Atmospheric band fitting coefficients derived from a self-consistent rocket-borne experiment

Mykhaylo Grygalashvyly1, Martin Eberhart5, Jonas Hedin4, Boris Strelnikov1, Franz-Josef Lübken1, Markus Rapp12, Stefan Löhle5, Stefanos Fasoulas5, Mikhail Khaplanov1,†, Jörg Gumbel4, and Ekaterina Vorobeva3

1Leibniz-Institute of Atmospheric Physics at the University Rostock in Kühlungsborn, Schloss-Str. 6, 18225 Ostseebad Kühlungsborn, Germany
2Deutsches Zentrum für Luft- und Raumfahrt, Institut für Physik der Atmosphäre, Oberpfaffenhofen, Germany
3Department of Atmospheric Physics, Saint-Petersburg State University, Universitetskaya Emb. 7/9, 199034, Saint-Petersburg, Russia
4Department of Meteorology (MISU), Stockholm University, Stockholm, Sweden
5University of Stuttgart, Institute of Space Systems, Stuttgart, Germany
†deceased

Correspondence: Mykhaylo Grygalashvyly (gryga@iap-kborn.de)

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Abstract. Based on self-consistent rocket-borne measurements of temperature, the densities of atomic oxygen and neutral air, and the volume emission of the atmospheric band (762 nm), we examined the one-step and two-step excitation mechanism of $O_2 (b^1 \Sigma_g^+ \rightarrow X^3 \Sigma_g^-)(0,0)$. This allows us to derive the atomic oxygen concentration from nighttime observations of $O_2 (b^1 \Sigma_g^+ \rightarrow X^3 \Sigma_g^-)(0,0)$. The derived empirical parameters can also be utilized for atmospheric band modeling. Additionally, we derived the fit function and corresponding coefficients for the combined (one- and two-step) mechanism. The simultaneous common volume measurements of all the parameters involved in the theoretical calculation of the observed $O_2 (b^1 \Sigma_g^+ \rightarrow X^3 \Sigma_g^-)(0,0)$ emission, i.e., temperature and density of the background air, atomic oxygen density, and volume emission rate, is the novelty and the advantage of this work.

1 Introduction

The mesopause region is essential to understanding the chemical and physical processes in the upper atmosphere because this is the region of coldest temperature (during summer at high latitudes) and highest turbulence in the atmosphere (e.g., Lübken, 1997), the region of formation of such phenomena as noctilucent clouds (NLCs) and polar mesospheric summer echoes (PMSEs) (e.g., Rapp and Lübken, 2004), the region of gravity wave (GW) breaking and the formation of secondary GWs (Becker and Vadas, 2018), and the region of coupling between the mesosphere and thermosphere. This region is characterized by different airglow emissions and, particularly, by the emissions of the atmospheric band, which is produced by the excited state of molecular oxygen $O_2 (b^1 \Sigma_g^+)$. Airglow observation in the atmospheric band is a useful method to study dynamical processes in the mesopause region. There have been a number of reports of GW detection in this band (Noxon, 1978; Vierreck and Deehr, 1989; Zhang et al., 1993). Planetary wave climatology has been investigated by the Spectral Airglow Temperature Imager (SATI) instrument (López-González et al., 2009). In addition, the parameters of tides have been reported from SATI (López-González et al., 2005).
Two main mechanisms of nighttime population (note that the daytime mechanisms are quite different; see, e.g., Zaboo et al., 2018) were proposed: the first is the direct population from a three-body recombination of atomic oxygen (e.g., Deans et al., 1976); the second is the so-called two-step mechanism, which assumes an intermediate excited precursor $O_2^+$ (e.g., Witt et al., 1984; Greer et al., 1981). It has been shown by laboratory experiments that the first mechanism alone has not explained observed emissions (Young and Sharpless, 1963; Clyne et al., 1965; Young and Black, 1966; Bates, 1988). The second mechanism entails a discussion about the precursor excited state and additional ambiguities in their parameters (e.g., Greer et al., 1981; Ogryzlo et al., 1984). Thus, Witt et al. (1984) proposed the hypothesis that the $O_2^+(e^1 \Sigma^-_u^+)$ state is, possibly, the precursor; López-González et al. (1992a) suppose that the precursor could be $O_2^+(\Delta^3 \Sigma^-_u)$; and Wildt et al. (1991) found through laboratory measurements that it could be $O_2^+(A^3 \Sigma^-_u)$. Hence, the problem of identification is still not solved. The essential step in this direction has been made after the ETON 2 (Energy Transfer in the Oxygen Nightglow) rocket experiment. The ETON 2 mission yielded empirical fitting parameters that allow us to either quantify the $O_2\left(\frac{1}{2} \Sigma^+_g\right)$ (and consequently volume emission) by known O or atomic oxygen by known volume emission values (McDade et al., 1986). Despite the significance of this work, the temperature and density of air (necessary for derivation) were taken from the CIRA-72 and MSIS-83 (Hedin, 1983) models. This leads to some degree of uncertainty (e.g., Murtagh et al., 1990). Thus, more solid knowledge on these fitting coefficients based on consistent measurements of atomic oxygen, the volume emission of the atmospheric band, and temperature and density of the background atmosphere is desirable. In this paper we present common volume measurements of these parameters performed in the course of the WADIS-2 sounding rocket mission. In the next section, we describe the rocket experiment and obtained data relevant for our study. In Sect. 3, to make the paper easier to understand, we repeat some theoretical approximations from McDade et al. (1986). The obtained results of our calculations are discussed in Sect. 4. Concluding remarks and a summary are given in the last section.

