Increase of Radiative Forcing through Midinfrared Absorption by Stable CO₂ Dimers?

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ABSTRACT: We performed matrix-isolation infrared (MI-IR) spectroscopy of carbon dioxide monomers, CO₂, and dimers, (CO₂)₂, trapped in neon and in air. On the basis of vibration configuration interaction (VCI) calculations accounting for mode coupling and anharmonicity, we identify additional infrared-active bands in the MI-IR spectra due to the (CO₂)₂ dimer. These bands are satellite bands next to the established CO₂ monomer bands, which appear in the infrared window of Earth’s atmosphere at around 4 and 15 μm. In a systematic carbon dioxide mixing ratio study using neon matrices, we observe a significant fraction of the dimer at mixing ratios above 300 ppm, with a steep increase up to 1000 ppm. In neon matrix, the dimer increases the IR absorbance by about 15% at 400 ppm compared to the monomer absorbance alone. This suggests a high fraction of the (CO₂)₂ dimer in our matrix experiments. In atmospheric conditions, such increased absorbance would significantly amplify radiative forcings and, thus, the greenhouse warming. To enable a comparison of our laboratory experiment with various atmospheric conditions (Earth, Mars, Venus), we compute the thermodynamics of the dimerization accordingly. The dimerization is favored at low temperatures and/or high carbon dioxide partial pressures. Thus, we argue that matrix isolation does not trap the gas composition “as is”. Instead, the gas is precooled to 40 K, where CO₂ dimerizes before being trapped in the matrix, already at very low carbon dioxide partial pressures. In the context of planetary atmospheres, our results improve understanding of the greenhouse effect for planets of rather thick CO₂ atmospheres such as Venus, where a significant fraction of the (CO₂)₂ dimer can be expected. There, the necessity of including the mid-IR absorption by stable (CO₂)₂ dimers in databases used for modeling radiative forcing, such as HITRAN, arises.

INTRODUCTION

Carbon dioxide is the preeminent anthropogenic greenhouse gas on Earth. In thermal equilibrium with the sun, the Earth’s surface temperature should be 255 K according to Stefan–Boltzmann’s law. The actual surface temperature, however, is 288 K, implying a warming of 33 K caused by its atmospheric greenhouse gases. While CO₂ is a trace gas on Earth, Venus has a much denser atmosphere composed of mainly 96.5% CO₂ and 92 bar of surface pressure. Hence, the greenhouse effect on Venus is about 15 times stronger than that on Earth. The Earth’s atmospheric CO₂ mixing ratio increased from 270 ± 10 ppm in 1750 to about 417 ppm in May 2020, where annual increases have reached 3 ppm in the past decade. Consequently, the temperature has risen by +2.50 ± 0.14 °C for Northern hemisphere landmasses comparing 2020 with 1884. By year 2100, CO₂ mixing ratios might climb to 830 ppm according to the SSP3 scenario that predicts an increase of global mean surface temperature by 4.1 °C, far above the Paris Agreement aims of 1.5 °C. This scenario seems to be a path closely resembling that of the global development in the past few years. A skyrocketing atmospheric CO₂ mixing ratio favors CO₂ dimerization and raises the question of what the impact of (CO₂)₂ on greenhouse warming might be. Up to now, no carbon dioxide dimers have been detected in Earth’s atmosphere. However, once dimers appear in significant amounts in planetary atmospheres, we expect increasing
radiative forcing due to additional infrared (IR) absorption and, thus, global warming beyond the greenhouse effect of the monomers. If there were carbon dioxide dimers in the troposphere, revision of contemporary climate models could be necessary because of potentially unaccounted \((\text{CO}_2)_2\) infrared absorptions. A step in this direction has been done by considering collision-induced absorption (CIA) due to unstable \((\text{CO}_2)_2\) dimers for dense \text{CO}_2 atmospheres.\(^8\) CIA has been included in the HITRAN database for the \((\text{CO}_2)_2\) dimers, together with a variety of other dimers and other collision complexes.\(^7\) However, dimerization may also result in stable dimers (or tightly bound dimers), which have been recently considered in broad-band spectra.\(^10\) In the mid-IR, such stable dimers would lead to permanent additional absorptions in the vicinity of the monomer’s IR absorption. Those additional IR bands are usually hard to distinguish from the rotational–vibrational spectrum of the pure monomer.

Thus, we investigate the IR spectrum of carbon dioxide using matrix isolation to quench rotational transitions and use anharmonic calculations to assign all observed vibrational transitions. Previously, we successfully assigned all \text{CO}_2 monomer bands,\(^11\) where we also provided a short review of the theoretical and experimental spectroscopy of carbon dioxide and its dimer. For a detailed review on computational studies on the \((\text{CO}_2)_2\) dimer and also recent calculations we refer to Maistroskyy et al.,\(^12\) who demonstrated that variational calculations using a tailor-made potential energy surface for the dimer on explicitly correlated coupled cluster theory yield excellent results. In the present work, we investigate the infrared absorption of the \((\text{CO}_2)_2\) dimer in the mid-IR and discuss its importance to different planetary atmospheres, specifically Mars, Venus, and Earth in comparison. On the basis of two matrix-isolation experiments, we demonstrate the characteristics of the mid-IR spectrum of the \((\text{CO}_2)_2\) dimer: (a) a mixing ratio series study of carbon dioxide isolated in neon matrix and (b) direct matrix isolation of laboratory air (containing about 417 ppm carbon dioxide). In addition to the experimental data, we provide \textit{ab initio} calculations of the vibrational spectrum of the monomer and dimer, relying on multimode potential energy surfaces\(^13\) and vibrational self-consistent field (VSCF) and configuration interaction (VCI) computations.\(^14,15\) This approach allows us to unequivocally identify dimer absorptions by minimizing the discrepancy between experiment and calculation based on the inclusion of mode coupling and anharmonicity.\(^16\) Finally, we calculate the equilibrium constant for the monomer–dimer equilibrium to estimate the fraction of dimers for different \text{CO}_2 partial pressures and temperatures.

