Multichannel quantum defect theory for cold molecular collisions with a strongly anisotropic potential energy surface

James F. E. Croft and Jeremy M. Hutson

Joint Quantum Centre Durham/Newcastle, Department of Chemistry, Durham University, South Road, Durham, DH1 3LE, United Kingdom
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We show that multichannel quantum defect theory (MQDT) can be applied successfully as an efficient computational method for cold molecular collisions in Li + NH, which has a deep and strongly anisotropic interaction potential. In this strongly coupled system, closed-channel poles restrict the range over which the MQDT Y matrix can be interpolated. We present an improved procedure to transform the MQDT reference functions so that the poles are removed from the energy range of interest. Effects due to very-long-range spin dipolar couplings are outside the scope of MQDT, but can be added perturbatively. This procedure makes it possible to calculate the elastic and inelastic cross sections, over the entire range of energies and fields needed to evaluate the feasibility of sympathetic cooling of NH by Li, using coupled-channel calculations at only five combinations of energy and field.

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I. INTRODUCTION

Cold and ultracold molecules provide an exciting doorway to new fields in physics and chemistry. They share the controllability and tunable interactions that have made cold-atom studies so fruitful. However, the richer structure of molecules makes them suitable for many new applications and research directions [1,2]. Polar molecules are of particular interest because their electric dipole moment allows the interactions between them to be controlled and manipulated by external fields. This strong tunable response may make it possible to develop a fully controlled chemistry [3] where every degree of freedom of the reaction can be tuned, providing fundamental insights into chemical reaction processes. Molecules such as KRb [4] and Cs₂ [5] have already been produced at submicrokelvin temperatures in their lowest-energy electronic, vibrational, rotational, and hyperfine state, by magnetoassociation followed by laser state transfer. Reactions of ultracold 40K⁸⁷Rb with itself and with K and Rb atoms have been studied [6] and it was seen that quantum statistics and quantum threshold laws play an important role in determining the rates of inelastic collisions.

The only ultracold molecules that are accessible with current methods are alkali-metal dimers. However, there is great interest in producing samples of other molecular species in the ultracold regime. The most promising route to this is first to cool and trap them in the cold regime (at temperatures below 1 K), using a method such as buffer-gas cooling [7] or molecular beam deceleration [8], and then to bring them to the ultracold regime using a second-stage approach such as evaporative cooling [9], sympathetic cooling [10], or laser cooling [11,12].

Evaporative and sympathetic cooling both rely on elastic collisions to thermalize the sample, but both can be prevented by inelastic collisions that release kinetic energy and lead to trap loss. Collisional properties are also key to methods for controlling ultracold atoms and molecules. Calculations on atomic and molecular collisions are therefore crucial to both the production and control of cold and ultracold molecules.

Such calculations require the solution of the set of coupled differential equations obtained from the Schrödinger equation.

There are various numerical methods for solving these coupled differential equations, of which the most commonly used is the full coupled-channel method. This propagates the matrix solution of the Schrödinger equation from short to long range and takes a time proportional to $N^3$, where $N$ is the number of coupled channels. The properties of completed collisions are described by the scattering matrix $S$, which is obtained by matching the propagated solutions to free-particle wave functions (Ricatti-Bessel functions) at long range [13].

Full coupled-channel calculations can be extremely expensive, particularly in applied electric and magnetic fields [14–16]. The expense is particularly great for systems with deep and strongly anisotropic potential wells [17], for molecule-molecule collisions [18], or when nuclear hyperfine interactions are included [19,20].

In cold collision studies, the scattering $S$ matrix is often a fast function of collision energy $E$ and magnetic field $B$, with extensive structure due to scattering resonances and discontinuous behavior at threshold. Calculations are thus required over a fine grid of energies and/or applied field, typically over an energy range of order 1 K from threshold, and for magnetic fields up to a few thousand gauss. This contrasts with the situation for collisions of ultracold atoms, where the energy range of interest is commonly a few microkelvins and the fields are typically a few hundred gauss.