2 Rocket experiment description

The WADIS (Wave propagation and dissipation in the middle atmosphere: Energy budget and distribution of trace constituents) sounding rocket mission aimed to simultaneously study the propagation and dissipation of GWs and measure the concentration of atomic oxygen. It comprised two field campaigns conducted at the Andøya Space Center (ASC) in northern Norway (69º N, 16º E). The WADIS-2 sounding rocket was launched during the second campaign on 5 March 2015 at 01:44:00 UTC under nighttime conditions. For a more detailed mission description, the reader is referred to Strelnikov et al. (2017) and the accompanying paper by Strelnikov et al. (2018).

The WADIS-2 sounding rocket was equipped with the CONE instrument to measure absolute neutral air density and temperature with high spatial resolution, an instrument for atomic oxygen density measurements (FIPEX; Flux Probe Experiment), and an airglow photometer for atmospheric band (762 nm) volume emission observation.

CONE (COMbined measurement of Neutrals and Electrons), operated by IAP (Leibniz Institute of Atmospheric Physics at Rostock University), is a classical triode-type ionization gauge optimized for a pressure range between $10^{-5}$ and 1 mbar. The triode system is surrounded by two electrodes: whilst the outermost grid is biased to +3 to +6 V to measure electron densities at a high spatial resolution, the next inner grid (~15 V) is meant to shield the ionization gauge from ionospheric plasma. CONE is suitable for measuring absolute neutral air number densities at an altitude range between 70 and 120 km. To obtain absolute densities, the gauges are calibrated in the laboratory using a high-quality pressure sensor, like a Baratron. The measured density profile can be further converted to a temperature profile assuming hydrostatic equilibrium. For a detailed description of the CONE instrument, see Giebeler et al. (1993) and Strelnikov et al. (2013). Molecular oxygen and molecular nitrogen are derived from CONE atmospheric number density measurements and partitioning according to the NRLMSISE-00 reference atmosphere (Picone et al., 2002).

The airglow photometer operated by MISU (Stockholm University, Department of Meteorology) measures the emission of the molecular oxygen atmospheric band around 762 nm from the overhead column, from which the volume emission rate is inferred by differentiation. For airglow measurements in general, a filter photometer is positioned under
the nose cone viewing along the rocket axis with a defined field of view (FOV). For WADIS-2, however, the FOV of ±3° was tilted from the rocket axis by 3° to avoid having other parts of the payload within the FOV and at the same time roughly view the same volume as the other instruments. The optical design is a standard radiometer-type system with an objective lens, a field lens, aperture, and stops, which provide an even illumination over a large portion of the detector surface (photomultiplier tube) and a defined FOV. At the entrance of the photometer there is an interference filter with a passband of 6 nm centered at 762 nm. During ascent, after the nose cone ejection, the photometer then counts the incoming photons from the overhead column (or actually the overhead cone). When the rocket passes through the layer the measured photon flux drops and above the emission layer over an overhead cone. When the rocket passes through the layer the 0–0 band of the O\(^{1}\Sigma_{g}^{+}\) atmospheric band system and the overlap of the interference filter passband. The profile is then smoothed and numerically differentiated with respect to altitude to yield the volume emission rate of the emitting layer. The data were sampled with a frequency of 100 Hz during the passage of the airglow layer (the velocity was \(\sim 800\) m s\(^{-1}\) at 95 km). However, because of the high noise level, the profile needed to be averaged to a vertical resolution of at least 3 km in order to get satisfactory results after the differentiation. A more detailed description and review of this measurement technique is given by Hedin et al. (2009).

The aim of the FIPEX developed by the IRS (Institute of Space Systems, University of Stuttgart) is to measure the atomic oxygen density along the rocket trajectory with high spatial resolution. It employs a solid electrolyte sensor, which has a selective sensitivity to atomic oxygen. A low voltage is applied between anode and cathode pumping oxygen ions through the electrolyte ceramic (yttria-stabilized zirconia). The current measured is proportional to the oxygen density. Sampling is realized with a frequency of 100 Hz and enables a spatial resolution of \(\sim 10\) m. Laboratory calibrations were done for molecular and atomic oxygen. For a detailed description of the FIPEX instruments and their calibration techniques, see Eberhart et al. (2015, 2018).

3 Theory

Here, we are repeating the theory of O\(_2\) \((b^{1}\Sigma_{g}^{+} - X^{3}\Sigma_{g}^{-})\) (0, 0) nighttime emissions following McDade et al. (1986) to make our paper more readable, using all nomenclature as in the original paper. All utilized reactions are listed in Table 1, together with corresponding reaction rates, branching ratios, quenching rates, and spontaneous emission coefficients. Some components have been updated according to modern knowledge, thus deviating from the work of McDade et al. (1986).

Assuming a direct one-step mechanism as the main one for the population and that O\(_2\) \((b^{1}\Sigma_{g}^{+})\) is in photochemical equilibrium, we can write its concentration as a ratio of production to the loss term:

\[
\left[\text{O}_2 \left( b^{1}\Sigma_{g}^{+} \right) \right] = \frac{\varepsilon k_1 [\text{O}^2 M]}{A_2 + k_2^O [\text{O}_2] + k_2^N [\text{N}_2] + k_2^O [\text{O}]}.
\]

where \(k_1\) is the reaction rate for the three-body recombination of atomic oxygen, \(\varepsilon\) is the corresponding quantum yield of O\(_2\) \((b^{1}\Sigma_{g}^{+})\) formation, \(A_2\) represents the spontaneous emission coefficient, and \(k_2^O\), \(k_2^N\), \(k_2^O\) are the quenching coefficients for reactions with O\(_2\), N\(_2\), and O, respectively. Then the volume emission, \(V_{at}\), is obtained by multiplying the O\(_2\) \((b^{1}\Sigma_{g}^{+})\) concentration by the spontaneous emission coefficient, \(A_1\), of Reaction (R5) (hereafter, nomenclature RX means the reaction X for Table 1).

