Pulsed, supersonic molecular beams and pulsed lasers are particularly well matched tools when combined in molecular reaction dynamics studies. Salient features of an experiment using two pulsed molecular beam sources, a pulsed ultra-violet laser for creating reactive atoms by laser ablation and a pulsed dye laser for performing laser-induced fluorescence detection of the products are described. Differences with steady-state molecular beam experiments are outlined with respect to the following points: facility of inverting the data, possibility of obtaining high signal-to-background ratios and wide ranges of collision energy. These points are illustrated with some results concerning the reactions: C(3P) + NO(X2Π) → CN(X2Σ+) + O(3P), C(3P) + N2O(X1Σ+) → CN(X2Σ+) + NO(X2Π) and Mg(1S0) + N2O(X1Σ+) → MgO(X1Σ+) + N(X4Σ+).

KEY WORDS: Pulsed lasers, laser ablation, laser-induced fluorescence, pulsed supersonic molecular beams, reactive scattering.

About ten years ago, it was stressed that bimolecular inelastic scattering experiments gained substantial advantages when performed with supersonic molecular beam sources, delivering pulses of very short duration and high peak intensity.\(^1-3\) In particular, the most severe limitation encountered in crossed, continuous molecular beam scattering studies, the signal versus background problem, was virtually eliminated due to higher instantaneous molecular beam intensities, which are not limited by differential pumping requirements, and time-of-flight discrimination of the background signal. Another major advantage of pulsed molecular beam sources suggested at that time, which became fully justified afterwards, was their facile interfacing with powerful pulsed ultra-violet lasers for generating high fluxes of reactive chemical species.

By laser photolysis of a stable precursor molecule, or by laser ablation of a solid target, many interesting reactive species, including free radicals and atoms, could be produced. Moreover, by focussing the laser beam in the hydrodynamic region or in the free molecular region of the pulsed molecular flow, or by varying the delay between the pulsed valve and the pulsed laser triggering, various degrees of cooling...
of the translational and internal degrees of freedom of the reactive species of interest could be obtained. Finally, other exotic species could be synthesised by laser-induced chemical processes resulting from interactions of photolyzed or photoablated products with reactive gases in the hydrodynamic region.

The highly efficient pulsed valve–pulsed ultra-violet laser combination launched reactive scattering experiments which could hardly have been performed by using the conventional continuous molecular beam approach. The first reaction dynamics studies of this new type were performed in our laboratory using the laser ablation technique for generating beams of ground state C and Al atoms. It is noteworthy that successful studies have since been reported by other groups. State-resolved differential cross sections have thus been obtained for the reaction \( \text{D} + \text{H}_2 \rightarrow \text{HD} + \text{H} \): the high potential energy barrier being overcome by the kinetic energy mainly due to the fast D atoms (2.2 eV in the laboratory frame) generated by ArF photolysis of D$_2$S in the collisionless region. The excitation function and the product state distribution have also been found for another isotopic exchange reaction: \( \text{CH} + \text{D}_2 \rightarrow \text{CD} + \text{HD} \), the supercooled CH\((^2\text{S}_1/2, \nu'' = 0, N'' = 1)\) radicals being created by ArF laser-induced chemistry in the hydrodynamic region of expanding CH$_3$I/Xe/H$_2$ mixtures.

CROSSED BEAM EXPERIMENTS

In our molecular beam experiments, reactions between ground-state atoms of more or less refractory elements and small oxidant molecules have been studied in the single collision regime, by crossing collimated, pulsed, supersonic molecular beams of short duration at right angles. Product quantum-state distributions and their collision energy dependence could be obtained by using pulsed laser-induced fluorescence (LIF) at the crossing point and by varying the velocity of the refractory atom beam. Such data can provide information about the nature of the potential energy hypersurface (PES) connecting reactants and products.