Approaches based on multichannel quantum defect theory (MQDT) avoid the repetition of the expensive propagation by representing the scattering properties in terms of a matrix $Y(E,B)$ that is a smooth function of $E$ and $B$ [21–25]. MQDT has proved highly successful for cold atomic interactions [24–30] and more recently it has been applied to collisions of cold and ultracold molecules [31–36]. MQDT defines the matrix $Y(E,B)$ at a matching distance $r_{\text{match}}$ at relatively short range in order to achieve this insensitivity to energy and field. The $Y$ matrix contains all the scattering dynamics inside $r_{\text{match}}$. The smooth variation of $Y$ allows it to be obtained once and then used for calculations over a wide range of energies and fields or obtained by interpolation from a few points. The computational cost of calculations at additional energies and fields is only proportional to $N$, not $N^3$. 
We have previously demonstrated the application of MQDT to cold molecular collisions for the moderately anisotropic system Mg + NH(2Σ−) [34,36]. In Ref. [36] we showed that the choice made for the phase of the MQDT reference functions is very important in producing a Y matrix that can be interpolated smoothly over a wide range of energy and field. The purpose of the present paper is to explore how the approach performs for a much more strongly anisotropic system with many more closed channels. For this purpose we choose Li + NH, which has been studied previously using full coupled-channel calculations by Wallis et al. [17]. As in Ref. [17], we focus on collisions between spin-polarized Li and NH, which occur on the quartet potential energy surface. This surface is deep and highly anisotropic, with a well depth about 1800 cm−1 at the Li-NH geometry, but only 113 cm−1 at the NH-Li geometry. With a small but important modification to the method of choosing phases described in Ref. [36], we find that we can obtain accurate results for elastic and inelastic cross section, over the entire range of energies relevant to sympathetic cooling, using only five coupled-channel calculations.

II. THEORY

The theory of MQDT is given in detail in Refs. [22–25]. Here we give only a brief description, following Refs. [34,36], which is sufficient to describe the notation we use.

MQDT makes the approximation that the multichannel Schrödinger equation is uncoupled at distances r > r_match, so its solution in this region may be written in the matrix form

\[ \Psi = r^{-1}[f(r) + g(r)Y]. \] (1)

Here f and g are diagonal matrices containing the functions f_i and g_i, which are linearly independent solutions of a reference Schrödinger equation in each asymptotic channel i,

\[ -\frac{\hbar^2}{2\mu} \frac{d^2}{dr^2} + U^\text{ref}_i(r) - E \] f_i(r) = 0, \] (2)

and similarly for g_i(r). Here \( \mu \) is the reduced mass and the reference potentials \( U^\text{ref}_i(r) \) approach the true potential at long range. They include the centrifugal terms \( \hbar^2L_i(L_i + 1)/2\mu r^2 \), where \( L_i \) is the partial-wave quantum number for channel i.

The phase of the short-range reference functions f_i and g_i is a disposable parameter of MQDT and may be chosen to generate a Y matrix that is smooth and pole-free over a wide range of energy and field [36]. Equation (1) shows that the Y matrix has a pole whenever the component of the propagated multichannel wave function \( \psi_i \) in any channel i is proportional to the reference function g_i and has no contribution from f_i, i.e., when \( g_i \) and the full coupled-channel solution have the same phase at \( r_\text{match} \). In Ref. [36] we proposed transforming the pair of reference functions f_i and g_i in channel i with a rotation angle \( \theta_i \), chosen so that the diagonal matrix element \( Y_{ii} \) is 0 at a particular reference energy \( E^\text{ref} \) and field \( B_{\text{ref}} \). This ensures that the reference function g_i and the full coupled-channel solution in channel i are perfectly out of phase at the chosen \( r_\text{match} \) and the resulting Y matrix is therefore pole-free close to \( E^\text{ref} \) and \( B_{\text{ref}} \) [36].

