Current state of modeling the photochemistry of Titan's mutually dependent atmosphere and ionosphere

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[1] In the context of recent observations, microphysical models, and laboratory data, a photochemical model of Titan's atmosphere, including updated chemistry focusing on rate coefficients and cross sections measured under appropriate conditions, has been developed to increase understanding of these processes and improve upon previous Titan photochemical models. The model employs a two-stream discrete ordinates method to characterize the transfer of solar radiation, and the effects of electron-impact, cosmic-ray deposition, and aerosol opacities from fractal and Mie particles are analyzed. Sensitivity studies demonstrate that an eddy diffusion profile with a homopause level of 850 km and a methane stratospheric mole fraction of 2.2% provides the best fit of stratospheric and upper atmosphere observations and an improved fit over previous Titan photochemical models. Lack of fits for C₃H₈, HC₃N, and possibly C₂H₃CN can be resolved with adjustments in aerosol opacity. The model presents a benzene profile consistent with its detection in Titan's stratosphere [Coustenis et al., 2003], which may play an important role in the formation of Titan hazes. An electron peak concentration of 4200 cm^{-3} is calculated, which exceeds observations by 20%, considerably lower than previous ionosphere models. With adjustments in aerosol opacities and surface fluxes the model illustrates that reasonable fits to existing observations are possible with a single eddy diffusion profile, contrary to the conclusions of previous Titan models. These results will aid in the receipt and interpretation of data from Cassini-Huygens, which will arrive at Titan in 2004 and deploy a probe into Titan's atmosphere in January 2005. INDEX TERMS: 5405 Planetology: Solid Surface Planets: Atmospheres-composition and chemistry; 6280 Planetology: Solar System Objects: Saturnian satellites; 5435 Planetology: Solid Surface Planets: Ionospheres (2459); 0335 Atmospheric Composition and Structure: Ion chemistry of the atmosphere (2419, 2427); 0305 Atmospheric Composition and Structure: Aerosols and particles (0345, 4801); KEYWORDS: Titan, photochemistry, composition, haze, ion chemistry, planetary atmospheres

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1. Introduction

[2] Titan, the only satellite in the Solar System with an extensive atmosphere, has been an object of considerable scrutiny for some time. Titan has a largely N₂ atmosphere (90-98%) [*Lindal et al.*, 1983; *Broadfoot et al.*, 1981] with methane (2-6%) [*Samuelson et al.*, 1997; *Hanel et al.*, 1981] being the most abundant minor constituent. Methane is dissociated through the transfer of UV solar radiation, photoelectrons, and magnetospheric electrons, which produce highly reactive radical species. These species along with the nitrogen atoms produced through N₂ dissociation react to form a bevy of hydrocarbons and nitriles that

characterize Titan's atmosphere. The presence of CO along with the likely influx of micrometeorites into Titan's atmosphere contributes to the collection of oxygen-bearing species that exist in Titan's atmosphere, as well.

[3] A brief historical perspective of relevant observations and previous modeling studies is given first in order to place the present work in proper perspective. The presence of most of the species known to exist in Titan's atmosphere was revealed by the Voyager flybys, which unveiled an object covered with orange-brown hazes. Beneath the haze region the surface was found to have a temperature of 94 K at a pressure of 1.5 bars and a radius of 2575 km [*Lindal et al.*, 1983]. Through these radio occultation measurements, *Lindal et al.* [1983] were able to infer the temperature profile up to 200 km, which included a tropopause region near 42 km with a temperature of 71 K and a stratospheric

HCO

H₂CO

formyl radical

formaldehyde

Table 1. List of Neutral and Ionic Compounds Used in the Model

 Table 1. (continued)

	Neutrals	Ions
Н	atomic hydrogen	H^+
H_2	molecular hydrogen	H_2^+
C	atomic carbon	H_3^+
CH	methylidyne	H_2O^+
CH ₂	excited-state methylene	H_3O^+
CH ₂	ground-state methylene	HCO ⁺
CH ₃	methyl radical	CH_3^+
CH ₄	methane	CH_4^+
22	molecular carbon	CH_5^+
C ₂ H	ethynyl radical	$C_2H_2^+$
C_2H_2	acetylene	$C_2H_3^+$
C_2H_3	vinyl radical	$C_2H_4^+$
C_2H_4	ethylene	C_2H_5
C_2H_5	ethyl radical	$C_2H_6^+$
$_{2}H_{6}$	ethane	c-C ₃ H ₃ ⁺
C ₃ H ₂ C ₃ H ₃	propadienylidene	C_3H_5 $C_2H_7^+$
лага Н ₃ С ₂ Н	propargyl radical methylacetylene	$\begin{array}{c} \mathrm{C_4H_2^+}\\ \mathrm{C_4H_3^+} \end{array}$
H_3C_2H H_2CCH_2	allene	$C_{6}H_{7}^{+}$
H2CCH2	allyl radical	N^+
$_{3}H_{6}$	propylene	N_2^+
3116 3H7	isopropyl radical	NH ⁺
³¹¹⁷ ³ H ₈	propane	N_2H^+
₄ H	r-spane	NH_2^+
$_{4}H_{2}$	diacetylene	NH ₃
$_{4}H_{2}^{4}$	excited-state diacetylene	NH4
4H ₂ 4H ₃		CN ⁺
4H4	vinylacetylene	HCN ⁺
4H5	1-butyn-3-yl radical	H_2CN^+
$_{4}H_{6}$	1,3-butadiene	$\tilde{C_xH_v^+}$
$_{4}H_{8}$	1-butene	$C_x H_y N_z^+$
$_{4}H_{10}$	n-butane	$C_x H_v O_z^+$
₆ H		N_x^+
C_6H_2	triacetylene	
C_6H_4	benzyne	
$-C_6H_4$	linear-C ₆ H ₄	
₆ H ₅	phenyl radical	
$-C_6H_5$	linear-C ₆ H ₅	
C_6H_6	benzene	
$-C_6H_6$	linear-C ₆ H ₆	
C_6H_7	cyclized-C ₆ H ₇	
$-C_6H_7$	linear-C ₆ H ₇	
$_{8}H_{2}$	tetraacetylene	
4s ~	ground-state atomic nitrogen	
2d	excited-state atomic nitrogen	
2	molecular nitrogen	
Н	imidogen	
H ₂	amino radical	
H ₃	ammonia	
₂ H ₂	diimide	
₂ H ₃	hydrazinyl radical	
$_{2}H_{4}$	hydrazine	
N	cyano radical	
CN	hydrogen cyanide	
2CN	methylene-amidogen radical	
HCN H CN	ovenomethyl radical	
H ₂ CN	cyanomethyl radical	
H ₃ CN	acetonitrile	
2N2 CaNa	cyanogen	
C ₂ N ₂ ₃ N	cyanoethynyl radical	
C ₃ N	cyanoacetylene	
$L_{2}C_{3}N$	cyanovinyl radical	
	acrylonitrile	
₂ H ₃ CN	dicyanoacetylene	
4N2 3p	ground-state atomic oxygen	
1d	excited-state atomic oxygen	
,)H	hydroxyl radical	
20	water	
0	carbon monoxide	
0 0 ₂	carbon dioxide	

	/	
	Neutrals	Ions
CH ₂ OH	hydroxymethyl radical	
CH ₃ O	methoxy radical	
CH ₃ OH	methanol	
CH ₂ CO	ketene	
CH ₃ CO	acetyl radical	
CH ₃ CHO	ethylene oxide	
C_2H_4O	oxirane	

rsion layer, increasing temperatures to 170 K at 200 km. her analysis of Voyager Infrared Spectrometer (IRIS) Voyager Ultraviolet Spectrometer (UVS) data yielded h of what is known about the distribution of constituents itan's atmosphere listed as shown in Table 1. However, ager did not uncover much about the middle and upper osphere. An exospheric temperature of 186 K was ined by Smith et al. [1982]. But, beyond some observaof CH₄ and C₂H₂ [Smith et al., 1982], little is known ut the vertical distribution of Titan's constituents ughout the atmosphere, largely due to the opacity vided by CH₄ and Titan's hazes. In fact, the hazes nselves remain largely a mystery. Voyager revealed an ue haze region consisting of a main haze layer around km and a detached haze layer at 300-350 km [Rages Pollack, 1983]. However, the precise source of this on and its composition is left to speculation.

