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Efficient, highly selective laser isotope separation of carbon-13

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ABSTRACT We recently demonstrated an original approach to highly selective laser isotope separation of carbon-13 that employs vibrational overtone pre-excitation of CF₃H together with infrared multiphoton dissociation [O.V. Boyarkin, M. Kowalczyk, T.R. Rizzo, J. Chem. Phys. **118**, 93 (2003)]. The practical implementation of this approach was complicated by the long absorption path length needed for the overtone excitation laser beam. In the present work, we employ a low overtone level for the pre-excitation that shortens this pathway, facilitating engineering of the process. We propose an optimal configuration of the isotope separation scheme and consider a realistic example of a separation unit for isotopic enrichment of carbon-13 to 94%–98%. The photon energy expenditure of 97 eV per separated atom is much lower than that of the current commercial laser technology, making this process economically feasible.

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1 Introduction

While the use of infrared lasers for molecularbased separation of stable isotopes has led to many interesting findings and promising demonstrations [1-7], few laser isotope separation schemes have reached the stage of commercial application [8]. In the case of schemes based on infrared multiphoton excitation, the limitation is their low isotopic selectivity. Physical methods of isotope separation, such as low temperature distillation of CO in the case of carbon, do not alter the working molecule, and this facilitates the implementation of a multi-cycle separation process [9]. Although the selectivity of each cycle is low, the repetition of many separation cycles makes it possible to achieve the high levels of selectivity demanded by the market. Because molecular laser isotope separation techniques that are based on infrared multiphoton dissociation chemically convert the parent molecule into a new species, repetitive cycles can be complicated and costly. Development of a single-stage, energy-efficient, laser separation technique that could produce the high levels carbon-13 enrichment required by the market (above 95% or above 99%) would fill an important niche.

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Over the last several years we have developed a highly selective separation scheme for carbon-13 that combines isotopically selective vibrational overtone pre-excitation (OP) of the CH-stretch vibration in ¹³CF₃H followed by selective IRMPD of the pre-excited molecules (Fig. 1) [10-12]. The CF₂ fragments combine via 3-body collisions to form C_2F_4 , which can be separated from the starting material. We have demonstrated selectivities up to 9000 (99% enrichment in carbon-13 starting from CF₃H in natural abundance) that can be enhanced at high sample pressure by V-V collisional relaxation [11]. Since our original discovery of this unexpected effect [11], we have taken several steps that significantly increase the yield of the process, bringing it closer to commercial relevance. In particular, overlapping the preexcitation laser pulse with a short (50 ns) dissociation pulse from a CO₂ laser allows working pressures of 100-250 mbar while maintaining a high relative dissociation yield and high isotopic selectivity [12].

Despite the high working pressure of the process, the low absorption cross section of the 3_1 vibrational band [14] that we originally employed for overtone pre-excitation makes the 1/e absorption length of the pre-excitation beam fairly long (e.g., > 200 m at 100 mbar), inhibiting the efficient use of photons. Shortening the absorption length can be achieved by pre-exciting lower CH-stretch bands, which have higher absorption cross-sections. Table 1 summarizes the spectroscopic data for the absorption bands of interest. Pre-excitation through the 2_1 and 2_2 bands are 6.7 and 4.4 times more efficient than use of the 3_1 band, and if the high level of selectivity could be maintained, this would lead to a significantly higher performance of the process. Moreover, generation of highenergy, high repetition rate pulses in the region of 1.7 µm (for the 2_1 and 2_2 bands) seems to be significantly easier than at 1.14 μ m (for the 3₁ band) using an OPO + OPA system pumped by a Nd: YAG laser, for example. Thus, use of either the 2_1 or 2_2 band would be highly desirable for simplifying the engineering of the process and increasing its practical relevance. There are, however, two factors that may complicate the use of these bands. First, the isotopic shift of the 2_1 and 2_2 bands is smaller than that of the 3_1 band, and this reduces isotopic selectivity of the pre-excitation step and thus may substantially reduce the overall selectivity of the process. Second, IRMPD of molecules pre-excited to the first overtone level should be less efficient than dissociation of molecules



