Natural Lifetimes of Excited-states of Neutral Nitrogen Determined By Time-resolved Laser Spectroscopy

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Natural lifetimes of excited states of neutral nitrogen determined by time-resolved laser spectroscopy

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Radiative lifetimes were determined for three quartet states of neutral nitrogen, and sequences of Rydberg states were studied using depletion spectroscopy. Free nitrogen atoms were generated by photodissociation of N₂O using frequency-tripled dye-laser radiation that was two-photon resonant with the 2p³ 3p S or D states. Further quartet states were reached by a subsequent single-photon absorption. We obtain τ(2p³ 3p D₁/₂) = 44(2) ns, τ(2p³ 3p S₁/₂) = 26.0(1.5) ns, and τ(2p³ 6s P₃/₂) = 41(7) ns.

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I. INTRODUCTION

Nitrogen is the most abundant atmospheric gas. Thus, nitrogen molecules and free atoms are of great importance in a multitude of processes, i.e., atmospheric physics and chemistry, combustion kinetics, and plasma processes. Furthermore, nitrogen is abundant in the sun and the stars. Therefore, it is of considerable interest to accurately study the energy structure and the radiative properties of these atoms. In the present paper we present the results of a laser-spectroscopic study of free nitrogen atoms. Little data from methods using selective excitation exist since considerable difficulties have to be overcome in the case of nitrogen. Free nitrogen atoms have to be produced by dissociation of a suitable molecule. The fact that the resonance lines in nitrogen fall in the vacuum-ultraviolet region is a further complication for laser spectroscopy.

The problems were first solved by Bischof, Perry, and Crosley [1] who dissociated N₂ atoms in a microwave discharge. The 2p³ 3p D state could be reached by two-photon absorption of 211-nm quanta and the lifetime of the state was measured by observing the decay to the 2p³ 3s 4P state. In a later investigation [2], the same group revised the previously obtained lifetime value by a factor of 1.6. The same optical scheme was used in our department for detecting nitrogen atoms [3,4], and earlier also oxygen atoms [5], in a flame. While a hot flame is a natural atomizer, the high ambient pressure results in a very unsuitable environment for studying radiative properties. We have demonstrated that, using a sufficiently high laser power in the two-photon excitation scheme, we could produce oxygen atoms by photodissociation of NO₂ molecules in a low-pressure environment. While thus the same uv laser beam was used for photodissociation and population of the 2p³ 3p 4P state, a further dye-laser beam was used to transfer the atoms into more highly excited states that in this way could be made available for studies of radiative properties [6]. In the present paper, we report on measurements of this kind on the nitrogen atom.

A relevant energy-level diagram for the nitrogen atom is shown in Fig. 1 [7], where transition wavelengths have also been indicated. In the present work two states were populated by two-photon absorption, the 3p 4D₁/₂ state studied previously [3,4] and the 3p 4S₃/₂ state that had not been studied before using selective excitation techniques. In the figure, second-step excitations from these platform states are also indicated. The transfer of platform-state atoms to such higher states was monitored as a decrease in the near-ir fluorescence light from the platform states. This “depletion-spectroscopy” technique was used in Ref. [6] and allows Rydberg-state and also autoionization-state spectroscopy without observing fluorescence light or photoelectrons from such states. We demonstrate such spectroscopy for a large number of nitrogen states.

II. EXPERIMENTAL ARRANGEMENT

A schematic diagram of the experimental arrangement used in the present experiments is shown in Fig. 2. A Quantel Datachrome 5000 laser system was employed as an excitation source. It consists of a YG 581c Q-switched Nd-YAG laser (YAG denotes yttrium-aluminum garnet) pumping a TDL 50 tunable dye laser,
which was operated on (DCM) [4-(dicyanomethylene)-2-methyl-6-(p-dimethyl-amino styryl)-4H-pyran] dye. The dye laser delivered pulses of tunable radiation of about 10-ns duration at a repetition rate of 10 Hz. The dye-laser radiation was frequency doubled in a KDP (potassium dihydrogen phosphate) crystal and tripled by mixing of the doubled frequency and the fundamental frequency in a BBO (β barium borate) crystal. In order to achieve the process, a mechanically compressed crystalline-quartz plate was used to rotate the fundamental and the doubled dye-laser radiation to obtain parallel polarization planes for the two waves before entering the BBO crystal. Starting with a dye-pulse energy of 50 mJ at 621 nm, we ended up with about 2 mJ at the 207-nm wavelengths required for two-photon excitation of the $3p^4S_{3/2}$ state. The tripled frequency was isolated from the unwanted wavelengths by a Pellin-Broca prism and was subsequently sent into the experimental chamber which was filled with a low-pressure atmosphere of N$_2$O.

N$_2$O was chosen because of its photophysical properties [8]. The processes

$$\text{N}_2\text{O} \rightarrow \text{N}_2 + \text{O}(3P),$$

$$\text{N}_2\text{O} \rightarrow \text{NO} + \text{N}(^4S)$$

are energetically possible for wavelengths below 250 nm. The first of the two processes is the most efficient one dominating over the second, desired one, by a factor of 50 (Ref. [8]). The ground-state nitrogen atoms, created in the second process, find themselves two-photon resonant with the strong uv field.

