

# A practical walk through formal scattering theory

**Connecting bound states, resonances, and scattering  
states in exotic nuclei and beyond**

## Contour rotation

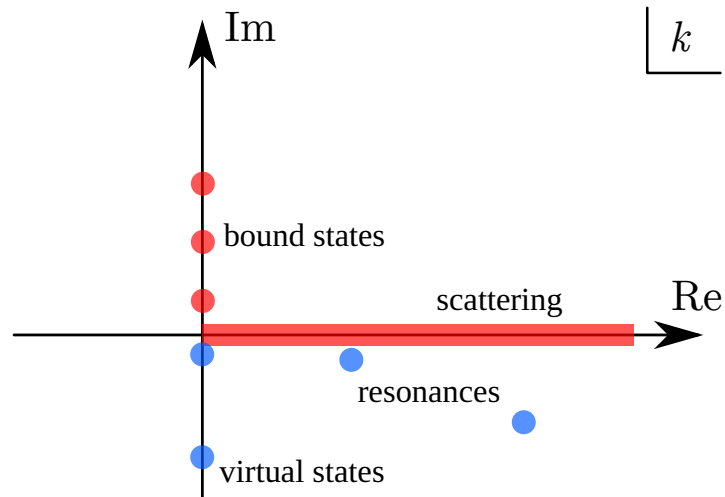
Sebastian König, NC State University



Theory  
Alliance

# Virtual states

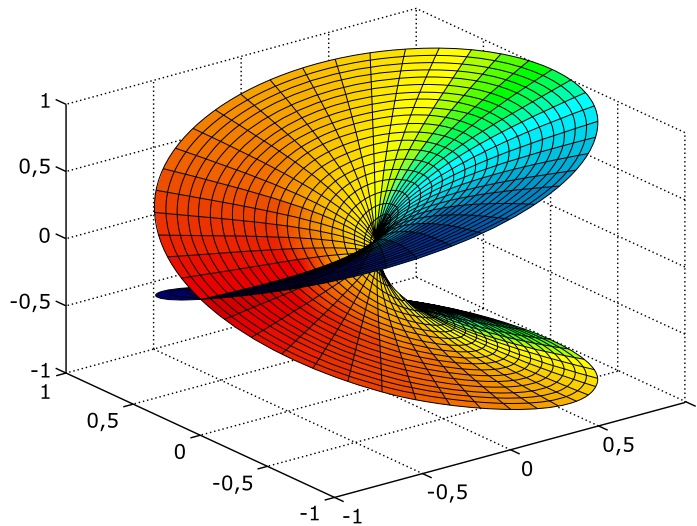
- as mentioned before, complex momenta  $k = -i\kappa$  also yield negative energies
- S-matrix poles at such positions in the complex  $k$  plane are called **virtual states** (or antibound states)



- as a function of energy, the S-matrix has **multiple branches**:  $S_l^I(E)$ ,  $s_l^{II}(E)$ 
  - **bound states** are poles of  $S_l^I(E)$  for negative  $E$ ,  $k = i\kappa$
  - **virtual (antibound) states** are poles of  $S_l^{II}(E)$  for negative  $E$ ,  $k = -i\kappa$
  - other poles of  $S_l^{II}(E)$  are resonances

