SINGLE-CHANNEL CALCULATION OF SCATTERING USING THE MANDELSTAM ITERATION

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ABSTRACT

The Mandelstam iteration with appropriate cutoff is shown to be a practical technique for the study of strong-interaction dynamics in the framework of the strip approximation. Comparison with known potential (nonrelativistic) scattering problems shows that the method is accurate enough to allow workable numerical calculations. Calculations of single channel relativistic $\pi\pi$ scattering with an elementary $\rho$ potential are reported.

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I. INTRODUCTION

In a previous paper\(^1\) the Mandelstam iteration technique was analyzed in the context of the strip approximation and shown to be, in principle, a workable method for the calculation of strongly interacting amplitudes. In this paper we describe some preliminary calculations which use this technique to study a single-channel zero-spin case (\(\pi - \pi\)), and which assures us that this approach is numerically feasible.

In Section II we describe the calculation method in detail. Section III is devoted to a comparison of solutions of potential problems obtained by the iteration method and by integration of Schrödinger's equation. Section IV describes some preliminary calculations in the fully relativistic \(\pi - \pi\) problem. These solutions are compared with solutions of the equivalent problem obtained by the N/D technique of the "New Strip Approximation."\(^2\)

II. CALCULATION METHOD

The iteration technique involves the integration of the pair of coupled equations,\(^+\)

\[
\rho^s(s,t) = \frac{g(s)}{2\pi q_s^2(s)} \int \int dt' dt'' \frac{M^*(t',s) M(t'',s)}{k^{1/2} [q_s^2(s); t, t', t'']}
\]

\[
M_t(t,s) = V_t^s(t,s) + \frac{1}{\pi} \int_{s_0}^{\infty} \frac{ds'}{s' - s} \rho^s(s',t),
\]

\(^+\) We use the notation of Ref. 1.
with
\[ K(q^2, y, y', y'') = y^2 + y'^2 + y''^2 - 2(yy' + yy'' + y'y'') - \frac{yy'y''}{q^2} \] (2.3)

and
\[ g(s) = q_s(s) \] (2.4)

for potential scattering, and
\[ g(s) = \frac{2q_s(s)}{(s)^\frac{1}{2}} h(s, s_1) \] (2.5)

for relativistic scattering, where
\[ q_s^2(s) = \frac{s^2}{4} - 1 \] (2.6)

Here \( h(s, s_1) \) is a cutoff function equal to unity below \( s_1 \) and rapidly going to zero above \( s_1 \). As usual \( t' \) and \( t'' \) integrals in Eq. (2.1) are carried over the region where \( K \) is positive.

A detailed study of the solution of a similar set of equations has been made by Bransden et al., and it is continued here.

A computer program designed to solve these equations has been written, and it operates as follows: given an initial potential discontinuity function \( V_t^S(t, s) \) for all \( s \) and \( t \), it can, using (2.1), compute \( p^S(s, t) \) for a limited range of \( t \). Equation (2.2) then allows one to compute \( M_t(t, s) \) for this same range of \( t \), which upon return to
Eq. (2.1), can be further extended. The trick is that due to the
nature of the region of integration in Eq. (2.1), to compute \( \rho^s(s,t) \)
for \( t \), say, equal to \( t_1 \), only values of \( M_t(t,s) \) for \( t \) less
than \( t_1 \) are required. This "iteration" process can, in principle,
be repeated indefinitely. However, after a sufficient number of
iterations, we can expect that the power behavior of the discontinuity
function \( M_t^s(t,s) \) will emerge, dominated by the leading Regge pole
in \( s \),

\[
M_t^s(t,s) \approx \beta(s) t^\alpha(s)
\]  

(2.6)

and it is unnecessary to proceed any further. The trajectory function
\( \alpha(s) \) and residue function \( \beta(s) \) can then be obtained from the relations,

\[
\ln |M_t^s(t,s)| = \ln |\beta(s)| + \text{Re} \; \alpha(s) \ln t ,
\]

(2.7)

\[
\text{arg} \; (M_t^s(t,s)) = \text{arg} \; (\beta(s)) + \text{Im} \; \alpha(s) \ln t ,
\]

by simple least-squares straight-line fit to \( \ln |M_t^s(t,s)| \) and
\( \text{arg}(M_t^s(t,s)) \) over a sufficiently large range of \( \ln t \). The functions
\( \alpha(s) \) and \( \beta(s) \) can now be used to define by analytic continuation
the scattering amplitude

\[
M^s(s,t) = \frac{1}{\pi} \int \frac{M_t(t,s)}{t' - t} \, dt'
\]

(2.8)
even if the above integral does not converge, as will be the case if a resonance is present.