The present paper aims at describing some salient differences with the steady-state approach to such experiments. The first consists of the production of intense beams of atoms. The second one concerns data reduction to extract quantitative information from the spectra, which requires the solution of the problem of density-flux transformation. The last two features discussed hereafter result from the characteristics of our pulsed atom beam source, i.e. the significant increase in the signal-to-background ratio, and the ability to scan a wide range of relative translational energy.

**Pulsed Atom Beam Source**

A schematic view of the experiment is given on Figure 1. As the apparatus has been the object of a recent paper describing all the experimental details, this section only relates the key points of the metal-atom beam source design.

The idea of the pulsed supersonic metal-atom beam came from the spectroscopy
experiments performed at Rice University on supersonic metal clusters. Laser ablation of a solid metal inside the throat of a pulsed nozzle using a pulsed laser (generally doubled Nd:Yag) was found so efficient and so promising that it literally induced an explosion of the metal cluster field. However, our own goal being to produce atom beams if possible free of any aggregates, the supersonic metal cluster source was redesigned for the production of monoatomic species.

In the supersonic metal cluster experiments cited above, optimum cluster growth for various elements such as Cu, Mo, W or Nb was found to occur by focusing 7–15 mJ in 6 ns pulses of 532 nm radiation (2.33 eV or 225 kJ mole$^{-1}$) onto a 0.5 mm target resulting in fluences of $0.4–0.85 \times 10^4$ J m$^{-2}$. Our own approach was to use higher photon energy together with higher fluence in order to induce complete dissociation of all the fragments ablated from the solid by multiphoton absorption. This was first achieved with spatially filtered radiation of a KrF laser operated with unstable resonator optics. Pulses of 248 nm radiation (5 eV, or 482 kJ mole$^{-1}$) with the following characteristics: 7 mJ of energy within 1.2 mrad and 15 ns full-width at half-maximum (FWHM), could be focussed onto the target as 0.4 mm spots, resulting in a fluence of $5.5 \times 10^4$ J m$^{-2}$. Such a fluence, the highest that could be obtained with the type of laser used (Lambda Physik EMG 101 E with EMG 70
unstable resonator), was very satisfactory for Al atom experiments but insufficient for some of the C atom studies. Indeed, unlike most of the other refractory elements, carbon gives small clusters having strong chemical bonds ($D_0(C_2) = 6.21$ eV, $D_0(C_3) = 7.31$ eV) which perturb the system for two reasons. Firstly, $C_2$ is an extremely reactive radical which could yield the same LIF detected product: a case study is the $C + NO \rightarrow CN + O$ system with the reaction $C_2 + NO \rightarrow CN + CO$ occurring simultaneously. Secondly, $C_3$ exhibits intense absorption bands ($A^1\pi_u - X^1\Sigma_g^+$ transitions), extending throughout the ultra-violet and visible regions due to an unusually low bending mode ($63$ cm$^{-1}$), which could blur out the LIF spectra of the scattered product. $C_2$ and $C_3$ densities in the carbon beam could be efficiently lowered in experiments performed up to ca. $20 \times 10^4$ J m$^{-2}$ fluence using 7 ns pulses of 266 nm radiation delivered by a quadrupled Nd : Yag laser (Quantel SA YG585 with temperature phase-matched quadrupler).

Following quasi-complete dissociation of the ablated species into atoms, cluster growth was limited by minimising three body recombination reactions. These collision processes occur in the sonic extension channel, between the vaporising point and the vacuum, and afterwards in the hydrodynamic region of the expansion of the carrier gas in vacuum. Cluster growth is therefore favoured by increasing the gas load of the pulsed valve and the extension-channel length. The supersonic cluster experiments at Rice were performed with a magnetically operated double-solenoid pulsed valve giving under typical conditions a total gas output of ca. 70 mm$^3$ per pulse (normal pressure and temperature conditions) and a sonic channel length extending up to 30 mm after the vaporising point. In our metal-beam source, the Gentry and Giese$^1$ pulsed valve model used (Beam-Dynamics VCD-1) gave a gas load less than 20 times this value, and the sonic channel length was reduced to 3 mm. These characteristics were nonetheless found to ensure efficient electronic quenching of the atom metastable states, reducing them to negligible amounts compared to the atom ground-state concentration.$^{15}$ Any ions produced were easily removed from the beam with an electrostatic field.