# Riemann sheets

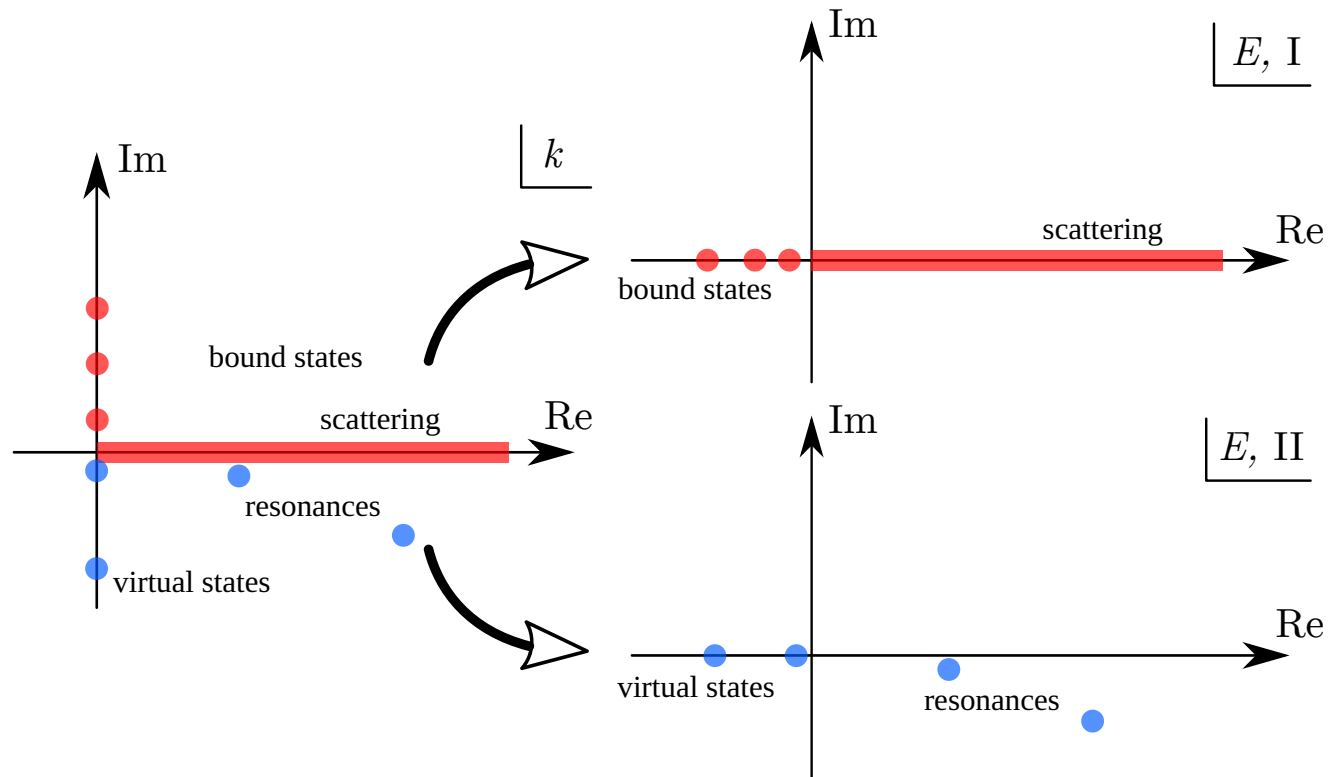
- recall that for  $E = k^2$ ,  $\sqrt{E}$  can equally well be defined as  $+k$  or  $-k$
- these are the two branches of the square root function
- typically, the **principal branch** is taken to be the positive solution
- both branches can be combined by defining  $\sqrt{E}$  on a **Riemann surface**
  - in this case, it is built out of two **Riemann sheets**
  - these are connected at the **branch cut**, chosen along the negative real axis



