

PHY 982 Homework 2

John Ash, Mengzhi Chen, Tong Li, Jason Surbrook

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1 Elastic scattering calculations of a nucleon on ^{208}Pb target

1.1 Coulomb scattering

In the case of point-Coulomb scattering, the Coulomb interaction between the projectile and the target with charges Z_1 and Z_2 is

$$V_c(R) = \frac{Z_1 Z_2 e^2}{R^2}, \quad (1)$$

where R is the distance between them. The Schrödinger's equation with this potential can be solved analytically. Related discussions are detailed in Ref.[1]. In sum, the angular distribution of point-Coulomb potential is described by the differential cross section

$$\sigma(\theta) = \frac{\eta^2}{4k^2 \sin^4(\theta/2)}; \quad \eta = \frac{Z_1 Z_2 e^2}{\hbar} \left(\frac{\mu}{2E} \right)^{\frac{1}{2}}, \quad (2)$$

where μ is the reduced mass and k is the wave number with the energy E . It worth noticing that the Eq. 2 is the same as the classical *Rutherford cross section*.

When the finite size of the target is included, due to different charge distributions, the interaction between the projectile and the target becomes more complicated.

Normally, these potentials are not analytically solvable. Sometimes, we can apply the *plane wave Born approximation* (PWBA) for the Coulomb scattering. Under PWBA, the differential cross section is

$$\sigma(\theta) = \frac{\eta^2}{4k^2 \sin^4(\theta/2)} |F(\theta)|^2, \quad (3)$$

where $F(\theta)$ is called the *form factor* related with the charge distribution as

$$F(\theta) = \int e^{i(\vec{k}_f - \vec{k}_i) \cdot \vec{r}'} \rho(r') d^3 r'. \quad (4)$$

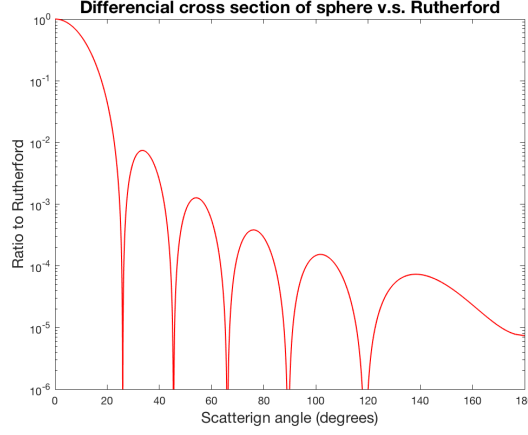


Figure 1: Differential cross sections for solid sphere charge distribution. Here we take $a = 1; k = 10$.

1.2 Form factor of a solid sphere

To see substantial influence of the form factor, we give an simple example. Suppose the charge distribution of the target with charge Z_2 is homogenous inside a solid sphere with radius a , which means

$$\rho(R) = \frac{3}{4\pi a^3} \text{ for } R < a \quad (5)$$

$$= 0 \text{ elsewhere.} \quad (6)$$

The form factor is then calculated from Eq. 4

$$F(\theta) = \frac{3}{qa} \left(\frac{\sin(qa)}{(qa)^2} - \frac{\cos(qa)}{qa} \right) = \frac{3j_1(qa)}{qa}. \quad (7)$$

where j_1 is spherical Bessel functions and $q = 2k \sin(\theta/2)$. In Figure 1, we show the differential cross sections' ratio between these two cases. We can see fluctuation appears due to diffraction of finite-size body.

In sum, we see the finite-size target can give cross sections different from the point-like case.

1.3 Description of elastic scattering using optical potentials

In this section, we study the elastic scattering between nucleons and the ^{208}Pb target with the help of FRESKO[2]. Their effective interaction is referred to as the optical potential. We take the global parameterizations from Ref. [3, 4] as the input for FRESKO.