### EXPERIMENTAL METHODOLOGY AND COMPUTATIONAL DETAILS

**Matrix-Isolation Infrared Spectroscopy of Carbon Dioxide in Neon.** We performed infrared spectroscopy measurements of mixtures of gases in frozen neon matrices, i.e., matrix-isolation infrared (MI-IR) spectroscopy. Former studies\(^17\) have shown that the composition of an atmosphere trapped in a frozen matrix resembles the gas phase composition. Thus, MI-IR spectroscopy is a suitable analytical technique for investigating short-lived and unstable species from the gas phase. As matrix isolation inhibits molecular rotation, i.e., quenches the rotational–vibrational transitions, the complexity of the MI-IR spectra is reduced compared to a gas-phase IR spectrum. Consequently, observation of pure vibrational features in MI-IR spectra facilitates the distinction of different molecules, conformers, and clusters as well as dimers and oligomers. For studying the carbon dioxide dimerization with increasing carbon dioxide mixing ratio, we employed MI-IR experiments using neon as a host. Compared to other rare gases as a matrix material, neon minimizes matrix effects such as matrix shifts or multiple trapping sites.\(^11\) Hence, MI-IR spectroscopy using neon as a host provides particularly clean spectra and facilitates studying carbon dioxide dimerization.

We used a \(^{12}\text{C}^{16}\text{O}_2\) sample of high purity (99.9995\%, Messer Austria, order no. 1290102114, lot 27531923) to deposit \text{CO}_2/\text{Ne} mixtures as a frozen, immobile solid on a cold mirror. The neon matrix containing carbon dioxide is then probed in the mid-infrared using a Vertex 80v Bruker spectrometer. The mixing ratio of carbon dioxide in neon, from now on denoted as \(\rho\), was adjusted and quantified by barometric monitoring. This setup is detailed elsewhere.\(^11\) In our main experiments, matrix isolation and IR spectroscopy were performed at 6 K. We accumulated the IR spectra (resolution, 0.3 cm\(^{-1}\); scans, 512) within roughly 30 min. To study the influence of initial gas temperature during mixing on the dimerization, the \text{CO}_2/\text{Ne} mixtures were equilibrated at either 25 °C (cf. the spectra in Figure S4) or 65 °C (cf. spectra in Figure S5). The relative band areas for the monomer and dimer at these different mixing temperatures (cf. Figure S6, parts b and c) show no significant difference.

As neon has a comparably high vapor pressure in comparison to, e.g., argon, we probed the thermal stability of our neon matrices. All gas mixtures were deposited at 6 K, and then slowly heated in steps of 0.5 K up to 12 K. In each step, we accumulated “quick” spectra (resolution, 0.3 cm\(^{-1}\); scans, 32) within about 1 min. Neon evaporation from the matrix commences at roughly 9 K. Afterward we further increased the temperature beyond 12 K by turning off the helium cryostat and keeping the heater on. In roughly 2 min steps, we accumulated spectra up to a temperature of about 100 K. Figure S7 shows for \(\rho = 500\) ppm that solid carbon dioxide residues remain after neon evaporation. We observed very similar behavior for the other carbon dioxide mixing ratios in our study.

**MI-IR Spectroscopy of Isolated Air (Carbon Dioxide in Nitrogen–Oxygen–Argon).** When isolating air at 12 K, all trace gases in the air, including carbon dioxide, are trapped in a solid matrix of \text{N}_2/\text{O}_2/\text{Ar}, with their volume mixing ratio of 78:21:1 in the atmosphere. For this experiment, we used a different high-vacuum setup, also employing a helium cryostat. The cryostat was used to cool a cesium iodide window to 12 K. By introducing air into the chamber through a needle valve, we could deposit air onto this window slowly. During the deposition, the window was monitored \textit{in situ}, where the entire vacuum chamber is placed inside the sample chamber of an IR spectrometer (Varian Excalibur 4500). The sample chamber of the spectrometer was constantly flooded with nitrogen to remove IR-active trace gases from the air, through which the beam passes. Over 800 spectra, with a resolution of 0.25 cm\(^{-1}\), were collected and combined to a single measurement. We isolated air on November 15, 2020 at Innsbruck 52c, Innsbruck, Austria. The deposition times were long enough to avoid significant heating of the air matrix.

