Title
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Permalink
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Publication Date
2001-12-02
Improved alternating gradient transport and focusing of neutral molecules

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PACS numbers: 39.10.+j, 33.15.Kr, 07.77.Gx

I. INTRODUCTION

A polar molecule has an intrinsic separation of charge on which an electric field gradient exerts a force. The force, $F_x$, in the (transverse) x direction is:

$$F_x = -\frac{\partial W}{\partial x} = -\frac{\partial W}{\partial E} \frac{\partial E}{\partial x}$$

where $W$ is the potential energy of the molecule in an electric field (Stark effect) of magnitude $E = (E_x^2 + E_y^2)^{1/2}$; and similarly for the force, $F_y$ in the y direction. The force and gradient are in opposite directions (weak-field seeking) for rotational states that become less tightly bound in an electric field ($\partial W/\partial E > 0$), while the force and gradient are in the same direction (strong-field seeking) for rotational states that become more tightly bound in an electric field ($\partial W/\partial E < 0$). The $J=0$ state is always strong-field seeking and all rotational states become strong-field seeking in the limit of strong electric field as shown in Fig. 1.

Focusing a beam of molecules, traveling in the z direction, is achieved using static two-dimensional ($x,y$) electric field gradients. We neglect end field effects and assume $E_z = 0$ inside the focusing lenses. Molecules in weak-field seeking states can be focused, in both directions, using quadrupole and/or sextupole fields that have a minimum, in the electric field, in both directions. Molecules in strong-field seeking states can be focused in only one transverse direction, while being defocused in the other, because it is not possible to have a maximum in the electric field in both dimensions (in free space). Alternating gradient lenses are used to transport and focus molecules in strong-field seeking states. While alternating gradient focusing has been used for molecules [1–7] and neutral atoms [8], it has been neither as successful, nor as widely used, as has quadrupole and sextupole focusing for molecules in weak-field seeking states [7, 9].

In this paper we show, by calculation and numerical simulation, how one may greatly improve the alternating gradient transport and focusing of molecules by optimizing the lens field geometries. We use this optimized multipole lens design, in a FODO lattice beam transport line and use lenses to match the beam transport line to the beam source and the final focus.

The remainder of this paper is organized as follows: Section II discusses alternating gradient focusing, transport lattices, matching lenses, and linear optics. Section III derives the formulae for designing linear focusing elements and presents examples of lenses. Section IV derives the equations for molecular beam transport and models both a simple 30 m-long FODO lattice and a complete 15 m-long transport line with entrance and exit matching lenses. Section IV G estimates the intensity of a methyl fluoride jet-source beam, transported through a beamline and focused. Section V discusses the use of strong-field seeking states, and possible applications of alternating grading focusing and transport through very long transport lines.

II. ALTERNATING GRADIENT BEAM TRANSPORT AND FOCUSING

A beam of charged particles can be focused and transported over almost unlimited distances by alternating F and D type magnetic quadrupole lenses. The F type lens focuses the beam in the horizontal ($x$) direction while defocusing it in the vertical ($y$) direction. The D type defocuses the beam in the horizontal direction while focusing it in the vertical direction.

A complete alternating gradient transport line begins with a beam source whose output is optically matched, by lenses, into the acceptance of a transport section which, in turn is matched, by lenses, to a final focus. The final focus can be,
used for asymmetric focal points (unequal strengths) and to match the transport line to the final focus, we use doublets (FD/DF) and triplets (FDF/DFD) type lens configurations. In the F/D doublet, a net focusing in \(x\) and \(y\) occurs because particles first focused in \(x\) are, in the second element, closer to the center axis and therefore less defocused. Particles first defocused in \(y\) are, in the second element, further from the axis and thus more strongly focused.

In all lenses, linear focusing is needed for optimum optical properties. Linear focusing requires that the force in each direction be linearly proportional to the displacement in that direction (\(|F_x/x| = \text{constant}\) and \(|F_y/y| = \text{constant}\)), and independent of the displacement in the other direction (uncoupled motion). Strong non-linearities in focusing elements will result in loss of beam, generation of beam halo, and growth in the transverse emittance (the product of the angular divergence and the spatial dimension) producing larger beam sizes.

### III. LINEAR OPTICS

#### A. Potential energy of a molecule in an electric field

To determine the lens shape that will produce the most linear force on the molecule, we need \(\partial W/\partial E\) for Eq. (1). This quantity will change with the \(J, |m_J|\) rotational state and electric field strength. If a number of rotational levels have a similar \(\partial W/\partial E\), then one lens design will be nearly optimum for all of them. In the limiting case of a strong electric field (large \(\omega\) in Fig. 1), \(\partial W/\partial E = -d_e\).

