The Role of Excited Oxygen Molecules in the Formation of the Secondary Ozone Layer at 87 to 97 km

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Abstract The secondary ozone layer is located at elevations of 87 to 97 km in the upper mesosphere – lower thermosphere. It overlaps with the ionospheric D-layer. Daytime intensive UV radiation is dissociating O₂ molecules to O atoms and photoexcitating O₂ molecules up to 11.07 eV level. Ozone photolysis between the wavelengths of 118.7–121.6 nm produces three oxygen atoms from one ozone molecule. Collision reactions of O₂(B³Σ_u−) and O₂(X³Σ_g−, ν≥26) with O₂(X³Σ_g−, ν=0) produce additional oxygen atoms. The number of oxygen atoms is maintained at such a high level that a small but significant ozone concentration survives. UV radiation weakens radically during the night. The number of O atoms shows no diurnal variation in the MLT. This leads to a ten-fold increase of ozone concentration over the course of the night. Dissociative recombination of O₂+ (entered via diffusion from above) and reactions of O(3P) atoms with excited O₂ molecules generate O(1S) atoms. The quenching of O(1S) → O(1D) emits the green nightglow. The reactions of O(1D) with ozone and O₂ absorption of UV nightglow produce O₂(c¹Σ_u+, A'³∆_u and A³Σ_u+). When these molecules relax, they emit the O₂ UV nightglows. The relaxations of O₂(a¹∆_g) and O₂(b¹Σ_g+) emit infrared nightglows.

Keywords Secondary Ozone Zone, Photoexcitation of O₂, Vibrational and Electronical Excitation of O₂, O₂+ Dissociative Recombination, Nightglows

1. Introduction

Before the 1970s, the mesospheric–lower thermospheric (MLT) layer was located too far from the ground for the purposes of obtaining good observation results. On the other hand, for rocket research it was located too near, as they were targeting objects that were further away. Thomas and Bowman [1] examined the diurnal variations of constituents in an oxygen-hydrogen atmosphere. They compared the results of available experimental data, and suggested that their laboratory tests demonstrated atmospheric oxygen chemistry in the Schumann–Runge band region as well.

Hays and Roble [2] found that stellar ultraviolet light near 2500 Å is attenuated in the Earth’s upper atmosphere in the 60- to 100-km region due to strong absorption in the Hartley continuum of ozone. They found that the night-time ozone number density has a bulge in its vertical profile with a peak of 1 to 2 x 10⁸ cm⁻³ at approximately 83km and a minimum near 75km.

Miller and Ryder [3] also measured the concentration of ozone in the mesosphere and lower thermosphere at sunset using the occultation technique. They found a minimum in the O₃ concentration near 80km and a secondary O₃ maximum around 90km.

Nowadays, there is a special Global Ozone Monitoring by Occultation of Stars (GOMOS) instrument on board the European Space Agency’s Envisat satellite. This instrument measures ozone using the stellar occultation method [5]. The number density of ozone has a strong maximum region around 90km (see Figure 1). This is known as the secondary ozone maximum [5, 6]. The night-time mixing ratio of ozone in this maximum zone is comparable to that found in the stratospheric maximum (around 8 to 9 ppm). Daytime mixing ratios are substantially smaller but are significantly higher than those observed elsewhere above or (immediately) below [5, 7].

Rogers et al. [8] measured the ozone in a layer of 80 to 104 km by the microwave line of 11.072454 GHz. The Gaussian distribution of the ozone mixing ratio was centred at 92km with the full width at maximum of 10km. The location of the secondary ozone maximum can be set between altitudes of 87 to 97 km. An important detail is that this ozone layer is a part of the ionospheric D-region as well [9].
2. Materials and Methods

2.1. Aims of the Study

The aim of this meta-study is to provide a basic understanding of the formation and dynamics of energetically excited oxygen molecules, and of their importance in the generation of the secondary ozone layer. The study is based on the available literature of excited O₂ molecules and excited O atoms in the secondary ozone layer. Data from the literature regarding atmospheric nightglows on Earth and Venus are used as crucial evidence.

2.2. Formulas Used in Calculations

The formation enthalpies of reactions are calculated by formula (1). If difference $\Delta H^0<0$, reaction is proceeding spontaneously, it is an exergonic reaction.

$$\begin{align*}
\text{(products)} - \text{(starting products)} & = \Delta H^0 \\
(\Delta H^0_1 + \Delta H^0_2) \text{ end products} - (\Delta H^0_1 + \Delta H^0_2) \text{ starting products} & = \Delta H^0
\end{align*}$$

Relationship between energy and wavelength of electromagnetic radiation (EMR) is calculated by formula (2) [10].

$$\lambda = \frac{hcN}{E}$$  

The formations of enthalpies (kJ/mol) are: for ozone 142.7, for O (⁴P) 249.2, for O (⁴D) 438.9 and for O (⁴S) 653. The relationships between energy units eV, kJ/mol and cm⁻¹ are 1 eV = 96.49 kJ/mol = 8064 cm⁻¹ [11].

3. Discussion

3.1. Number Densities of Oxygen Atom in the MLT

According to Lin et al. [12] at an altitude of about 101 km the model-dependent peak of atomic oxygen density is 3.6 ($\pm$1.9) × 10¹¹ cm⁻³ (or in terms of the number of atomic oxygen 4.5 x 10¹¹). In a later study by Hedin et al. [13] the maximum concentration of atomic oxygen (4.7 x 10¹¹ atoms in cubic cm) was achieved at an altitude of 95 km.