**Monomer–Dimer Thermodynamic Equilibrium.** The free energy of the dimer dissociation \((\text{CO}_2)_2 \leftrightarrow 2\text{CO}_2\) is computed as \(\Delta G (\text{kJ mol}^{-1}) = 2G_{\text{f}}^{\infty} - G_{\text{f}}^{\infty}(\text{CO}_2)_2\) where \(G_{\text{f}}^{\infty}\)
and $G_{\text{CO}_2}$ are the free energy of formation of the dimer and the monomer. First, we calculated the electronic energies and the harmonic frequencies of the monomer and dimer at the CCSD(T)-F12 level of theory with the VTZ-F12 basis set using the MOLPRO software package.20 Second, we estimated the free energies of formation within the rigid rotor harmonic oscillator (RRHO) approach using the KiSTheLP program.21 This approach computes thermodynamic properties from partition functions based on previously calculated electronic energies, harmonic frequencies, rotational constants, as well as the molecular symmetry and mass. Note that calculated thermodynamics and the equilibrium constants for temperatures close to 0 K may potentially be inaccurate as the approximations break down. However, for the temperature range we are interested in (above 30 K) the RRHO approximations can be expected to be valid.

**In Vacuo VSCF/VCI Calculations of Carbon Dioxide and Its Dimer.** Our present band assignment to monomer or dimer vibrational modes relies on VSCF and VCI computations using the MOLPRO software package.20 Starting from the global equilibrium structure of the carbon dioxide dimer, which has $C_{2v}$ symmetry,22 we performed a harmonic frequency calculation using the distinguishable cluster approach with a double-$\xi$ basis (DCSD-F12a/VDZ-F12)23,24 to obtain the normal modes of the dimer. The multimode potential energy surface (PES) is expanded up to 3D terms, i.e., three mode couplings, using the XSURF algorithm,25 as implemented in MOLPRO 2020. Hereby, intermolecular as well as intramolecular modes are coupled. Within this approach we used DCSD-F12a/cc-pVDZ-F12 single points to construct the grid representation of the PES. To estimate whether addition of diffuse functions improves the description of the CO$_2$ dimer, we also computed a PES relying on DCSD-F12a/aug-cc-pVDZ single points (cf. Table S2). However, the results were very similar and will not be considered in detail here. The grid representation of the PES is transformed into an analytic representation using polynomials.26 On the basis of this polynomial representation, anharmonic vibrational frequencies are computed using the vibrational self-consistent field and configuration interaction (VSCF/VCI) approach27 as implemented in MOLPRO. In the VCI approach, single to quadruple (SDTQ) excitations are considered. Vibrational intensities are computed within the same approach using a multimode dipole moment surface (DMS) at the HF/VTZ-F12 level of theory. For the carbon dioxide monomer in $D_\infty h$ symmetry, we followed the same procedure using up to three mode couplings. Our previous studies on the CO$_2$ monomer using the same computational approach show excellent agreement between VCI computations and MI-IR spectra.11

The (CO$_2$)$_2$ dimer can be distinguished from the monomer by consideration of its structure and symmetry. The linear CO$_2$ monomer (Figure 1a) comprises four normal modes ($3N-5$...
with \( N = 3 \) and is in the \( D_{2h} \) point group. Only three modes are in the IR-active irreducible representation \( A_{1u} \) or \( E_{1u} \), namely, the antisymmetric stretch \( \nu_3(A_{1u}) \) and the bending \( \nu_2(E_{1u}) \). The bending occurs as two degenerate normal modes; hence, only two distinguishable IR-active vibrations are considered in total. Considering the structure of the \((\text{CO}_2)_2\) dimer, Kalugina et al.\(^{22}\) suggested the slipped-parallel 60–60–0 conformer as the global minimum. Other conformers are transition states (T-shaped and crossed dimer) or higher in energy (linear dimer). For the assignment of our MI-IR spectra, the consideration of the mentioned slipped-parallel dimer is sufficient; hence, we omit any anharmonic calculations of the other conformers.

The slipped-parallel \((\text{CO}_2)_2\) dimer (Figure 1b) is in the \( C_{2h} \) point group and has 12 vibrational modes (\( 3N - 6 \) with \( N = 6 \)). Six out of 12 modes are in the IR-active irreducible representation \( A_a \) or \( B_a \), namely, the in-phase antisymmetric stretch \( \nu_6(B_a) \), the in-phase symmetric stretch \( \nu_{10}(B_a) \), the in-phase out-of-plane bending \( \nu_{7}(A_a) \), the in-phase in-plane bending \( \nu_{11}(B_a) \), and two intermolecular modes \( \nu_8(A_a) \) and \( \nu_{12}(B_a) \). While the intermolecular modes are outside the atmospheric IR window in the far-infrared region, the remaining four are inside the atmospheric window. The \( \nu_{10}(B_a) \) vibration in the atmospheric window is too weak to be detected in the present work. That said, an IR spectrum containing \((\text{CO}_2)_2\) dimers comprises three additional mid-IR bands compared to that of a pure monomer spectrum. In the following experimental spectra, we make use of these differences to elaborate on the impact of the \((\text{CO}_2)_2\) dimer spectral features relative to the \(\text{CO}_2\) monomer features.