In the first step, we take two lab energies 5 and 50 (40) MeV for proton (neutron) in our calculations. The results together with experimental data are shown in Figure 2. It

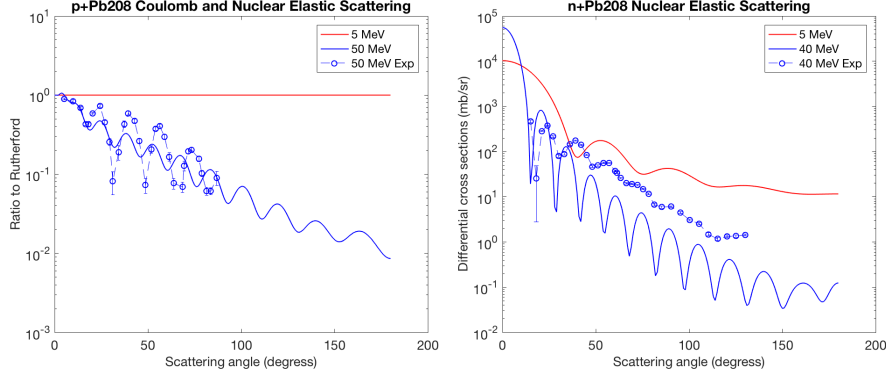


Figure 2: Differential cross sections for nucleons scattering with ^{208}Pb at 5 and 50 MeV. Left panel is proton and right panel is neutron

gives the angular distributions in center of mass for protons and neutrons. For the 5 MeV proton, the curve agrees well with the *Rutherford cross section*. Given that the radius parameter of optical potential is $r_w \sim 1.2 \text{ fm}$, we can estimate the Coulomb barrier for a proton to overcome. It should be

$$E_{Coul} \approx \frac{Z_1 Z_2 e^2}{r_w + R_2} \approx 15 \text{ MeV}. \quad (8)$$

We can see that 5 MeV is much lower than the Coulomb barrier. Thus, this scattering can be referred as a point-like Coulomb scattering properly. The agreement with the *Rutherford cross section* also verifies this point. For neutrons, there is no Coulomb interaction, hence the value of the neutron cross section being listed as fm^2 . At 5 MeV the cross section hints at a broad diffraction pattern.

Since 50 MeV is strong enough to overcome the Coulomb barrier, the effective interaction plays a role and induces a diffraction pattern in Figure 2. The neutron cross section at 40 MeV exhibits a similar behavior, speaking to the dominance of the nuclear interaction at this energy. We can also see that our curves basically obey same trends as experiments. However, they deviate in values, especially for neutron. It is because we use the global parameterizations. We will include the qualitative analyses and better fittings in the next section. In order to explore the influence of the optical potential, we will keep using these energies in following calculations.

In the next step, we would like to see the influences of the depth of optical potential's imaginary part of the S-matrix. Fixing other parameters, we scale the depth of imaginary parts V_i (including volume, surface derivative and spin-orbit potentials) a factor of 1.5, 0.5 and 0. The modules of S-matrix for the proton and the neutron are shown in Table 1 and 2.

We can see that as we lower the depth, $|\mathbf{S}|^2$ fluctuates and eventually converges to one which indicates an elastic scattering. It inspires us that the imaginary parts can describe

Proton partial waves					
L	J	$ \mathbf{S} ^2(\bar{V}=1.5)$	$ \mathbf{S} ^2(\bar{V}=1.0)$	$ \mathbf{S} ^2(\bar{V}=0.5)$	$ \mathbf{S} ^2(\bar{V}=0.0)$
0	0.5	6.45E-04	2.33E-04	1.56E-02	1.0
1	0.5	9.17E-04	1.17E-03	1.63E-02	1.0
2	1.5	8.33E-04	1.28E-04	1.34E-02	1.0
1	1.5	9.19E-04	1.23E-03	1.74E-02	1.0
2	2.5	7.97E-04	1.17E-04	1.52E-02	1.0
3	2.5	1.27E-03	1.64E-03	1.81E-02	1.0
4	3.5	1.51E-03	2.89E-04	1.02E-02	1.0
3	3.5	1.27E-03	1.85E-03	2.11E-02	1.0
...