In the upper atmosphere the absorption of solar EUV ation produces an ionosphere that interacts with Saturn's netosphere and the solar wind. This interaction is not stant as variations in solar wind pressure cause a comsion of the magnetosphere, resulting in Titan being ide of the magnetosphere part of the time. However, n Voyager 1's arrival, Titan was inside Saturn's magnehere, and Voyager observed a Venus-like interaction veen Titan and the magnetosphere with no bow shock ubauer et al., 1984; McNutt and Richardson, 1988], wing Titan to mainly interact with precipitating magneheric electrons along with photoelectrons produced in Titan's ionosphere. Early analysis revealed a variation lectron peak concentration with derived upper limits of 0 cm^{-3} at the evening terminator and 5000 cm⁻³ at the ning terminator [Lindal et al., 1983]. Bird et al. [1997] alyzed Voyager data to obtain an electron peak density $400 \pm 1100 \text{ cm}^{-3}$ at $1180 \pm 150 \text{ km}$ at the evening inator at a solar zenith angle near 90°.

The data presented by the Voyager flybys provided the etus for the construction of photochemical models that attempted to investigate how chemical species are ibuted in Titan's atmosphere. The first extensive photonical model after the Voyager flybys was developed by g et al. [1984], which made use of a large reaction set and analyses of Voyager data [Hanel et al., 1981; Maguire l., 1981; Kunde et al., 1981; Samuelson et al., 1983]. blanc et al. [1995] and Lara et al. [1996] took advantage nproved analysis of Voyager observations [Coustenis et 1989, 1991] and subsequent ground-based millimeter ervations [Tanguy et al., 1990; Hidayat et al., 1997] to different profiles of the eddy diffusion coefficient. blanc et al. [1995] included a Monte Carlo treatment he transfer of solar radiation in the atmosphere, while Lara et al. [1996] included physically based water ablation profiles to account for the oxygen source from micrometeorites. *Keller et al.* [1992] and *Fox and Yelle* [1997] used neutral photochemical models as a basis for the construction of ionosphere models. *Galand et al.* [1999] used the *Toublanc et al.* [1995] photochemical model to investigate diurnal effects on Titan's ionosphere, while *Banaszkiewicz et al.* [2000] combined the *Lara et al.* [1996] model with an investigation of charged particle chemistry to construct a coupled model of Titan's atmosphere and ionosphere.

[6] Despite these investigations many questions regarding Titan chemistry still remain unanswered. Photochemical models have been unable to simultaneously fit the vertical profiles of observed species. Furthermore, although many microphysical models have been developed [e.g., *Rannou et al.*, 1995], the chemical sources of Titan's haze layer are still quite unknown.

[7] The low temperatures which characterize outer planetary atmospheres, and Titan, in particular, have prompted many measurements of reaction rates and cross sections at low temperatures which were not included in previous Titan photochemical models. Microphysical modeling and analysis of Voyager photometric and polarimetric observations and albedo data have suggested the fractal nature of Titan haze whose scattering effects differ than those considered in previous one-dimensional Titan photochemical models (the two-dimensional model of Lebonnois et al. [2001] does consider fractal haze particles). With this in mind and in the context of preparation for the retrieval and interpretation of data from the upcoming Cassini-Huygens investigation of Titan, a one-dimensional steady state photochemical model has been constructed. This model computes the mole fraction for 80 neutrals and 33 ions, shown in Table 2, from the surface to 1600 km, while self-consistently calculating the total number density as a function of the mean molecular weight. The model incorporates a two-stream discrete ordinates scattering model, differing from the scattering treatment of previous Titan models, and includes dissociation via photoelectrons from 15-1000 eV, magnetospheric electrons, and cosmic rays, along with solar radiation from 50-3000 Å. A multicomponent treatment of molecular diffusion is included along with sensitivity studies of various eddy diffusion profiles. The sensitivity studies include the analysis of constituent profiles generated by 100 eddy diffusion profiles, increasing through the stratosphere in varying degrees, with homopause levels ranging from 600 km to 1150 km.

[8] This paper discusses the details of the photochemical model, including the equations and inputs used to calculate constituent densities in sections 2 and 3. A discussion of the important free parameter of eddy diffusion is provided, detailing the basis for the profiles used in the sensitivity studies regarding Titan's homopause level. The chemical mechanisms, which govern the distribution of hydrocarbon, oxygen, and nitrile neutrals as well as ions in Titan's atmosphere, are discussed in section 4. Sensitivity studies regarding eddy diffusion and aerosol opacity are examined in section 5, along with a discussion on constituent profiles. Finally, conclusions are detailed in section 6.

2. Model Description

[9] To examine the physical and chemical processes that shape the distribution of constituents with altitude in Titan's atmosphere, a one-dimensional photochemical model is developed. This photochemical model solves the steady state altitude-dependent continuity-diffusion equation in spherical coordinates

$$P_i - L_i = \frac{1}{r^2} \frac{\partial (r^2 \Phi_i)}{\partial r} \tag{1}$$

where P_i is the chemical production rate of species *i*, L_i is the chemical loss rate, the radius $r = (R_0 + z)$, where R_0 is the radius of Titan and *z* is the altitude, and Φ_i is the vertical flux, which can be expressed as

$$\Phi_{i}(z) = -D_{i} \left[n \frac{\partial \xi_{i}}{\partial z} + \xi_{i} \frac{\partial n}{\partial z} + \left[1 + \left[1 - \xi_{i} \right] \alpha_{i} \right] \frac{n \xi_{i}}{T} \frac{\partial T}{\partial z} + \frac{n \xi_{i}}{H_{i}} \right] - Kn \frac{\partial \xi_{i}}{\partial z}$$

$$(2)$$

where *n* is the total number density, ξ_i is the mole fraction, D_i is the molecular diffusion coefficient, *K* is the eddy diffusion coefficient, α_i is the thermal diffusion coefficient, *T* is the temperature, and H_i is the scale height

$$H_i = \frac{RT}{m_i g}, g = \frac{GM}{r^2} \tag{3}$$

with R as the gas constant, m_i as the molecular weight of species *i*, *g* as the gravity, *G* as the gravitational constant, and *M* representing Titan's mass. The total number density as a function of mean molecular weight can be determined from the equation of hydrostatic equilibrium,

$$\frac{dp}{dz} = -\rho g \tag{4}$$

where *p* is the pressure and ρ is the mass density = *nm*, where $m = \sum_{i} m_i \xi_i(z)$, the mean molecular weight at altitude *z*, and the gas law

$$p[f+1] = nkT,\tag{5}$$

where k is Boltzmann's constant and f is a factor marking the departure from gas ideality, as Titan's atmosphere exhibits non-ideal characteristics near its surface [Lindal et al., 1983]. Atmospheric parameters that are inputs into the model are the surface pressure $p_0 = 1496$ mb [Lindal et al., 1983] and the non-ideality profile and temperature profile (Figure 1a), modeled by Yelle et al. [1997] and used for all studies related to the Huygens probe.