The Laboratory to Centre of Mass Transformation

In crossed molecular beam experiments, the data obtained in the laboratory must be transposed in the centre-of-mass (CM) system before any interpretation of the results is made. This is common practice in reactive scattering experiments using an ionisation mass detector scanning the range of accessible laboratory angles: in general the laboratory intensity distribution is strongly weighted by those contributions which have low relative velocities. When LIF is performed at the beam crossing point, it yields laboratory quantum-state product densities which must also be transformed into fluxes in the CM frame. A simple density to flux conversion taking the recoil velocity vectors into account was introduced with the pioneering experiments in this field.$^{18}$ Its application, however, requires the angular distribution to be known, which is generally not the case.

The simple transformation invoked above is also constrained to a limited domain of validity. It cannot be applied when performing experiments with pulsed beams of
short physical length because a steady state is not reached when the LIF detection process is triggered. A misuse of this conversion also arises when experiments have been done with a laser beam not irradiating all the reaction zone but only its central part, which is very common in practice. In replacement, a mathematical model has been developed in our laboratory for reactive atom-diatomic systems. Product densities within the laser irradiated volume are calculated from the outcome of all the reactive events taking place in the beam overlap region up to the detection time. The model shows that the LIF detection efficiency can become extremely dependent not only on the internal energy state but also on the CM scattering angle. Moreover, changing the geometric arrangement of the experiment without changing the collision energy can drastically shift the maxima and minima of detection efficiency towards other values of the internal energy and recoil angle; in other words, for sharply peaked scattering angle distributions, the apparent rovibrational distributions of the excitation spectra can look totally different.

Typical results for the reaction:

$$C(3P_1) + NO(X^2\pi_r) \rightarrow CN(X^2\Sigma^+) + O(3P_1), \Delta\varepsilon_0 = -1.27 \text{ eV},$$

are displayed on the axonometric plots of Figure 2 which give the conversion function or LIF detection efficiency of the CN radical as a function of the CM recoil energy and the CM scattering angle. For example, point A on Figure 2a corresponds to radicals with no internal energy scattered forwards, point B to radicals at the excitation limit (almost no recoil energy) scattered backwards. The product flux in any given internal energy state can be obtained when dividing the intensity of a corresponding LIF rovibrational line by the integral of the conversion function over the scattering angle distribution.

Effects of beam collimation and pulse duration are clearly seen on these plots. It is worth noting that operating conditions can be found where the LIF detection efficiency is almost independent of the scattering angle (Figure 2b), simply by selecting the diameters of the collimators, which is of great interest when the angular distribution is unknown. Gross features of the scattering can also be deduced by performing experiments with different collimators (Figures 2a and 2b), the LIF signal behaving differently in the case of forward, backward or symmetric scattering. While these possibilities exist with pulsed, supersonic beam sources because the size and location of skimmers, which essentially act as collimators, are not critical parameters due to the absence of shock structure, they are, however, unrealistic in continuous beam experiments. The latter experiments can also have the drawback of giving very peculiar figures of the LIF detection efficiency (Figure 2c), unsuitable for correct determination of the internal energy partitioning. Such a very unfavourable case as in Figure 2c cannot be found using even mildly collimated pulsed beams of short duration because the overlap (both in space and time) of the beams away from the scattering centre is limited.