In the experiments with a second-step excitation, an extra beam was diverted from the Nd:YAG laser to pump a Quanta-Ray PDL-1 dye laser operated with Rhodamin 6G or DCM dye to cover the wavelength regions 550–580 and 600–670 nm, respectively. The red laser light generated by the second dye laser was sent into the experimental chamber close to and parallel with the uv beam. Inside the vacuum chamber the two beams were focused by an $f = 10$-cm lens. Producing an overlap at the focus of the uv beam was simplified since the chromatic lens focused the uv light earlier than the visible light of the second-step beam. The detection of the fluorescence was made through a window perpendicular to the laser beam with a Hamamatsu R943-02 photomultiplier tube preceded by different interference filters. In the lifetime measurements, the signal was recorded by a Tektronix DSA 602 transient digitizer, while in the depletion spectroscopy a Stanford Research Instruments model 265 boxcar integrator was used. Both were connected to an IBM-compatible computer for data processing. When using the boxcar integrator, the intensity of the third-harmonic laser light was monitored with a diode in order to compensate for pulse-to-pulse fluctuations. Compensation for detector nonlinearities was made by a signal normalization routine used by Wolf and Tiemann [9]. Both boxcar integrator and transient digitizer were triggered by a diode, which detected a reflection of the red light from the first-step dye laser.

The vacuum chamber used for the experiments was cubic with a side of about 20 cm. After evacuation, N$_2$O could be introduced through a needle valve. The chamber was surrounded by coils for compensation of the earth's magnetic field. The gas pressure in the chamber was measured with a Pirani vacuum gauge, which was calibrated for N$_2$O in the pressure region of interest.

For depletion spectroscopy, calibration of the second-step dye laser was made by letting a beam reflection hit a neon-filled hollow-cathode lamp. When the wavelength coincided with certain neon transitions, a weak current transient was generated in the lamp discharge current. This signal could be sampled and used for wavelength calibration during a depletion-spectroscopy scan.

**III. MEASUREMENTS AND RESULTS**

The correct tuning of the first excitation step laser (two-photon transition) was normally monitored by observing the near-ir fluorescence light from the platform state. In an early stage of the experiments, we tried to use the presence of stimulated emission on this line in the forward direction for ensuring correct tuning. Strong, stimulated emission was detected with a photodiode as has been reported in different investigations at this department [3], but only at gas pressures far too high for the experiments of interest in the present study. We also investigated the possibility of using the photoionization current expected when the platform states were two-photon resonant with the laser. Strong photoionization was detected using metal-plate electrodes close to the interaction volume, but, unfortunately, the atomic signal was mostly blurred by a nonresonant molecular background. This is a common problem in photoionization monitoring when not using a time-of-flight spectrometer or some other type of mass filter. Photoionization detection involves the presence of light scattering objects close to the interaction volume that are undesirable when detecting low levels of laser-induced fluorescence.
The $3p\,^4S_{3/2}$ state was excited by two-photon absorption at 206.7 nm. Fluorescence was detected in the decay to the $3s\,^4P_{3/2,1/2}$ states (742–747 nm) at pressures ranging from 0.05 to 0.5 mbar. The light from the decaying nitrogen atoms was averaged on the transient digitizer for multiple laser pulses. The decay constant was obtained by computer fitting of an exponential for times long enough to be outside the duration of the laser pulse. Under certain experimental conditions, more accurate data can be obtained by fitting a convolution of an exponential with a system response function to the experimental curve. However, such a method is not applicable in our case with multiphoton interaction and an unknown degree of saturation. The natural radiative lifetime unperturbed by collisions is obtained by linear extrapolation of the decay constants versus pressure to zero pressure. Normally, the collisional cross sections can be inferred from the slope of such Stern-Vollmer plots. This interpretation is not valid in our case. Using the previously described method for producing free atoms, several kinds of atoms, molecules, and ions are present in the interaction region. Hence, the collision partner of the nitrogen atoms is not well defined.

In corresponding measurements on the $3p\,^4D_{7/2}$ state, the wavelength for two-photon absorption was 210.8 nm and fluorescence light was detected at 868 nm down to the $3s\,^4P_{3/2}$ state. Decay curves for both investigated platform states are shown in Fig. 3. Several measurement series were taken and the corresponding Stern-Vollmer plots were constructed as shown in the lower part of Fig. 3. From the axis intercepts of the fitted lines, the radiative lifetimes were evaluated. The results are given in Table I, where the error bars include possible systematic errors as well as the statistical scattering in the data, resulting in an uncertainty in the extrapolation to zero pressure. Although other $J$ components of the platform state were observed, only the state providing the strongest signal was selected for an accurate determination of the radiative lifetime.

Depletion spectra for transitions originating in the two platform states were recorded by scanning a second-step dye laser. By proper tuning of the first-step laser, well-defined fine-structure levels of the platform state could be selected. Every time the wavelength of the second-step laser coincided with a transition to an upper level,
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