Table 1: The modules of S-matrix for proton partial waves with varying imaginary potentials, where \bar{V} is the scaled depth defined as $V_i(new)/V_i(initial)$.

absorptions in scattering processes. It also worth noticing the fluctuation of $|\mathbf{S}|^2$ with varying V_i in each partial wave. For this reason, there is no simple monotonous relations between the relative depth \bar{V} and the total absorptive cross section.

Moreover, we are interested in the effects brought by different radius parameters. In analogy with the Fresnel circular disk diffraction, as distance between disk and screen increases, the rings become denser. We can also check the special situation in Eq. 7, since $j_1(x)$ has fixed zero points, then, as radius parameter increases, we expect more compact diffraction patterns. We don't scale the Coulomb interaction with optical potentials at same time, so we should see stronger up-down shift for proton caused by it.

Again, fixing other parameters, we repeat our calculations by scaling the radius parameters r (including volume, surface derivative and spin-orbit potentials) a factor of 0.2, 0.5, 2, 3 and 4. The results are shown in Figure 3. For the proton, we observe a dramatic up-down shifts and denser diffraction patterns with increasing r . It's the synergistic effect by the Coulomb and optical potentials. As for neutron, the amplitude changing of pattern is relative small as it only feels the optical potential. Although the interactions are more complicated than the solid sphere, the trends in these two figures agree well with our anticipations.

The total reaction cross section and the absorptive cross section are equal in a single channel elastic scattering. For neutron, we extract them according to the equation

$$\sigma_e = \frac{\pi}{k^2} \sum_J (2J+1)(1 - |\mathbf{S}_J|^2). \quad (9)$$

The total reaction and absorptive cross sections are 3.764×10^3 mb. The number deviates little from the value 3.796×10^3 mb given directly by FRESCO for slightly different physical constants used.

Neutron partial waves					
L	J	$ \mathbf{S} ^2(\bar{V}=1.5)$	$ \mathbf{S} ^2(\bar{V}=1.0)$	$ \mathbf{S} ^2(\bar{V}=0.5)$	$ \mathbf{S} ^2(\bar{V}=0.0)$
0	0.5	7.88E-04	1.40E-03	2.16E-02	1.0
1	0.5	5.24E-04	4.68E-05	1.75E-02	1.0
2	1.5	8.94E-04	1.62E-03	2.29E-02	1.0
1	1.5	5.24E-04	4.68E-05	1.75E-02	1.0
2	2.5	8.94E-04	1.62E-03	2.29E-02	1.0
3	2.5	7.27E-04	1.66E-05	1.67E-02	1.0
4	3.5	1.15E-03	1.87E-03	2.51E-02	1.0
3	3.5	7.27E-04	1.66E-05	1.67E-02	1.0
...

Table 2: The modules of S-matrix for neutron partial waves with varying imaginary potentials, where \bar{V} is the scaled depth defined as $V_i(new)/V_i(initial)$.

2 Fit optical potentials for elastic scattering of a nucleon on ^{208}Pb target

In this section we will do χ^2 data fitting to obtain optical potentials for the elastic scattering of a proton or neutron on ^{208}Pb . Experimental data are taken from Ref. [5] for proton and [6] for neutron. The beam energies of proton and neutron are 49.35 MeV and 40.0 MeV, respectively. In addition, we choose the optical parameters employed in Sec. 1 (Ref. [3, 4]) as the starting point of our fitting. All the results discussed in this section are generated by SFRESCO (Ref. [2]).