The Obtention of High Signal-to-Background Ratios

The signal-to-background ratio problem, which remains the major limitation in
Figure 2 Conversion function or LIF detection efficiency $Y(e_{\text{tr}}, \theta)$: z axis, versus CM recoil energy $e_{\text{tr}}$; y axis and CM scattering angle $\theta$; x axis, of the CN($X^2\Sigma^+$) product from the C($^3P_J$) + NO($X^2\pi_r$) reaction at $e_c = 0.234$ eV (beam velocities: $v_c = 2140$ ms$^{-1}$ and $v_{\text{NO}} = 830$ ms$^{-1}$), in the case of a laser beam irradiating the central part of the collision volume. a: pulse durations FWHM: $\delta t_c = 4.3$ $\mu$s and $\delta t_{\text{NO}} = 35$ $\mu$s, beam diameters FWHM at the collision zone: $d_c = 17$ mm and $d_{\text{NO}} = 14$ mm; b: same as a except $d_{\text{NO}} = 6$ mm; c: $\delta t_c$ and $\delta t_{\text{NO}} \to \infty$, $d_c = 17$ mm and $d_{\text{NO}} = 6$ mm.

molecular beam scattering experiments, is well illustrated with the case of the reaction:

$$C(^3P_J) + N_2O \rightarrow \text{CN}(X^2\Sigma^+) + \text{NO}(X^2\pi_r), \Delta e_0 = -2.78 \text{ eV}$$

Carbon is one of the most refractory elements and obtaining a carbon beam with a metal-oven effusive source can only be achieved by operating the oven at an extremely high temperature (up to 3500 K). At such a temperature, the effusive carbon beam contains roughly equal amounts of C, C$_2$ and C$_3$ species. On the one hand, complete time-of-flight discrimination of these species, which have different velocity distributions, by using a multiple chopper-disk velocity selector is hardly conceivable without loosing most of the C flux. On the other hand, leaving a small fraction of the highly rovibrationally excited C$_3$ radical in the C($^3P_J$) beam, would result in a background LIF signal throughout the whole spectral region of the CN
CN LIF spectra previously obtained in the early stages of development of our apparatus were not completely free of C3 lines.\(^6\)\(^7\) This problem was finally solved as stressed above by increasing the fluence used for the laser ablation of graphite. The CN(X\(^2\Sigma^+\)) excitation spectrum of Figure 3a, obtained this time at larger nozzle-crossing point distances and with beams collimated to 10° FWHM, has a larger signal-to-noise ratio and therefore contains dynamical informations of higher quality. The high rovibrational excitation results in prominent bandheads up to \(v'' = 6\). The \((6-7)\) band is not clearly observed although it should also exhibit a head in the vicinity of the \((5-6)\) one. However, due to the spectral congestion in this region, it is difficult to ascertain whether the rovibrational manifold is populated above \(v'' > 6\) or not. Synthetic spectra\(^1\),\(^2\) in fair agreement with the experimental one could be obtained by including a vibrational distribution only up to \(v'' = 6\) (Figure 3b and 3c).

It must be stressed at this level that the relative translational energy for a given CN quantum-state is not as well defined as in the C + NO \(\rightarrow\) CN + O reaction. It can, rather, take any discrete value allowed by the energy conservation law, depending upon the partitioning of the remaining available energy between fragment relative translation and NO internal degrees of freedom. As a consequence, some assumptions are needed to compute the CN LIF detection efficiency. Synthetic spectra have thus been computed assuming either similar internal energy distributions for CN and NO, or no NO internal energy at all (Figures 3b and 3c, respectively).

The following dynamical trends, independent of the hypotheses, can be derived from the inspection of the spectra of Figure 3. The rotational excitation is high: the distributions are definitely non-Boltzmann; however, the mean energies correspond to "rotational temperatures" as high as 11 000 K for CN (\(v'' = 1\)). The CN vibrational distribution is bell shaped, peaking at \(v'' = 3\) and exhibiting a sharp drop for \(v'' > 6\). This apparent excitation limit, roughly at half the total energy available to the products, strongly suggest that an important fraction of the reaction energy is deposited in NO. Such behaviour is consistent with the existence of a deep well in the PES\(^1\) which may favour a balanced energy partitioning between two vibrating rotators having almost identical vibrational and rotational constants.