For polar molecules in weaker fields, we calculate the interaction energy, in the rigid rotor approximation, following the approach of von Meyenn [10]. The Hamiltonian operator is \(H = BJ^2 - d_eE \cos \theta\) where \(B\) is the rotational constant, and the direction cosine matrix elements, which couple \((J, m_J)\) with \((J + 1, m_J)\) and \((J - 1, m_J)\), are taken from Townes and Schawlow [11]. We diagonalize the matrix for terms through \(J = 30\). The first few levels are shown in Fig. 1. As \(\omega = d_e/E\) and \(B\) determine the Stark effect for each \(J, |m_J|\), it is straightforward to construct a simple function for the Stark effect for any \(J, |m_J|\). For the \(J = 0\) state of a diatomic molecule or symmetric top (\(K = 0\)) molecule, a satisfactory approximation [12] is:

\[
W(E) = \frac{C_1 \omega^2 B}{1 + C_2 \omega} = \frac{C_1 d_e^2 E^2}{B + C_2 d_e E}
\]  

(2)

with \(C_1 = -0.2085\) and \(C_2 = 0.2445\). This expression works best for small and intermediate values of \(\omega\).

From Eq. (2), for the \(J = 0\) state the field derivative of the potential is:

\[
\frac{\partial W}{\partial E} = \frac{W}{E} \left[ \frac{2B + C_2 d_e E}{B + C_2 d_e E} \right]
\]

(3)

Similar expressions may be found for other rotational levels. A perturbation expression for \(W\) can be used for rotational states in weak electric fields [11]. In some applications, using \(\partial W/\partial E\) from either the strong field limit or the weak field limit will be sufficient to design a linear lens.

Finally, we note that all atoms and molecules, including nonpolar molecules, are polarizable with an interaction energy for example, at an experimental target or at a matching point for injection into a (storage) ring lattice. A complete transport line, for molecules, starting with a jet source and skimmer, is shown in Fig. 2. For long-distance beam transport, the lens system providing the highest beam acceptance and requiring the lowest focusing strength, is a lattice of identical FODO-cells, in which the F/D- lenses are separated by drift spaces (O).

To match the molecular beam source to the transport line, and to match the transport line to the final focus, we use doublet (FD/DF) and triplet (FDF/DFD) type lens configurations. These lens configurations generally require higher focusing strengths than do the FODO cells. Doublets are typically used for asymmetric focal points (unequal \(x\) and \(y\) dimensions) while triplets are used for symmetric ones. In the F/D doublet,
\[ W_0 = -\frac{1}{4} \alpha E^2, \] where \( \alpha \) is the dipole polarizability [13, 14]. For laboratory electric fields this interaction is much smaller than the interaction with a molecular electric dipole moment, but it can be used to focus [8] or decelerate atoms [15].

**B. Electric field gradient of a focusing lens**

Any lens electrostatic potential \( \Phi \) can be expressed as the following multipole expansion in cylindrical coordinates \( r, \theta \):

\[
-\Phi(r, \theta) = E_0 r_0 \left[ \sum_{n=1}^{\infty} \frac{b_n}{n} \left( \frac{r}{r_0} \right)^n \cos(n\theta) + \sum_{n=1}^{\infty} \frac{a_n}{n} \left( \frac{r}{r_0} \right)^n \sin(n\theta) \right]
\]

where \( E_0 \) is the central field \( E = E_0(b_1^2 + a_1^2)^{1/2} \), \( r_0 \) is a scaling length, and \( b_n \) and \( a_n \) are the dimensionless constants of the \( 2n \)-pole strengths for normal (skew) elements.

Setting \( b_n = a_n \) for simplicity, converting to Cartesian coordinates, and retaining only the \( a_1 \), \( a_3 \), and \( a_5 \) terms (see section III C), the lens potentials for normal multipoles are:

\[
-\Phi_N(x, y) = E_0 x \left[ a_1 + \frac{a_3(x^2 - 3y^2)}{3r_0^2} + \frac{a_5(x^4 - 10x^2y^2 + 5y^4)}{5r_0^4} \right]
\]

(4)

or for skew multipoles are:

\[
-\Phi_S(x, y) = E_0 y \left[ a_1 + \frac{a_3(3x^2 - y^2)}{3r_0^2} + \frac{a_5(5x^4 - 10x^2y^2 + y^4)}{5r_0^4} \right]
\]

(5)

where \( E_0 \) is the field on axis for \( a_1 = 1 \).