FIGURE 1 Schematic energy level diagram of the overtone pre-excitation infrared multiphoton dissociation method for highly selective carbon-13 isotope separation using CF_3H as the parent molecule

from the 3_1 level, because the former have less vibrational energy. In this paper we present the results of our experiments applying the OP-IRMPD approach for separation of carbon-13 in CF₃H, employing the 2_1 and 2_2 bands for pre-excitation. We compare these results with those obtained for the 3_1 band and evaluate the overall performance of the isotope separation scheme. We also verify that the process can benefit from reducing the intensity of the dissociation laser pulse at fixed fluence, as we have previously suggested [12]. Finally, we estimate the quantum efficiency of the process and consider a potential example of its practical implementation.

2 Experimental approach

The experimental set-up, shown in Fig. 2, is identical to that used in the earlier experiments where we employed the 3₁ band for pre-excitation [12], except for the method of generating tunable infrared laser radiation. The current system uses difference frequency mixing (DFM) in a nonlinear crystal to generate near IR light and optical parametric amplification (OPA) to achieve high pulse energy. A fraction of the fundamental beam from a single-mode, Nd: YAG laser (Spectra-Physics GCR-250) is frequency-doubled to pump a dye laser (Lambda Physik Scanmate 2E). The 30–40 mJ output of the dye laser is mixed with 150 mJ of the Nd: YAG laser fundamental in a LiNbO₃ crystal to generate 2–4 mJ of tunable IR radiation. The output of the DFM stage is further amplified in a two-stage OPA pumped by 200–300 mJ



FIGURE 2 Optical layout of the experiments

Band in ¹³ CF ₃ H	Transition wavenumber ^a cm^{-1}	Isotopic shift ^b $v(^{12}C) - v(^{13}C),$ cm ⁻¹	Absorption cross-section at $T = 300 \text{ K}^c$, cm^2
31	8753	39.7	1.8×10^{-21}
2_{1}	5936.6	22.8	12×10^{-21}
2_{2}	5680.9	29.5	7.9×10^{-21}

^{*a*} From [15]; ^{*b*} Calculated from [14] and [15]; ^{*c*} Calculated considering integral absorption intensities from [14] and our measurements of absorption spectra

TABLE 1 Spectroscopic data for selected CH-stretch overtone bands of $^{13}\text{CF}_3\text{H}$

of the infrared fundamental of the same Nd:YAG laser. This results in 120 mJ, 6 ns pulses of tunable around 1.7 μ m radiation with 0.1 cm⁻¹ bandwidth. The pre-excitation laser beam is slightly focused by a telescope before entering the reaction cell through a BaF₂ window at the Brewster angle. The dissociating CO₂ laser beam is focused by an F = 100 cm lens and enters the reaction cell from the opposite side. The 50 ns long pulses of this laser overlap the 6 ns pre-excitation pulses. The isotopic composition of the C₂F₄ dissociation product is measured by a computer controlled quadruple mass-spectrometer (Balzers, QMS-422) as previously described [12].

3 Results

3.1 *Pre-excitation to the first* CH-stretch overtone

The *Q*-branches of the 2_2 and 2_1 CH-stretch overtone transitions in room temperature ¹³CF₃H appear as prominent peaks – an order of magnitude more intense than their