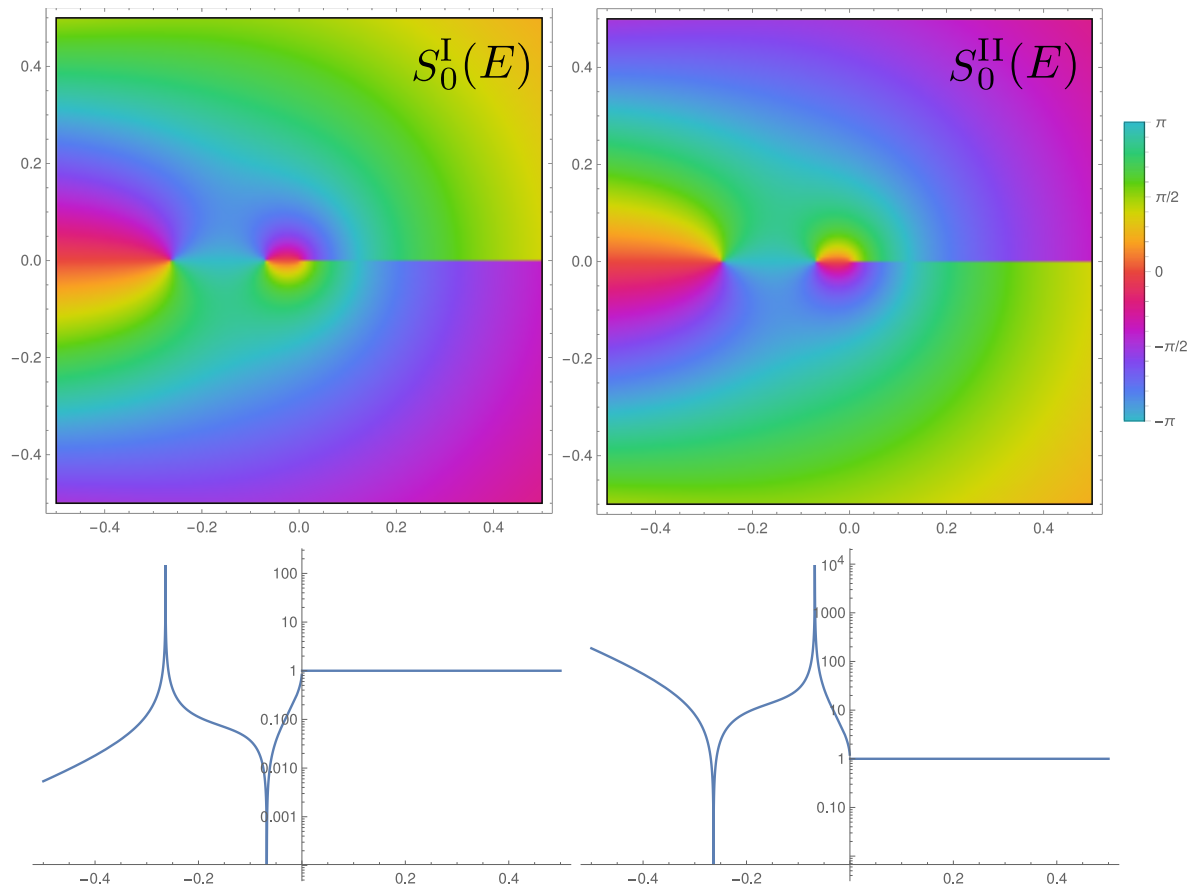
Leonid 2, via Wikimedia commons

# Analytic structure of the S-matrix

- from the square-root structure it follows that the **two sheets** of the S-matrix as a function  $E$  correspond to the **upper and lower half planes** as a function of  $k$



# Example



calculation by Nuwan Yapa

# Riemann sheets of the T-matrix

- consider now the (partial-wave projected) Lippmann-Schwinger equation in momentum space:

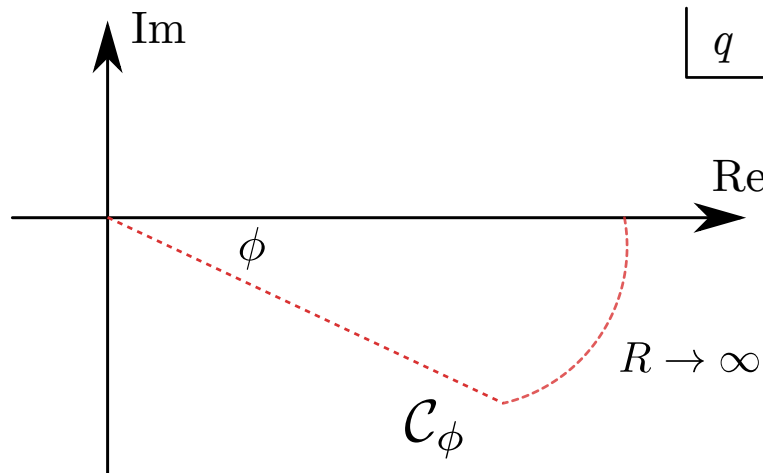
$$T(E; p, p') = V(p, p') + \int_0^\infty \frac{dq q^2}{2\pi^2} V(p, q) G_0(E; q) T(E; q, p') \quad (1)$$

- we have written this in **full off-shell form**, with the energy  $E$  a free parameter not associated with either  $p$  or  $p'$
- just like the S-matrix, the T-matrix has two Riemann sheets, which in the following we denote by  $T^{\text{I}}$  and  $T^{\text{II}}$ , and Eq. (1) is the equation for  $T = T^{\text{I}}$
- that means, even if we choose  $E$  complex, we do not leave the first sheet

**How then can we obtain  $T^{\text{II}}$ ?**

# Contour rotation

- recall that the **scattering cut** connects the first and second Riemann sheets
  - it runs along the positive real axis
  - this is precisely where we integrate in the Lippmann-Schwinger equation:  $\int_0^\infty dq$
  - for scattering calculations, we use  $i\varepsilon \rightarrow 0$  to approach the upper rim of the cut
- let us now **deform this integration contour** by rotating it into the lower half plane



- the contribution from the arc can be neglected if both  $V(p, q)$  and  $T(E; q, p')$  fall off sufficiently fast for  $q \rightarrow \infty$