Both of these potentials give the same total electric field \( E = -\nabla \Phi \).

\[
E(x, y) = (E_x^2 + E_y^2)^{1/2} = E_0 \left[ a_1^2 + \frac{2a_1a_3(x^2 - y^2)}{r_0^2} + \frac{a_3^2(x^4 + 2x^2y^2 + y^4)}{r_0^4} + \frac{2a_1a_5(x^4 - 6x^2y^2 + y^4)}{r_0^4} + \ldots \right]^{1/2}
\]

(6)

which has the electric field gradients:

\[
\frac{\partial E}{\partial x} = \frac{2a_1E_0^2 G_x x}{E} \quad \frac{\partial E}{\partial y} = -\frac{2a_1E_0^2 G_y y}{E}
\]

(7)

where

\[
G_x(x, y) = a_1 + \frac{a_3(x^2 + y^2)}{r_0^2} + \frac{a_3(x^2 - 3y^2)}{a_3^2 r_0^4} + \ldots
\]

(8)

\[
G_y(x, y) = a_1 - \frac{a_3(x^2 + y^2)}{r_0^2} + \frac{a_3(3x^2 - y^2)}{a_3^2 r_0^4} + \ldots
\]

C. Force on the molecule due to an electric field gradient

The force on a polar molecule in the \( x \) or \( y \) direction can now be calculated using Eq’s. (1, 7, and 8) with a suitable expression for \( \partial W / \partial E \) from section III A.

Non-linear forces, inside the focusing lenses limit the maximum beam size that one can transport without suffering beam losses, emittance (size) growth, and beam halo. Non-linear forces generally arise from higher-order multipole components of the lens electrostatic potential. Cylindrical electrodes, which are two-wire field lenses (with \( r_0 = \) half-gap), shown in Fig. 3 (a), have long been used to focus molecules in strong-field seeking states [1–7]. They contain skew multipoles of all odd orders and of equal strengths ( \( a_1 = 1, a_3 = -1, \) etc.)
a_5 = 1\ldots). As we will see below, its strong decapole field ($a_5 = 1$) reduces the area of the lens over which the focusing is linear (dynamic aperture). A beam transport line using these lenses will be limited to a smaller diameter beam compared to a similar transport line using optimized multipole lens such as the one shown in Fig. 3 (b).

To produce a larger linear region, than that of the two-wire field lens, we try to limit the multipole fields to those that are essential for the optics. We include a dipole field ($a_1$) to give a non-zero field on axis, without which strong-field seeking molecules defocus in both transverse directions; a sextupole field ($a_3$) to provide the linear focusing force (of order $r^3$); and a decapole field ($a_5$) to correct the non-linear forces (of order $r^5$) produced by the sextupole field. We omit the quadrupole field ($a_2$), because it bends the beam, and we omit the octupole field ($a_4$), because it introduces stronger non-linear forces (of order $r^7$) than those of the sextupole/decapole fields. All other multipole strengths are set to zero. In a real lens, however, the electrodes may be truncated equipotentials, with the consequence that small residual higher order multipoles will remain.

Since the lens potentials, as defined by Eqs. (4, 5), have two free parameters, we chose $a_1 = 1$, making $E_0$ the central field and we choose $|a_3| = 1$, making it easier to compare the optimized multipole lens to a two-wire field lens. The remaining choice is the decapole field strength $a_5$, which we use to optimize the lens optics. We do this in two ways: first by calculating, for specific molecules, the forces $F_x$ and $F_y$ inside a lens and comparing their linearity in $x$ and $y$ for different $a_5$, and second, in section IV, by simulation of beams in model transport lines.

If we calculate the horizontal force, $F_x$, on a molecule whose potential energy is $W = -d_x E$ (strong field limit), we find that it is most linear on-axis ($y = 0$) for no decapole field ($a_5 = 0$). For this example, the constant-force contours for $F_x$ in the $x, y$ plane are shown, for the two wire field lens, in Fig. 4 (a) and, for the zero-decapole lens, in Fig. 4 (b). The zero decapole lens is seen to have far better linearity than the two-wire field lens. (This is also true for $F_y$ which is not shown here.) In a similar way, we find that the optimized multipole lens outperforms the two-wire field lens in the low electric field limit (quadratic Stark effect), where the most linear field on axis has $a_5 = -\frac{1}{3}$.

D. End effects

Real lenses have finite length and the two-dimensional fields inside become three-dimensional fields at the ends. There are three effects. First, the $z$-component of the field gradient affects the transverse motion. This will be small if the lens aperture is small compared to the physical length of the lens.