In the case of known temperature, volume emission, and concentrations of O, O\(_2\), N\(_2\), and \(M\), the quantum yield of O\(_2\) \((b^{1}\Sigma_{g}^{+})\) formation can be calculated as follows:

\[
\varepsilon = \frac{V_{at}}{A_2 + k_2^O [\text{O}_2] + k_2^N [\text{N}_2] + k_2^O [\text{O}]} \frac{A_1 k_1 [\text{O}^2 M]}{A_2^b}. \tag{2}
\]

In the case of the two-step mechanism, the unknown excited-state O\(_2^*\) is populated at the first step from Reaction (R7). Then, it can be deactivated by quenching (Reaction R9), spontaneous emission (Reaction R10), or producing O\(_2\) \((b^{1}\Sigma_{g}^{+})\) by Reaction (R8). Note that Reaction (R8) is one pathway of the overall quenching Reaction (R9).

In the second step, O\(_2^*\) is transformed into O\(_2\) \((b^{1}\Sigma_{g}^{+})\), which can be deactivated by quenching (Reactions R2–R4) and by spontaneous emission (Reaction R6). Assuming photochemical equilibrium for O\(_2^*\) and, as before, for O\(_2\) \((b^{1}\Sigma_{g}^{+})\), the volume emission in the case of O\(_2\) \((b^{1}\Sigma_{g}^{+} - X^{3}\Sigma_{g}^{-})\) (0, 0) is

\[
V_{at} = \frac{A_1 \varepsilon k_1 [\text{O}^2 M] k_2^O [\text{O}_2]}{(A_2 + k_2^O [\text{O}_2] + k_2^N [\text{N}_2] + k_2^O [\text{O}])}.
\]

where the quantum yield of O\(_2^*\) formation is \(\alpha\), the quantum yield of O\(_2\) \((b^{1}\Sigma_{g}^{+})\) formation is \(\gamma\), the spontaneous emission coefficient is \(A_3\), and \(k_2^O\), \(k_3^N\), \(k_3^O\) are unknown quenching rates of O\(_2^*\). Note that the assumption about photochemical equilibrium for O\(_2^*\) and O\(_2\) \((b^{1}\Sigma_{g}^{+})\) is valid because the
curs via both channels: one-step and two-step. We discuss calculated them based on our measurements and will discuss precursors is assumed true, but solid evidence is absent. We mention on the basis of available information about potential $k_2$ for all quenching partners ($N_2$) independence is weak) or have the same temperature dependence. This means that the reaction rates are assumed that the coefficients are temperature independent (or $k_2 \propto \frac{1}{T}$). Such derivation as-

$$C_{O_2}[O_2] + C^O(O) = \frac{A_1 k_1 [O_2] M_{O_2}}{V_a (A_2 + k_2 [O_2] + k_2 [N_2] + k_2 [O] + k_2 [O])},$$

where $C_{O_2} = \left(1 + k_3 [N_2] / k_3 [O_2] \right) / \alpha \gamma$ and $C^O = k_3 / \alpha \gamma k_3 O_2$ are the fitting coefficients that can be calculated by the least-squares fit (LSF) procedure. Such derivation assumes that the coefficients are temperature independent (or temperature dependence is weak). This means that the reaction rates $k_3$ are assumed to be temperature independent (dependence is weak) or have the same temperature dependency for all quenching partners ($N_2$, $O_2$, $O$). Currently, this statement on the basis of available information about potential precursors is assumed true, but solid evidence is absent. We calculated them based on our measurements and will discuss the results in the following section.

In a more general case the population of $O_2 \left(b^1 \Sigma_g^+ \right)$ occurs via both channels: one-step and two-step. We discuss such processes in Sect. 4.3 and derive an expression for the corresponding fit function in Appendix A.

### 4 Results and discussion

Figure 1 shows input data for our calculations: temperature from the CONE instrument (Fig. 1a), number density of air (Fig. 1b), atomic oxygen concentration measured by FIPEx (Fig. 1c), and volume emission at 762 nm from the photometric instrument (Fig. 1d). A temperature minimum of $\sim 158$ K was observed at 104.2 km. A local temperature peak was measured at 98.9 km with values of 204.5 K. The secondary temperature minimum was visible at 95.4 km and amounted to $\sim 173$ K. The atomic oxygen concentration (Fig. 1c) had a peak of $\sim 4.7 \times 10^{11}$ [cm$^{-3}$] at 97.2 km and approximately coincided with the secondary temperature peak. The peak of volume emission was detected between 95 and 97 km with values of more than 1700 [phot. cm$^{-3}$ s$^{-1}$]; this is slightly beneath the atomic oxygen corresponding maximum and slightly above the secondary temperature minimum. Note that this points to the competition of temperature and the atomic oxygen concentration in processes of atomic oxygen excited-state $O_2 \left(b^1 \Sigma_g^+ \right)$ formation. Independently of the mechanism of atmospheric band emission (Eq. 1 or Eq. 3), the numerator is directly proportional to the square of the atomic oxygen concentration and inversely propor-