P and R-branches [15]. Figure 3 shows the photoacoustic absorption spectra of naturally abundant CF₃H in the regions of the 2_2 and the 2_1 band *Q*-branches of the carbon-13 species. Because of its larger isotopic shift, the 2_2 *Q*-branch of the carbon-13 species is overlapped with high J transitions of the *P*-branch in ¹²CF₃H. Due to the splitting of high *J*-states of the 2_2 level by weak perturbations [14], these transitions appear unresolved at our moderate spectral resolution of 0.08 cm^{-1} , forming a nearly continuous background. Given the degree of spectral overlap, the maximum isotopic selectivity that one could expect in the pre-excitation step using the 2_2 band is around 90, which is substantially lower than the selectivity of 400 previously measured using the 3_1 band [11]. Because of its smaller isotopic shift, the Q-branch of the 2_1 band in ¹³CF₃H falls into a region of more intense, but resolved, transitions of the carbon-12 species. Here the 2_1 *Q*-branch in ¹³CF₃H is adjacent to a single transition. The isotopic selectivity using the 2_1 level, estimated by fitting the spectrum in Fig. 3, could be up to 100 if a narrow-bandwidth laser is used for pre-excitation. Because of the particular overlap pattern of these transitions, excitation at the maximum of the $2_1 Q$ -branch of ¹³CF₃H will not necessary yield the highest isotopic selectivity - some optimization of the pre-excitation frequency is required.

Figure 4a and b shows the isotopic selectivity and relative IRMPD yield (i.e., fraction of the pre-excited molecules that have been dissociated) as a function of CF₃H pressure for the OP-IRMPD process with pre-excitation to the 3_1 and 2_2 bands. In the latter case, the dissociation fluence has been increased to 4.5 J/cm² in order to achieve a relative IRMPD yield approximately equal to that obtained with 3 J/cm^2 when the 31 band is pre-excited. This allows a convenient comparison of the two experiments using a single parameter, isotopic selectivity, which is less ambiguous than the simultaneous comparison of two parameters (selectivity and yield). When extrapolated to zero pressure, the isotopic selectivity with pre-excitation through the 2_2 band is about three-fold lower than with pre-excitation through the 31 band. However, already at sample pressures above 50 mbar, the selectivity in both experiments becomes almost equal, and further increase in pressure does not influence the ratio between them. This suggests that the enhancement of selectivity by collisions is higher in the case of pre-excitation to the lower energy level (2_2) compared to that achieved using the 3_1 level. Thus, using pre-excitation via the 2_2 band and increasing the fluence of the dissociation laser by a factor of 1.5, one can achieve approximately the same overall isotopic selectivity and fraction dissociated in the IRMPD step as with the 31 band for sample pressures above 50 mbar. However, because the absorption cross-section of the 2_2 band is 4.4 times higher than of the 3_1 band, the absolute yield of the processes per irradiated unit volume increases by this same factor (Fig. 4c). Therefore, for the same (short) absorption pathway, the overall productivity of the process increases a factor of 4.4 using 2_2 versus 3_1 as a pre-excitation level.

The productivity is even higher with pre-excitation through the 2_1 band. At 100 mbar we observe an increase in the per volume dissociation yield by a factor of 1.5 compared to the 2_2 band, which reflects the ratio of the 2_1 to 2_2 absorption cross sections. While there is a decrease in the pre-excitation



FIGURE 4 Isotopic selectivity (**a**), relative IRMPD yield (**b**) and the absolute yield (**c**) as a function of CF₃H pressure in the case of pre-excitation to the 2₂ level (*filled markers*) and the 3₁ level (*open markers*). The absolute yield is given in units of ¹³C atoms per cm³ and per J/cm² pre-excitation fluence. The dissociation fluence is 4.5 J/cm^2 and 3 J/cm^2 for the 2₂ and 3₁ bands respectively. The pre-excitation and dissociation pulses are overlapped in time

selectivity of less than 10% in changing the level from 2_2 to 2_1 , it can be regained by increasing the sample pressure, although with 10% decrease of dissociation yield. Thus, for a fixed isotopic selectivity and a fixed laser beam pathway, the process yield with pre-excitation through the 2_1 band is 1.35 times higher than pre-excitation through the 2_2 band and about six times higher than with the 3_1 band employed previously. In what follows we thus concentrate on experiments employing pre-excitation through the 2_1 band.