Second, the lens field acts over a distance that is different than the physical length of the electrodes because of (extending) fringe fields at the ends. For linear design optics we approximate the real lens by a lens of constant central field $E_0$ over an effective length, different than the physical length of the lens. The effective length is determined by integrating the actual central field (found by numerical calculation or measurement) through the lens including the ends and dividing by $E_0$.

Third, the $z$-component of the electric field gradient, present at the ends of the lens, has an effect on the $z$-component ( lon-
The non-relativistic Hamiltonian (neglecting gravity) for a molecule of mass \( m \) and total velocity \( v \) is:
\[
H = \frac{1}{2}mv^2 + W.
\]
The Hamiltonian is conserved if the electric field is static. Consequently, a molecule in a strong-field seeking (weak-field seeking) state will gain (lose) kinetic energy entering the lens and then lose (gain) the same amount of kinetic energy upon exit [15, 16]. For the central \((x = y = 0)\) trajectory, the longitudinal velocity, \( v_{zE} \), of the molecule, in the lens is:
\[
v_{zE}^2 = v_{z0}^2 - W/2m
\]
where \( v_{z0} \) is the velocity of the molecule in the drift spaces. For most thermal or jet-source molecules in laboratory electric fields, the change in velocity is a small effect. (A beam of 560 m/s methyl fluoride molecules in the \( J = 0 \) state increases its velocity by 0.2 m/s upon entering an electric field of 10 MV/m.)

IV. BEAM TRANSPORT

A. Equations of motion

To track the trajectories of molecules passing through combinations of lenses, we use the transverse nonlinear equations of motion for a molecule traveling in the \( z \) direction, given from Eq’s. (1, 7, 8):
\[
x'' + \frac{2a_3}{r_0^2} \frac{E_0^2}{mv_{ZE}^2} \frac{G_z x}{E} \frac{\partial W}{\partial E} = 0
\]
\[
y'' - \frac{2a_3}{r_0^2} \frac{E_0^2}{mv_{ZE}^2} \frac{G_y y}{E} \frac{\partial W}{\partial E} = 0
\]
with trajectory gradients defined as \( x' = dx/dz = v_x/v_{ZE} \) and \( y' = dy/dz = v_y/v_{ZE} \).

We start the design optics using completely linear lenses \((G_z = G_y = 1, \text{ and } E = E_0)\), for which Eq. (9) reduces to:

\[
x'' + K_0x = 0 \quad y'' - K_0y = 0
\]
where \( K_0 \) is the lens linear focusing strength:
\[
K_0 = \frac{2a_3}{r_0^2} \frac{E_0}{mv_{ZE}^2} \frac{\partial W}{\partial E}
\]
\[
= \frac{2a_3}{r_0^2} \frac{dE_0}{mv_{ZE}^2} \quad \text{for } W = -d_xE
\]
\[
= \frac{a_3}{r_0^2} \left( \frac{\partial E_0}{mv_{ZE}^2} \right)^2 + \frac{1}{2} \frac{\partial E_0^2}{mv_{ZE}^2} \quad \text{for } W = -d_xE^2
\]
\[
= \frac{2a_3}{r_0^2} \frac{d^2E_0^2}{mv_{ZE}^2} \frac{C_2(2i+C_0dE_0)}{(B+C_0dE_0)} \quad \text{for } J = 0, \text{ from Eq. (21)}
\]

B. Example of methyl fluoride

For a realistic beam transport simulation, we assume a beam of methyl fluoride \((\text{CH}_3\text{F})\), in the \( J = 0 \) rotational state, having a longitudinal velocity \( v_{z0} = 560 \text{ m/s} \). (This is the approximate velocity of a beam produced by seeding methyl fluoride in an argon jet source with a reservoir temperature of 300 K (See section IV G). The electric field derivative of the potential energy (of the \( J = 0 \) state), is given by Eq. (3) with \( d_x = 6.25 \times 10^{-30} \text{ J/V/m} \) (1.86 Debye) and rotational constant \( B = 1.76 \times 10^{-22} \text{ J} \) (0.88 cm\(^{-1}\)).

To study long distance transport, we model a FODO lattice consisting of a sequence of identical F and D lenses separated by drift spaces \((O)\). Then, to study a complete beam line, we add an upstream lens section, to match the beam from the jet source into the FODO lattice, and a downstream section for a final focus of the beam (see Fig. 2).