| Reaction | Coefficient | Reference |
|----------|-------------|-----------|
| (R1) $O + O + M \rightarrow \frac{z \lambda}{k_1} \rightarrow O_2 \left(b^1 \Sigma_g^+ \right) + M$ | $k_1 = 4.7 \times 10^{-13} (300/T)^2$ | unknown Campbell and Gray (1973) |
| (R2) $O_2 \left(b^1 \Sigma_g^+ \right) + O_2 \rightarrow \frac{k_2}{k_1} \rightarrow$ products | $k_2 = 7.4 \times 10^{-17} T^{0.5} e^{-\frac{104.7}{T}}$ | Zagidullin et al. (2017) |
| (R3) $O_2 \left(b^1 \Sigma_g^+ \right) + N_2 \rightarrow \frac{k_3}{k_1} \rightarrow$ products | $k_3 = 8 \times 10^{-20} T^{1.5} e^{\frac{503}{T}}$ | Zagidullin et al. (2017) |
| (R4) $O_2 \left(b^1 \Sigma_g^+ \right) + O \rightarrow \frac{k_4}{k_1} \rightarrow$ products | $k_4 = 8 \times 10^{-14}$ | Slanger and Black (1979) |
| (R5) $O_2 \left(b^1 \Sigma_g^+ \right) \rightarrow \frac{k_5}{k_1} \rightarrow A_1 \rightarrow O_2 + h\nu (762 nm)$ | $A_1 = 0.0834$ | Newnham and Ballard (1998) |
| (R6) $O_2 \left(b^1 \Sigma_g^+ \right) \rightarrow \frac{k_6}{k_1} \rightarrow A_2 \rightarrow O_2 + h\nu$ (total) | $A_2 = 0.088158$ | Yankovsky et al. (2016) |
| (R7) $O + O + M \rightarrow \frac{ak_1}{k_1} \rightarrow O_2^* + M$ | $\alpha$ | unknown |
| (R8) $O_2^* + O_2 \rightarrow \frac{k_1}{k_1} \rightarrow O_2 \left(b^1 \Sigma_g^+ \right) + O_2$ | $\gamma$ | unknown |
| (R9) $O_2^* + O_2 + N_2 \rightarrow \frac{k_1}{k_1} \rightarrow O_2 \left(b^1 \Sigma_g^+ \right) + O_2 + N_2$ | $k_3^2, k_3^3, k_3^4$ | unknown |
| (R10) $O_2^* \rightarrow \frac{k_1}{k_1} \rightarrow O_2 + h\nu$ | $A_3$ | unknown |

$O_2 \left(b^1 \Sigma_g^+ \right)$ radiative lifetime is less than 12 s and all potential candidates for $O_2^*$ have lifetimes less than several seconds (e.g., López-González et al., 1992a, b, c; Yankovsky et al., 2016, and references therein).

Collecting all known values on the right-hand side (RHS), all unknown summands on the left-hand side (LHS), and omitting emissive summand $A_3$ as noneffective loss (McDade et al., 1986). Eq. (3) can be rearranged as follows:

$$C_{O_2}[O_2] + C^O(O) = \frac{A_1 k_1 [O_2] M_{O_2}}{V_a (A_2 + k_2 [O_2] + k_2 [N_2] + k_2 [O] + k_2 [O])},$$

where $C_{O_2} = \left(1 + k_3^2 [N_2] / k_3^2 [O_2] \right) / \alpha \gamma$ and $C^O = k_3^2 / \alpha \gamma k_3^2 O_2$ are the fitting coefficients that can be calculated by the least-squares fit (LSF) procedure. Such derivation assumes that the coefficients are temperature independent (or temperature dependence is weak). This means that the reaction rates $k_3$ are assumed to be temperature independent (dependence is weak) or have the same temperature dependency for all quenching partners ($N_2$, $O_2$, $O$). Currently, this statement on the basis of available information about potential precursors is assumed true, but solid evidence is absent. We calculated them based on our measurements and will discuss the results in the following section.

In a more general case the population of $O_2 \left(b^1 \Sigma_g^+ \right)$ occurs via both channels: one-step and two-step. We discuss
tional to the third power of the temperature (via reaction rate \( k_1 \) and \( M \), considering the ideal gas law). Our rocket experiment shows an essential difference of emissions between ascending and descending flights (see Strelnikov et al., 2018). It also demonstrates significant variability in other measured parameters, including neutral temperature and density as well as atomic oxygen density (Strelnikov et al., 2017, 2018). This suggests that, in the case of the ETON 2 experiments, the temporal extrapolation of atomic oxygen for the time of the emission measurement flight (which was approximately 20 min earlier) may lead to serious biases in estimations because, as one can see from Eqs. (1) and (3), volume emission depends on the atomic oxygen concentration quadratically. Since the best-quality data were obtained during the descent of the WADIS-2 rocket flight, we chose this data set for our analysis (Strelnikov et al., 2018). The region above 104 km is subject to auroral contamination. In the region below 92 km, negative values may occur in the volume emission profile as a result of self-absorption in the denser atmosphere below the emission layer. Hence, we considered the region near the emission peak between 92 and 104 km as most appropriate for our study. The comparisons of our measurements with other observations, as well as with the results of modeling, are presented in several papers (e.g., Eberhart et al., 2018; Strelnikov et al., 2018).

4.1 One-step mechanism

Figure 2 shows the quantum yield of \( \text{O}_2 \left( b^1 \Sigma_g^+ \right) \) formation \( \varepsilon \) calculated according to Eq. (2), which is necessary to form \( \text{O}_2 \left( b^1 \Sigma_g^+ \right) \) under the assumption that the direct three-body recombination of atomic oxygen is the main mechanism. The uncertainties for this figure (as well as for other figures) were calculated according to a sensitivity analysis (von Clarmann, 2014; Yankovsky and Manuilova, 2018, their Appendix 1; Vorobeva et al., 2018), for which the errors represent error propagation from the experimental data. Calculated values of \( \varepsilon \) are placed in the range [0.07; 0.13], which is in good agreement with the values derived by McDade et al. (1986). The averaged value amounts to 0.11 ± 0.02. The range of values, taking into account both the variance and the error range, amounts to [0.02; 0.22]. By the physical nature of this value, the quantum yield of \( \text{O}_2 \left( b^1 \Sigma_g^+ \right) \) formation should not depend on altitude. Figure 2 shows some altitude dependence of central values of \( \varepsilon \), but considering the large error range, there is no clear altitude dependence. The variability of the data points is much smaller than the errors of the individual points. Hence, in light of the analysis of our rocket experiment, we may not reject the direct excitation mechanism.