The range of the pole-free region is dependent on where the matching occurs. When matching is in the classically allowed region, the phases of both the reference functions and the propagated coupled-channel solutions vary approximately linearly with energy and setting the diagonal Y matrix elements to zero is effective: The relative phase of the reference functions and the coupled-channel solution is a slow function of energy. For a closed channel where matching is carried out in the classically forbidden region, however, there is a resonance structure in both the coupled-channel solutions and the reference functions, so the phase of each is a fast (and nonlinear) function of energy. Fortunately, the energies at which the resonance structure occurs depend on the choice of \( \theta_i \). In the present work we show that a more sophisticated choice of \( \theta_i \) than that of Ref. [36] can produce a larger pole-free region for closed channels.

A. Basis set and quantum numbers

We construct the collision Hamiltonian in the fully uncoupled basis set \( |nm_n, s_{\text{NH}}m_{\text{NH}}⟩|s_i,m_{s_i}⟩|L,M_L⟩ \), where the quantum numbers \( n \) and \( s_{\text{NH}} \) describe the rotation and electron spin of the NH molecule and \( s_{Li} \) describes the electron spin of the Li atom. The corresponding \( m \) quantum numbers are the projections onto the space-fixed magnetic field axis. Hyperfine structure is neglected. The matrix elements required for the coupled equations are the same as for scattering of NH from a closed-shell atom [16], with the addition of the anisotropic intermolecular spin-spin dipolar interaction [17].

The coupled-channel calculations may in principle be carried out in any sufficiently complete basis set. However, the Y and S matrices are expressed in a basis set of eigenfunctions of the field-dressed monomer Hamiltonians. At low field the states of the NH molecule are approximately described by quantum numbers \( j \) and \( m_j \), where \( j \) is the resultant of \( n \) and \( s \). We label elements of Y and S by subscripts \( n,j,m_j,m_{s_i},L,M_L \rightarrow n',j',m_j',m_{s_i}',L',M_L' \). For diagonal elements we suppress the second set of labels.

B. Numerical methods

The coupled-channel calculations required for both MQDT and the full coupled-channel approach are carried out using the MOLSCAT package [37], as modified to handle collisions in magnetic fields [16]. The coupled equations are solved numerically using the hybrid log-derivative propagator of Alexander and Manolopoulos [38], which uses a fixed-step-size log-derivative propagator in the short-range region \( r_{\text{min}} \leq r < r_{\text{mid}} \) and a variable-step-size Airy propagator in the long-range region \( r_{\text{mid}} \leq r \leq r_{\text{max}} \). As in Ref. [17], the full coupled-channel calculations use \( r_{\text{min}} = 1.8 \text{ Å}, r_{\text{mid}} = 12.5 \text{ Å}, \) and \( r_{\text{max}} = 600 \text{ Å} \) (where 1 Å = \( 10^{-10} \text{ m} \)). MQDT requires coupled-channel calculations only from \( r_{\text{min}} \) to \( r_\text{match} \) (which is less than \( r_{\text{mid}} \)), so only the fixed-step-size propagator is used in this case.

The initial MQDT reference functions and quantum defect parameters are obtained as described in Ref. [34], using the renormalized Numerov method [39] to solve the one-dimensional Schrödinger equations for the reference potentials. In this paper all MQDT calculations use the reference
potential

\[ U_{i}^{\text{ref}}(r) = V_0(r) + \frac{\hbar^2 L_i(L_i + 1)}{2 \mu r^2} + E_i^\infty, \]  

where \( V_0(r) \) is the isotropic part of the interaction potential and \( L_i \) is the partial-wave quantum number for channel \( i \). The reference potential contains a hard wall at \( r = r_{\text{wall}}^{\text{ref}} \), so \( U_{i}^{\text{ref}}(r) = \infty \) for \( r < r_{\text{wall}}^{\text{ref}} \). In the present paper we choose \( r_{\text{wall}}^{\text{ref}} = 4.0 \text{ Å} \). This reference potential has been shown to produce quantitatively accurate results for Mg + NH [34,36].