C. FODO lattice

We chose a simple FODO lattice consisting of identical FODO cells (see Section II). The optics of a FODO cell starts in the center of an F lens (or D lens) and ends at the center of the next F lens (or D lens). At the ends of the FODO cell the beam is at a waist (defined as zero slope in the beam size) with a maximum size in one plane and a minimum size in the other plane.

For this study, we use 10 cm-long F/D lenses with scaling length \( r_0 = 6 \text{ mm} \) and separated by 40 cm drift spaces, giving a FODO cell length of 100 cm. This design leaves 80% of the cell unfilled. It is economical to build but does not have as large an acceptance as designs that filled more of the lattice (see section V).

In a FODO lattice the motion of a particle is periodic in phase-space. The phase-advance measures how far along the period it has proceeded from its initial starting point. The transverse linear optics are characterized by the phase advance in the FODO cell \( \mu_\perp \) (under focussed 0 < \( \mu_\perp < \pi \) over focused).

For our cell we choose \( \mu_\perp = \pi/3 \) rad for which all particles return to their initial phase-space position after 6 FODO cells. This specifies a central field of \( E_0 = 3.23 \text{ MV/m} \), which is not close to breakdown, as well as, a beam size that is nearly the minimum possible.
D. Modeling beam transmission

1. beam distribution in phase space

We take the molecular beam to be continuous (unbunched) and monoenergetic, in which case, it is completely specified by its density in the \((x, x', y, y')\) phase space (distribution function). The beam size in a transport line depends, not on the density, but, on the four-dimensional volume \((V_{4D})\) occupied by the beam. This can be defined, experimentally, in a number of ways: (a) the volume containing some fraction of the beam, (b) the volume calculated from the root-mean-square (rms) beam sizes, or (c) the volume defined by a set of collimators. These four dimensional volumes can be related to an equivalent theoretical distribution function of Kapchinsky and Vladiminsky (KV) [17]. With the KV distribution function, we can calculate the linear focusing beam sizes \((a_x, a_y)\) along the beam transport line and, in so doing, model the design optics.

The KV distribution consists of a uniform density of particles on a hyper-ellipsoid in four-dimensional phase space. At a beam waist \((a_x^0 = a_y^0 = 0)\), the hyper-ellipsoid is given by:

\[
\left(\frac{x}{a_x}\right)^2 + \left(\frac{y}{a_y}\right)^2 + \left(\frac{x'}{a_x'}\right)^2 + \left(\frac{y'}{a_y'}\right)^2 = 1
\]  

(12)

and has the volume, \(V_{KV} = \frac{4}{3}\pi a_x a_y a_x' a_y'\). Typically we have beam waists at the beam source (minimum), at any focus (minimum), and inside the focusing lenses (minimum or maximum).

The beam sizes along the beam transport line are given by the uncoupled envelope equations, which for the equations of motion [Eq. (10)] are:

\[
\begin{align*}
a_x'' + \kappa_0 a_x &= \frac{\varepsilon_{Kx}^2}{a_x} \\
a_y'' - \kappa_0 a_y &= \frac{\varepsilon_{Ky}^2}{a_y}
\end{align*}
\]

where \(\varepsilon_{Kx}\) and \(\varepsilon_{Ky}\) are the invariant transverse emittances which at a waist [see Eq. (12)] are simply given by \(\varepsilon_{Kx} = a_x a_{x'}\) and \(\varepsilon_{Ky} = a_y a_{y'}\).

If we project the KV distribution onto the \((x, x')\) plane, we obtain a uniform density of particles inside an ellipse of constant area \(\pi \varepsilon_{Kx}\). For an arbitrary beam distribution, the equations of motion, for the rms beam sizes, have the same form as Eq. (12). This defines the rms-equivalent KV distribution sizes as \(a_x = 2\sigma_x, a_{x'} = 2\sigma_{x'}, a_y = 2\sigma_y,\) and \(a_{y'} = 2\sigma_{y'}\). Under linear forces, a molecule remains on the same KV surface on which it started, with the shape of the ellipse changing but its volume remaining constant.

Since the non-linear forces inside a lens become stronger, the further the molecule is from the central axis, lenses may be evaluated by computing the increase in non-linear effects with increasing beam size. For this we use nested KV distributions of increasing volume, each of which is characterized by its volume, \(V_{KV}\). The simulation results are then independent of

![FIG: Emittance versus beam envelope size of the shell in the Kapchinsky-Vladiminsky (KV) equilibrium beam distribution.](image)

FIG. 5: Emittance versus beam envelope size of the shell in the Kapchinsky-Vladiminsky (KV) equilibrium beam distribution.