At an altitude of 105km, the number density of O atoms is 2.8 x 10¹¹, at an altitude of 120km it is 8 x 10¹⁰ and at an altitude of 150km it is 8 x 10⁹ [14]. The number of O atoms reaches their atmospheric maximum in the MLT. At an altitude of 120km the number density of O₂ molecules is 6 x 10¹⁰ [14], which is less than the number density of O atoms. At an altitude of 91km the number of atomic oxygen is 1.4 x 10¹¹ during night and day [12].

3.2. Absorption of UV, Visible and IR Radiation by Oxygen Species

In the thermosphere, solar UV radiation is absorbed via photoionization by hydrogen, oxygen and nitrogen atoms and molecules and helium atoms. It leads to almost complete absorption of UV radiation shorter than 102nm in the thermosphere above the MLT layer.

In order to be photodissociated or photoexcititated by a
non-ionizing photon, a molecule must absorb on the wavelength of the photon. These absorptions define the absorptive optical thickness of the atmosphere towards the UV radiation.

Oxygen atoms absorb UV radiation at wavelengths between 10nm and 91nm \[15\]. An oxygen atom has the ground state O(\(^3\)P), which is a triplet state. Higher energy forms, singlet oxygen atoms, are formed if the electrons pair up, like in the first excited state O(\(^1\)D) and second excited state O(\(^1\)S). Radiative lifetime of O(\(^1\)S) is only 0.84s and that of O(\(^1\)D) is 114s \[16\].

Oxygen molecule absorbs UV radiation at wavelengths from 58 to 260 nm. The absorption cut-off of UV photons by O2 molecules is 260nm in 1 atm. air pressure and 250nm in zero pressure \[17\].

For dissociation, a photon must have so much energy that it exceeds the dissociation energy of the molecule. Due to the properties of the electron configuration, the excited O2 molecules have individual dissociation ranges within which they are more or less metastable. If the UV photon raises an electron to such a high level in which it is in the dissociation range of a certain excited state, the O2 molecule is photoexcitated rather than dissociated.

The spectrum of O2 from 102 to 125 nm is very complex (see Figure 2) \[18\]. Lyman β line at 102.57nm is known to penetrate down to about 86 km level \[19\]. The ionization potential of oxygen molecules is 12.07eV, which is equal to the wavelength of 102.78nm. O2 molecules absorb weakly at the Lyman β line \[20\] so the ionization of O2 is possible in the entire MLT layer as well.
Absorption by ozone is a much studied topic, there remains disagreement regarding the absorption minimum between the Huggins and Chappuis bands (350–425 nm), and especially near 380 nm due to low absorption coefficients and the difficulties of preparing pure ozone for study [27].

3.3. Basic Notations and Rules in Quantum Chemistry

3.3.1. Quantum Chemistry Notations

The spectroscopic state of atoms and molecules is conveniently summarized by the use of spectroscopic term symbols. The Greek capital letters Σ, Π or Λ are used as the principal symbol for the state of the molecule. It tells the resultant angular momentum of all the electrons with respect to the internuclear axis Λ, which may have values 0, 1 and 2. When |Λ| is 0, the letter is Σ; when |Λ| is 1, the letter is Π and when |Λ| is 2, the letter is Λ. In atoms, the notation is analogous to the roman letters S, P and D. The multiplicity (singlet = 1, triplet = 3) is denoted by a superscript number to the left side of the letter [28, 29].

A closed molecular group, whether σ or π, must give a 1Σ term, as the resultant angular momentum and the multiplicity must both be zero. A superscript + or − on the right side indicates whether the total wave function is symmetric or antisymmetric with respect to the reflection in a plane containing the principal axis. A subscript g or u on the right side is given corresponding to the gerade or ungerade molecular symmetry. The product of any number of g functions or an even number of u functions results in a g function [28].

The lowest state of the molecule (either Σ, Π or Λ) is labelled X. The excited states are then labelled alphabetically, starting with Σ, Π and so on to indicate the first excited state, second excited state, and so on. Uppercase letters are used for states with the same multiplicity as the ground state, while lowercase letters are used for states with a different multiplicity. Occasionally, new states are found which lie in between previously assigned states. These states are labelled A′ or B′. Thus, for O₂, the lowest state is X 3Σ⁻, followed, in terms of increasing energy, by a 1Δg, b 1Σ⁺, A 3Σ⁺, and B 3Σ⁻ [28, 30].

3.3.2. Quantum Chemical Rules: Obeyed and Disobeyed

by O₂ Molecules and Atoms

Generally, according to Wigner’s spin selection rules [31], chemical reactions between triplet and singlet reactants are forbidden. So the transitions Σ→Δ, Δ→Σ, Σ→Π and Σ→Σ are forbidden. In practice, all other elements more or less obey these rules, except excited O₂ molecules. Transitions (and vice versa) of O₂(a' Δg→X 3Σg⁻), O₂(A 3Σ⁺→X 3Σg⁻) and O₂(c' 3Σg−→X 3Σg−) take place in atmosphere (see section 3.10.3).

Ground state oxygen molecules have the triplet electronic configuration whereas most natural compounds have the singlet multiplicity. The triplet multiplicity of O₂ (X 3Σg⁻) is the reason why most reactions of oxygen
molecules with organic substances do not, despite being exergonic, proceed spontaneously at room temperature. Instead, heating or a catalyst is needed [32]. The ground state O₂ molecule follows strictly this quantum chemical rule. It is of vital importance for life on Earth.