The uncoupled basis functions used to solve the coupled-channel equations are not eigenfunctions of the Hamiltonians of the separated monomers. The log-derivative matrix obtained from the coupled-channel calculations at a distance \( r_{\text{match}} \) is therefore transformed into a basis set that diagonalizes the asymptotic Hamiltonian [40]. The MQDT \( Y \) matrix is then obtained by matching to this log-derivative matrix at \( r_{\text{match}} \) using Eq. (13) of Ref. [34]. All channels with \( n \geq 2 \) are treated as strongly closed and thus not included in the MQDT part of the calculation, but are included in the log-derivative propagation.

III. RESULTS AND DISCUSSION

In a magnetic field, the lowest Li-NH threshold \((n = 0, \Delta n \text{Li} = 1, \Delta n \text{NH} = \frac{1}{2})\) splits into six Zeeman sublevels. We consider collisions between Li atoms and NH molecules that are both initially in their magnetically trappable low-spin-seeking states \( m_{\text{Li},1} = 1/2 \) and \( m_{\text{NH},1} = 1 \). This corresponds to the highest of the six thresholds.

Figure 1 shows the variation of the representative element \( Y_{1,2,-2,-1/2,7/4} \), as a function of the matching distance and energy, when the phases \( \theta_i \) are chosen to make all diagonal \( Y \) matrix elements zero at collision energy \( E_{\text{ref}} = 0.5 \text{ K} \) and field \( B_{\text{ref}} = 10 \text{ G} \). It may be seen that there are poles (where \( \text{arctan} \frac{Y_{ii}}{\pi} \) passes through \( \pm 1/2 \)) whose positions depend strongly on \( r_{\text{match}} \). Other \( Y \) matrix elements are quantitatively different but have poles in the same places. The basis set used for Fig. 1 includes all functions up to \( n_{\text{max}} = 3 \) and \( L_{\text{max}} = 8 \). This un converged basis set was used due to the substantial computational cost of performing a full coupled-channel calculation at every energy in the figure. The outer turning point of the \( n = 1 \) reference potential is at 6.1 Å and it may be seen that, for values of \( r_{\text{match}} \) inside this, the \( Y \) matrix is pole-free over many kelvin. However, MQDT with such small values of \( r_{\text{match}} \) does not produce accurate results because it neglects all channel couplings that exist outside \( r_{\text{match}} \). It is therefore essential to place \( r_{\text{match}} \) far enough out that some channels are closed [34]. When \( r_{\text{match}} > 6.1 \text{ Å} \), however, poles start to enter the \( Y \) matrix in the energy range of interest. As \( r_{\text{match}} \) increases further, the poles move and at some values of \( r_{\text{match}} \) can come within 0.1 K of \( E_{\text{ref}} \).

A contour plot such as Fig. 1 requires coupled-channel calculations at every energy and producing it thus sacrifices most of the computational savings that MQDT is designed to achieve. Furthermore, we need a procedure for choosing the phases \( \theta_i \) that will guarantee a large pole-free region for any choice of \( r_{\text{match}} \). In the remainder of this paper, we perform calculations at only a single value of \( r_{\text{match}} = 6.5 \text{ Å} \), deliberately chosen to be in a region where Fig. 1 shows that there are poles in \( Y \) inconveniently close to \( E_{\text{ref}} \). In addition, the remaining calculations use a basis set including all functions up to \( n_{\text{max}} = 6 \) and \( L_{\text{max}} = 8 \), except where stated otherwise.