### RESULTS AND DISCUSSION

Figure 2 depicts the calculated temperature dependence of the equilibrium constant \( K_{eq} \) for the \((\text{CO}_2)_2\) dimer dissociation

![Figure 2](image-url)

**Figure 2.** Temperature-dependent equilibrium constant \( K_{eq} \) for \((\text{CO}_2)_2 \rightarrow 2\text{CO}_2\) between 30 and 1000 K calculated within the rigid rotor harmonic oscillator (RRHO) approximation in the KISTheIP program by Canneaux et al. (ref 21) and relying on the electronic energy and the harmonic frequencies at the CCSD(T)-F12/VTZ-F12 level of theory.

reaction. For temperatures above 50 K, the monomer is thermodynamically favored (cf. thermochemistry data in Figure S1), and the equilibrium constant \( K_{eq} \) is on the \(\text{CO}_2\) monomer side. This is the case for various planetary atmospheres. While the equilibrium constant \( K_{eq} \) increases steeply for temperatures above 300 K (like Earth’s atmosphere), it reaches a maximum at around 700 K (like Venus’s atmosphere). The dimer dissociation, however, depends also on the partial pressure of carbon dioxide \( p(\text{CO}_2) \) in the respective environment (cf. the pressure dependence of the degree of dissociation in Figure S2). On Earth and Mars, the partial pressure of carbon dioxide is very low \((p(\text{CO}_2) \ll 1 \text{ bar})\), leading to high degrees of dissociation. On Venus, however, the high partial pressure \((p(\text{CO}_2) \approx 90 \text{ bar})\) leads to a lower degree of dissociation \( \alpha \), although the high temperature favors dissociation.

For Venus, the very dense atmosphere composed of 96.5% carbon dioxide results in a significant fraction of dimers. According to our thermodynamic estimations, the ratio of monomers to dimers is approximately 12:1, resulting in a partial pressure of \((\text{CO}_2)_2\) dimers of 6.98 bar on Venus (7.6 vol %). Mars has an atmosphere that is composed of 95.1% carbon dioxide but is about 15 000 times thinner than that of Venus (6.36 mbar vs 92 bar). Consequently, the dimer is only a trace species on Mars, despite the much colder climate on Mars (210 vs 737 K). Hence, the monomer-to-dimer ratio is about \( 5.4 \times 10^4 \) and the \((\text{CO}_2)_2\) partial pressure is only 0.111 mbar (17 ppmv). By contrast to Mars and Venus, \(\text{CO}_2\) is only a trace species in our atmosphere (410 ppmv in a 1 bar atmosphere). This leads to a very high monomer-to-dimer ratio of \( 1.3 \times 10^4 \) and a \((\text{CO}_2)_2\) partial pressure of only 0.317 mbar (312 pptv) in the boundary layer above Earth’s surface at 288 K.

If \((\text{CO}_2)_2\) exists in the troposphere, then it is merely at a mixing ratio on the order of parts per trillion (ppt). At higher altitude, e.g., in the tropopause at ~20 km or the mesopause at ~80 km, temperatures are much lower, namely, about 200 and 150 K, respectively. Partial pressures are also lower, namely, by a factor of 10^2 and 10^3 at such altitudes. While the lower temperatures favor dimerization, the lower partial pressure disfavoring dimers is much stronger. This leads to much smaller dimer fractions in the higher layers of our atmosphere. Consequently, it is a huge challenge in terms of detection limit and sensitivity to spectroscopically observe \((\text{CO}_2)_2\) dimers on Mars and Earth. For comparison, our estimated mixing ratio of \((\text{CO}_2)_2\) of 312 pptv on Earth is on the same scale as that of the OH radical\(^2\) or prominent ozone-depleting halocarbons, currently in between ~80 pptv (CCl\(_3\)) and ~490 pptv (CFC-12) according to measurements at Manua Loa, Hawaii within the NOAA/GML in situ halocarbons program.

In both our neon matrix and the air-freezing experiments detailed below, we in fact do observe a considerable amount of \((\text{CO}_2)_2\) dimers. However, as we will also show below, the monomer/dimer ratio in our matrix experiments does not represent the ratio at 25 °C (or 65 °C). Rather than that, the gas precools to 40 K prior to being trapped in the matrix. Cooling of the gas from room temperature (or above) to 40 K shifts the equilibrium massively to the side of the dimers. That is, in our matrix experiment we trap an atmosphere enriched in dimers, where the monomer/dimer ratio is about 8:1 (see Table 1).