The program to carry out this calculation is reasonably straightforward if a bit complicated. Accuracy in the integrals is of great importance if stable solutions are to be obtained. Particular care has to be exercised in Eq. (2.1) close to the boundaries of the \( t' \) and \( t'' \) integrals, as the denominator vanishes there like an inverse square root.

As it is evident from this description of the calculation, only the leading trajectory is detected. In principle, once this trajectory were known, its effect could be subtracted, and lower trajectories could then be calculated. At present it seems unlikely that the overall accuracy of the method is enough to allow this subtraction to be carried out successfully.

III. POTENTIAL PROBLEM

The potential scattering problem involving the exchange of spin zero particles can be solved by this method without any cutoff. This allows us to check the accuracy and reliability of the iteration solution by comparing it with solutions obtained by direct integration of Schroedinger's equation. For this purpose, an attractive potential with the discontinuity

\[
V^S_{t'}(t,s) = \frac{A \varepsilon}{(t - t_R)^2 + \varepsilon^2} \quad \text{for} \quad t > \frac{1}{m^2},
\]

\[
= 0 \quad \text{for} \quad t \leq \frac{1}{m^2}
\]  

(3.1)
was chosen. It corresponds to a superposition of Yukawa potentials of
range close to $1/(t_R)^{1/3}$ and it attempts to model the exchange of a
spin zero particle of width $\epsilon$. Although a single Yukawa potential
would perhaps have been preferable, its corresponding discontinuity
is a $\delta$ function which makes its numerical treatment awkward.

The same potential can then be used to integrate Schroedinger's
equation. As we are mainly interested in comparing trajectory and
residue functions, this integration can be best performed numerically,
using a modified version of P. G. Burke and C. Tate's TREGGE program.

In Figures 1 through 6 we exhibit $\alpha(s)$ and $\beta(s)$ for the
iterative and the Schroedinger solution of this problem for different
values of the width $\epsilon$, and strength $A$. It is seen that the
agreement is in general quite good throughout the ranges of $s$
explained. In particular, the iterative calculation seems to give
reasonable residue functions $\beta(s)$, which are usually more difficult
to calculate than the trajectory functions $\alpha(s)$.

As it can be expected, the agreement is poorer for narrower
or stronger potentials, the errors arising mainly from inaccuracies
in the (2.1) integration. Also it was noted that the residue functions
of potentials whose trajectories did not rise much above zero were
rather poorly determined. This is probably due to error buildup in
$\rho(s,t)$, which in these case does not increase much as a function of $t$.

All trajectories shown were obtained at $t = 19600 \text{ m}_\pi^2$. It
is necessary to go that far in $t$ to eliminate oscillations which
appear in $M_b(t,s)$ from interference with lower trajectories.
All these calculations were performed in a CDC 6600 computer, and required about 7 minutes per set.

IV. RELATIVISTIC $\pi - \pi$ SCATTERING

Having ascertained the accuracy of the iteration procedure in nonrelativistic problems, we turn to the interesting case, relativistic $\pi\pi$ scattering. As pointed out in Ref. 1, the major difference between the potential and relativistic problems is the necessity of introducing a cutoff, as otherwise the integral in Eq. (2.2) cannot be performed. The cutoff procedure adopted is the one suggested in Ref. 1, which has the advantage of both being mathematically tractable and at the same time modeling closely the "strip" structure assumed for the amplitude. To this end, the function $h(s,s_1)$ in Eq. (2.5) was set to

$$h(s,s_1) = \frac{1}{1 + \exp[(s - s_1)/\Delta]}.$$  

The solutions of the relativistic problem can be expected to depend rather critically on $s_1$, as it presumably represents the extremely complicated higher $s$ structure of $\rho^S(s,t)$ arising from the increasing number of inelastic channels open to the reaction. However, if the strip approximation is a sensible one, the dependence on $\Delta$ should not be too severe.

In these preliminary calculations, the iteration technique was used to calculate the trajectory and residues of the $\rho$ and Pomeranchuk ($I = 1$ and $I = 0$) trajectories in $\pi - \pi$ scattering with an
"elementary" $\rho$ exchanged in the $u$ and $t$ channel as potential. No attempt to obtain self-consistent or "bootstrap" solutions was made, as the presumably important Pomeranchuk repulsion\textsuperscript{5} was entirely neglected. These calculations are not expected to reproduce too closely the physical values of the position and widths of the resonances involved.