Due to excitation, O₂ molecules have extra energy, with the aid of which they are able to go against the quantum chemical rules of forbidden transitions/reactions for state O₂ molecule follows strictly this quantum chemical rule. It is of vital importance for life on Earth.

For ozone in the MLT layer than that which was observed in the 1980s chemical models predicted a lower density due to excitation, O₂ molecules have extra energy, with the aid of which they are able to go against the quantum chemical rules. Of course, transitions/reactions may be slower and yields may subsequently be smaller than in the allowed reactions. Nature has overcome this issue by the set-up of two ozone layers in the atmosphere. The outcome is that no harmful UV radiation is able to enter the troposphere through both of them. Only that part of UV radiation that helps our body to produce D-vitamin, and which is so vital for human life, is able to enter [33]. O₂ is called a life supporting and saving molecule for a reason.

In the 1980s chemical models predicted a lower density for ozone in the MLT layer than that which was observed [34]. One reason for this may have been that the quantum chemical rules of forbidden transitions/reactions for oxygen species were taken too literally in the chemical models of those times.

3.4. Importance of O₂ Vibrational Energies

Molecular vibrations are caused by thermal infrared wavelengths and molecular rotations are caused by microwave and far-IR wavelengths. The vibrational energy levels E(υ) within one electronic state (Eel) are schematically described by the Morse potential energy curve [35]. Vibrational energies are typically much smaller than electronic energies, and rotational energies E(J) are much smaller than vibrational energies.

However, the E(υ) levels of O₂ molecules contain relatively much energy. O₂(X³Σg⁻) possibly has up to 29 vibrationally excited states [36]. Slangen and Cosby [37] have determined the vibrational energy content of 14 of them. For example, at the level of υ = 1 the energy content of O₂(X³Σg⁻) is 1553 cm⁻¹ (18.6 kJ/mol), at the level of υ = 8 it is 11803 cm⁻¹ (141.2 kJ/mol) and at the level of υ = 14 it is 19693 cm⁻¹ (235.6 kJ/mol).

Energy of O₂ (a¹Δg, υ = 0) is 94.3 kJ/mol and that of O₂ (a¹Δg, υ = 1) is 112.3 kJ/mol. So vibrational energy is as much as 19% from the Eel of the O₂ (a¹Δg, υ = 1) molecule. The energy of O₂ (a¹Δg, υ = 10) is 19695 cm⁻¹ or 235.7 kJ/mol [37]. It exceeds 2.5 times the Eel of the υ = 0 molecule. So in case of O₂ molecules, it is necessary to deal with vibrational energies as well. For O₂ molecules, E (J) is much smaller than E (υ). Rotational energies are important in advanced research work regarding, for example, the identification of the emissions from differently excited oxygen molecules [38].

3.5. Electronically Excited States of O₂ Molecules

Paul Krupenie [39] lists several electronically excited states of neutral oxygen molecules and O₂⁺ ions. The electronic configurations and the basic energy (υ=0 levels) of 20 of them are presented in Table 1.

Table 1. Some excited O₂ molecules, their formation of enthalpies and dissociation energies [39]

| Excited state | Formation enthalpy cm⁻¹/eV | Dissociation energy D^0 cm⁻¹/eV |
|---------------|-----------------------------|-------------------------------|
| X³Σg⁻         | 0                           | 41260/5.12                    |
| a¹Δg          | 7882/0.98                   | 33378/4.14                    |
| b¹Σ⁺          | 13121/1.63                  | 28139/3.49                    |
| c¹Σ⁻          | 32664/4.05                  | 8596/1.07                     |
| A¹ÎΔg         | 34319/4.26                  | 6941/0.86                     |
| A¹Σ⁺          | 35007/4.34                  | 6253/0.77                     |
| B¹Σ⁺          | 49358/6.12                  | 7770/0.96                     |
| B¹Σ⁺          | 75450/9.35                  |                               |

Cooper et al. [40] have determined the wavelengths and enthalpies for transitions/emissions of several excited O₂ molecules to the ground state.

3.5.1. The Excited States of O₂(X³Σg⁻, a¹Δg and b¹Σ⁺)

The electron configuration which gives rise to the ground state for O₂(X³Σg⁻), and also the first excited states of O₂(a¹Δg) and O₂(b¹Σ⁺), is [28, 39]:

\[
(1σ_g^1s)^2 (1σ_u^1s)^2 (2σ_g^2p)^2 (2σ_u^2p)^2 (3σ_u^2p)^2 < (1σ_u^2p)^2 < (1π_u^2p)^2
\]

Or, expressed as the number of electrons: 2, 2, 2, 2, 2, 4 and 2, 1 (σ_g^1s)^2 and (1σ_u^1s)^2 are inner (and virtually atomic) orbitals, usually marked with the letter K. Antibonding orbitals are marked with an asterisk while the others are bonding orbitals. (2σ_g^2p)^2 and (2σ_u^2p)^2 are actually modified (2σ_u)^2-orbitals. An O₂ molecule has 12 valence electrons, eight of which are bonding MOs and four are antibonding MOs.

All other orbitals no longer resemble their separated (atomic form). O₂(X³Σg⁻, a¹Δg and b¹Σ⁺) molecules differ only in the spin and occupancy of oxygen’s two antibonding (1π_u)^2 MOs, which are degenerate (equal in energy). The energies of O₂(a¹Δg) and b¹Σ⁺) are 94.3 kJ/mol and 157 kJ/mol respectively.