[10] These equations are finite differenced (see Appendix A) and solved through the reduced Jacobian solver method [*Sandilands and McConnell*, 1997] where the species in Table 2 are divided into two families, representing the neutrals and the ions. The structure of the model is laid out in Figure 2. The primary input parameters - the solar flux and cross sections combined with the thermal profile - are used to calculate the next group of parameters located in the first box. The calculation of the radiation field then follows, producing photolysis rates for absorbing species. The continuity-diffusion equation is then inverted

	Table 2.	Observations	of	Constituents	in	Titan's	Atmosphere
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Species	Altitude, km	Instrument	Observation	References
	1400		$20 \pm 2\%$	Strobel et al. [1992]
H_4^a	1130	Voyager UVS	$8 \pm 3\%$	Smith et al. [1982]
	1000		$6 \pm 1\%$	Smith et al. [1982]
	>825		1-2%	Smith et al. [1982]
	725	Voyager UVS	0.1-0.3%	
2H2 ^{a,b}	$300_{-60}^{+80} \\ 180_{-30}^{+50}$		$NP^{b}:4.7^{+3.5}_{-2.1} \times 10^{-6}$	
2 2	180^{+50}_{-30}	Voyager IRIS	$NP^{b} 2 3^{+1.6} \times 10^{-6}$	Coustenis et al. [1991]
	125^{+50}_{-40}	vojugor nuo	$EQ^{b}:2.2^{+0.7}_{-0.9} \times 10^{-6}$	Coustenis et al. [1989]
	180^{+50}_{-30}		$NP^{b}:3.0^{+2.8}_{-2.1} \times 10^{-6}$	Coustenis et al. [1991]
$_{2}H_{4}^{a}$	125^{+55}_{-35}	Voyager IRIS	$EQ^{b}:9.0^{+3}_{-5} \times 10^{-8}$	Coustenis et al. [1989]
	300^{+80}_{-60}		$NP^{b}: 1.5^{+2.6}_{-0.9} \times 10^{-5}$	
	180^{+50}_{-30}	Voyager IRIS	$NP^{b} \cdot 10^{+1.4} \times 10^{-5}$	Coustenis et al. [1991]
₂ H ₆ ^a	125^{+50}_{-40}	voyager ners	$\begin{array}{c} \mathrm{NP^{b}:}1.0^{+1.4}_{-0.6}\times10^{-5}\\ \mathrm{EQ^{b}:}1.3^{+0.5}_{-0.7}\times10^{-5} \end{array}$	Coustenis et al. [1989]
	125_{-40} 105-300	IRHS-IRTF	$8.8 \pm 2.2 \times 10^{-6}$	
	103-300 200+80	IKH3-IKI F	$NP^{b}:6.2^{+4}_{-2.5} \times 10^{-8}$	Livengood et al. [2002]
псп	300^{+80}_{-60}	V IDIC	NP $:0.2_{-2.5} \times 10^{-8}$	Coustenis et al. [1991]
H_3C_2H	180^{+50}_{-30}	Voyager IRIS	NP ^b :2.0 ^{+1.1} _{-0.8} × 10 ⁻⁸	
	105^{+55}_{-30}		$EQ^{b}:4.4^{+1.7}_{-2.1} \times 10^{-9}_{-9}$	Coustenis et al. [1989]
	180^{+50}_{-30}	Voyager IRIS	$NP^{b}:5.0^{+4}_{-3.5} \times 10^{-7}_{-7}$	Coustenis et al. [1991]
$_{3}H_{8}$	105^{+55}_{-30}		$EQ^{b}:7.0^{+4}_{-4} \times 10^{-7}_{-7}$	Coustenis et al. [1989]
	90-250	TEXES-IRTF	$6.2 \pm 1.2 \times 10^{-7}$	<i>Roe et al.</i> [2003]
	$\begin{array}{c} 300 \substack{+80\\-60}\\ 180 \substack{+50\\-30}\\ 105 \substack{+55\\-30}\end{array}$		NP ^b :4.2 ^{+3.3} _{+2.1} × 10 ⁻⁸	Coustenis et al. [1991]
$_{4}\text{H}_{2}^{a}$	180^{+50}_{-30}	Voyager IRIS	$NP^{b}:2.7^{+2.1}_{-1.2} \times 10^{-8}_{-0.2}$	
	105^{+55}_{-30}		$EQ^{b}:1.4^{+0.6}_{-0.7} \times 10^{-9}$	Coustenis et al. [1989]
20	400	ISO	$8.0^{+6}_{-4} \times 10^{-9c}_{-9c}$	Coustenis et al. [1998]
20	>40	150	4.0×10^{-10}	constents et ut. [1996]
	350		$\begin{array}{c} 4.8^{+3.8}_{-1.5} \times 10^{-6} \\ 2.4^{+0.5}_{-0.5} \times 10^{-5} \\ 2.9^{+0.9}_{-0.5} \times 10^{-5} \end{array}$	
	175	IRAM – Pico Veleta, Spain	$2.4^{+0.5}_{-0.5} \times 10^{-5}$	Hidayat et al. [1998]
0	60		$2.9^{+0.9}_{-0.5} \times 10^{-5}$	
	200 - 300		$5.2 \pm 1.2 \times 10^{-3}$	Gurwell and Muhleman [2000]
	40 - 200	Owens Valley millimeter array	$5.2 \pm 0.6 imes 10^{-5}$	
	180^{+50}_{-30}	Voyager IRIS	NP ^b : $<7.0 \times 10^{-9}$	Coustenis et al. [1991]
O ₂	~ 180		$EO^{b,c}$: 1.4 × 10 ⁻⁸	
- 2	$105^{+55}_{-30} \\ 300^{+80}_{-60}$		$EQ^{b}:1.4^{+0.3}_{-0.5} \times 10^{-8}$	Coustenis et al. [1989]
	300^{+80}	Voyager IRIS	$NP^{b} \cdot 2 \cdot 3^{+1.8} \times 10^{-7}$	
	180^{+50}	tojuger mus	$NP^{b} \cdot 4 0^{+2.8} \times 10^{-7}$	Coustenis et al. [1991]
	$180^{+50}_{-30} \\ 125^{+55}_{-40}$		$\begin{array}{c} NP^{b}:2.3^{+1.8}_{-1.4} \times 10^{-7} \\ NP^{b}:4.0^{+2.2}_{-2.2} \times 10^{-7} \\ EQ^{b}:1.6^{-0.6}_{-0.6} \times 10^{-7} \\ \end{array}$	Coustenis et al. [1989]
	350		$37^{+1.8} \times 10^{-7}$	Hidayat et al. [1907]
CN ^a	300		$3.7^{+1.8}_{-1.2} \times 10^{-7} \\ 5.2^{+6.6}_{-3.9} \times 10^{-6}$	Tanguy et al. [1997]
CN	200		$2.5^{+1.2} \times 10^{-7/6} 2^{+1.9} \times 10^{-7}$	Hidayat et al. [1997]/Tanguy et al. [1990]
		IRAM – Pico Veleta, Spain	$\begin{array}{c} 3.5^{+1.2}_{-1.1} \times 10^{-7} / 6.2^{+1.9}_{-2.1} \times 10^{-7} \\ 2.0^{+0.3}_{-0.4} \times 10^{-7} / 3.3^{+0.9}_{-0.8} \times 10^{-7} \end{array}$	Hidayal el al. [1997]/languy el al. [1990]
	170	· •	$2.0_{-0.4} \times 10^{-3.5} \times 10^{-8} \times 10^{-8}$	
	110		$5.0_{-0.9} \times 10^{-8}$	Hidayat et al. [1997]
	100		$7.5^{+8.0}_{-3.0} \times 10^{-8}_{-8.0}$	Tanguy et al. [1990]
II ON	320		1.0×10^{-8c}	D/ 1 1 [1002]
H ₃ CN	250	IRAM – Pico Veleta, Spain	3.5×10^{-9c}	<i>Bézard et al.</i> [1993]
	180		1.5×10^{-9c}	
	300^{+80}_{-60}		NP ^b : $2.5^{+1.1}_{-1} \times 10^{-7}_{-7}$	Coustenis et al. [1991]
C_3N^a	180^{+50}_{-30}	Voyager IRIS	NP ^b :8.4 ⁺³ _{-3.5} × 10 ⁻⁸	
	105^{+55}_{-30}		$EQ^{b}: \le 1.5 \times 10^{-9}$	Coustenis et al. [1989]
	300^{+80}_{-60}		NP: $1.6^{+2.6}_{-1} \times 10^{-8}$	Coustenis et al. [1991]
$C_2 N_2^{a}$	180^{+50}_{-30}	Voyager IRIS	NP: $5.5^{+5}_{-2.2} \times 10^{-9}$	
	105^{+55}_{-30}		EQ ^b : $\leq 1.5 \times 10^{-9}$	Coustenis et al. [1989]