2.1 Fit process and results

All the intermediate and final results of our fitting are summarized in Table 3 (for proton) and Table 4 (for neutron). For simplicity, we neglect spin-orbit components at the beginning, and only volume real and volume imaginary components are included. Keeping the volume imaginary component unchanged, we fit the volume real part (index 2 in Table 3 and 4), which cannot reduce χ^2/N to a low value. Then, we vary both volume real and imaginary parts simultaneously (index 3 in Table 3 and 4) and achieve a significant improvement. However, for proton the diffuseness parameter av becomes too small to be considered physical.

Another choice of optical potential is to use a surface imaginary component instead of a volume one. Starting from parametrization of index 4 in Table 3 and 4, we fit the volume real part and surface imaginary part (index 5 in Table 3 and 4). For both proton and neutron, a surface imaginary component gives a lower χ^2/N , and thus we will add

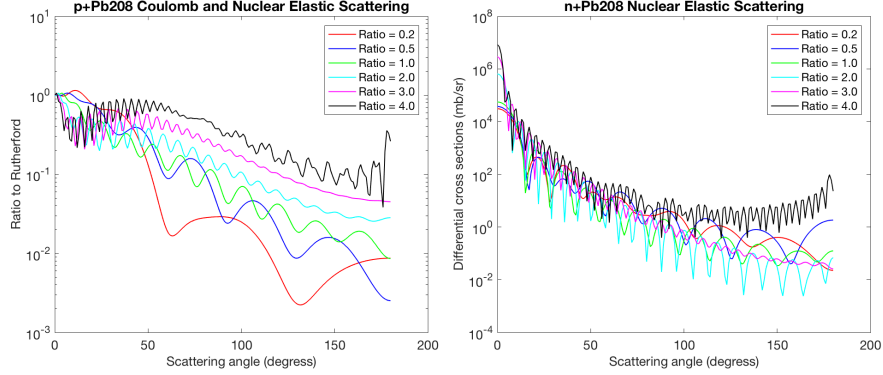


Figure 3: Differential cross sections for nucleons scattering with ^{208}Pb with different radius parameters at 50MeV, where ratio is defined as $r_{\text{new}}/r_{\text{initial}}$. Left panel is proton and right panel is neutron.

spin-orbit terms onto this optical potential form.

As shown in the last four lines in Table 3 and 4, we gradually add and fit different parameters in the spin-orbit term. As for proton, the description can hardly be improved by only adding a real spin-orbit potential (index 6 in Table 3). If we only add a imaginary spin-orbit component, the diffuseness parameter “awso” will become negative and thus unphysical (index 8 in Table 3), so we abandon this set of parameters. If both real and imaginary components of spin-orbit potential are included, χ^2/N will slightly decrease from 4.860 to 3.611, (index 9 in Table 3), which indicates that the spin-orbit term is not very important for the description of elastic scattering of a proton on ^{208}Pb .

As for neutron, the radius parameter “rvso” will become negative and unphysical when only a real spin-orbit component is included (index 6 in Table 4), so we abandon this set of parameters. By only adding an imaginary spin-orbit component, χ^2/N is slightly lowered from 4.638 to 3.310 (index 8 in Table 4). When both real and imaginary spin-orbit parts are varied, χ^2/N can be even lower (index 9 in Table 4). But in this case the diffuseness “avso” will become less than radius “rvso”, which is unphysical and unacceptable. As no significant improvement is seen from adding spin-orbit terms, we think that the spin-orbit term is not important for the description of elastic scattering of a neutron on ^{208}Pb .

Table 3: Fitting results of scattering of a proton on ^{208}Pb . The unit of V_v , W_v , W_s , V_{so} and W_{so} is MeV, and the unit of r_v , a_v , r_{wv} , a_{wv} , r_{ws} , a_{ws} , r_{vso} , a_{vso} , r_{wso} and a_{wso} is fm. Unphysical values are highlighted in red.