Although the population via the one-step mechanism alone is, generally speaking, possible, it is improbable because laboratory experiments show that direct excitation alone may not explain observed emissions (Young and Sharpless, 1963; Clyne et al., 1965; Young and Black, 1966; Bates, 1988). This conclusion is in agreement with the conclusion from McDade et al. (1986), which stated that the one-step excitation mechanism is not sufficient to explain the \( \text{O}_2 \left( b^1 \Sigma_g^+ \right) \) population. Hence, in the following, we check the second energy transfer mechanism.

4.2 Two-step mechanism

Figure 3 depicts the altitude profile of the RHS of Eq. (4) and the profile calculated by the LSF. The fitting coefficients, \( C^{\text{O}_2} \) and \( C^\text{O} \), resulting from this fit amount to \( 9.8_{-5.4}^{+5.3} \) and \( 2.1_{-0.6}^{+0.3} \), respectively. The uncertainties were calculated, as is common for LSF (Bevington and Robinson, 2003), based on error propagation from the RHS as provided in Fig. 3. Our \( C^{\text{O}_2} \) coefficient is partially, within the error range, in agreement with \( C^{\text{O}_2} \) coefficients given in McDade et al. (1986), which amount to \( 4.8 \pm 0.3 \) and \( 6.6 \pm 0.4 \) for temperature from CIRA-72 and MSIS-83, respectively. The \( C^\text{O} \) coefficient is approximately 1 order lower. There are several possible reasons for the large discrepancy in \( C^\text{O} \), for example the temperature dependence of the O quenching or that, in the case of ETON 2 experiments, mean temperature profiles from the models CIRA-72 and MSIS-83 were utilized, which does not reproduce any short-time dynamical fluctuations, solar cycle conditions, etc. In the framework of our analysis, we may not identify the reason for the large discrepancy in \( C^\text{O} \) more precisely. Fitting coefficients defined in such a way (Eq. 4) do not have a direct physical meaning. However, they have a physical meaning in several limit cases. If the quenching coefficients of a precursor with molecular nitrogen are much smaller than those with molecular oxygen (\( k_3 N_2 \ll k_4 \)), then \( \alpha \eta = 1/C^{\text{O}_2} \). The assumption that the quenching of the precursor with \( N_2 \) is much slower than quenching with \( \text{O}_2 \) is just a working hypothesis, which is commonly used for the analysis of possible precursors (e.g., McDade et al., 1986; López-González et al., 1992a, b; and references therein). It is true for such potential precursors as \( \text{O}_2 \left( A^1 \Sigma_g^+ \right) \) (Kenner and Ogryzlo, 1983b), but generally, there is no evidence for or against that. If it is not true, any definite conclusion on precursors by known \( C^{\text{O}_2} \) is not possible. In our case \( \alpha \eta = 0.102_{-0.041}^{+0.120} \). In other words, in the case of the two-step formation of \( \text{O}_2 \left( b^1 \Sigma_g^+ \right) \) with energy transfer agent \( \text{O}_2 \), the total efficiency \( \eta = \alpha \eta \) amounts to 10.2 %, which is the lowest amongst known values. Based on rocket experiment data analysis (ETON), Witt et al. (1984) obtained \( \alpha \eta = 0.12 \). According to McDade et al. (1986) for the case with \( k_4^2 = 8 \times 10^{-14} \), the total efficiencies are 0.15 and 0.21 for temperature profiles adopted from MSIS-83 and CIRA-72, respectively. The analyses of López-González et al. (1992a, c), which adopted \( \text{O}_2 \), \( N_2 \), and temperature profiles from the model (Rodrigo et al., 1991), showed a total efficiency of 0.16. In contrast to our work, all investigations mentioned above utilized temperature and atmospheric
Measurements of (a) temperature (CONE), (b) number density of air (CONE), (c) atomic oxygen concentration (FIPEX), and (d) volume emission at 762 nm (photometer).

density from models that describe a mean state of the atmosphere. This is a possible reason for discrepancy in the results. Total efficiency $\eta$ may serve as an auxiliary quantity to identify the precursor. According to the physical meaning of efficiency, it may not be larger than 1. Hence, $\alpha$ and $\gamma$, as well as the total efficiency, are smaller than 1. Consequently, $\gamma = \eta / \alpha < 1$, and we can examine potential candidates for $O_2^*$ with this criterion. From an energetic point of view, only four bound states of molecular oxygen can be considered as an intermediate state for the $O_2 \left( b^1 \Sigma^+_g \right)$ population: $O_2 \left( A^3 \Sigma_u^+ \right)$, $O_2 \left( A^3 \Delta_u \right)$, $O_2 \left( c^1 \Sigma_u^- \right)$, and $O_2 \left( 5 \Pi_g \right)$ (Greer et al., 1981; Wraith, 1982; Witt et al., 1984; McDade et al., 1986; López-González et al., 1992c). For better readability, we will partially repeat a table from López-González et al. (1992b, c) with known $\alpha$ in our work (Table 2). From Table 2, it can be seen that only $O_2 \left( A^3 \Delta_u \right)$ and $O_2 \left( 5 \Pi_g \right)$ fit the criterion of $\gamma = 0.102 / \alpha < 1$. At a lower limit of uncertainty ($\gamma = 0.061 / \alpha < 1$) $O_2 \left( A^3 \Delta_u \right)$ and $O_2 \left( 5 \Pi_g \right)$ satisfy the criterion and, considering the upper limit ($\gamma = 0.222 / \alpha < 1$), only $O_2 \left( 5 \Pi_g \right)$ may serve as a precursor.