^a*Vervack* [1997] and *Vervack et al.* [2003] conducted Voyager UVS reanalysis of upper atmosphere observations for these constituents. See text. ^bVoyager IRIS observations taken at NP: north polar region, EQ: equatorial region.

^cValue obtained by linear fitting of a model-calculated density profile.

for a given chemical family before proceeding back to the recalculation of box 1 parameters with the new abundance values. This cycle continues until a solution is reached, defined by the convergence level

$$\left(\Delta\xi,\Delta n\right)_{conv} = \begin{cases} \frac{\Delta\xi}{\xi} < 10^{-4}\\ \frac{\Delta n}{n} < 10^{-4} \end{cases}.$$

The resultant nominal total number density profile is shown in Figure 1b.

3. Model Parameters

3.1. Radiation Field

3.1.1. Solar Flux Input

[11] During the Voyager 1 flyby of Titan in November 1980, the solar flux output was near the maximum of its 11-year cycle, amplifying the effect of solar radiation on the

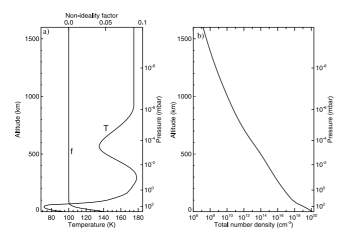


Figure 1. a) Temperature profile and non-ideality factor from *Yelle et al.* [1997]. b) Calculated total number density in Titan's atmosphere.

atmosphere. However, most of the stable constituents in Titan's atmosphere have lifetimes longer than one solar cycle in the stratosphere and are thus less sensitive to the changes in solar flux over the course of that cycle in that region. Thus the solar flux for moderate solar conditions obtained by The Solar-Stellar Irradiance Comparison Experiment (SOLSTICE) [Woods et al., 1996] on the Upper Atmosphere Research Satellite (UARS) from 1150-3000 Å and an EUV flux, calculated using the EUVAC model [Richards et al., 1994] for solar conditions associated with a 10.7 cm radio solar flux of 130 ($\times 10^{-22}$ W m⁻² Hz⁻¹), is used to calculate neutral densities in the nominal model. However, in the upper atmosphere above $\sim 0.01 \ \mu bar$, lifetimes for most constituents are considerably shorter. Sensitivity to solar flux for many of these constituents is discussed in section 5.2, in which simulations are run for solar minimum, solar maximum, and moderate solar conditions. Due to the short ion chemical lifetimes, profiles for charged particles in the solar maximum case are considered

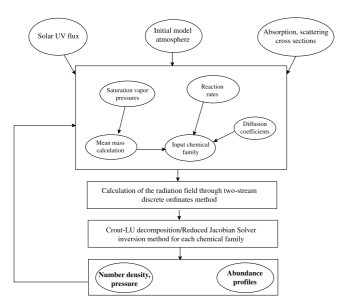


Figure 2. Layout of the structure of the model.

for the nominal model. In the solar maximum case, an EUV flux for F10.7 = 233 and the 81-day average 10.7 cm radio solar flux (F10.7A) equal to 211.9, which matches the solar conditions during the Voyager flyby, is used to calculate the distribution of species. The incident solar flux is calculated with a zenith angle of 58°, corresponding to the zenith angle representing a globally averaged incident flux in November 1980.

[12] The scattering contribution to the solar flux is calculated through a two-stream discrete ordinates algorithm detailed by *Edgington et al.* [1998]. The contributions of solar radiation, direct and scattered flux, for three characteristic wavelengths are shown in Figure 3. In the EUV region, scattering does not play a major role, shown in the 800 Å example, as nitrogen absorption prevents the penetration of photons to deeper regions where scattering would have a larger effect. In the 1700 Å example, radiation penetrates further into the atmosphere, allowing some backscattering to take place. In the denser regions, the effect

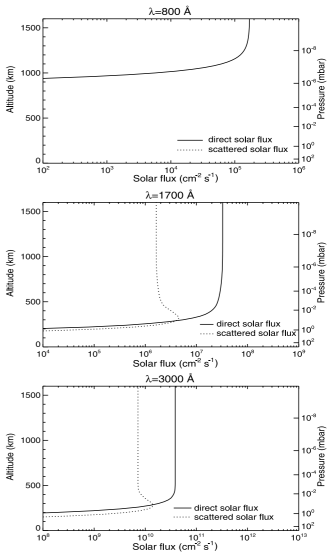


Figure 3. The direct solar flux and diffuse contribution as a function of altitude for 800 Å, 1700 Å, and 3000 Å intensities.

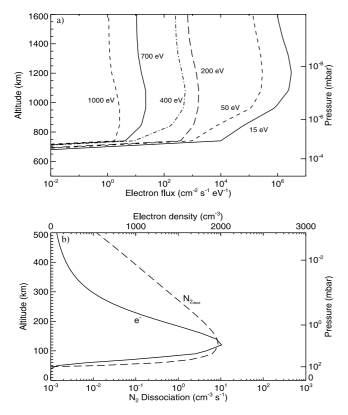


Figure 4. a) Radial electron fluxes from T. Cravens (personal communication, 1998) for various electron energies. b) Electron density profile in Titan's stratosphere (solid line) and molecular nitrogen dissociation as a result of cosmic ray deposition (long dashed line).

of scattering increases. At 3000 Å, the high single scattering albedo results in similar levels of direct and diffuse radiation throughout the atmosphere, with scattering dominating the solar input below 300 km.