2. beam survival

We calculate the survival of a 560 m/s methyl fluoride beam in the \(J = 0\) state, through a 30-m FODO lattice, as a function of the decapole strength \((a_5)\). The lattice is described in section IV C, and we assume that the beam is already matched to the lattice. As molecular beam jet sources are usually axisymmetric, we take the emittance, \(\varepsilon_{KV}\), to be the same in both transverse planes. Then, the initial matched beam sizes and divergences, for the \(\pi/3\) phase-advance FODO cell, are:

\[
\begin{align*}
a_{\text{max}} &= a_x = (\varepsilon_{KV}\beta_{\text{max}})^{1/2} \\
a_{t'} &= (\varepsilon_{KV}\beta_{\text{max}})^{1/2} \\
a_y &= (\varepsilon_{KV}\beta_{\text{min}})^{1/2} \\
a_{t'} &= (\varepsilon_{KV}\beta_{\text{min}})^{1/2}
\end{align*}
\]

where \(\beta_{\text{min}} = 0.587\) m and \(\beta_{\text{max}} = 1.703\) m. As \(\beta\) is independent of the emittance, we use the initial maximum beam size \(a_{\text{max}}\) as our KV distribution size parameter:

\[
V_{KV} = \frac{1}{2}\pi \varepsilon_{KV} = \frac{\pi^2 a_{\text{max}}^2}{2\beta_{\text{max}}^2}
\]

The relation between the beam size and the emittance is shown in Fig. 5.

The trajectories were numerically integrated along the FODO lattice using a 4th-order Runge-Kutta algorithm. We take as lost those particles whose transverse displacement in the beam transport line in \(x\) and/or \(y\) became too large. The transported particles, which started on a zero-thickness KV-surface, finished up on a smeared-out fuzzy one, producing emittance growth and halo.

From the simulations, we found the fraction of the initial methyl fluoride beam surviving as a function of position along
FIG. 6: Calculated beam survival for 560 m/s methyl fluoride in the $J = 0$ state in a 30 m-long FODO lattice constructed of (a) two wire lenses or (b) optimized multipole lens with $a_5 = 0$. In each case $r_0 = 6$ mm and the central field, $E_0 = 3.2$ MV/m. Survival is plotted as a function of (KV) beam size. For the 100% beam survival, the beam intensity scales as KV beam size to the fourth power (see text). Consequently, the FODO lattice of optimized multipole lenses will transport about a factor of ten more beam than the same FODO lattice of two-wire field lenses.

the beamline. We did this for two different lens designs (two- wire field lens and optimized multipole) and for selected initial KV beam sizes. The results are shown in Figs. 6 (a) and 6 (b).

For the two-wire field lens, the largest KV beam that can be transmitted without loss is 1.4 mm. For the optimized multipole lens the largest lossless KV beam is 2.5 mm. If the initial phase-space density of the beams is approximately constant, the relative beam intensity, is given by the ratio of the lossless phase-space volumes. This is a factor of $(2.5/1.5)^4 \approx 10$ improvement by using the optimized multipole lenses in place of the two-wire field lenses in this FODO lattice.

Figures 6 (a) and 6 (b) also show that most of the beam losses occur in the first 5 m and by 30 m the losses are essentially complete. This suggests that extending the beam transport line to much longer distances will not further reduce beam survival.

To find the most appropriate values of the lens decapole strength, for other molecules, we also studied beam survival for the two limiting cases of strong electric field (linear Stark effect) and weak electric field (quadratic Stark effect). A KV beam size of $a_{max} = 2.5$ mm was used and the results, along with the results for methyl fluoride, are shown in Fig. 7.

For the weak field limit, as well as for the methyl fluoride example, we obtain the highest transmission for $a_5 = 0$. In the weak electric field limit the optimum value of $a_5$ is shifted to $a_5 \approx -0.2$, this nonlinearity in $\partial E/\partial x$ compensating the nonlinearity in $\partial W/\partial E$. Thus, an $a_5$ near zero would be a good choice for CsF molecules in low-lying rotational states, and an $a_5$ near -0.2 would be a good choice for ground-state atoms.

E. Velocity dependence of the beam transmission

So far we have considered only a monoenergetic beam. To look at the tolerance of the different lenses to energy deviations, we calculated the transmission for initially identical KV distributions but with different energies (560 m/s being the matched velocity.)