The second expression that helps to clarify the choice of the precursor is the ratio of quenching rates. In the limit of low quenching with molecular nitrogen ($k_{N_2} \ll k_{O_2}$), the ratio of fitting coefficients equals the ratio of the quenching rates of atomic and molecular oxygen ($C^O / C^{O_2} = k^O_3 / k^{O_2}_3$). An analysis from the ETON 2 rocket experiment yields values for the quenching coefficient ratios of potential precursors of 3.1 and 2.9 for temperature from CIRA-72 and MSIS-83, respectively. This is close to the value from Ogryzlo et al. (1984), who found $k^O_3 / k^{O_2}_3 = 2.6$ by laboratory measurements; however, as was noted in their work, substitution of these values into the equation for emission yields 16% of the observed emission (Ogryzlo et al., 1984). These findings point to the possibility of a too-
Table 2. Efficiencies $\alpha$ of the different excited states of O$_2$.

| State of O$_2$ | $\alpha$ | $\alpha$ | $\alpha$ | Reference |
|---------------|---------|---------|---------|-----------|
| O$_2$(c$^1\tilde{X}_u$) | 0.03 | 0.12 | 0.04 | 0.66 | Wraight (1982), Smith (1984) |
| O$_2$(A$^3\tilde{X}_u$) | 0.04 | 0.18 | 0.06 | 0.5 | Bates (1988) |
| O$_2$(A$^3\tilde{X}_u$) | 0.03 | 0.18 | 0.06 | 0.52 | López-González et al. (1992a, b, c) |

Figure 2. Quantum yield of O$_2$(b$^1\Sigma_g^+$) formation $\varepsilon$ for the case of the one-step mechanism.

Figure 3. RHS (dots) and least-squares fit of LHS (black line) of Eq. (4).

Figure 4. Atomic oxygen concentration: FIPEX (black line); model MSIS00 (red line); derived from emission observation with McDade et al. (1986) coefficients (blue line); calculated with newly derived fitting coefficients for the two-step mechanism (green line).

High measured ratio $k_{3}^{O}/k_{3}^{O_2}$ as a result of too-strong quenching of the precursor by atomic oxygen. Our value of quenching ratios $k_{3}^{O}/k_{3}^{O_2}$ amounts to $0.21^{+0.32}_{-0.12}$. There is not enough information on measured values for bound states of molecular oxygen. Laboratory measurements for O$_2$ (A$^3\tilde{X}_u$) ($v = 0 − 4$), O$_2$ (A$^3\tilde{X}_u$) ($v = 2$), and O$_2$ (c$^1\tilde{X}_u$) infer the values of the $k_{3}^{O}/k_{3}^{O_2}$ ratio to be $30 \pm 30$, $100 \pm 15$, and $200 \pm 20$, respectively (Kenner and Ogrzyzlo, 1980, 1983a, b, 1984). On the other hand, Slanger et al. (1984) found that the lower limit of O$_2$ (A$^3\tilde{X}_u$) ($v = 8$) quenching by O$_2$ must be $\geq 8 \times 10^{-11}$. If the results from Slanger et al. (1984) were applied to the results from Kenner and Ogrzyzlo (1980, 1984) for k$_3^{O_2}$, then the ratio of $k_{3}^{O}/k_{3}^{O_2}$ would be 2 orders lower. This short discussion illustrates a strong scattering of this ratio. For our two potential candidates (O$_2$ (A$^3\tilde{X}_u$) and O$_2$ (A$^3\tilde{X}_u$)), there is information about the $k_{3}^{O}/k_{3}^{O_2}$ ratio for only O$_2$ (A$^3\tilde{X}_u$). Through the comprehensive analysis of known rocket experiments, López-González et al. (1992a, b, c) inferred that the upper limit of the ratio amounts to 1. Hence, our value of $k_{3}^{O}/k_{3}^{O_2} = 0.21^{+0.32}_{-0.12}$ agrees with this result. Consistent information from laboratory experiments on the ratio for O$_2$ (A$^3\tilde{X}_u$) is absent. Thus, we can propose as potential candidates for
precursors both \( \text{O}_2 \left( A^3 \Delta_u \right) \) and \( \text{O}_2 \left( ^3 \Pi_g \right) \); however, we are not able to identify which of these two is more preferable.

In order to illustrate the application of the newly derived fitting coefficients we compare in Fig. 4 the atomic oxygen concentration from FIPEX (black line), from the NRL MSISE-00 reference atmosphere model (Picone et al., 2002) (red line) calculated with McDade et al. (1986) coefficients (blue line), and our fitting coefficients for the two-step mechanism (green line). In the region 94–98 km, i.e., at atomic oxygen peak and volume emission peak (see Fig. 1d), fitting coefficients from this paper reproduce observed values better than the McDade coefficients (MSIS-83 case). Our fitting coefficients and the fitting coefficients of McDade give a similar approximation above the atomic oxygen peak (\( \sim 98–104 \) km). The shape of the calculated profiles appears slightly different, with the peak maximum at a higher altitude than the observed. In this, our result resembles the McDade results, probably because in both cases the ratio of two reaction rates is derived, but not the rates themselves. In the lower part our results and those of McDade differ because our \( \text{CO}_2 \) value is larger and the term with molecular oxygen dominates. Nevertheless, the atomic oxygen retrieved with our fitting coefficients satisfactorily reproduces measurements, especially at the peak.