The dependence of a diagonal \( Y \) matrix element on the phase of the reference functions (in any channel handled by MQDT) is

\[ Y_{ii}(E) = \tan[\theta_i + \delta_i(E)], \]

where \( \delta_i(E) \) is the phase shift between the unrotated reference function \( f_i \) and the propagated multichannel wave function in channel \( i \). For a closed channel that is capable of supporting resonances, the phase shift around a resonance has a Breit-Wigner form

\[ \delta_i(E) = \delta_i(E_0) + \text{arctan} \left( \frac{\Gamma_i/2}{E - E_i^{\text{ref}}} \right), \]

where \( \delta_i \) is a slowly varying (nonresonant) background term. The resonant part of this function is shown in the bottom panel of Fig. 2 for values of the parameters appropriate to one of the channels in Li + NH. It may be seen that choosing a value of \( \theta_i \) that makes \( Y_{ii} \) zero (shown by the dashed vertical line) does not guarantee a large pole-free region in the case where \( E_{\text{ref}} \) is close to \( E_i^{\text{ref}} \). A much better choice in this case is to set \( \theta_i \) to the value shown by the solid vertical line. In the following we will show how this can be achieved.

A basic problem of MQDT in coupled-channel problems is that a pole in \( Y \) that originates in any channel causes a pole in every channel. We refer to this as the contamination of one channel by another. The top panel of Fig. 2 shows arctan \( Y_{ii}/\pi \) for the single matrix element \( Y_{1,2,-2,-1/2,7/4} \) obtained from coupled-channel calculations as a function of \( \theta_i \) and collision energy \( E \). The phases \( \theta_i \) in all other channels \( i' \) are set to the values that produce \( Y_{i'i'} = 0 \) at \( E_{\text{ref}} = 0.5 \text{ K} \). The broad horizontal sweep around 0 K arises from a resonance in channel \( i \), while the narrower sweeps near \(-1.8, -1.6, -1.2, 1.4, \) and \(2.0 \text{ K} \) are poles due to contamination from other channels. Setting \( Y_{i'i'} \) to zero in all these other channels has shifted these contamination effects to energies either above

FIG. 1. (Color online) arctan \( Y_{ii}/\pi \) for a single representative diagonal \( Y \) matrix element, as a function of collision energy \( E \) and \( r_{\text{match}} \), with the phases \( \theta_i \) set so that \( Y_{ii} = 0 \) in all channels for energy \( E_{\text{ref}} = 0.5 \text{ K} \) and field \( B_{\text{ref}} = 10 \text{ G} \).
about 1.4 K or below about −1.2 K, leaving a region of about 2.6 K uncontaminated by other channels. For the specific circumstances shown in Fig. 2, it is seen that choosing $\theta_i$ so that $Y_{ii} = 0$ results in $\theta_i/\pi \approx 0.33$ (dashed vertical line). This produces a pole in $Y_{ii}$ itself at about −0.1 K, which is inconveniently close to $E_{\text{ref}}$. However, a choice of $\theta_i/\pi = 0.63$ (solid vertical line) would produce a much larger pole-free range, limited only by poles due to contamination effects in other channels. If improved values of $\theta_i$ can also be obtained for the channels responsible for the contamination poles, then there is clearly the prospect of achieving a much improved pole-free region.

The pole structure in channel $i$ when uncontaminated by other channels is given by Eqs. (4) and (5). In order to use these equations to obtain a better choice of $\theta_i$, we need values for $E_{\text{ref}}^i$, $\Gamma_i$, and $\delta_i$. To obtain these we first optimize the phases as in Ref. [36], transforming the reference functions so that $Y_{ii} = 0$ in all channels at energy $E_{\text{ref}}$. This provides at least a small region where $Y_{ii}$ is uncontaminated by poles in other channels. We then carry out coupled-channel calculations at two additional energies near $E_{\text{ref}}$ and use Eqs. (4) and (5) to obtain the three parameters $E_{\text{ref}}^i$, $\Gamma_i$, and $\delta_i$ numerically, neglecting the slow variation of $\delta_i$ with $E$. The optimum pole-free region for this channel is then achieved by setting $\theta_i = \pi/2 - \delta_i$.