The results are shown in Fig. 8, for KV beam size of $a_{max} = 1.5$ mm, for both the two-wire field lens and for the optimized multipole lens. Both do quite well. At energies from 0.65 to 1.2 times the nominal energy of 640 K (velocity 560 m/s), transmission in the FODO lattice, of two-wire field lenses, is 50% or more of its maximum value. The FODO lattice of optimized multipole lenses does even better, as we would expect, since it has lossless transmission up to a KV size of $a_{max} = 2.5$. One should note that Fig. 8 does not represent the energy acceptance of a complete transport line, which will be limited by the source and matching optics.
from the jet source. (If needed, the beam can be focused, at the end, to an even smaller size.)

Finally, we note that much higher performance beam transport lines can be designed. If one uses stronger electric fields, shorter FODO cells, fills more of the beamline with focusing elements, and increases the size of the final focus, most of the solid angle from the skimmer can be accepted. Similar measures would allow one to efficiently transport and focus faster beams of molecules, such as those seeded in a helium jet source.

V. APPLICATIONS

Improving alternating gradient transport and focusing will make it easier to use molecules in strong-field seeking states for experiments, for beam transport, or to focus molecules for easier detection. In beam resonance experiments, molecules, prepared in a strong-field seeking state, can be detected, after a transition to a weak-field seeking state as a flop-in resonance.

The optimized multipole lenses can greatly reduce the problem of Majorana transitions [19, 20]. These are transitions that arise because different $m_J$ levels belonging to the same $J$ are degenerate in zero field. In very weak electric fields, a time-varying component caused, for example, by the motion of the molecule through the lens, can induce a transition to a different $m_J$ state with a very different Stark effect. This leads to beam loss, or loss of signal and a large background in sensitive

G. Transmitted intensity

If the characteristics of the initial molecular beam are known, the transverse and longitudinal acceptances of the full beam transport line may be used to calculate the beam intensity at the final focus. As an example, consider the beamline in section IV F, which has a skimmer radius of 0.75 mm and a $\pm 3.3$ mrad angular acceptance. From the point of view of an observer at the source, the entire beam transport line intercepts the same solid angle as would a 2 mm diameter collimator, located 0.3 m from the skimmer with no lenses. The intensity of an unfocused beam at this location can often be calculated or measured. From this, and a knowledge of the rotational state population fraction and the velocity spread, we can estimate the intensity at the final focus of the beamline.

For the methyl fluoride example, we assume a (seeded) jet source beam temperature of 3.5 K. (This is based upon the equations in Ref. [18] for a 0.0035 cm diameter source orifice, a source pressure of $1.9 \times 10^5$ Pa (1400 torr) of Ar at a reservoir temperature of 300 K, and a methyl fluoride seed of 5%. The source could be either pulsed or continuous.) The 3.5 K, results in a $J = 0$ population of about 30%, based upon a Maxwell Boltzmann distribution, and a kinetic energy spread of 11.4% which implies a longitudinal acceptance of about 60%.

Thus an intensity, equivalent to 18% of a unfocused methyl fluoride beam passing through a 2 mm dia. collimator 0.3 m from the skimmer, would reach the final focus some 14.5 m from the source. And since the beam transport line acceptances demonstrate transmission without loss, the same intensity should also reach a final focus much further from the source.

We complete the model transport line by adding matching optics upstream and downstream of the FODO lattice. This matches the source to the lattice and the lattice to the final beam focus. The matching, in both cases, is achieved using a doublet- and a triplet- lens configuration as shown in Fig. 2. As the matching lenses require stronger focusing, producing more non-linearities, we raise the lens scaling length in the matching lenses to $r_0 = 12$ mm. And since most of the beam losses occur early in the FODO lattice, we shorten the lattice to 15 m to reduce the computing time necessary for the simulation. The beam envelope in $x$ and $y$, and the placement of the lenses, is shown in Fig. 9, for an initial beam which is close to the linear focusing limit in the lenses.

For our example, of a 560 m/s beam of methyl fluoride in the $J = 0$ state, we assume that the jet source has a very small orifice and a skimmer of 1.5 mm diameter is placed 25 cm upstream of the first lens. An angular spread of $\pm 3.3$ mrad is fixed by collimators (see Fig. 2). This gives an initial beam emittance of 2.5 mm-mrad in both transverse planes, and a maximum beam size of 2.1 mm in the FODO lattice. In addition, the initial beam is taken to have a Gaussian energy spread of $\pm 10\%$ (rms). The calculated beam transport, through this beamline, is 84% of the entering beam, the losses being due to the energy spread. Thus, we can transmit and focus most of the velocity distribution from the jet source. (If needed, the beam can be focused, at the end, to an even smaller size. However in this case, we have to increase the size of the beam in the doublet/triplet lenses, where the non-linear forces will produce emittance growth and beam halo. If we are not limited by the electric field strength, we can correct this by using lenses with a larger scaling length.)