O₂(a¹Δg, υ = 0 and 1) are photoexcited by IR photons of 1060 nm and 1269 nm [41]. The photoexcitation of
0₂(b′Σ_u⁻) takes place with visible and IR photons at the A-, B- and Y bands of O₂ (see Figure 3) in which the photon energy is more than 157 kJ/mol.

3.5.2. The Excited States of O₂ (c¹Σ_u⁻, A¹Δ_u, A³Σ_u⁺ and B³Σ_u⁻)

These molecules are produced when one electron is moved from bonding (1π_g)² MO to antibonding (1π_u)² MO. The number of electrons on the last three MOs becomes (3σ_g)²(2, (1π_u)³=3 (39)]. Compared to O₂ (A³Σ_u⁺), the excitation of O₂ (B³Σ_u⁻) requires 171.6 kJ/mol more energy.

Copeland et al. [22] photoexcited O₂(X³Σ_g⁻) with a laser beam between the wavelengths 242 to 252 nm to O₂ (A³Σ_u⁺) at vibrational levels of v = 6 to the highest level of v = 11. This proves that an important formation mechanism of the excited states of O₂ (A³Σ_u⁺), the O₂ (B³Σ_u⁻) and 25–50 seconds for O₂ (a¹Σ_u⁻) are formed by direct photoexcitation: 1) UV photons of λ > 211–566 nm. They report measurements of 17 3Σ_u⁻, v = 0, 1 and 2). However, there is no explanation for why these states relax by radiating, but when v increases > 2, the relaxation would be radiationless.

Predissociation does not affect the energy content of O₂ (B³Σ_u⁻). Within its 7700 cm⁻¹ (92.1 kJ/mol) dissociation energy, there is room for 19 different vibronic states of O₂ (B³Σ_u⁻) molecules [40]. So in the supposed radiationless predissociation, the O₂ (B³Σ_u⁻, v > 2) molecules lose no energy.

When O₂(X³Σ_g⁻, v = 0) absorbs a UV photon in the Schumann–Runge bands, its internuclear distance (R) increases from the value 1.2075358 Å to the value 1.6042799 Å for O₂ (B³Σ_u⁻) [39]. The double bond between the oxygen atoms in the O₂ (B³Σ_u⁻) molecule probably becomes weaker. Considering the great E_u and E (υ, v), the O₂ (B³Σ_u⁻, v > 2) molecules would become more exposed towards all possible collision reactions, supposing its radiative lifetime is sufficiently long.

Sick et al. [46] have actually demonstrated that O₂ (B³Σ_u⁻) can undergo collisional reactions. It may be that due to the predissociation phenomenon (lack of rotation) the molecule internally strengthens against radiative relaxation. This would allow the O₂ (B³Σ_u⁻, v > 2) molecules to be destroyed in collisions. In this case, no emission spectra would be generated.

An increase of internuclear distance, predissociation via tunneling effect and the fact that transition O₂ (B³Σ_u⁻, v > 2) molecules lose no energy. The double bond of the oxygen atoms in the O₂ (B³Σ_u⁻) molecule probably becomes weaker. Considering the great E_u and E (υ, v), the O₂ (B³Σ_u⁻, v > 2) molecules would become more exposed towards all possible collision reactions, supposing its radiative lifetime is sufficiently long.

3.5.3. The Excited States of O₂ (B³Σ_u⁻, a¹Σ_u⁺, 1Δ_u, 2Σ_u⁺, 1Π_u and 3Π_u)

The molecules are formed when a electron is moved from (2σ_g)² MO to (3σ_u)² MO. The number of electrons on the last four MOs becomes 2σ_g=1, (1π_u)³=4, (1π_g)²=2 and 3σ_u=1. In 1Δ_u and 3Π_u, one electron is moved from 1π_u MO to 3σ_u MO [39].

The excited oxygen molecules in this category belong to the energy range of 9.35 to 11.07 eV (see Table 1). They are formed by direct photoexcitation: 1) UV photons of λ < 131 nm produce O₂ (B³Σ_u⁻) and O₂ (a¹Σ_u⁺) and 2) UV photons of λ < 112 produce 2Σ_u⁺, 1Δ_u and 1Π_u.

Englund et al. [47] have made a comprehensive vibronic assignment of the O₂(3Π_u) states in the region 104 to 120 nm. They report measurements of 17 3Σ_u⁻ bands in the 85,800–93,000 cm⁻¹ (10.64–11.53 eV) region of the photoabsorption spectra of ¹⁶O₂ and ¹⁸O₂. Lewis and Gibson [38] have measured the wavelength of the transition O₂ (B³Σ_u⁻, v = 3) → O₂ (X³Σ_g⁻) to be 123.4 nm.
When dissociated, these O₂ molecules are energetically able to produce O (¹S) and O (¹D) atoms.

3.5.4. Ionic O₂⁺ Molecules

The first of the O₂⁺ ions (with the lowest energy of 12.07 eV), is X²Πg which has lost one 1π⁺ electron. The O₂(a³Πg) and O₂(A³Πu) molecules have lost one 1π⁺ electron. The O₂(b³Σg⁺), O₂(C³Δg) and O₂(B³Σu⁺) molecules have lost one 3σg electron. The O₂(c³Σu⁻) molecule has lost one 2πa electron. [39] The energy range of ionic O₂ molecules is between 12.07-24.56 eV, which is equal to the wavelength range 102.7 nm-50.5 nm.