3.1.2. Electron Input

[13] The solar flux while ionizing molecules also creates photoelectrons in the process. These photoelectrons play a significant role in the radiation field by providing an additional source of energy through which chemistry is induced. In their study of Titan's ionosphere, Gan et al. [1992] calculated photoelectron fluxes through Titan's atmosphere as a function of electron energy. On the basis of this study, the radial fluxes of photoelectrons from 15-100 eV, for a solar zenith angle of 60 degrees is adopted for altitudes from 725-1600 km (T. Cravens, personal communication, 1998). The radial fluxes for some of these electrons are shown in Figure 4a. Considering the input from Saturn's magnetosphere, Keller et al. [1992] calculated the ion production rates of N_2^+ , N^+ , and CH_4^+ , the ions that begin Titan ion chemistry, due to magnetospheric electron impact. These rates are also considered in the model.

3.1.3. Cosmic Rays

[14] The deposition of cosmic rays in Titan's stratosphere can be an important source of nitrile formation through cosmic ray-induced N_2 dissociation. Cosmic ray input in Titan's atmosphere is treated using the model of local energy deposition described by *Atreya et al.* [1995], assum-

ing zero magnetic rigidity for Titan, under moderate solar conditions. The cascading penetration of energy deposition analyzed by *Capone et al.* [1983] is neglected, amounting to a possible underprediction of N₂ dissociation of a factor ~2, according to *Lellouch et al.* [1994]. The total ionization rate calculated is assumed to equal the N₂ ionization rate and is scaled for the dissociation and ionization rate of N₂ and other species by a factor relating the peak absorption cross section via electron impact of that species with the N₂ absorption cross section at that wavelength. Thus, for $X + e^- \rightarrow j^{(+)}$, where j is a neutral or ion,

$$P_j^X = \frac{\sigma_X \left(E_j^{peak} \right) q_j}{\sigma_{N_2} \left(E_j^{peak} \right) q_{ion}^{N_2}} \mathcal{Q}_{N_2} \xi_X = J_j n \xi_X \tag{6}$$

where, P_j^X is the production rate of species j from destruction of X, σ_j is the electron impact absorption cross section of species j, E_j^{peak} is the electron energy of peak absorption for species j, q_j is the quantum yield of species j from destruction of X, $q_{ion}^{N_2}$ is the total ionization quantum yield of N₂ for electron impact, Q_{N_2} is the calculated total ionization rate of N₂ from cosmic ray impact, and J_j is the coefficient of production from destruction of X.

3.2. Photochemistry

[15] The chemical production term in the continuity equation (1) is given by

$$P_i = \sum_j \sum_k k_{jk} n^2 \xi_j \xi_k + \sum_l J_l^i n \xi_l \tag{7}$$

where k_{jk} is the rate coefficient of the reaction in which species *j* and species *k* react to form species *i*, while

$$J_l^i(z) = \int\limits_{\lambda} \sigma_l^{abs} I(z,\lambda) q_i(\lambda) d\lambda \tag{8}$$

where $q_i(\lambda)$ is the quantum yield, representing the probability that the photolysis of *l* will produce *i*. The expression $k_{jk}n^2\xi_j\xi_k$ represents the production rate of *i* resulting from the two-body reaction of *j* and *k*, and $J_in\xi_l$ is the production rate of *i* resulting from the photolysis of *l*. The chemical term for species *i*, on the other hand, is dependent on the mixing ratio of *i*, as the loss rate of a species is affected by its abundance

$$L_i = \sum_j k_{ij} n^2 \xi_i \xi_j + J_i n \xi_i.$$
(9)

 k_{ij} is the rate coefficient of the reaction of reactant *i* with some reactant *j*, with $k_{ij}n^2\xi_i\xi_j$ expressing the reaction rate of *i* and *j*. The photolysis rate of *i* is $J_in\xi_i$.

[16] For three-body pressure dependent reactions, where the two reactants react with the background atmosphere, k_{jk} is often given in terms of a low-pressure rate k_0 and a high-pressure rate k_{∞} where

$$k_{jk} = \frac{k_0 k_\infty n}{(k_0 n + k_\infty)}.$$
(10)

Meanwhile, the production rate due to the deexcitation of an excited constituent is $k_j n \xi_j$, where k_j is the deexcitation rate of constituent *j*.

[17] The set of reactions used in the model is displayed in Table 3. The reaction rates chosen are the rates calculated or measured under the conditions most representative of Titan's atmosphere. The most important of these conditions is temperature. The rate of a reaction can vary by many orders of magnitude over the span of 100 K in temperature, which can have a large effect on the overall chemistry of the atmosphere. The bulk of chemistry in Titan's atmosphere occurs in the region where atmospheric temperatures range between about 130 K and 180 K. So, rate coefficient measurements that are applicable at or near these temperatures are favored over rate measurements that are applicable at room temperature, for example. The pressure at which the rate was measured and the bath gas used is also taken into consideration.

[18] The absorption cross sections and quantum yields which are used in (7) to calculate photolysis rates are referenced in Table 4. Many of these cross sections can also vary significantly with temperature, and thus low temperature cross sections, where available, are used. For Rayleigh scattering, the cross section is

$$\sigma_i^{Ray} = \frac{32\pi^3 \alpha_i^2 \left[\frac{6+3\delta_i}{6-7\delta_i}\right]}{3\lambda^4},\tag{11}$$

where $\alpha_{CH_4} = 25.6 \times 10^{-25}$, $\delta_{CH_4} = 0$, $\alpha_{N_2} = 3.96 \times 10^{-25}$, and $\delta_{N_2} = 0.03$ [Allen, 1976].

[19] Microphysical models matched with geometric albedo observations [e.g., *McGrath et al.*, 1998] have suggested that Titan aerosols in the main haze layer are likely fractal in nature. Aerosol opacities for fractal particles were taken from *Lebonnois et al.* [2001], assuming aerosol single scattering albedos calculated from scattering and extinction efficiency factors from *Rannou et al.* [1995]. Haze opacity profiles for Mie particles from *Rannou et al.* [1995] were also tested.

[20] Charged particles are assumed to be governed solely through chemical processes with the assumption of charge neutrality. The electron temperature is taken from *Keller et al.* [1992] model A.