### Figure 8: Calculated transmission, as a function of relative beam kinetic energy, of a beam of methyl fluoride in the $J = 0$ state through a 30-m FODO lattice. The nominal energy is 640 K (560 m/s) and the KV beam size is $\sigma_{\text{max}} = 1.5$ mm. Points for the FODO lattice using two-wire field lenses and optimized multipole lenses (with decapole constant $a_5 = 0$) are shown as squares and circles, respectively.
FIG. 9: Lens placement of a complete transport line with matching lenses and a 15-m FODO lattice (center line), and the calculated rms beam envelope, in $x$ and $y$, for 560 m/s beam of methyl fluoride in the $J = 0$ state. We model the beam from a skimmed jet source and focus it to a 2 mm diameter spot. A schematic of the beam transport line is shown in Fig. 2. With a central field of 3.2 MV/m, scaling lengths of $r_0 = 6$ mm for the FODO lattice lenses, and 12 mm for the matching lenses, the beam transport line has an acceptance of 2.5 mm - mrad in both transverse planes. The longitudinal acceptance is about 84% for a ± 10 % rms energy spread.

resonance experiments.

The optimized multipole lenses reduce this problem by allowing one to have F and D lenses with their central electric fields in the same direction. This is done by changing the sign of $a_3$, the hexapole strength (and $a_5$) while leaving the dipole strength ($a_1$) unchanged. This eliminates field direction changes in going between F lenses and D lenses. If a weak dipole field is added to the region between the lenses, the molecules may never be in a rapidly changing weak field region.

The problem of Majorana transitions will be greatest for molecules in weak-field seeking states focused by pure quadrupole and/or sextupole lenses. These lenses have a vanishing field at the center and rapid changes in the field direction at their entrances and exits [7]. The problem will be diminished if the molecules are focused in alternating gradient fields using optimized multipole lenses with their central electric fields in the same direction and with alternating positive and negative values of $a_3$. These lenses have a non-vanishing field everywhere and can be optimized for a quadratic Stark effect by choosing $|a_5| = 0.2$. Again a small dipole bias field can be used between lenses.

The problem of Majorana transitions can be eliminated by choosing the $J = 0$ state; which is non-degenerate, always strong-field seeking, has the largest Stark effect of any rotational level, and is highly populated in a cold jet source beam.

The $J = 0$ and other strong-field seeking states have unique and useful properties that can be exploited in experiments. Within a rotational level, J, the $|m_J| = J$ states are strong-field seeking. States that are strong-field seeking in weak fields remain strong-field seeking in stronger fields, but weak-field seeking states will become strong-field seeking in the limit of strong fields. Thus, for strong field seeking states, there is no restriction on the size of the electric field that can be used to focus them. This is an advantage for molecules with small rotational constants and large dipole moments where the weak field-seeking states in low rotational levels become strong-field seeking in modest electric fields. CsF (Fig. 1) and other heavy alkali halides are good examples.

Long distance beam transport which can exceed 100 m or more has a number of applications. Since monatomic carrier gasses, clusters, and many contaminants will not focus through the beam transport line, it can be used to clean up a beam. For hazardous and radioactive molecules, a long beamline allows one to separate the source material and reservoir from the experiment and allows one to use radioactive detection in a lower background environment.

The long transit time (54 ms for the 30 m beam line in our example) corrected, if necessary for the small longitudinal velocity changes in the focusing elements, can be used for time-of-flight measurements with pulsed sources or a beam chopper. The different velocities will focus at slightly different longitudinal positions which may be exploited for position sensitive detection. Alternatively, by using a pulsed beam and ramping the electric field in the final focus lenses, all molecules may be brought to a focus at different times but at the same position.

The long flight path may be useful for colinear laser excitation of weak transitions. If the colinear laser excitation is combined with time-of-flight measurement, the Doppler spread from the velocity distribution may yield information about the absorption profile. Long transit time also allows for the decay of some long-lived states. Molecular beams may be run in both directions to form a very long colliding beam apparatus. The reservoir temperatures of the beam sources may be adjusted to equalize the focusing strengths of different molecules.

Acknowledgments

We thank Jason Maddi for performing the Stark effect calculation used in Fig. 1 and, along with Daniel Schwan, for early assistance with this work. We thank Swapan Chattopadhyay for helping to get the work started. This work was sup-