3.2 Dissociation by a temporally modified CO₂ laser pulse

The output beam of our free-running CO_2 laser appears as a train of 1-2 ns pulses separated by the cavity



FIGURE 5 Oscillograms of the regular (a) and the "doubled" (b) dissociation CO_2 laser pulses obtained with 2 ns resolution

round trip time of 10 ns (Fig. 5a) as a result of self modelocking [16]. This spiky structure of the CO_2 laser pulse imposes a certain time structure on the multiphoton absorption process, which competes with collisions. Considering the competition between up-pumping and collisional V-V relaxation [12], there should be a predominance of up-pumping during a laser intensity spike and of relaxation in the short time between the spikes. Thus, the entire process of multiphoton excitation consists of a number of short, repetitive subprocesses that limit the possibility of increasing the sample pressure with simultaneous increase of dissociation laser fluence. Indeed, if the collisional relaxation rate is too high, during the time-delay between two laser spikes the pre-excited molecules may relax below the energy where efficient IR multiphoton pumping can occur. Increasing the dissociation laser fluence to compensate this will increase the probability for dissociation of both the relaxed and the ground state carbon-12 species. It is clear that for the efficient use of the competition between up-pumping and collisional V-V relaxation during the CO_2 laser pulse, the distance between the adjacent spikes should not considerably exceed the average time between two subsequent collisions. We previously measured a value of $5-7 \,\mu s^{-1} mbar^{-1}$ for the vibrational relaxation rate constant of the first CH-stretch overtone level in CF₃H [11], which is close to the recently calculated value of $5 \,\mu s^{-1} mbar^{-1}$ [13]. Given this rate constant, one can see that already at CF₃H pressures above 20 mbar, the spiky structure of the dissociation pulse has a considerable destructive influence on the performance of the process. A smooth temporal pulse shape would thus be highly desirable. This can be achieved by forcing the dissociation laser to oscillate on a single longitudinal mode by one of several means: injection seeding; using a combination of a high-pressure discharge region with a low pressure gain section; or by using an intra-cavity cell with a selective absorber, each of which would require rebuilding our CO₂ laser. To demonstrate the influence of laser pulse shape, we have chosen a simple approach that modifies the shape of our laser pulses outside of the laser cavity. Although this approach cannot be used in a large-scale apparatus, its simplicity and pulse-to-pulse reproducibility makes it convenient for laboratory experiments with a small working volume. The CO₂ laser beam is split to two beams of nearly equal intensity by an ZnSe beam splitter. One of the two beams is optically delayed by 5 ns with respect to the other split beam. Both beams are then focused by the same F = 100 cm ZnSe lens and overlap in the 2 cm long reaction cell. This has the effect of interposing spikes in the pulse of one beam precisely between those of the other, shortening the delay between spikes that the molecules experience from 10 ns to 5 ns. Figure 5 shows the temporal profile of a pulse emanating from the CO₂ laser and the profile of the "doubled" pulse, measured using a Ge photon drag with 1 ns time resolution. Although the spiky structure in such "doubled" pulses has not been completely smoothed, we still can draw conclusions about the potential advantages of using a truly "smooth" dissociation laser pulse in our isotope separation process.

The yield and selectivity of the OP-IRMPD process using 2_1 pre-excitation is shown in Fig. 6 as a function of the sample pressure for the regular and "doubled" dissociation laser pulses at a dissociation laser fluence of 3 J/cm^2 . As the figure illustrates, the "doubled" pulses cause a slower increase of isotopic selectivity and slower decrease of dis-



