the well, past the branch point where the two poles coalesced, produces two anti-bound states. Before proceeding, we point out that even though after the coalescence of the resonance and virtual poles to two anti-bound states left no resonance pole no longer exists, the peak is still apparent at \( k^{AB} \). The remarkable agreement between the above formula and the numerically exact transmission can be witnessed in figure 3.

The resonance profile in equation (2) was developed under the assumption of a symmetric one-dimensional potential. We now discuss briefly the generalization of this result. One rather straightforward extension can be made for s-wave resonances. These resonances are the Siegert solutions to a radial 1D potential. The expansion of the outgoing Green’s function for such a single radial channel has been given elsewhere [7]. Following the same procedure described above, one can readily find that the corresponding isolated low-energy resonance profile is identical to the one given in equation (2). Note that the low-energy restriction is necessary for the BW profile as well [1]. A second extension concerns asymmetric 1D potentials. Here the situation is somewhat more complicated since the available product formula is of limited use. Still, many tested examples show that resonance profiles in the transmission through asymmetric potentials are proportional to the resonance profile given in equation (2). Further study is underway.

In summary, we have demonstrated that a local theory of resonances can still be used even in cases previously thought impossible. Our analysis allows one to unequivocally connect the resonance peak in the transmission to specific poles of the scattering matrix throughout various changes of the scattering potential. Even for wide overlapping resonances, one can connect each transmission unity with a single resonance pole.

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