The O₂⁺ ion has 11 valence electrons, of which eight are bonding electrons and three antibonding electrons. Thus, its bond order is ½(8-3) = 2½. The experimental dissociation energy and bond length of the O₂ (X²Πg) ion are 643 kJ/mol and 1.123 Å, respectively [39]. In terms of dissociation energy, losing one electron makes the O₂(X²Πg) molecule “stronger” than O₂(X³Σg⁻). However, the ionic O₂ molecules have a “weak” side – absorption of an electron changes their bond order from 2.5 to 2.0. So a single electron with its 32.8 kJ/mol energy [48] can dissociate ionic O₂(X²Πg) molecule in the dissociative recombination reaction (DR) [49]:

\[ O₂(X²Πg) + e → O(¹D) + O(¹S); \Delta H^0 = -105.5 \text{ kJ/mol} \] (4)

The lifetime of O₂⁺ ions in the night depends mostly on the available electron concentration. O₂⁺ ion concentrations peak at 110 km [23]. It is quite possible that some O₂(X²Πg) molecules are produced in the MLT layer and that at 110 km, more O₂(X²Πg) and other O₂⁺ ions enter into the MLT layer via diffusion. During the night the reaction (4) may have importance, as a continuous additional source of O(¹S) atoms.

3.6. Reactions of Vibrationally and Electronically Excited O₂ Molecules

3.6.1. Vibrationally Excited O₂(X³Σg⁻)

Kinetic studies by Rogaski et al. [50] and Miller et al. [51] suggest that O₂(X³Σg⁻, v ≥ 26) is formed in ozone photolysis via the triplet channel in the ozone photodissociation in the Hartley band with UV photons of λ < 233 nm:

\[ O₃ + hν → O₂(X³Σg⁻, v ≥ 26) + O(¹P); \Delta H^0 = -2 \text{ kJ/mol} \] (5)

These vibrationally excited ground state oxygen molecules have energies ≥ 4.2 eV (405 kJ/mol). It is possible that the O₂(X³Σg⁻, v ≥ 26) molecules enter vibrational-to-electronic (v→E) energy transfer reactions when colliding with a ground state O₂ molecule [50]:

\[ O₂(X³Σg⁻, v ≥ 26) + O₂(X³Σg⁻) → O₂(c³Σu⁺) + O₂(X³Σg⁻); \Delta H^0 = -14.2 \text{ kJ/mol} \] (6)

A theoretical problem here is that this reaction is forbidden by quantum chemical rules. However, it is known that the Herzberg II UV nightglow is due to the relaxation of O₂(c³Σu⁺ → X³Σg⁻). The reaction (6) would be an important night-time source of O₂(c³Σu⁻) for the Hertzberg II nightglow.

A second possible channel is that O₃ and O are formed [50, 51]:

\[ O₂(X³Σg⁻, v ≥ 26) + O₂(X³Σg⁻) → O₃ + O(¹P); \Delta H^0 = -13.1 \text{ kJ/mol} \] (7)

In this channel the result would be production of two unpaired oxygen species. This would be important for maintaining both the daytime and night-time ozone and oxygen atom concentration.

3.6.2. Electronically excited O₂(B³Σu⁻)

The Schumann–Runge bands have been observed in hydrogen, carbon monoxide and ammonia flames. This suggests that there may be a reaction common to all flames such as [52]:

\[ O(¹P) + O(¹D) + M → O₂(B³Σu⁻) + M; \Delta H^0 = -97.6 \text{ kJ/mol} \] (8)

The prerequisite of reaction (8) is the participation of O (¹D) atoms, generated either via the heat of flame at the surface or via quenching of O (¹S) or via various reactions in the MLT layer.

Copeland et al. [22] have shown that O atoms are generated efficiently in the collision of vibrationally excited O₂(A³Σu⁺, v ≥ 9) with (vibrationally excited) ground state O₂ (reaction 9). Collision reactions of O₂ (B³Σu⁻) with ground state O₂ molecules (reaction 10) is also allowed by quantum chemical rules, and is so even more likely. If O₂(X³Σg⁻) is vibrationally excited (v ≥ 26) it is energetically possible that O (¹S) is formed (reaction 11):

\[ O₂(A³Σu⁺, v ≥ 9) + O₂(X³Σg⁻, v ≥ 1) → O₂(X³Σg⁻) + 2O(¹P); \Delta H^0 = -5.7 \text{ kJ/mol} \] (9)

\[ O₂(B³Σu⁻) + O₂(X³Σg⁻) → O₂(X³Σg⁻) + 2O(¹P); \Delta H^0 = -92.1 \text{ kJ/mol} \] (10)

\[ O₂(B³Σu⁻) + O₂(X³Σg⁻, v ≥ 26) → O(¹S) + O(¹P) + O₂(X³Σg⁻); \Delta H^0 = -93.3 \text{ kJ/mol} \] (11)

Reaction (8) contributes to the amount of O₂ (B³Σu⁻) molecules both in the daytime and in night-time in the MLT layer. The reaction (11) may have importance to the night-time O (¹S) budget. Reactions (9) to (11) are important for the stability of oxygen atom concentration during the night.