Let us now turn to our experimental observations obtained for \(\text{CO}_2/\text{Ne}\) mixtures after deposition as matrices at 6 K. Figure 3a shows the IR spectra recorded in transmission—reflection for carbon dioxide mixing ratios \( p = 100–700 \text{ ppm} \). Other spectra for \( p = 700–4000 \text{ ppm} \) are shown in Figures S4 and S5. At \( p = 100 \text{ ppm} \), the well-known \(\text{CO}_2\) monomer bands dominate the spectra. In the antisymmetric stretch region (Figure 3c), we observe the \(\text{CO}_2\) \(\nu_3(A_{1u})\) transition at 2348.3 cm\(^{-1}\). In the bending region (Figure 3d), we observe the \(\text{CO}_2\) \(\nu_2(E_{1u})\) transition as a doublet at 668.5 and 667.9 cm\(^{-1}\). Note that doublet splitting was observed in earlier studies for both the antisymmetric stretch and the bending region for carbon...
Table 1. Estimation of the $(CO_2)_2$ Partial Pressure and Monomer/Dimer Ratio*

|          | temp (K) | $K_{eq}$ | $p_{CO_2}$ | $p((CO_2)_2)$ | ratio (monomer/dimer) |
|----------|----------|----------|------------|---------------|-----------------------|
| Mars     | 210      | 325      | $6.00 \times 10^{-3}$ bar ($\sim 95.1$ vol %) | $0.111 \times 10^{-6}$ bar ($\sim 95.1$ ppmv) | 54 157:1 |
| Earth    | 288      | 572      | $0.426 \times 10^{-3}$ bar ($\sim 426$ ppmv)  | $0.317 \times 10^{-9}$ bar ($\sim 312$ pptv) | 1 344 231:1 |
| Venus    | 737      | 958      | $88.8$ bar ($\sim 96.5$ vol %) | $6.98$ bar ($\sim 7.6$ vol %) | 12:1 |
| “matrix” | 40       | 0.00273  | $0.400 \times 10^{-3}$ bar | $0.460 \times 10^{-6}$ bar ($\sim 46$ ppmv) | 8:1 |

*Equilibrium constants $K_{eq}$ for a given temperature are taken from the calculation shown in Figure 2. Calculation of the dimer fraction $p((CO_2)_2)$ and monomer/dimer ratio is detailed in the Supporting Information. The characteristic temperatures and carbon dioxide partial pressures $p_{CO_2}$ of Mars, Earth, and Venus are taken from ref 29.

Figure 3. Matrix isolation of $(CO_2)_2$ dimers from gas-phase CO$_2$/Ne mixtures at 6 K. (a) The mid-IR spectrum between 4000 cm$^{-1}$ (2.5 $\mu$m) and 500 cm$^{-1}$ (20 $\mu$m) exhibits three major absorption regions: (b) the Fermi resonance overtone region, which we do not consider in detail, (c) the antisymmetric stretch region, containing the $(CO_2)_2$ $\nu_9(B_u)$ transition, and (d) the bending region, containing the $(CO_2)_2$ $\nu_8(A_u)$ and $\nu_{11}(B_u)$ transitions. The assignment relies on frequencies calculated in vacuo, shown as colored lines. To better compare with the experiment, we scale all calculated frequencies by a factor of 0.9989 in panel c and 0.9924 in panel d and the calculated intensities by 0.2397.

Table 2. Monomer–Dimer Shift of the Stretching and Bending Regions*

|          | ref | $(CO_2)_2$ $\nu_9(B_u)$ | shift $\rightarrow$ | $(CO_2)_2$ $\nu_{11}(A_u)$ | shift $\rightarrow$ | $(CO_2)_2$ $\nu_8(B_u)$ | shift $\rightarrow$ |
|----------|-----|--------------------------|---------------------|--------------------------|---------------------|--------------------------|---------------------|
| VCI      | this study | 2353.2 | 2.2 | 2351.0 | 677.0 | 3.9 | 673.1 | $-3.9$ | 669.2 |
| air $\text{(N}_2/\text{Ar/O}_2)$ | this study | 2350.9 | 2.2 | 2348.7 | 663.5 | 1.2 | 662.3 | |
| neon     | this study | 2350.2 | 1.9 | 2348.3 | 670.3 | 1.8 | 668.5 | $-3.9$ | 664.6 |
| nitrogen $\text{N}_2$ | 30 | 2345.8 | 1.3 | 2344.5 | 664.1 | 0.7 | 663.4 | $-3.5$ | 659.9 |
| argon    | 30 | 2346.5 | 2.0 | 2344.5 | 664.2 | 0.8 | 663.4 | $-3.8$ | 659.6 |
| 31 | 2346.7 | 2.7 | 2345.0 | 664.4 | 0.7 | 663.7 | $-3.8$ | 659.9 |
| 36 | 2342.5 | 2.0 | 2340.5 | 662.2 | 1.0 | 661.2 | $-2.6$ | 658.6 |
| krypton | 31 | 2336.0 | 1.5 | 2334.5 | 661.2 | 1.2 | 660.0 | $-1.5$ | 658.5 |
| xenon   | 31 | 2344.0 | 663.3 | 1.2 | 665.1 | $-2.1$ | 663.0 |
| deuterium $\text{D}_2$ | 33 | 2334.5 | 663.5 | 1.2 | 665.0 | $-2.1$ | 663.0 |

*Monomer–dimer shifts derived from absolute (unscaled) frequencies as assigned to the CO$_2$ monomer and $(CO_2)_2$ dimer in various matrix-isolation experiments from literature and in the matrix-isolation experiments and VCI computations from this study. All values are in wavenumbers (cm$^{-1}$).
dioxide in argon and krypton matrixes but not in xenon, nitrogen, and deuterium matrixes.\textsuperscript{30–36} These doublets were identified as a matrix effect, reflecting that the monomer, of linear geometry, was in two different surroundings referred to as matrix cages. While we reproduced this splitting in argon matrixes, in neon matrixes we observe only a splitting for the CO\textsubscript{2} \nu\textsubscript{2}(E\textsubscript{1u}) transition.\textsuperscript{11} It has been shown that the matrix environment makes the two degenerate bending vibrations distinguishable, leading to the observed splitting of the band that corresponds to the bending vibration.