3.3. Condensation

[21] Due to the very low temperatures reached in Titan's tropopause and lower stratosphere, many gases become saturated and proceed to condense according to their saturation vapor pressures. The saturation laws governing condensing species are taken from *Allen and Nelson* [1998] for CH₄ above 90 K, C₂H₂, and C₂H₄, *Moses et al.* [1992] for other hydrocarbons, *Washburn* [1924] for H₂O, and *Sagan and Thompson* [1984] for HC₃N. Vapor pressures for other compounds are determined by fitting the lowest two points in the vapor pressure table from *Weast et al.* [1987] for the expression $\ln p(z) = A + \frac{B}{T(z)}$. This process is taken into account in the chemical side of equation (1) where for condensing species,

$$P_i - L_i - \gamma_i = \frac{1}{r^2} \frac{\partial (r^2 \Phi_i)}{\partial r}$$
(12)

where γ_i is the condensation factor of species *i*. The condensation factor is assumed to dominate in regions of condensation, forcing the constituent mixing ratio profile to follow the profile associated with its saturation vapor pressure. This is done by setting the condensation factor

$$\gamma_i = \left[An\left[\xi_i - \xi_i^{sat}\right]\right]^p \tag{13}$$

where ξ_i^{sat} equals the saturation mole fraction, $A \gg P_i$, L_i , and p is a constant ≈ 1 to smooth the transition between condensation and non-condensation regions.

[22] As evident in equation (4) the total number density is dependent on the mean mass of the atmosphere, which will be affected by the distribution of the most abundant constituents. Thus condensation must be taken into account in the mean mass calculation in a similar fashion as the mole fractions in equation (13), by forcing $m(z) = m^{sat}(z)$ in condensing regions, where

$$m^{sat}(z) = \sum_{i} m_{i} \min\left(\xi_{i}, \xi_{i}^{sat}\right).$$
(14)

3.4. Boundary Conditions

[23] The boundary conditions for the model are presented in Table 5. At the lower boundary, the mole fractions for H_2 and CO are set to their observed quantities in the stratosphere, while CH₄ assumes a mixing ratio profile consistent with the saturation model of Samuelson et al. [1997], with a surface mole fraction of 5.6% and a supersaturation of 1.37. ξ_{N_2} follows as $1 - \sum \xi_i$, $i \neq N_2$ throughout the model. The total number density is set to 1.2×10^{20} cm⁻³ at the lower boundary, associated with a pressure of 1496 mb, a temperature of 94 K, and a non-ideality factor of 0.0347. At the upper boundary, $\Phi_i = 0$ is assumed for most species, although $\Phi_{H_2O} = -5.0 \times 10^6$ molecules cm⁻² s⁻¹ is adopted for H₂O, accounting for the influx of water molecules arising from micrometeorites [Feuchtgruber et al., 1997]. For H and H₂, the escape flux is dependent on the density of these constituents that are being solved. Thus the boundary condition accounting for escape of H, H_2 is expressed in terms of velocity

$$w_i(\tilde{r}_N) = -D_i \left[\frac{1}{\xi_i} \frac{\partial \xi_i}{\partial z} + \frac{1}{n} \frac{\partial n}{\partial z} + [1 + \alpha_i] \frac{1}{T} \frac{\partial T}{\partial z} + \frac{1}{H_i} \right] - K \frac{1}{\xi_i} \frac{\partial \xi_i}{\partial z}.$$
(15)

The top of the model is assumed to be the exobase, where the effusion velocity from Jeans escape is given as

$$w_{e} = \sqrt{\frac{kT}{2\pi m}} [1 + E_{e}] e^{-E_{e}}$$
(16)

where $E_e = \frac{r_e}{H_e}$. Thus $w_H = 2.72 \times 10^4$ cm s⁻¹ and $w_{H_2} = 7000$ cm s⁻¹.

3.5. Vertical Transport

3.5.1. Molecular Diffusion

[24] Molecular diffusion coefficients are often provided as measurements of diffusivity in a medium consisting of two constituents. Atmospheres, however, are not binary