The input potential was taken to be

$$V_t(s,t) = \frac{2\beta^{'\Pi}}{3} \left[ \frac{t_R}{(t - t_R)^2 + \Gamma^2 t_R} \right] \left( \frac{t_R}{t_R - 4} \right)^{\frac{1}{2}} \left[ \frac{\Gamma^2 t_R}{(t - t_R)^2 + \Gamma^2 t_R} \right],$$

where $t_R$ and $\Gamma$ are the mass and the width of the input $\rho$ particle, and $\beta^{'\Pi}$ is the familiar $\pi - \pi$ crossing matrix. The leading factor of 2 is introduced to take into account the effect of the potential in both the $t$ and $u$ channels. The parameters $\Gamma$, $s_1$, and $\Delta$ were then adjusted to obtain reasonable output trajectories, consistent with the physical situation. It was found, however, that $\Delta$ has little effect over the lower part of the trajectory, which is mainly controlled by $\Gamma$ and $s_1$. The cutoff point $s_1$ is expected to be anywhere from about 200 to 600 $m_\pi^2$, the width of the resonance region which characterizes the strip approximation,\textsuperscript{2} and $\Gamma$ is known experimentally to be 0.9 $m_\pi^6$.

For parameters in this region, it is possible to obtain a continuum of solutions which yield a trajectory with $Re \alpha(2\beta) = 1$ in the $I = 1$ partial wave, including one for $\Gamma$ slightly higher than the physical width of the $\rho$ meson. The real and imaginary parts of $\alpha$ for two such examples are shown in Figs. 7 to 10 corresponding to
(a) \( \Gamma = 1.1 \, m_\pi \) and \( s_1 = 400 \, m_\pi^2 \), and (b) \( \Gamma = 1.6 \, m_\pi \) and \( s_1 = 256 \, m_\pi^2 \). For case (b) we also show the effect of changing the parameter \( \Delta \) from 30.0 \( m_\pi^2 \) to 100.0 \( m_\pi^2 \). As it can be readily seen, this has little effect for low positive energies, but becomes more important towards the high end of the strip and at large negative energies, where this calculation is not expected to be accurate anyway. Although neither set of trajectories is very close to the physical one, the second one seems to be the better, as it is initially steeper, more in accordance with the experimentally determined trajectories.\(^7\) In case (a) the \( I = 0 \) trajectory rises up to \( J = 2 \) to produce the \( f^0 \) resonance while in case (b) the trajectory does not reach \( J = 2 \) for a real value of \( s \) but probably does for a slightly complex one. Also, as expected, no trajectory above \( J = 0 \) is observed for \( I = 2 \), as in this case the \( \rho \) potential is repulsive.

The width of the output \( \rho \) meson is in both cases too large: 3.7 \( m_\pi \) in case (a) and 2.7 \( m_\pi \) in case (b). This is not surprising, as we have not yet included the effect of the Pomeranchuk trajectory, which can be expected to narrow this resonance.

As we approach the high end of the strip all trajectories bend downwards, as they can be expected to do if they satisfy a dispersion relation of the form

\[
\alpha(s) = \alpha(\infty) + \frac{1}{\pi} \int_{s_0}^{\infty} \frac{\text{Im} \alpha(s')}{s' - s} \, ds',
\]

as the imaginary part of \( \alpha(s) \) should go to zero outside the strip.
We also show for comparison the $0 < s$ part of the trajectories calculated using the New Form of the Strip Approximation and the N/D method. It is seen that for this particular potential, it is not very different from the iterative solution. However, this is quite reasonable, as we are dealing with a purely attractive elementary potential, where the N/D solution can be expected to perform rather well. Even so, the trajectories are flatter than the corresponding iterative ones, indicating that it will probably be easier to obtain steeper trajectories required by experiment with the new technique, once a better input potential is used. The flatness of the N/D trajectories persists for $s > 0$, where their effect can be seen by calculating cross sections. This lack of slope in turn implies that the N/D method will require considerably stronger potentials than the iterative method to give the correct mass to the $\rho$ and $f^0$ resonances. Thus, for $s > 0$ the N/D calculation yields less binding than the iterative one.