# Analytic continuation

- to rotate the contour in the first place, we need to assume of course that the potential is actually defined for **complex momenta**
  - for **short-range local potentials** this is just fine because the integral

$$V_l(p, k) = 4\pi \int_0^\infty dr r^2 j_l(pr) V(r) j_l(kr)$$

converges for all  $p$  and  $k$

- so-called **separable potentials**, i.e., potentials that factorize as

$$V(p, k) \sim g(p)g(k)$$

are also no problem provided the "form factor"  $g(p)$  is an analytic function of  $p$

- after rotating the contour, we can pick  $E$  with  $q_0 = \sqrt{2\mu E}$  such that  $-\arg q_0 < \phi$  and write down the **Lippmann-Schwinger equation on the second sheet** as

$$T^\Pi(E; p, p') = V(p, p') + \int_{\mathcal{C}_\phi} \frac{dq q^2}{2\pi^2} V(p, q) G_0(E; q) T^\Pi(E; q, p') \quad (2)$$



# Rotation reversed

- the contour-rotation method is strikingly simple, but it introduces the angle  $\phi$  as an **additional parameter in the calculation**
- note now that the free Green's function for has a **pole at  $q = q_0 = \sqrt{2\mu E}$** :

$$G_0(E; q) = \frac{2\mu}{q_0^2 - q^2} \quad (3)$$

- if we want to rotate the contour back to the real axis, we will **sweep across this pole**
- this means that we will **pick up a residue contribution**

for a more detailed discussion, see W. Glöckle, The Quantum Mechanical Few-Body Problem, Springer, 1983

# Full circle

- let us retrace our steps so far:
  1. without specifying the energy explicitly, we rotated the  $dq$  integral
  2. we then chose the energy  $E$  in the accessible part of the second sheet
  3. after fixing  $E$ , we rotate the integral back and pick up a residue
- this leads to the following equation:

$$T^{\text{II}}(E; p, p') = V(p, p') + \int_0^\infty \frac{dq q^2}{2\pi^2} V(p, q) G_0(E; q) T^{\text{II}}(E; q, p') - \frac{i\mu q_0}{\pi} V(p, q_0) T^{\text{II}}(E; q_0, p') \quad (4)$$

- for the new amplitude  $T^{\text{II}}(E; q_0, p')$  we need a supplementary equation:

$$T^{\text{II}}(E; q_0, p') = V(q_0, p') + \int_0^\infty \frac{dq q^2}{2\pi^2} V(q_0, q) G_0(E; q) T^{\text{II}}(E; q, p') - \frac{i\mu q_0}{\pi} V(q_0, q_0) T^{\text{II}}(E; q_0, p') \quad (5)$$

# Second-sheet kernel

- in numerical calculations, where we discretize the  $dq$  integral, we can combine the two equations (4) and (5) by adding  $q_0$  as an extra mesh point
- this is similar to our numerical treatment of the principal-value integral that we encountered for scattering calculations
- a yet simpler equation can be obtained by eliminating  $T^{\text{II}}(E; q_0, p')$  explicitly:

$$T^{\text{II}}(E; p, p') = \tilde{V}(q_0; p, p') + \int_0^\infty \frac{dq q^2}{2\pi^2} \tilde{V}(q_0; p, q) G_0(E; q) T^{\text{II}}(E; q, p'), \quad (6)$$

with

$$\tilde{V}(q_0; p, p') = V(p, p') - V(p, q_0) \frac{i\mu q_0 / \pi}{1 + i\mu q_0 V(q_0, q_0) / \pi} V(q_0, p') \quad (7)$$

- this **modified kernel for the second sheet** allows us to search for virtual states and resonances
- note that in all these equations, we have  $q_0 = \sqrt{2\mu E}$

# Second-sheet S-matrix poles

- in order to actually search for virtual states and resonances, we need to identify **poles of the S-matrix on the second sheet**
- as for bound states, the poles actually are poles of the T-matrix
- to find these poles, we **proceed exactly as we did for bound states**
- assuming the existence of simple pole at energy  $E^*$ , the second-sheet T-matrix factorizes at the pole position:

$$T^{\text{II}}(E) \sim \frac{|R\rangle\langle R|}{E - E^*} \quad \text{for } E \rightarrow E^* \quad (8)$$

- we use  $R(p) = \langle p|R\rangle$  here to denote the vertex function
- inserting this into the second-sheet Lippmann-Schwinger equation (6) yields the **homogeneous equation**

$$R(p) = \int \frac{dq q^2}{2\pi^2} \tilde{V}(q_0; p, q) G_0(E^*; q) R(q), \quad (9)$$

where now  $q_0 = \sqrt{2\mu E^*}$