For \( \rho \geq 200 \) ppm, satellite bands appear, shifted from the assigned monomer bands by just a few wavenumbers. Specifically, two bands in the antisymmetric stretch region are shifted by +1.9 and −1.8 cm\(^{-1}\) compared to the CO\textsubscript{2} \nu\textsubscript{3}(A\textsubscript{1u}) transition (Figure 3c), and two bands in the bend region are shifted by +1.8/+2.4 and −3.3/−3.9 cm\(^{-1}\) compared to the CO\textsubscript{2} \nu\textsubscript{2}(E\textsubscript{1u}) transition (Figure 3d). As the shift between the satellite bands and the established monomer bands agrees with our VCI-calculated monomer–dimer shifts, we assign the satellite bands to the (CO\textsubscript{2})\textsubscript{2} dimer. Our VCI calculations of the slipped-parallel dimer in the C\textsubscript{2h} point group reveal monomer–dimer shifts that are very similar to the experiment, namely, +2.2 cm\(^{-1}\) for the antisymmetric stretch and +3.9 and −3.9 cm\(^{-1}\) for the bend (cf. the line spectra in Figure 3, parts c and d). On the basis of these calculations, we assign the \text{(CO\textsubscript{2})\textsubscript{2}} as follows: the band at 2350.2 cm\(^{-1}\) to the \text{(CO\textsubscript{2})\textsubscript{2}} \nu\textsubscript{3}(A\textsubscript{1u}) and \nu\textsubscript{11}(B\textsubscript{2u}) transitions, respectively.

The present VCI calculations systematically overestimate the experimental frequencies by approximately 5 cm\(^{-1}\). In parts c and d of Figure 3, we scale the VCI frequencies of the monomer in each the antisymmetric region by a factor of 0.9989 and the bending region by 0.9924 to match our neon monomer in each the antisymmetric region by a factor of and d of Figure 3, we scale the VCI frequencies of the band. For this reason, calculated and observed shifts agree best with gas-phase spectra, i.e., the matrix effect in neon is the smallest one among all matrixes,\textsuperscript{4} which is why we decided to study the CO\textsubscript{2} dimerization in neon matrixes.

The present VCI calculations allow for a correct assignment as they generally reproduce the positive monomer–dimer shift in the antisymmetric stretch region (Figure 3e) and one positive and one negative monomer–dimer shift in the bending region (Figure 3d). In the latter region, the calculated monomer–dimer shift from CO\textsubscript{2} \nu\textsubscript{2}(E\textsubscript{1u}) to \nu\textsubscript{11}(B\textsubscript{2u}) agrees with the experiment, whereas the shift from CO\textsubscript{2} \nu\textsubscript{2}(E\textsubscript{1u}) to \nu\textsubscript{1}(A\textsubscript{1u}) is slightly overestimated. Further improvement in our VCI calculations may be achieved by an extension of the multimode potential energy surface to couple more than three modes. However, we do not expect such calculations to improve the assignment; hence, we consider them as beyond the scope of the present work. Note that VSCF and VCI calculations including up to three mode couplings have been previously performed by Maystrovsky et al.\textsuperscript{12} Considering the mbCO2/grid/VCI calculations in ref 12, the monomer–dimer shift from CO\textsubscript{2} \nu\textsubscript{1}(E\textsubscript{1u}) to \nu\textsubscript{11}(B\textsubscript{2u}) would be −4.5 cm\(^{-1}\), the shift from CO\textsubscript{2} \nu\textsubscript{1}(E\textsubscript{1u}) to \nu\textsubscript{11}(B\textsubscript{2u}) would be +13.7 cm\(^{-1}\), and the shift from CO\textsubscript{2} \nu\textsubscript{1}(E\textsubscript{1u}) to \nu\textsubscript{11}(B\textsubscript{2u}) would be +4.7 cm\(^{-1}\). These shifts do not agree with our or any previous MI-IR experiments,\textsuperscript{30–36} which is why we rely on our present VCI calculations for the final assignment. The discrepancy of the monomer–dimer shifts from ref 12 is most likely rooted in the VCI results of the dimer, where the authors correctly stress that the VCI eigenvectors are very sensitive to small changes in the computation\textsuperscript{12} and emphasize the value of additional investigations. The assigned monomer–dimer shifts observed from our and previous MI-IR experiments may be a reference for future VCI investigations.