All trajectories shown were calculated at $t = 10000 \text{ m}^2 \pi$. The time required to perform these calculations in a CDC 6600 was about 4 minutes per value of the isotopic spin. The time needed to solve the N/D equations for an equivalent range of $J$ is about 1.7 minutes, so the iterative method cannot be said to be much more complicated than the N/D.

As a final point, we would like to indicate that these results are substantially different from those obtained by Bransden et al. in a similar calculation using almost the same input potential. Bransden et al. were unable to obtain trajectories rising up to $J = 1$ in the
I = 1 partial wave for a pure  $^0$ input, and were forced to include an elementary $f^0$ in the potential.*

The main differences between our calculation and theirs are improved accuracy and the different cutoff scheme. Their solution involves the smooth cutoff the the potential $V^S(t,s)$ past a given $s_\perp$, and introduces no cutoff in $\rho^S(s,t)$. It can be easily checked that both these differences play an important role in the discrepancy between the two calculations. It is our impression that our cutoff procedure is the more natural one, as it does not interfere with the power blowup of the potential in the $s$ direction, and also allows simple mathematical justification, as seen in Ref. 1.

To summarize, we can say that the above calculations seem to show that the Mandelstam iteration technique is indeed a feasible one from the computational point of view. It is quite able to produce reasonable output trajectories from a simple elementary particle input potential, and it offers many advantages over the more usual N/D approach without an outrageous increase in the necessary computations.

At present attempts are being made at calculating fully Reggeized input potentials which will include Pomeranchuk repulsion effects. Also a more ambitious self-consistent scheme is being considered whereby the output $\rho(s,t)$ function obtained after the above iterations have been completed is used to compute a new potential $V^S(t,s)$ by means of crossing. This potential could then be used in a "macro-iteration" to restart the whole calculation.

* This point is rather questionable, as has been pointed out by Chew: the inclusion of the $f^0$ as an elementary particle vastly exaggerates its effect.
V. ACKNOWLEDGMENTS

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FOOTNOTES AND REFERENCES


7. C. Chiu (Lawrence Radiation Laboratory), private communication.


9. G. F. Chew, Supplement of the Progress of Theoretical Physics, Extra Number, 118 (1965).
FIGURE CAPTIONS

Fig. 1. Real and imaginary parts of $\alpha$ for $A = 15.0$, $\epsilon = 3.0$, $t_R = 6.0$, --- Iterative --- Schroedinger.

Fig. 2. Real and imaginary parts of $\beta$ for $A = 15.0$, $\epsilon = 3.0$, $t_R = 6.0$, --- Iterative --- Schroedinger.

Fig. 3. Real and imaginary parts of $\alpha$ for $A = 35.0$, $\epsilon = 3.0$, $t_R = 6.0$, --- Iterative --- Schroedinger.

Fig. 4. Real and imaginary parts of $\beta$ for $A = 35.0$, $\epsilon = 3.0$, $t_R = 6.0$, --- Iterative --- Schroedinger.

Fig. 5. Real and imaginary parts of $\alpha$ for $A = 50.0$, $\epsilon = 3.0$, $t_R = 6.0$, --- Iterative --- Schroedinger.

Fig. 6. Real and imaginary parts of $\beta$ for $A = 50.0$, $\epsilon = 3.0$, $t_R = 6.0$, --- Iterative --- Schroedinger.

Fig. 7. Real part of $\alpha$, $\pi \pi$ problem, case (a), --- $I = 1$; --- $I = 0$; --- N/D solution.

Fig. 8. Imaginary part of $\alpha$, $\pi \pi$ problem, case (a), --- $I = 1$; --- $I = 0$.

Fig. 9. Real part of $\alpha$, $\pi \pi$ problem, case (b), --- $I = 1$, $\Delta = 30.0$; $I = 0$, $\Delta = 30.0$; $\cdots$ $I = 1$, $\Delta = 100.0$; --- $I = 0$, $\Delta = 100.0$; --- N/D solution.

Fig. 10. Imaginary part of $\alpha$, $\pi \pi$ problem, case (b), --- $I = 1$, $\Delta = 30.0$; $I = 0$, $\Delta = 30.0$; $\cdots$ $I = 1$, $\Delta = 100.0$; --- $I = 0$, $\Delta = 100.0$. 
Fig. 1.
Fig. 3.
Fig. 5.
Fig. 6.
Fig. 8.
Fig. 9.
Fig. 10
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