4.3 Combined mechanism

In the most general case, the \( \text{O}_2 \left( b^1 \Sigma^+_g \right) \) population passes through two channels: directly and via a precursor. In fact, theoretical calculations from Wraight (1982) and laboratory measurements from Bates (1988) predicted a direct population with efficiencies of 0.015 and 0.03, respectively, which is not sufficient to explain the observed emissions (Bates, 1988; Greer et al., 1981; Krasnopolsky, 1986). A similar value, \( \varepsilon = 0.02 \), was shown in the analysis by López-González et al. (1992b, c). We investigated a combined mechanism based on the LSF calculation and fit function (derivation in Appendix A):

\[
\frac{\text{[O}_2\text{] + } D_1 \text{[O]}}{D_2 + \tilde{\varepsilon} \left( 1 + D_1 \text{[O]} / \text{[O}_2\text{]} \right)} = \frac{A_k k_1 [\text{O}]^2 M [\text{O}_2]}{V_{at} \left( A_2 + k_2^O \text{[O}_2\text{]} + k_2^N \text{[N}_2\text{]} + k_2 [\text{O}] \right)}, \tag{5}
\]

where hereafter tildes denote these values for the combined mechanism and do not equal the values for one-step or two-step mechanisms (Sect. 4.1 and 4.2): \( D_1 = \tilde{\varepsilon} \text{[O]} / \tilde{\varepsilon} \text{[O}_2\text{]} \) and \( D_2 = \tilde{\alpha} \tilde{\gamma} \) are the fitting coefficients, which refer to the ratio of quenching rates and \( \tilde{\gamma} \equiv \tilde{\alpha} \tilde{\gamma} \) total efficiency for the two-step channel, respectively. The fitting coefficients were calculated for two limit cases, \( \varepsilon = 0.015 \) (Wraight, 1982) \( \tilde{\varepsilon} = 0.03 \) (Bates, 1988), and for the averaged case \( \tilde{\varepsilon} = 0.022 \).

The results for the best fit in each case are listed in Table 3. Analogously to the two-step mechanism (Sect. 4.2), for the case of the combined mechanism \( \gamma = \tilde{\gamma} / \tilde{\alpha} < 1 \); hence, the precursor should satisfy \( \tilde{\alpha} > 0.08^{+0.12}_{-0.04} \) (see Table 3). For central values of \( \tilde{\alpha} \), only \( \text{O}_2 \left( A^3 \Delta_u \right) \) and \( \text{O}_2 \left( ^3 \Pi_g \right) \) satisfy this criterion (see Table 2). At a lower limit of uncertainty (\( \tilde{\alpha} > 0.04 \)) \( \text{O}_2 \left( A^3 \Delta_u \right) \), \( \text{O}_2 \left( A^3 \Sigma^+_u \right) \), and \( \text{O}_2 \left( ^3 \Pi_g \right) \) satisfy the criterion and, considering the upper limit (\( \tilde{\alpha} > 0.2 \), only \( \text{O}_2 \left( ^3 \Pi_g \right) \) may serve as a precursor. The upper limit of the ratio \( k_3^O / k_5^O < 1 \) for \( \text{O}_2 \left( A^3 \Delta_u \right) \), derived by López-González et al. (1992a, b, c), is in agreement with our calculations \( \left( 0.231^{+0.358}_{-0.142} \right) \). As noted above, the ratio for \( \text{O}_2 \left( ^3 \Pi_g \right) \) is unknown. Consequently, taking into account both conditions, only \( \text{O}_2 \left( A^3 \Delta_u \right) \) and \( \text{O}_2 \left( ^3 \Pi_g \right) \) may serve as precursors.

Figure 5 illustrates a sanity check for volume emissions derived (black lines) with the fitting coefficients of McDade et al. (1986) for the MSIS-83 (Fig. 5c) case, the CIRA-72 case (Fig. 5d), and with our newly derived fitting coefficients for the two-step (Fig. 5a) and combined (Fig. 5b) mechanisms (Fig. 5b) in comparison with the measured one (red lines). All of the derived volume emission profiles (black lines) were calculated based on the temperature, concentration of surrounding air, and concentration of atomic oxygen from our rocket launch. The calculations with the combined mechanism (Eq. 5) and two-step energy transfer mechanism (Eq. 4) give almost identical results. The results obtained with the new fitting coefficients are in satisfactory agreement with the measured volume emissions at the peak and above, whereas the McDade coefficients related to temperature from CIRA-72 give better approximations below the volume emission peak (92 km). The coefficients of McDade related to temperature from MSIS-83 are in better agreement with our results and are almost identical above the volume emission peak. More independent common volume in situ measurements are necessary to validate these results.

5 Summary and conclusions

Based on the rocket-borne common volume simultaneous observations of atomic oxygen, atmospheric band emission (762 nm), and density and temperature of the background atmosphere, the mechanisms of \( \text{O}_2 \left( b^1 \Sigma^+_g \right) \) formation were analyzed. Our calculations show that one-step direct excitation alone is less probable for the reasons discussed above (Sect. 4.1).

For the case of the two-step mechanism, we found new coefficients for the fit function in accordance with McDade et al. (1986) based on self-consistent temperature, atomic oxygen, and volume emission observation. These coefficients amounted to \( C^O_2 = 9.8^{+5.3}_{-6.5} \) and \( C^O = 2.1^{+0.3}_{-0.6} \). The \( C^O_2 \) coefficient is partially, within the error range, in agreement with
Table 3. Fitting coefficients for the combined mechanism (Eq. 5) at different efficiencies.

|            | Low $\bar{\varepsilon}$ | High $\bar{\varepsilon}$ | Averaged $\bar{\varepsilon}$ |
|------------|--------------------------|---------------------------|-----------------------------|
|            | Wraight (1982)           | Bates (1988)              | (this work)                 |
| $\bar{\varepsilon}$ | 0.015                    | 0.03                      | 0.022                       |
| $D_1 = \frac{k_{O^3}^O}{k_{O^2}^3}$ | 0.211 ± 0.355 − 0.136    | 0.397 ± 0.22 − 0.282       | 0.231 ± 0.358 − 0.142       |
| $D_2 = \bar{\alpha} \bar{\gamma}$    | 0.087 ± 0.12 − 0.041      | 0.073 ± 0.119 − 0.042      | 0.08 ± 0.12 − 0.04          |

Figure 5. Volume emissions: photometer (red line); derived from atomic oxygen (black line) with (a) newly derived fitting coefficients for the two-step mechanism, (b) with fitting coefficients for the combined mechanism, (c) with McDade et al. (1986) coefficients that correspond to MSIS-83 temperature, and (d) with McDade et al. (1986) coefficients that correspond to CIRA-72 temperature.