**The Variation of the Relative Translational Energy**

The possibility of scanning the collision energy over a wide range of relative translational energies is another specification of a crossed pulsed supersonic beam apparatus. A study of the reaction:

\[
\text{Mg}^{(1S_0)} + \text{N}_2\text{O}(X^1\Sigma^+) \rightarrow \text{MgO}(X^1\Sigma^+) + \text{N}_2(X^1\Sigma_g^+), \Delta \varepsilon_0 \approx -1.5 \text{ eV},
\]

presently in progress illustrates its use.

Previous works by several groups have always remained at a qualitative level. The reaction was not observed with a Broida-type apparatus at 300 K.\(^2\)\(^2\) LIF detection of MgO(X\(^1\Sigma^+)\) failed at a first attempt in beam-gas configuration with a magnesium-oven effusive source operated at 1400 K and the scattering gas at 300 K but succeeded...
Figure 3  LIF spectrum of CN(\(X^2\Sigma^+\)) product from the C(\(^{3}\text{P}_j\)) + N\(_2\)O(\(X^1\Sigma^+\)) reaction at 0.28 eV relative translational energy. \(\text{CN}(B^2\Sigma^+ - X^2\Sigma^+)\) transition, \(\Delta v = -1\) sequence; \(v' - v''\): vibronic assignment. a: experimental spectrum, b: synthetic spectrum computed when assuming similar internal distributions for CN and NO, with the following relative vibrational populations, \(N_{v''} = 0.26, 0.67, 1.00, 0.66, 0.57, 0.28\) for \(v'' = 1-6\), respectively, c: synthetic spectrum computed when assuming no NO internal energy, with \(N_{v''} = 0.21, 0.60, 1.00, 0.75, 0.74, 0.27\) for \(v'' = 1-6\), respectively.

in a second experiment due to an improved detection sensitivity.\(^{23-24}\) Successful LIF detection was also reported in a flow experiment at 520 K but no spectrum was published.\(^{25}\) Finally MgO(\(X^1\Sigma^+)\) LIF was obtained with high signal-to-noise ratio by mixing Mg vapor from an oven at 1100 K with N\(_2\)O preheated at this temperature.\(^{26}\) Clearly, the apparent increase of reactivity with collision energy of the Mg(\(^1\text{S}_0\)) + N\(_2\)O system can be ascribed to the presence of a high potential energy barrier in the entrance channel of the ground-state PES which has been predicted by \textit{ab initio} calculations.\(^{27}\)
An excitation spectrum of the MgO(X^1Σ^+) product taken at ε_tr = 0.9 eV is presented on Figure 4. Several experiments were performed at a set of lower values of ε_tr obtained by varying the velocity of the Mg beam, keeping constant conditions on the N₂O beam, and recording alternatively the LIF intensities of the MgO(0–0) bandhead and of the Mg(^1P_1–^1S_0) resonance transition at 285.21 nm. The variation of the relative reactive cross section with collision energy could thus be extracted with the assumption that the internal energy partitioning and the angular distribution of the MgO product did not change markedly in this energy range. First, quantitative

![Figure 4](image-url)  
**Figure 4** LIF spectrum of MgO(X^1Σ^+) product from the Mg(^1S_0) + N₂O(X^1Σ^+) reaction at 0.9 eV relative translational energy. MgO(B^1Σ^+ − X^1Σ^+) transition, Δν = 0 sequence; ν′ − ν′: vibronic assignment.

![Figure 5](image-url)  
**Figure 5** Relative reactive cross-section of the Mg(^1S_0) + N₂O → MgO(X^1Σ^+) + N₂(X^1Σ_g^+) product channel, as a function of the relative translational energy.
dynamical information on the reaction is shown with the excitation function (Figure 5) which exhibits a sharp threshold near $\varepsilon_{tr} = 0.5$ eV. This translational energy threshold represents an upper limit of the energy barrier located on the entrance channel of the PES.

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