3.6.3. Electronically Excited O₂(a³Σ⁺, a¹Σ⁺, a³Σ⁺, ¹Δu, ¹Πu)

It is feasible that the collision of these high energy O₂ molecules with ozone or O₂ (X³Σg⁻) produce O (¹S) atoms (reactions 12 and 13).

\[ O₂(¹Δu) + O₃ → O(¹S) + 2O₂(X³Σg⁻); \Delta H^0 = -546.3 \text{ kJ/mol} \] (13)

\[ O₂(¹Δu) + O₂(X³Σg⁻) → O(¹S) + O(¹P) + O₂(X³Σg⁻); \Delta H^0 = -154.4 \text{ kJ/mol} \] (12)
3.7. Ozone Photolysis and Reactions Thereof

The ozone molecule is a kind of primus motor in many of the reactions in the MLT layer. Its readiness to react is enhanced by the participation of excited ozone molecules [53]. Wayne [54] has calculated limiting wavelengths for the formation of energetically different combinations of O + O2 in ozone photolysis (see Table 2).

When ozone is photolyzed by photons of visible light, \( \lambda < 611\text{nm} \), are produced \( \text{O}_2(a^3\Delta_g^+) + \text{O}(^3\text{P}) \). With UV photons \( \lambda < 310 \text{nm} \), the products are \( \text{O}_2(b^3\Sigma_g^+) + \text{O}(^3\text{D}) \), and when of \( \lambda < 266 \text{nm} \), the products are \( \text{O}_2(b^3\Sigma_g^+) + \text{O}(^3\text{D}) \). Similarly at UV wavelengths \( \lambda < 230 \text{nm} \), and \( \lambda < 170 \text{nm} \), \( \text{O}_2(A^3\Sigma_u^+) + \text{O}(^3\text{P}) \) and \( \text{O}_2(B^3\Sigma_u^+) + \text{O}(^3\text{P}) \) are formed, respectively. When these excited molecules are dissociated by UV photons \( \lambda < 242 \text{nm} \) two O atoms are formed. The net effect is the production of three O atoms from one ozone molecule.

### Table 2. Long wavelength limits (nm) for the production of various electronic states of \( \text{O}_2 \) and \( \text{O} \) in the primary step of ozone photolysis [54]

| O atom | Electronic state of \( \text{O}_2 \) molecules |
|--------|----------------------------------|
| \(^3\text{P}\) | \(1180\) |
| \(^1\text{D}\) | \(411\) |
| \(^1\text{S}\) | \(234\) |

Tu and Nee [26] suggested that when a UV photon at the wavelength \( \lambda = 121.6 \text{nm} \) dissociates ozone, the result is direct formation of three oxygen atoms from one ozone molecule:

\[
\text{O}_3 + h\nu (\lambda = 121.6 \text{nm}) \rightarrow \text{O}_2^* + \text{O} \quad (14)
\]

\[
\text{O}_2^* + \text{O} \rightarrow \text{O} + \text{O} + \text{O} \quad (15)
\]

The \( \Delta H^0 \) of the starting products (ozone and energy of UV photon) of the reaction (14) is \(142.7 \text{kJ/mol} \) + 983.8 kJ/mol = 1126.5 kJ/mol. When the O atom is \( \text{O}(^3\text{P}) \), then the energy content of the \( \text{O}_2^* \) molecule is 877.3 kJ/mol. It exceeds the dissociation energy of \( \text{O}_2(B^3\Sigma_u^+) \) (682.6 kJ/mol) and do not reach the formation enthalpy of \( \text{O}_2(b^3\Sigma_g^+) \) (902.2 kJ/mol). At the wavelength of \( \lambda = 118.7 \text{nm} \) the energy content of the \( \text{O}_2^* \) is 901.3 kJ/mol which is just below the formation enthalpy of \( \text{O}_2(b^3\Sigma_g^+) \). So between the wavelengths of 118.6 to 121.6 nm there is a window in which three oxygen atoms are formed from one ozone molecule.

3.8. Reactions of \( \text{O}(^1\text{D}) \) with \( \text{O}_2(X^3\Sigma_g^-) \)

In the reaction \( \text{O}(^1\text{D}) + \text{O}_2 \) is formed two \( \text{O}_2 \) molecules. At least one of these \( \text{O}_2 \) molecules produced is excited (reaction 16). This excited \( \text{O}_2 \) molecule is capable of decomposing additional \( \text{O}_3 \), yielding another \( \text{O}(^1\text{D}) \) atom (reaction 17) [55, 56]:

\[
\text{O}(^1\text{D}) + \text{O}_3 \rightarrow \text{O}_2(A^3\Sigma_u^+) + \text{O}_2(X^3\Sigma_g^-); \quad \Delta H^0 = -62.8 \text{kJ/mol} 
\quad (16)
\]

\[
\text{O}_2(A^3\Sigma_u^+) + \text{O}_3 \rightarrow \text{O}(^1\text{D}) + 2\text{O}_2(X^3\Sigma_g^-); \quad \Delta H^0 = -122.6 \text{kJ/mol} 
\quad (17)
\]

Reactions (16) and (17) constitute a chain reaction with reported chain lengths as large as 16 [55]. This allows \( \text{O}(^1\text{D}) \) to fuel the dynamics during the night. The reaction (16) is an important night-time source of \( \text{O}_2(A^3\Sigma_u^+) \) as well. It is 162.8 kJ/mol exothermic, so one mole of \( \text{O}_2(a^1\Delta_g) \) or \( \text{O}_2(b^3\Sigma_g^+) \) could also be produced.