Index	Fit from	V_v	r_v	a_v	W_v	r_{wv}	a_{wv}	W_s	r_{ws}	a_{ws}	V_{so}	r_{vso}	a_{vso}	W_{so}	r_{wso}	a_{wso}	χ^2/N
1	-	67.2	1.244	0.646	16.6	1.244	0.646	-	-	-	-	-	-	-	-	-	78.285
2	1	46.1	1.262	0.574	16.6	1.244	0.646	-	-	-	-	-	-	-	-	-	27.134
3	2	44.6	1.254	1.87E-04	4.89	1.726	0.171	-	-	-	-	-	-	-	-	-	5.479
4	-	46.1	1.262	0.574	-	-	-	19.5	1.246	0.615	-	-	-	-	-	-	11.335
5	4	45.3	1.229	0.528	-	-	-	13.2	1.295	0.713	-	-	-	-	-	-	4.860
6	5	45.2	1.232	0.489	-	-	-	12.7	1.295	0.732	0.14	1.07	0.55	-	-	-	4.838
7	-	45.2	1.232	0.489	-	-	-	12.7	1.295	0.732	-	-	-	-3.1	1.08	0.57	12.026
8	7	45.1	1.242	0.507	-	-	-	12.0	1.279	0.748	-	-	-	-2.1	0.66	-0.028	4.068
9	7	45.2	1.276	0.540	-	-	-	15.7	1.239	0.710	-6.9	0.84	0.55	-8.0	1.03	0.53	3.661

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Table 4: Fitting results of scattering of a neutron on ^{208}Pb . The unit of V_v , W_v , W_s , V_{so} and W_{so} is MeV, and the unit of r_v , a_v , r_{wv} , a_{wv} , r_{ws} , a_{ws} , r_{vso} , a_{vso} , r_{wso} and a_{wso} is fm. Unphysical values are highlighted in red.

index	Fit from	V_v	r_v	a_v	W_v	r_{wv}	a_{wv}	W_s	r_{ws}	a_{ws}	V_{so}	r_{vso}	a_{vso}	W_{so}	r_{wso}	a_{so}	χ^2/N
1	-	50.6	1.244	0.646	15.6	1.244	0.646	-	-	-	-	-	-	-	-	-	261.394
2	1	40.3	1.245	0.903	15.6	1.244	0.646	-	-	-	-	-	-	-	-	-	46.553
3	2	41.9	1.154	0.762	6.598	1.383	0.824	-	-	-	-	-	-	-	-	-	7.900
4	-	40.3	1.245	0.903	-	-	-	13.8	1.246	0.510	-	-	-	-	-	-	98.423
5	4	36.4	1.269	0.585	-	-	-	14.4	1.097	0.500	-	-	-	-	-	-	4.638
6	5	36.1	1.277	0.578	-	-	-	15.7	1.108	0.498	8.3	-4.3	3.2	-	-	-	3.084
7	-	36.4	1.269	0.585	-	-	-	14.4	1.097	0.500	-	-	-	-3.1	1.08	0.57	35.163
8	7	36.4	1.273	0.593	-	-	-	15.2	1.109	0.519	-	-	-	-3.1	1.09	0.43	3.310
9	8	35.8	1.266	0.592	-	-	-	14.9	1.107	0.513	4.9	1.03	1.23	-3.0	1.06	0.81	2.258

2.2 Fit Sensitivity to Initialization

Here we study the sensitivity of SFRESCO fitting to initial parameters. We used volume real and surface imaginary components of the optical potential only. Although there is improvement in the neutron scattering fit by inclusion of a spin-orbit imaginary term, optimal fits without this term are roughly 40% of the fit with this additional term ($\chi^2/N = 3.310, 4.638$). Optimal initial parameters were determined by successive fitting and reinitializing output parameters until a stable minimum was achieved with parameters comparable to the initial global potential fits from [3].