$C^O_2$ coefficients given in McDade et al. (1986), and the $C^O_2$ coefficient is approximately 1 order lower. The general implication of these results is the parameterization of volume emission in terms of known atomic oxygen. This can be utilized either for atmospheric band volume emission modeling or for the estimation of atomic oxygen by known volume emission. We identified two candidates for the intermediate state of $O_2^*$. Our results show that $O_2 (A^3 \Delta_u)$ or $O_2 (\Sigma^+_g)$ may serve as a precursor.

Taking into account both channels of $O_2 (b^1 \Sigma^+_g)$ formation, we proposed a combined mechanism. In this case, atomic oxygen via volume emission or volume emission based on known atomic oxygen can be calculated by Eq. (5). The recommended fitting coefficients amounted to $D_1 = 0.231 ± 0.358$ and $D_2 = 0.08 ± 0.12$ with the efficiency of the direct channel as $\bar{\varepsilon} = 0.022$. These coefficients have a mean total efficiency ($\bar{\alpha} \bar{\gamma} = 0.08 ± 0.12$) and a ratio of quenching coefficients ($\frac{k_{O^3}^O}{k_{O^2}^3} = 0.231 ± 0.358$) for the two-step chan-
nel. The analysis of their values indicates that \( \text{O}_2 \left( A^3 \Delta_u \right) \) and \( \text{O}_2 (\tilde{5} \Pi_p) \) may serve as possible precursors for the two-step channel in the combined mechanism. In the context of our rocket experiment, we do not have the possibility to figure out which mechanism is true. Nevertheless, we consider the combined mechanism as more relevant to nature because it has a higher generality. This conclusion does not contradict the current point of view that the two-step mechanism is dominant because \( \tilde{c} \) is assumed to be 1.5\%–3\%. Moreover, it is possible that in reality the mechanism is much more complex and it has a multichannel or more than two-step nature. Undoubtedly, more common volume simultaneous observations of the atmospheric band and atomic oxygen concentrations would be desirable to confirm and improve these results.

Data availability. The rocket-borne measurements and calculated data shown in this paper are available via the IAP ftp server at ftp://ftp.iap-kborn.de/data-in-publications/GrygalashvylyACP2018 (last access: 25 January 2019).
Appendix A

We consider photochemical equilibrium for the nighttime $\text{O}_2 \left( b^1 \Sigma_g^+ \right)$ concentration. If $\text{O}_2 \left( b^1 \Sigma_g^+ \right)$ is produced via both channels, the equilibrium concentration is given by the following expression:

$$\left[ \text{O}_2 \left( b^1 \Sigma_g^+ \right) \right] = \frac{\tilde{\epsilon} k_1 [\text{O}]^2 M + \tilde{\gamma} k_3^O [\text{O}_2] [\text{O}_2^*]}{A_2 + k_2^O [\text{O}_2] + k_2^N [\text{N}_2] + k_3^O [\text{O}]}, \quad (A1)$$

where the tilde denotes the combined mechanism, $A_1$, $k_1$, $k_2^O$, $k_2^N$, $k_3^O$, $k_3^N$ are the ratios for the corresponding processes (see Table 1), and $\text{O}_2^*$ is the unknown precursor.

Considering this precursor in photochemical equilibrium, we can obtain the following expression for its concentration:

$$[\text{O}_2^*] = \frac{\tilde{\alpha} k_1 [\text{O}]^2 M}{A_3 + k_3^O [\text{O}_2] + k_3^N [\text{N}_2] + k_3^O [\text{O}]}, \quad (A2)$$

where efficiency is $\tilde{\alpha}$, $A_3$ is the unknown spontaneous emission coefficient of $\text{O}_2^*$, and $k_3^O$, $k_3^N$, $k_3^O$ are the unknown quenching rates for $\text{O}_2^*$.

Substituting (A2) into (A1) and into the expression for volume emission we obtain

$$V_{at} = A_1 \left[ \text{O}_2 \left( b^1 \Sigma_g^+ \right) \right] = \frac{A_1 k_1 [\text{O}]^2 [\text{O}_2] M}{A_2 + k_2^O [\text{O}_2] + k_2^N [\text{N}_2] + k_3^O [\text{O}] \left( \frac{1}{[\text{O}_2]} + \frac{\tilde{\gamma} k_3^O}{A_3 + k_3^O [\text{O}_2] + k_3^N [\text{N}_2] + k_3^O [\text{O}]} \right)}. \quad (A3)$$

We assume that, in analogy with a two-step mechanism, a spontaneous emission $A_3$ of $\text{O}_2^*$ is much smaller than the quenching, and we utilized a traditional assumption about low quenching with molecular nitrogen ($\tilde{k}_3^N \ll \tilde{k}_3^O$), which is commonly used to analyze a potential precursor. In this case, Eq. (A3) can be rearranged as follows:

$$\frac{[\text{O}_2] + \frac{\tilde{k}_3^O}{k_3^O} [\text{O}]}{\tilde{\alpha} \tilde{\gamma} + \tilde{\epsilon} \left( 1 + \frac{\tilde{k}_3^O}{k_3^O} [\text{O}] / [\text{O}_2] \right)} = \frac{A_1 k_1 [\text{O}]^2 M [\text{O}_2]}{V_{at} \left( A_2 + k_2^O [\text{O}_2] + k_2^N [\text{N}_2] + k_3^O [\text{O}] \right)} \quad (A4)$$

We defined unknown fitting coefficients $D_1 \equiv \frac{\tilde{k}_3^O}{k_3^O}$ and $D_2 \equiv \tilde{\alpha} \tilde{\gamma}$. Expression (A4) was utilized to calculate them with LSF.
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