The three-body reaction (18) produces excited \( \text{O}_2(b^3\Sigma_u^+) \) molecules and \( \text{O}(^3\text{P}) \) atoms. The energy transfer via \( \text{O}(^3\text{P}) \) to \( \text{O}_2 \) is an important source of \( \text{O}_2(b^3\Sigma_u^+) \) at vibrational levels \( \nu = 0 \) and 1 in Earth’s atmosphere [57].

\[
\text{O}(^1\text{D}) + \text{O}_2(X^3\Sigma_g^-) + \text{M} \rightarrow \text{O}_2(b^3\Sigma_u^+) + \text{O}(^3\text{P}) + \text{M} 
\quad \Delta H^0 = -32.7 \text{kJ/mol} 
\quad (18)
\]

3.9. Formation of \( \text{O}(^1\text{S}) \) Atom

In the photolysis of ozone UV photons at wavelengths 230 < \( \lambda < 234 \text{ nm} \) produce \( \text{O}(^1\text{S}) \) and \( \text{O}_2(X^3\Sigma_g^-) \) (see Table 2). This reaction is feasible at night as well. Krasnopolsky and Levfèvre [58] suggest that reaction (19) would be important in producing \( \text{O}(^1\text{S}) \).

\[
\text{O}(^3\text{P}) + \text{O}_2(A^3\Sigma_u^+ \nu \geq 6) \rightarrow \text{O}(^1\text{S}) + \text{O}_2(X^3\Sigma_g^-); 
\quad \Delta H^0 = -64.4 \text{kJ/mol} 
\quad (19)
\]

Considering the higher energy, even the \( \text{O}_2(B^3\Sigma_u^-, \nu = 0) \), would enter the same reaction, and produce a vibrationally excited ground state \( \text{O}_2 \) molecule as well (reaction 20).

\[
\text{O}(^3\text{P}) + \text{O}_2(B^3\Sigma_u^-, \nu = 0) \rightarrow \text{O}(^1\text{S}) + \text{O}_2(X^3\Sigma_g^-, \nu \leq 6); 
\quad \Delta H^0 = -79.2 \text{kJ/mol} 
\quad (20)
\]

The highly excited molecules \( \text{O}_2(A^3\Sigma_u^+, \nu = 6) \) contain so much energy that their reaction with \( \text{O}(^1\text{P}) \) atoms would produce \( \text{O}(^1\text{S}) \) and \( \text{O}_2(B^3\Sigma_u^-, \nu > 5) \).

\[
\text{O}(^3\text{P}) + \text{O}_2(A^3\Sigma_u^+, \nu = 6) \rightarrow \text{O}(^1\text{S}) + \text{O}_2(B^3\Sigma_u^-, \nu > 5); 
\quad \Delta H^0 = -12.6 \text{kJ/mol} 
\quad (21)
\]

By reacting with \( \text{O}(^3\text{P}) \), the \( \text{O}_2(B^3\Sigma_u^-, \nu > 5) \) produce \( \text{O}(^1\text{S}) \) and \( \text{O}_2(X^3\Sigma_g^-, \nu \leq 9) \) (see reaction 20). This cascading process (reactions 20 and 21) emphasizes the importance of highly excited neutral \( \text{O}_2 \) molecules in producing \( \text{O}(^1\text{S}) \) and \( \text{O}_2(B^3\Sigma_u^-) \) especially during the night-time in the MLT layer.

3.10. Nightglows as Evidences of the Night-time Dynamics of \( \text{O} \) and \( \text{O}_2 \) Species in the MLT Layer

Nightglows are the emission of UV, visible or IR light by excited atoms and molecules when they are relaxing to the lower excited states or ground states in the upper atmosphere. Nightglows in the MLT layer last throughout the night with only a little falling off [59]. Understanding the nightglow dynamics is vital to understanding the dynamics of oxygen species during the night in the secondary ozone zone.
3.10.1. IR and Visible Nightglows

These nightglows are due to the relaxing of excited molecules $O_2(a'\Delta_u)$ and $b'\Sigma_g^+$. The transitions $O_2(a'\Delta_u, v=0, 1) \rightarrow O_2(X^2\Sigma_g^-)$ are called the IR atmospheric system. IR photons are emitted at wavelengths at 1060 nm and 1269 nm. The transition $O_2(b'\Sigma_g^+, v = 0$ and 1) $\rightarrow O_2(X^2\Sigma_g^-)$ is called the Atmospheric system. The emitting wavelengths are 761.9 nm and 864.5 nm [60].

3.10.2. Nightglows Due to the Oxygen Green Line and Red Line Emissions

The green nightglow of the $O(\text{I}S\rightarrow\text{I}D)$ transition is at a wavelength of 557.7 nm and is located at two elevations. The first peaks at an altitude between 300 and 160 km in the thermosphere [61] and the second one in the MLT layer at an altitude of 94 km [62].

The transition of $O(\text{I}D \rightarrow O(\text{I}P)$ produces the red nightglow at 630 nm and at 636.4 nm at an altitude of 300 to 160 km. Its intensity is 1000 R at the close of the day but diminishes to 50 R by midnight [61]. The green line nightglow at the 170 km diminishes soon after sunset from 1000 R to 300 R and by the midnight more [63]. (1 rayleigh equals $10^8$ photons per cm$^2$ column per second [61]).

In the altitude range of 300 to 160 km the most important sources of $O(\text{I}S)$ are $O_2^+$ ions are generated during the daytime, and are then exhausted by midnight by electrons and N atoms. The formation of $O_2^+$ ions by the EUV from geocorona, the formation of $O(\text{I}S)$ by the three-body reaction of $O+O+O$ atoms and the contribution from the galactic background [63] continues to provide so much $O(\text{I}S)$ that the falling-off stops at the 50 R level.