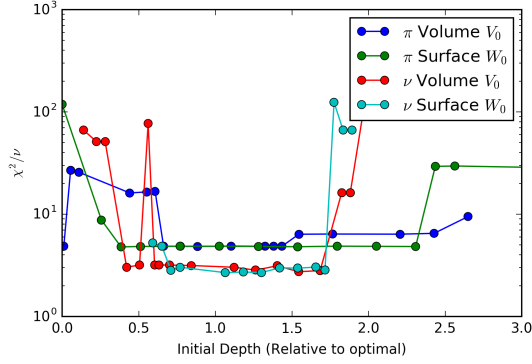
In Figure 4 below, we plot the SFRESCO χ^2/N fit error against modifications of a single parameterization. To overlay dissimilar parameter values, the plotted input parameter is normalized by its determined optimized value. χ^2/N as a choice of parameter of interest (as opposed to the output value of the varied parameter) was from the observation that χ^2/N correlates strongly with optical potential parameters approaching the optimal fit values within a few percent. From Fig 4a, we see that initializations to both depth parameters for protons and neutrons arrive at optimum for roughly $\pm 50\%$ of optimal values. From Fig. 4b, While not as centered around $\frac{P_{in}}{P_{optimal}} = 1$ as the depth parameters, the radius parameters appear to converge for roughly $\pm 25\%$ of optimal values. In Fig. 4c, we demonstrate that diffuseness is potentially the most convergent parameter (see neutron surface imaginary converge for $\pm 100\%$ of the optimal value), diffuseness is the least consistent parameter. The worst demonstrated convergence is the proton surface imaginary diffuseness, which converges for $\frac{P_{in}}{P_{optimal}} = 1.0 - 0.0 + 0.5$.

Two peculiarities were discovered while spanning these parameters.

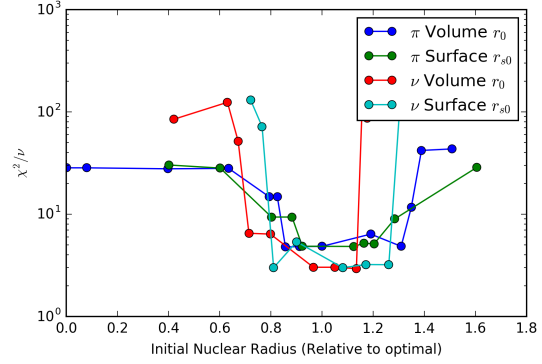
1. Multiple initializations regions of the proton real volume depth lead to optimal parameterization. Fig. 4a shows how $V_0 \approx 0$ gives results consistent with $V_0 \approx V_{optimal}$ while the region $0.1 \leq \frac{V_0}{V_{optimal}} \leq 0.6$ returns a significantly worse fit.
2. There appears to be a numerical instability for the neutron real volume depth $\frac{V_0}{V_{optimal}} = 0.55 \pm < 0.05$. This instability sits in a region of efficient optimal parameter finding, and has width of < 3 MeV. Given the approximate nature of the global potentials that were used to arrive at optimal fitting parameters, this narrow instability is concerning and so a range of initial values (centered around the global parameter fit) ought to be considered.

References

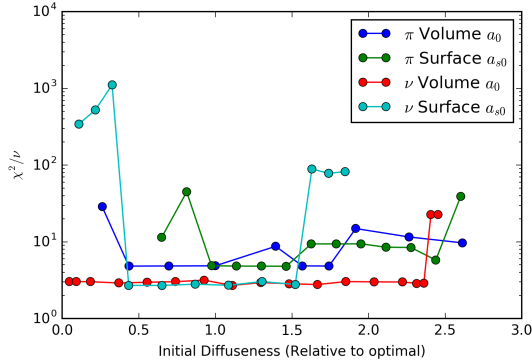
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(a) Optical potential depth parameters



(b) Nuclear radius parameters



(c) Nuclear diffuseness parameters

Figure 4: Measure of error of SFRESCO fits for different initial parameterization of the depth part of the optical potential for 49.35 MeV protons (π) and 40 MeV neutrons (ν) scattering on ^{208}Pb . Plotted on the x-axis is the ratio of $\frac{P_{\text{initial}}}{P_{\text{optimal}}}$.

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