In addition to the band positions, the band intensities are also of interest. In the region of 1400–1200 cm\(^{-1}\), we do not observe the (CO\textsubscript{2})\textsubscript{2} \nu\textsubscript{10}(B\textsubscript{2u}) transition. This agrees with the calculated IR intensities, where the intensity of \nu\textsubscript{10}(B\textsubscript{2u}) is 3–4 orders of magnitude weaker than the intensities of the other transitions, so we are not sensitive enough in our spectroscopy experiment to detect this very weak absorption. The observed intensities allow us to give a first estimate of the monomer/dimer ratio in our experiment. To match the monomer-to-dimer intensity ratio as observed for \( \rho = 700 \) ppm (black trace in Figure 3, parts c and d), the VCI-calculated intensities of the dimer bands in Figure 3, parts c and d, need to be scaled by 0.2397. In other words, for \( \rho = 700 \) ppm, the monomer is the dominant species with a ratio of 4:1 compared to the dimer. A better quantification of the monomer/dimer ratio from the
experiment is possible by investigation of the band areas, as explained below. On the basis of our assignment, we proceed to the central task of the present study, namely, quantifying the additional IR absorption of the \((\text{CO}_2)_2\) dimers from the experimental spectrum itself.

On the basis of the assigned dimer and monomer bands, we retrieve the band areas by numeric integration, as described in the Supporting Information. Figure S4 details this numerical integration of the monomer and dimer bands. Figure S6 shows the absolute areas of monomer and dimer bands individually in Figure S6a, with relative areas given in Figure S6, parts b and c. This quantification enables us to (1) estimate the monomer/dimer ratio in the matrix experiment and (2) quantify the increase in IR absorption (and thus radiative forcing) by the occurrence of dimers. This estimate of the dimer abundance based on band areas is more rigorous and accurate than the rough estimate based on band intensities detailed above.

We may compare the monomer/dimer ratio from our experiment with the estimation we gave in Table 1 from our thermochemical calculation. For \(\rho = 400\) ppm in our neon matrix experiment, we obtain a monomer-per-dimer ratio of 0.83/0.15 = 5.5 in the asymmetric stretch region or 0.87/0.13 = 6.7 in the bending region (cf. the relative dimer band areas in Figure S6b). For \(\rho = 400\) ppm, our thermochemical calculation predicts a monomer/dimer ratio of 8 at a temperature of 40 K. This implies that the composition in the neon matrix does not correspond to the gas-phase composition at room temperature, but to the gas-phase composition at roughly 40 K. While ideally we would like to trap the ambient-temperature gas-phase composition “as is”, this is apparently not the case in the matrix-isolation studies. We explain this observation by gas-phase molecules bouncing back from the matrix instead of being trapped right away. The gas-phase molecules are precooled during this hitting of the matrix and bounce back before eventually being trapped, apparently to 40 K in our experimental setup.

The analysis of band areas also allows us to quantify the impact of dimers on radiative forcing and greenhouse warming. Assuming an equal absorption coefficient for the monomer and dimer, we can provide an estimate of how much the radiative forcing of carbon dioxide is increased by the occurrence of dimers. The (relative) dimer contribution to radiative forcing is then calculated from the increase of the total band area due to dimerization \(\Omega = \frac{a(\text{dim})}{a(\text{mono})} \times 100\%\), where \(a(\text{dim})\) is the area under the dimer bands and \(a(\text{mono})\) is the area under the monomer band. Figure 4 shows \(\Omega\) in red for the antisymmetric stretch and blue for the bending bands. For example, at \(\rho = 400\) ppm \(\Omega\) is 18% in the asymmetric stretch region. In general, both spectral regions provide two central observations: (1) \(\Omega\) increases with increasing carbon dioxide mixing ratio \(\rho\). (2) \(\Omega\) increases steeply at lower \(\rho\) and less steeply at higher \(\rho\) (beyond 1000 ppm). In particular, \(\Omega\) increases steeply from approximately 3–6% at \(\rho = 100\) ppm to 15–20% at \(\rho = 400\) ppm, and further to 22–26% at \(\rho = 700\) ppm. Beyond \(\rho = 1000\) ppm, \(\Omega\) increases flatly from 25% to almost 50% at 4000 ppm.

Our observation of significant absorption due to dimers in neon matrices even at \(\rho = 400\) ppm prompted us to study the MI-IR spectrum of ambient air. This was done by very slow deposition of air onto a cesium iodide window at 15 K, where the lab was well-vented with outdoor air right before the deposition. In this experiment the matrix is composed of air \((\text{N}_2/\text{O}_2/\text{Ar})\) itself, as opposed to neon in the mixing ratio series described above. Figure 5 depicts a deposition-time sequence of spectra in the antisymmetric stretch and bend regions. Figure S8 covers additional spectral regions, where we observe, besides \(\text{CO}_2\) bands arising from \(\text{H}_2\text{O}\), \(\text{NO}\), and \(\text{NO}_2\) (most likely from pollution) as well as clusters, e.g., \((\text{H}_2\text{O})_2\) or \(\text{CO}_2-\text{N}_2\), in the matrix. As shown in Table S3, the deviation of the band positions of these species from literature data measured in a \(\text{N}_2\) matrix is less than 1 cm\(^{-1}\).