FIGURE 6 Isotopic selectivity (**a**) and relative IRMPD yield (**b**) as a function of CF₃H pressure in the case of pre-excitation to the 2_1 level by the regular (*opened markers*) and smoothed (*filled markers*) dissociation pulses. The pre-excitation fluence is 1 J/cm^2 , and the dissociation fluence is 3 J/cm^2 . The pre-excitation and dissociation pulses are overlapped in time

sociation probability upon increase of CF₃H pressure compared to the regular CO₂ laser pulses. The effect is similar to what one would observe by increasing the fluence of a regular pulse [12]. A certain value of isotopic selectivity achieved either with the smoothed, "doubled" pulse or with the intense, regular pulse corresponds to the same relative IRMPD yield. To achieve, for example, isotopic selectivity of 2000 (95.4% of carbon-13 in the final product), one can use either the regular CO₂ laser pulses and a pressure of 65 mbar or the smoothed pulses and 100 mbar. In both cases approximately 18% of pre-excited molecules are dissociated. However, working at higher pressure is advantageous from point of view of absolute productivity of the process. For the fixed selectivity of 2000, the absolute productivity per irradiated volume unit (per volume productivity) with our smoothed pulse is 1.5 times higher than for the regular (selfmode locked) CO_2 laser pulse. We believe that the use of a truly smooth pulse shape would increase the productivity even further.

3.3 Absorption of CO₂ laser radiation by pre-excited molecules

The dissipation of vibrational energy of preexcited carbon-13 species by collisions implies that the average number of absorbed CO_2 laser photons per molecule is higher than energetically necessary for dissociation (19 photons from the 2₁ level, for example). Moreover, some of the pre-excited molecules may not absorb enough CO_2 laser photons to dissociate. This will further increase the average number of photons absorbed per dissociated molecule. The number of photons actually absorbed is a key to evaluate the minimum energy expenditure per separated atom and the characteristic absorption length of the dissociation laser beam, the latter of which should be matched to that of the preexcitation laser beam. This number thus highly influences the practical design of the process and, ultimately, the productivity of a separation unit.

In order to estimate the average number of CO_2 laser photons absorbed per pre-excited molecule under the typical conditions of the separation process, we measured the CO₂ laser energy absorbed by CF₃H molecules pre-excited to the 2_1 level. In this experiment, a near collimated CO_2 laser beam of $2-3 \text{ J/cm}^2$ fluence is counter-propagated with a pre-excitation beam of larger diameter in the 50 cm long cell, which is filled with 100 mbar of CF₃H. A small fraction of the dissociation beam is split after the cell and sent to a pyroelectric detector. The CO_2 laser is tuned to the 9P(14)line at 1052.2 cm^{-1} , where the absorption cross-section of ground state CF₃H has an absorption minimum [17]. The fluence of the pre-excitation laser is 1 J/cm^2 , which promotes $\sim 10\%$ of the irradiated ¹³CF₃H to the 2₁ level. Absorption of the dissociation laser beam by the pre-excited molecules is measured as the difference between the detector signals with and without the pre-excitation pulse. The detected induced absorption is $\sim 2\%$, which corresponds to a characteristic absorption length of 25 m. Taking into account the concentration of pre-excited ¹³CF₃H in the irradiated volume, we estimate the average number of CO₂ laser photons absorbed by each pre-excited molecule to be 36 ± 6 .

Discussion

4

The experimental results reported here and in the previous paper [12] enable us to optimize the configuration of our OP-IRMPD scheme for highly selective separation of carbon-13 using CF_3H . We evaluate below the maximum possible productivity and the minimum necessary energy expenditure per separated atom for an ideal separation unit based on this configuration.

The optimal configuration of the separation scheme depends largely on the required level of isotopic enrichment. Although we have demonstrated carbon-13 enrichment levels as high as 99% [11], it seems that the most appropriate use of this method would be for enrichment levels of 94%–98%. To maximize the per volume productivity at these enrichment levels, one should use the 2_1 level for pre-excitation. A short (few tens of ns), single-mode dissociation laser pulse should overlap the pre-excitation pulse in time. As we observed earlier [12], such temporal arrangement significantly increases the dissociation yield at high sample pressure. A dissociation laser fluence of approximately 3 J/cm² gives a reasonably high yield, but is still lower than the damage threshold of many IR materials, enabling the use of unfocussed, collimated beams. This is essential, since it allows irradiation of large volumes over a long path length. For optimal use of laser photons, the absorption lengths for the two laser beams should be nearly equal. The required selectivity essentially determines the working pressure of the process and, therefore, the absorption length of the pre-excitation beam. However, the absorption length of the dissociation beam is also determined by the fraction of pre-excited molecules – that is, by fluence of the pre-excitation laser. This puts some constraints on the latter.