The pole-free region for the entire $Y$ matrix is optimized by applying this procedure in all closed channels where there is resonant structure close to the reference energy. We first calculate the numerical second derivative of the diagonal matrix elements $Y_{ii}$ with respect to energy. We then select the channel with the largest second derivative, apply the procedure described above, and use the new set of phases to recalculate the three $Y$ matrices. Because of channel mixing, this in principle changes all the diagonal matrix elements. If it reduces $\sum_i |d^2 Y_{ii}/dE^2|$ then we accept the new value of $\theta_i$. If not, we move on to the next channel and apply the same procedure. We loop over the channels in this manner until there is no channel for which changing $\theta_i$ to $\pi/2 - \delta_i$ reduces $\sum_i |d^2 Y_{ii}/dE^2|$. This is an inexpensive procedure, as it uses the same three coupled-channel calculations as before. Only the closed channels need to be included in the loop since only these channels have resonance structure. In physical terms, the procedure shifts the bound states supported by the closed-channel reference potentials so that the rapidly varying phase in the full coupled-channel solution in channel $i$ is matched by the phase of the corresponding reference function.

Figure 3 compares the final matrix elements $Y_{ii}$ in all the channels included in the MQDT procedure, obtained with the two optimization schemes. The dashed red lines show the result of choosing $\theta_i$ so that $Y_{ii}$ is zero in every channel, as in Ref. [36], while the solid black lines show the result of optimizing $\theta_i$ as described above. Both calculations use $r_{\text{match}} = 6.5$ Å and optimize $\theta_i$ at $E_{\text{ref}} = 0.5$ K and $B_{\text{ref}} = 10$ G. It may be seen that taking account of closed-channel resonances significantly increases the pole-free range of $Y$. Furthermore, it produces $Y$ matrix elements that are considerably more linear between 0 and 1 K and may thus be interpolated more accurately.

Figure 4 compares diagonal $T$-matrix elements $|T_{ii}|^2$ (where $T_{ij} = \delta_{ij} - S_{ij}$) obtained from full coupled-channel calculations at the two energies near $E_{\text{ref}}$.
calculations from [17] on the unscaled potential energy surface. Coupled-channel have therefore used MQDT to repeat the calculations of Ref. (required to optimize the phases) and two magnetic fields. We plot from coupled-channel calculations at just three energies the contour plots given in Ref. [17] actually included coupled-channel calculations at 204 combinations of energy and field. These calculations were very expensive because they required calculations on a fine grid of energies and fields. Wallis et al. [17] have investigated Li + NH collisions using coupled-channel calculations and produced contour plots that show the ratio $\gamma$ as a function of collision energy $E$ and magnetic field $B$. These calculations were very expensive because they required calculations on a fine grid of energies and fields. The contour plots given in Ref. [17] actually included coupled-channel calculations at 204 combinations of energy and field.

MQDT offers the possibility of producing the entire contour plot from coupled-channel calculations at just three energies (required to optimize the phases) and two magnetic fields. We have therefore used MQDT to repeat the calculations of Ref. [17] on the unscaled potential energy surface. Coupled-channel calculations from $r_{\text{min}}$ to $r_{\text{match}}$ were carried out at collision energies $E = 0.01, 0.05$, and $0.1$ K at magnetic field $B = 10$ G to optimize the phases and at $E = 0.01$ and $0.1$ K at $B = 1000$ G to allow linear interpolation in $B$ and $E$. These calculations used a basis set with $n_{\text{max}} = 6$ and $L_{\text{max}} = 8$ to allow direct comparison with Ref. [17]. The resulting contour plot of $\gamma$ is compared with the results of Ref. [17] in the top panel of Fig. 5: It may be seen that there is good agreement at both high collision energies ($E > 0.01$ K) and high fields ($B > 100$ G), but that MQDT by itself breaks down when both $E$ and $B$ are small [41].

The inaccuracy in MQDT at low energy and low field occurs because, in this region, the inelastic cross sections are dominated by long-range inelasticity involving the magnetic dipole interaction between the spins of Li and NH. As described by Janssen et al. [18], there are long-range avoided crossings between the effective potential curves for the incoming channel and for inelastic channels with larger values of $L$. These crossings usually occur outside the centrifugal barriers and even coupled-channel calculations must be propagated to very long range (hundreds of angstroms) to capture their effects. They are thus outside the scope of MQDT, which neglects couplings outside $r_{\text{match}}$.