In the MLT layer $O(\text{I}S)$ is steadily formed throughout the night, so the intensity of the green line nightglow emission stays at 337 R [64].

3.10.3. Ultraviolet Nightglows

Figure 4 presents the entire UV nightglow radiation of Earth [65]. It is generated

1) by H atoms (102.6 and 121.6 nm) by and by O atoms from geocorona

2) from relaxation of excited NO molecules, the wavelengths of $\delta$ band are 190–240 nm and those of the $\gamma$ band are 225–270 nm.

3) via Herzberg systems [39]:
   - Herzberg I system $O_2(A^3\Sigma_u^+ \rightarrow X^3\Sigma_g^-)$, emitting wavelengths are 243–488 nm.
   - Herzberg II system $O_2(c^3\Sigma_u^- \rightarrow X^3\Sigma_g^-)$, emitting wavelengths are 254–271.5 nm and 449–479 nm
   - Herzberg III system $O_2(A^1\Delta_u \rightarrow X^3\Sigma_g^-)$, emitting wavelengths are 257–263 nm.

4) via Chamberlain system $O_2(A^1\Delta_u \rightarrow a'\Delta_g)$, emitting wavelengths are 370–438 nm [39].

5) via the Schumann–Runge system $O_2(B^1\Sigma_u^- \rightarrow X^3\Sigma_g^-)$, emitting wavelengths are 211–566 nm [39, 45].

Figure 4. The entire composite UV nightglow [65]

Nightglows have been found on Venus as well [67, 68]. The UV nightglows due to $O_2(c^3\Sigma_u^- \rightarrow X^3\Sigma_g^-)$ and $O_2(A^3\Sigma_u^+ \rightarrow a'\Delta_g)$ emissions have a greater intensity in rayleighs on Venus than they do on Earth [68]. The ozone layer on Venus at 100 kilometres above the planet’s surface is considerably less dense compared to the MLT ozone layer at night [69].

So a likely conclusion is that on Earth during the night, the MLT ozone layer absorbs part of the UV photons emitted during nightglow. This absorbed UV radiation may have an important auxiliary role in the night-time dynamics of excited oxygen molecules and ozone.

3.10.4. Nightglow Dynamics: Economic Use of Available Energy

Energy reserved in the MLT layer during the daytime in the excited $O_2$ molecules and their replenishment by diffusion from above may not be enough to allow the nightglows to run throughout the night with steady intensity. Additional energy is provided by EUV from geocorona (producing highly excited $O_2$ molecules), UV from the nightglows of NO $\delta$ (producing $O_2(B^3\Sigma_u^-)$) and NO $\gamma$ bands (producing $O_2(A^3\Sigma_u^+)$), and UV generated by the nightglows of $O_2(A^3\Sigma_u^+, c^1\Sigma_g^-$ and $A^1\Delta_u)$ molecules (causing photolysis of ozone).

Visible photons from the oxygen green line produce $O_2(a'\Delta_g)$ via the photolysis of ozone Quenching of $O(\text{I}S)$ provides $O(\text{I}D)$ which reactions with ozone produce $O_2(A^3\Sigma_u^+, c^1\Sigma_g^-$ and $A^1\Delta_u)$ molecules.

In the MLT ozone layer the use of energy is highly efficient. Energy-rich compounds are upgraded from “waste” energy, and with no cost to the “middle man of ozone”, as these reactions provide also O atoms, which produce ozone.
4. Conclusions

During the daytime in the MLT layer the EUV photons (102.8 to 124 nm) are photoexciting O2 molecules up to 11.07 eV level. O2 (B3Σu+, ν = 0–19) molecules are formed via photoexcitation in the Schumann–Runge bands. Dissociation of O2 molecules in the Schumann–Runge and Herzberg continuums produces O atoms. In the photolysis of ozone there is a window between the wavelengths of 118.7 to 121.6 nm in which these oxygen atoms are formed from one ozone molecule. Vibrationally excited O2(X3Σg−, ν ≥ 26) molecules are formed via ozone photolysis. Both O2(B3Σu+, ν < 26) and (O2(X3Σg−, ν = 0) to produce O and O3. The O atom concentration reaches its atmospheric layer.

Due to the abundance of O atoms, a small but significant ozone concentration (as compared to the regions above or immediately below) remains during the daytime, even though ozone is then rapidly destroyed by photolysis.

During the night, solar UV radiation is radically reduced. Ozone, being the “weak” oxygen species, gains the most. The number of O atoms shows no diurnal variation in the MLT layer. This leads to a ten-fold increase of ozone concentration over the course of the night.

The MLT ozone zone coincides with the ionospheric D-ring, so there is a small concentration of electrons at night as well.

At an altitude of 110 km, O(S) ions enter the MLT layer via diffusion. Highly energetic neutral oxygen molecules (9.35 to 11.07 eV) generated during the daytime in the MLT layer are also replenished by diffusion from above. O1(S) atoms are produced via the BR reaction, via reactions of excited oxygen molecules (Ee ≥ 9.35 eV) with O2, O3, or O, and via reaction of O2(B3Σu+) + O(P) These reactions produce enough O1(S) atoms to keep the green line nightglow in the MLT layer running at a rather steady state throughout the night.

O(1D) quenched from O1(S), and produced in collision reactions of excited oxygen molecules and ozone is important for keeping the overall O atom concentration at a highly reactive level during the night as well.

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