Let us consider a particular example. Suppose the targeted level of enrichment is 95.4%, which corresponds to selectivity of 2000. Figure 6 suggests that with a smooth dissociation pulse of 3 J/cm²; this selectivity requires about 100 mbar of CF₃H. The characteristic absorption length for pre-excitation beam L_p is given as

$$L_{\rm p} = \frac{1}{n_{13}\sigma_{13}^{\rm p}(1+99/S_{\rm p})},\tag{1}$$

where n_{13} is concentration of ${}^{13}\text{CF}_3\text{H}$, σ_{13}^p is absorption crosssection for the *Q*-branch of the 2₁ band and $S_p = 85$ is the measured selectivity of pre-excitation. At 100 mbar and with $\sigma_{13}^p = 1.2 \times 10^{-20}$ (Table 1), (1) yields a characteristic absorption length for the pre-excitation beam of 14.3 m. Attenuation of the dissociation beam to the 1/e level by the pre-excited molecules occurs over length L_{d}^{+} , given as

$$L_{\rm d}^{\bigstar} = \frac{(\rm e-1)\,\Phi_{\rm d}\,\lambda_{\rm d}}{\langle N_{\rm d}\rangle\,n_{13}\,\sigma_{13}^{\rm p}\,\Phi_{\rm p},\lambda_{\rm p}} \tag{2}$$

Here, $\Phi_d = 3 \text{ J/cm}^2$, $\lambda_d = 9.5 \,\mu\text{m}$, Φ_p , $\lambda_p = 1.7 \,\mu\text{m}$ are the fluence and the wavelength of the dissociation and the preexcitation beams respectively, and $\langle N_d \rangle = 36$ is the average number of photons absorbed per each pre-excited molecule. We also have to take into account the absorption of this beam by the ground state molecules,

$$L_{\rm d}^0 = \frac{1}{n\sigma^{\rm d}};80{\rm m}\tag{3}$$

where L_d^0 is the characteristic absorption length, *n* is the total concentration of CF₃H and $\sigma^d = 0.47 \times 10^{-22}$ is the absorption cross-section of the ground state CF₃H at the dissociation wavenumber (1052.2 cm⁻¹) [17]. Within the typical pressure and fluence ranges used in our experiments, this absorption remains a linear function of these parameters [17].

The required balance between absorption lengths for the two laser beams implies

$$\frac{1}{L_{\rm p}} \cong \frac{1}{L_{\rm d}^0} + \frac{1}{L_{\rm d}^{\bigstar}}.\tag{4}$$

Solving (2)-(4) relative to the pre-excitation fluence yields its optimal value of 1.4 J/cm^2 . We assume that the preexcitation beam with pulse energy of 1.4 J and 1 cm² crosssection is overlapped with a similar dissociation beam of 3 J pulse energy inside a 15 m long reactor (or a shorter one with multiple passes). If the beams are slightly focused such that their respective fluence remains constant along their pathways, then the maximum irradiated volume of the reactor will be around 400 cm³. These optimal fluences result in pre-excitation of nearly 14% of irradiated CF₃H molecules, of which nearly 18% will be dissociated (Fig. 6b). At a pressure of 100 mbar, the productivity of such a separation unit per pair of laser pulses can reach 2.8×10^{17} atoms of carbon-13 in the chemical form of C₂F₄ at 95.4% isotopic purity. The main parameters of the considered separation unit are summarized in Table 2.