articl	e)		
Rxn	Reactions	Rate Coefficients	References and Comments
1	$H + H + M \rightarrow H_2 + M$	$1.5 \times 10^{-29} \text{ T}^{-1.3}$	Tsang and Hampson [1986]; bath gas N ₂
2	$C + C_2H_2 \rightarrow products$	2.6×10^{-10}	Guadagnini et al. [1998]
3	$CH + H_2 + M \rightarrow CH_3 + M$	$k_0 = 4.7 \times 10^{-26} T^{-1.6}$	Brownsword et al. [1997]
	3011	$k_{\infty} = 2.5 \times 10^{-10} \text{ T}^{-0.08}$	
4	$CH + H_2 \rightarrow {}^{3}CH_2 + H$	$3.1 \times 10^{-10} e^{-1650/T} 1.4 \times 10^{-11} $	Becken et al [1000]
4 5	$CH + H \rightarrow C + H_2$ $CH + CH_4 \rightarrow C_2H_4 + H$	$3.96 \times 10^{-8} \text{ T}^{-1.04} \text{ e}^{-36.1/\text{T}}$	Becker et al. [1989] Canosa et al. [1997]
6	$CH + C_{2}H_{4} \rightarrow C_{2}H_{4} + H$ $CH + C_{2}H_{2} \rightarrow C_{3}H_{2} + H$	$1.59 \times 10^{-9} \text{ T}^{-0233} \text{ e}^{-16/\text{T}}$	Canosa et al. [1997]
7	$CH + C_2H_2 \rightarrow CH_3C_2H + H$ $CH + C_2H_4 \rightarrow CH_3C_2H + H$	$3.87 \times 10^{-9} \text{ T}^{-0.546} \text{ e}^{-29.6/\text{T}}$	Canosa et al. [1997]
	$CH + C_2H_4 \rightarrow CH_2CCH_2 + H$	$3.87 \times 10^{-9} \text{ T}^{-0.546} \text{ e}^{-29.6/\text{T}}$	
8	$\mathrm{CH} + \mathrm{C_2H_6} \rightarrow \mathrm{C_2H_4} + \mathrm{CH_3}$	$1.9 \times 10^{-8} \text{ T}^{-0.859} \text{ e}^{-53.2/\text{T}}$	Canosa et al. [1997]
0	$CH + C_2H_6 \rightarrow C_3H_6 + H$	$1.9 \times 10^{-8} \text{ T}^{-0.859} \text{ e}^{-53.2/\text{T}}$	
9	$CH + C_4H_8 \rightarrow \text{products}$		Canosa et al. [1997]
10	$^{3}\mathrm{CH}_{2} + \mathrm{H} + \mathrm{M} \rightarrow \mathrm{CH}_{3} + \mathrm{M}$	$k_0 = 5.1 \times 10^{-10}$ e $k_\infty = 1.5 \times 10^{-10}$	Gladstone [1983]
	$^{3}\mathrm{CH}_{2} + \mathrm{H} \rightarrow \mathrm{CH} + \mathrm{H}_{2}$	$4.7 \times 10^{-10} e^{-3/0/1}$	Zabarnick et al. [1986]
11	$^{3}\mathrm{CH}_{2} + ^{3}\mathrm{CH}_{2} \rightarrow \mathrm{C}_{2}\mathrm{H}_{2} + \mathrm{H}_{2}$	$2.0 \times 10^{-11} e^{-400/T}$	Baulch et al. [1992]
	$^{3}CH_{2} + ^{3}CH_{2} \rightarrow C_{2}H_{2} + 2H$	$1.8 \times 10^{-11} e^{-400/T}$	
12	$^{3}\text{CH}_{2} + \text{CH}_{3} \rightarrow \text{C}_{2}\text{H}_{4} + \text{H}$	7.0×10^{-11}	Tsang and Hampson [1986]
13	$^{3}CH_{2} + C_{2}H_{2} \rightarrow C_{3}H_{3} + H$	$1.5 \times 10^{-11} e^{-3332/T}$	Bohland et al. [1986]
14	$^{3}CH_{2} + C_{2}H_{3} \rightarrow CH_{3} + C_{2}H_{2}$	3.0×10^{-11} 3.0×10^{-11}	Tsang and Hampson [1986]
15 16	${}^{3}\text{CH}_{2} + \text{C}_{2}\text{H}_{5} \rightarrow \text{CH}_{3} + \text{C}_{2}\text{H}_{4}$ ${}^{1}\text{CH}_{2} + \text{H}_{2} \rightarrow {}^{3}\text{CH}_{2} + \text{H}_{2}$	1.26×10^{-11}	Tsang and Hampson [1986] Langford et al. [1983]
10	$^{1}CH_{2} + H_{2} \rightarrow CH_{2} + H_{2}$	9.24×10^{-11}	
17	$^{1}\mathrm{CH}_{2}^{2} + \mathrm{CH}_{4} \rightarrow {}^{3}\mathrm{CH}_{2} + \mathrm{CH}_{4}$	1.2×10^{-11}	Bohland et al. [1985]
	$^{1}CH_{2} + CH_{4} \rightarrow CH_{3} + CH_{3}$	6.0×10^{-11}	
18	$^{1}\text{CH}_{2} + \text{C}_{2}\text{H}_{2} \rightarrow \text{C}_{3}\text{H}_{3} + \text{H}$	3.6×10^{-10}	Guadagnini et al. [1998]
19	$^{1}\text{CH}_{2} + \text{N}_{2} \rightarrow ^{3}\text{CH}_{2} + \text{N}_{2}$	$\begin{array}{l} 2.36 \times 10^{-14} \mathrm{T} \\ \mathrm{k}_0 = 4.0 \times 10^{-29} \end{array}$	Ashfold et al. [1981]
20	$\mathrm{CH}_3 + \mathrm{H} + \mathrm{M} \to \mathrm{CH}_4 + \mathrm{M}$	$k_0 = 4.0 \times 10$ $k_\infty = 4.7 \times 10^{-10}$	Brouard et al. [1989]
		$F_{-} = 0.902 - 1.03 \times 10^{-3} T$	
21	$\mathrm{CH}_3 + \mathrm{CH}_3 + \mathrm{M} \to \mathrm{C}_2\mathrm{H}_6 + \mathrm{M}$	$k_0 = 8.76 \times 10^{-6} T^{-7.03} e^{-1390/T}$	Slagle et al. [1988]; Rate used is $10 \times$ measured rate - see text
		$k_{ee} = 1.5 \times 10^{-6} T^{-1.18} e^{-329/T}$	
		$F_c = 0.381 e^{-T/37.2} - 0.619 e^{-T/1180}$	
	$CH_4 + C_2 \rightarrow C_2H + CH_3$	$5.05 \times 10^{-11} e^{-297/T}$ $1.2 \times 10^{-11} e^{-491/T}$	Pitts et al. [1982]
	$CH_4 + C_2H \rightarrow C_2H_2 + CH_3$ $CH_4 + C_4H \rightarrow C_4H_2 + CH_3$	$1.2 \times 10^{-11} e^{-491/T}$ $1.2 \times 10^{-11} e^{-491/T}$	<i>Opansky and Leone</i> [1996a] estimating that C_4H reaction rates are equal to their C_2H analogues;
24	$CH_4 + C_4 \Pi \rightarrow C_4 \Pi_2 + C\Pi_3$		Kiefer and von Drasek [1990]
25	$CH_4 + C_6H \rightarrow C_6H_2 + CH_3$	$1.2 \times 10^{-11} e^{-491/T}$	estimating that C_6H reaction rates are equal to their C_2H analogues;
			Kiefer and von Drasek [1990]
	$C_2 + H_2 \rightarrow C_2 H + H$	$1.77 \times 10^{-12} e^{-1469/T}$	Pitts et al. [1982]
27	$C_2H + H + M \rightarrow C_2H_2 + M$	$k_0 = 1.26 \times 10^{-18} \text{ T}^{-3.1} \text{ e}^{-721/\text{T}}$	Tsang and Hampson [1986]
20	$C_2H + H_2 \rightarrow C_2H_2 + H$	$k_{\infty} = 3.0 \times 10^{-10}$ $9.2 \times 10^{-18} \text{ T}^{2.17} \text{ e}^{-478/\text{T}}$	Openative and Leone [1006b]
	$C_2H + H_2 \rightarrow C_2H_2 + H$ $C_2H + C_2H_2 \rightarrow C_4H_2 + H$	$9.2 \times 10^{-11} e^{30.8/T}$ $9.53 \times 10^{-11} e^{30.8/T}$	Opansky and Leone [1996b] Chastaing et al. [1998]
	$C_2H + C_2H_2 \rightarrow C_4H_2 + H$ $C_2H + C_2H_4 \rightarrow C_4H_4 + H$	$4.6 \times 10^{-11} e^{24.6/T}$	Opansky and Leone [1996b], Chastaing et al. [1998]
	$C_2H + C_2H_4 \rightarrow C_2H_2 + C_2H_3$	$4.6 \times 10^{-11} e^{24.6/T}$	
	$C_2H + C_2H_6 \rightarrow C_2H_2 + C_2H_5$	$3.5 \times 10^{-11} e^{2.9/1}$	Opansky and Leone [1996b]
	$C_2H + C_3H_6 \rightarrow products$	$1.47 \times 10^{-10} e^{65.3/T}$	Chastaing et al. [1998]
	$C_2H + C_3H_8 \rightarrow C_2H_2 + C_3H_7$	$7.8 \times 10^{-11} \text{ e}^{3/\text{T}}$ $9.53 \times 10^{-11} \text{ e}^{30.8/\text{T}}$	Hoobler et al. [1997]
	$\begin{array}{l} C_2H+C_4H_2\rightarrow C_6H_2+H\\ C_2H+C_6H_2\rightarrow C_8H_2+H \end{array}$	$9.53 \times 10^{-11} e^{30.8/T}$ $9.53 \times 10^{-11} e^{30.8/T}$	<i>Chastaing et al.</i> [1998]; estimated from <i>Kiefer and von Drasek</i> [1990] <i>Chastaing et al.</i> [1998]; estimated from <i>Kiefer and von Drasek</i> [1990]
	$C_2H + C_6H_2 \rightarrow C_8H_2 + H$ $C_2H + C_8H_2 \rightarrow polymer$	$9.53 \times 10^{-11} e^{50.8/1}$	Chastaing et al. [1996]; estimated from Kiefer and von Drasek [1996] Chastaing et al. [1998]; estimated from Kiefer and von Drasek [1990]
	$C_2H_2 + H_2 + M \rightarrow C_2H_3 + M$	$k_0 = 3.3 \times 10^{-30} e^{-740/1}$	Baulch et al. [1992]
		$k_{\perp} = 1.4 \times 10^{-11} e^{-1300/1}$	
	$C_2H_2 + C_4H \rightarrow C_6H_2 + H$	$9.53 \times 10^{-11} e^{30.8/T}$	Chastaing et al. [1998]; estimated from Kiefer and von Drasek [1990]
	$C_2H_2 + C_6H \rightarrow C_8H_2 + H$	$9.53 \times 10^{-11} e^{30.8/T}$	Chastaing et al. [1998]; estimated from Kiefer and von Drasek [1990]
40	$\begin{array}{l} C_2H_3+H\rightarrow C_2H_2+H_2\\ C_2H_3+H+M\rightarrow C_2H_4+M \end{array}$	7.6×10^{-11} $k_0 = 5.76 \times 10^{-24} \text{ T}^{-1.3}$	k ₀ estimate; <i>Monks et al.</i> [1995]
	$C_2\Pi_3 + \Pi + \Pi_1 \rightarrow C_2\Pi_4 + \Pi_1$	$k_{\rm m} = 8.0 \times 10^{-11}$	
41	$C_2H_3+H_2 \rightarrow C_2H_4+H$	$5.01 \times 10^{-20} \text{ T}^{2.63} \text{ e}^{-4298/\text{T}}$	Tsang and Hampson [1986]
		$k_{min} = 1.0 \times 10^{-23}$	Isang and Handson [1900]
42	$C_2H_3+CH_3\rightarrow C_2H_2+CH_4$	3.4×10^{-11}	Fahr et al. [1991]
	$C_2H_3+CH_3+M \rightarrow C_3H_6+M$	$k_0 = 8.76 \times 10^{-6} T^{-7.03} e^{-1390/T}$	k_0 estimated from $k_0(CH_3 + CH_3)$; Fahr et al. [1991]
10		$k_{\infty} = 1.2 \times 10^{-10}$	
	$C_2H_3 + CH_4 \rightarrow CH_3 + C_2H_4$ $CH_4 + CH_4 \rightarrow CH_4 + H_4$	$ \begin{array}{c} & & & \\ 2.4 \times 10^{-24} \text{ T}^{4.02} \text{ e}^{-2754/\text{T}} \\ 3.32 \times 10^{-12} \text{ e}^{-2516/\text{T}} \end{array} $	Tsang and Hampson [1986]
44	$C_2H_3 + C_2H_2 \rightarrow C_4H_4 + H$ $C_2H_3 + C_2H_2 + M \rightarrow C_4H_5 + M$	$k_0 = 3.3 \times 10^{-29} e^{-740/T}$	Fahr and Stein [1988] k_0 estimated as $10 \times k_0(C_2H_3 + H)$; Weissman and Benson [1988]
	$C_{2113} + C_{2112} + W \rightarrow C_{4115} + W$	$k_0 = 3.3 \times 10^{-19} \text{ C}$ $k_{\infty} = 4.17 \times 10^{-19} \text{ T}^{1.9} \text{ e}^{-1058/\text{T}}$	R_0 commuted as 10 \wedge $R_0(C_2H_3 + H)$, reasonant and Denson [1700]
45	$\mathrm{C_2H_3} + \mathrm{C_2H_3} \rightarrow \mathrm{C_2H_4} + \mathrm{C_2H_2}$	2.4×10^{-11}	Fahr et al. [1991]