Unlike the neon matrix spectra in Figure 3a–d, the mixing ratio of carbon dioxide in ambient air is constant (\(\rho = 417\) ppm). Hence, we do not perform a mixing ratio series here, yet we increase the thickness of the air matrix with deposition time in Figure 5 to improve the signal-to-noise ratio. The interaction between \(\text{CO}_2\) and the matrix material \(\text{air (N}_2/\text{O}_2/\text{Ar})\) is stronger than the interaction with the matrix material neon. This leads to a significant broadening of bands, a change of band intensities, and a more pronounced matrix shift. Despite the broadening, we see direct evidence for the presence of \((\text{CO}_2)_2\) in isolated air, namely, shoulders at 2350.9 and 663.5 cm\(^{-1}\). The shoulder assignment relies on the assignment of the monomer and dimer shown in Figure 3, parts c and d, as well as earlier literature on matrix isolation of carbon dioxide in nitrogen matrices in ref 30 and in ref 36. Similar to the neon matrix spectra, the dimer bands are shifted by +2.2 and +1.2 cm\(^{-1}\) compared to the monomer bands. However, we do not observe the downshifted bending band as it is of very low intensity in \(\text{N}_2\) matrices (and hence also in air matrices). The downshifted band in the stretch region is missing, too. Thus, we assume that trimers and higher oligomers of \(\text{CO}_2\) are not present in the air matrix, as opposed to the neon matrix.

We have decomposed the bands shown in Figure 5 using three Gaussian functions to obtain more information on the monomer/dimer ratio in this matrix experiment. Figure S9 depicts this decomposition, where the black line represents the
measured band, the red line the sum of three Gaussian functions, and the green lines the individual Gaussian peaks. To correctly reproduce the bending region, we used two Gaussian functions for the dimer and one for the monomer. For the stretch region, we need two Gaussian functions for the monomer, which is probably a consequence of the cage geometry: a cage splitting occurs for the monomer in the stretch region and a splitting for the dimer in the bending region. The splitting amounts to 1.3 and 0.4 cm⁻¹, with very similar band areas and intensities for the Gaussian functions, indicating the cage splitting. In this analysis, each dimer band area is about 6–10 times smaller than the monomer band areas. We estimate that the IR absorption increases by about 13 ± 4% due to the dimers, which compares very well with the estimation from our neon experiments (see Figure 4). Also in this matrix experiment, the dimers are very likely present because the air is precooled before being trapped at the spectroscopy window. Upon precooking, a large fraction of monomers associate to produce dimers. The presence of such large fractions of carbon dioxide dimers in the air at ambient temperature is ruled out based on thermodynamics calculations.

CONCLUSION

We here provide a systematic matrix-isolation study of carbon dioxide with increasing mixing ratios up to 4000 ppm. High-level VCI calculations that account for mode coupling and anharmonicity allow us to clearly identify monomer and dimer bands. As expected, while we barely observe any dimers at mixing ratios of 100 ppm, we observe a strong increase of the dimer fraction at higher mixing ratios. Similarly, we observe carbon dioxide dimers by trapping laboratory air (mainly N₂/O₂/Ar) containing approximately 417 ppm of CO₂. On the basis of calculations of the equilibrium constants for the carbon dioxide monomer–dimer equilibrium, we conclude that the matrix composition represents the gas phase in thermodynamic equilibrium at 40 K. Accordingly, carbon dioxide is precooled before being trapped in the neon and air matrices. Upon precooking, dimers and higher oligomers form. In the matrix experiment, the IR absorption due to carbon dioxide dimers is significant. The mid-IR absorption compared to that of monomers alone increases between 18% (at 400 ppm) and 35% (at 4000 ppm) due to dimerization. Although it is a rather weakly bound system, the (CO₂)₂ dimer can introduce additional mid-IR bands.

The fraction of carbon dioxide dimers in the current atmosphere of Earth (~288 K) is in the parts-per-trillion range (see Table 1). In the coming 20 years, the carbon dioxide mixing ratio in the atmosphere will likely be doubled in comparison to preindustrial times and reach approximately 500 ppm. Although the increasing carbon dioxide mixing ratio will lead to a higher fraction of dimers, more sensitive methods will be necessary to identify dimers in Earth’s atmosphere spectroscopically. In contrast, we roughly estimate a 12:1 monomer/dimer ratio in the Venus atmosphere at 737 K and 88.8 bar of carbon dioxide. Consequently, climate models for Venus based on monomer bands alone would severely underestimate radiative forcing. Considering temperature and pressure broadening effects as the only mechanisms for IR band broadening would be insufficient. Hence, to properly account for band broadening, the occurrence of dimer satellite bands also ought to be considered. We tentatively propose that the (CO₂)₂ dimer is an underestimated driver for the Venus greenhouse effect that has previously not been recognized. On Earth and Mars, the (CO₂)₂ dimer fraction is too small to considerably increase IR absorption and act as a driver for the greenhouse effect of these planets.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jpca.2c00857.

Additional text explaining the results of the calculations and experiments in more detail, calculated frequencies of the monomer and dimer, experimentally observed bands in solid air, thermochemistry of the dimer dissociation, pressure dependence of the dissociation and dimer fraction, MI-IR spectra and calculated spectra without...
scaling, band integration of all MI-IR spectra of the dilution series in neon matrix (25 and 65 °C, gas phase), results of the band integration (absolute and relative area under the peak), heating experiment, complete MI-IR spectrum of isolated air, band integration of the MI-IR spectrum of isolated air, and additional references (PDF)

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**Notes**

The authors declare no competing financial interest.

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