An important characteristic of any isotope separation process is the energy expenditure per separated atom. This value gives a lower limit for energy consumption and, thus, the minimum running cost of the process. The photon energy expenditure by the pre-excitation laser per separated carbon-13 atom, calculated as a ratio of the pulse energy of this laser to the per pulse productivity of the separation unit (Table 2) is 31 eV. Similar calculation yields 66 eV for the energy expenditure by the dissociation laser. The minimum energy expenditure using our process to achieve isotopic selectivity of 2000 is, thus, 97 eV per carbon-13 atom. This can be compared to the energy expenditure in the 'classical' laser isotope separation of carbon-13 by IRMPD of CF₂HCl. Enrichment to a similar (96%) level of carbon-13 requires 760 eV of laser energy per carbon atom [3]. Clearly, for high levels of isotopic enrichment our laser isotope separation process is by far superior to the 'classical' IRMPD of CF₂HCl from

Parameter	Value
Pre-excitation wavenumber, cm ⁻¹	5936.6
Pre-excitation energy, J	1.4
Dissociation wavenumber, cm^{-1} (CO ₂ laser line)	1052.2 (9P)
Dissociation energy, J	3
CF ₃ H pressure, mbar	100
Length of the dissociation volume, m	15
Dissociation volume, cm ³	400
Productivity, ¹³ C atoms per pair of laser pulses	2.8×10^{17}
Energy expenditure per separated ¹³ C atom, eV	97

 TABLE 2
 The principal parameters of the separation cell

the point of view of the minimum possible photon energy expenditure.

Enrichment of carbon-13 by low-temperature distillation of CO, currently the main commercial production technology consumes up to 1500 eV/C – atom of energy in the form of liquid nitrogen [18]. Whether the technology based on our approach will consume less energy depends primarily on the efficiency of the lasers employed. The electrical efficiency of pulsed CO₂ lasers is typically 10%, bringing the energy consumption by the dissociation laser in our process to 660 eV/atom. Thus, to be competitive with the distillation process, the energy consumption by the pre-excitation laser has to be below $900 \, \text{eV}/\text{atom}$, which means that the efficiency of this laser should be above 3.5%. The development of an efficient, high repetition rate, high-energy near-IR laser system seems challenging, but currently realistic. This could be an all solid-state system based on a diodepumped Nd:YAG laser and an OPO/OPA wavelength converter. The efficiency of such laser system can approach 5%–10%. At 10%, this would imply that our laser isotope separation approach for carbon-13 would be approximately 1.5 times more efficient than low temperature distillation of CO.

5 Conclusions

We have further developed our collisionally assisted overtone pre-excitation-IRMPD approach to highly selective laser isotope separation of carbon-13 using CF₃H as a starting material. Employing the 2_1 vibrational level for pre-excitation, which has 6.7 times higher absorption crosssection than the previously used 3_1 level, allows for a significant increase of per volume absolute dissociation yield. Although the isotopic selectivity using pre-excitation of CF₃H to the former level is several times lower than that for the latter, collisional V-V relaxation quickly compensates this difference upon increase of pressure to above 50 mbar. Smoothing the spiky structure of pulses delivered by our CO₂ laser allows for the further increase of the per volume absolute dissociation yield.

We suggest an optimal configuration of the process that includes pre-excitation to the 2₁ level by a short (10–30 ns) pre-excitation pulse and a slightly longer single-mode dissociation pulse, overlapped in time. The investigated approach is most appropriate for carbon-13 enrichment to the level of 94%–98%. An example of an ideal separation unit illustrates the attainable performance of the process. The estimated productivity at an enrichment level of 95.4% is 2.8×10^{17} carbon-13 atoms per pair of laser pulses using 1.4 J/pulsefor the pre-excitation laser and 3 J/pulse for dissociation laser.

The evaluated energy expenditure, 97 eV per separated carbon-13 atom, is several times less than for the classical laser separation technique using IRMPD of CF₂HCl. The main challenge for practical implementation of our process is the development of the required pre-excitation laser system with at least 3.5% of photon/electrical power efficiency. With such a laser and when properly engineered, the process should be competitive with the current carbon-13 separation technologies.

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