The long-range couplings may however be included perturbatively at very little expense. Janssen et al. [42] have developed both a simple Born approximation and a distorted-wave Born approximation (DWBA) for calculating...
the inelastic cross sections due to these long-range couplings. In this approach the incoming and outgoing waves are written in terms of spherical Bessel functions, which represent the wave function accurately at very long range where the avoided crossings neglected by MQDT occur. The simple Born approximation is stable to evaluate, but can give results up to a factor of 2 in error for cross sections for initial L = 0 because it does not take account of the phase shift due to short-range interactions. The DWBA, by contrast, is quite accurate for initial L = 0 but can be unstable when L and L' are both nonzero. However, in the latter case, short-range effects are unimportant. We have therefore used a hybrid Born approximation, made up of the DWBA for initial L = 0 and the simple Born approximation for initial L > 0. When we add the resulting inelastic cross sections to those from MQDT, we obtain the contour plot for the ratio γ shown in the bottom panel of Fig. 5. It may be seen that this gives essentially complete agreement with the full coupled-channel results.

The MQDT approach makes it feasible to use a larger basis than was possible in Ref. [17] and to carry out the calculations on a much finer grid of energies and fields. Figure 6 shows the results obtained from MQDT with perturbative long-range corrections for a converged basis set with \( n_{\text{max}} = 10 \) and \( L_{\text{max}} = 8 \), with cross sections calculated on a 51 × 51 grid of energies and fields. The resulting coupled-channel basis sets contains 1887 basis functions, as compared to 937 functions for the smaller basis set used in Fig. 5, and each coupled-channel calculation is therefore a factor of 8 more expensive.

As described in Ref. [17], the elastic and inelastic cross sections are a strong function of both potential scaling and basis set size because they depend sensitively on the positions of near-dissociation levels. Because of this, calculations on a single potential do not give quantitative predictions for the ratio of elastic and inelastic cross sections and it is essential to explore the potential dependence of the results. We found that using the unscaled potential with a converged basis set gave a highly atypical contour plot because it has an accidentally near-zero scattering length and therefore a very small elastic cross section. The calculations in Fig. 6 used a potential with an overall scaling factor of 0.995, which produces a much more typical contour plot. Exploring the dependence of the results on the scaling factor confirmed the conclusions of Ref. [17] that sympathetic cooling of NH by Li is likely to succeed if the molecules can be precooled to a temperature around 20 mK.

V. CONCLUSION

We have demonstrated that MQDT can provide quantitatively accurate cross sections for cold and ultracold elastic and inelastic collisions in magnetic fields for a strongly coupled molecular system Li + NH. However, the choice of the phases of the MQDT reference functions is crucial. For Mg + NH it was sufficient to choose the phases so that \( \gamma_{ij} = 0 \) in every channel included in MQDT [36]. For Li + NH, however, this does not guarantee that all closed-channel poles are moved far away in energy and the poles can cause problems in interpolation. In the present paper we have developed an improved approach for optimizing the phases that ensures that closed-channel poles are far away from a reference energy.

We have been able to reproduce the results of coupled-channel calculations across the entire range of energies and field relevant to sympathetic cooling of NH by Li using our new version of MQDT combined with perturbative corrections for long-range inelasticity caused by the magnetic spin dipolar interaction. The MQDT results required coupled-channel calculations at only five combinations of energy and field, whereas the coupled-channel calculations [17] required 204 combinations. MQDT thus has enormous potential as an efficient computational method for molecular collisions.

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This transformation was neglected in Refs. [34,36], where the partially coupled basis set used for the propagation was asymptotically very nearly diagonal. However, it is important in the decoupled basis set used here.

For consistency of comparison, we have performed MQDT calculations on the same grid of energies and fields as was used for the coupled-channel calculations in Ref. [17]. This actually included only four field values across the range shown. Including extra fields in the MQDT calculations produces small but visible changes in the contours.

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