Table 3 (Representative Sample). Rate Coefficients Used in the Model (The full Table 3 is available in the HTML version of this article)

Table 4. Absorption Cross Sections and Quantum Yields Used in	ntum Yields Used in the Model		
Pathways	Branching Ratios	Cross Sections	Quantum Yields
	1.0 see Quantum Yields 1.0.7160 Å)	Paler Chan et al. [1992]; Samson and Haddad [1994] Dan	Palenius et al. [1976] Ford et al. [1975] Darbas et al. [1973]
$\begin{bmatrix} J_{0,2} \\ J_{1,1} \end{bmatrix} CH_4 + h_V \rightarrow a) CH_2 + H_1 \\ b) (CH_2 + H_2 \\ c) ^1 CH_2 + 2H \\ d) ^3 CH_2 + 2H \\ d) ^3 CH_2 + 2H \\ c) CH + H_2 + H \\ c) CH_1 + H_2 + H_2 \\ d) CH_3 + H + e^- \\ b) U^+ + U + U^- + e^- \\ d \end{bmatrix}$	$\begin{array}{l} q_{\rm neu}^{a} (<\!950 \ {\rm \AA}); 1.0, (2100 \ {\rm A}); 0.41 \ (\rm Ly \ \alpha) \\ 0.28 \ (\rm Ly \ \alpha) \\ 0.0 \\ 0.10 \ (\rm Ly \ \alpha) \\ 0.10 \ (\rm Ly \ \alpha) \\ see \ Quantum \ Yields; Au \ et \ al. \ [1993] \end{array}$	Au et al. [1993]; Mount et al. [1977], Mount and Moos [1978] at 200 K	Au et al. [193]; Romani [1996]; Au et al. [1989]; see text Samson et al. [1989]; see text
	0.3 × q _{neu} ^a (<1100 Å); 0.3 (1100–1660 Å); 0.06 (1660–1900 Å); 0.3 (1900–2400 Å); 0.1 × q _{neu} ^a (<1100 Å); 0.1 (1100–1660 Å); 0.1 (1660–1900 Å); 0.1 (1900–2400 Å); 0.6 × q _{neu} ^a (<1100 Å); 0.6 (1100–1660 Å); 0.84 (1660–1900 Å); 0.6 (1900–2400 Å) see Quanturn Yields; <i>Zheng and Srivastava</i> [1996]	Wu and Judge [1985]; Cooper et al. [1995]; Wu et al. [2001] at 150 K; Seki and Okabe [1993]	50–1100 Å quantum yields are taken from electron impact quantum yields Zheng and Srivustava [1966]; Okabe [1981, 1983]; Seki and Okabe [1993]
ا <i>ی</i> :	$\begin{array}{c} 0.58 \times q_{neu}{}^{a} (<\!1180 \mathring{A}\!); 0.58 (1180-1750 \mathring{A}\!); \\ 0.73 (1750-2100 \mathring{A}\!) \\ 0.42 \times q_{neu}{}^{a} (<\!1180 \mathring{A}\!); 0.42 (1180-1750 \mathring{A}\!); 0.27 (1750-2100 \mathring{A}\!) \\ \text{see Quantum Yields; } 7ian and 7idal [1998a] \end{array}$	Fahr Holland et al. [1997]; Zelikoff and Watanabe [1953]; Schmitt and Brehm [1966]; F. Raulin (personal communication, 1995)	 Fahr et al. [1998] 500-1200 Å quantum yields are taken from electron impact quantum yields <i>Tian and Vidal</i> [1998a]; 995) <i>Vidal</i> [1998a]; Holland et al [1997]; Chang et al. [1998]
[J8] $C_2H_5 + h\nu \rightarrow \breve{C}H_3^{-+}^{1}CH_2$ [J9] $C_2H_6 + h\nu \rightarrow a) C_2H_4 + H_2$ b) $C_2H_4 + 2H$ c) $C_2H_2 + 2H_2$ d) $CH_4 + {}^{1}CH_2$ e) $2CH_3$ f) $C_2H_5^{+} + e^{-}$ g) $C_2H_5^{+} + e^{-} + \dots$ h) $C_2H_5^{+} + e^{-} + \dots$ h) $H_7^{+} + e^{-} + \dots$ h) $H_7^{+} + e^{-} + \dots$ m) $H_3^{+} + e^{-} + \dots$ m) $CH_3^{+} + e^{-} + \dots$	$\begin{array}{l} 1.0 \\ 0.56 \times q_{neu}^{a} (< 1130 \ {\rm \AA}); 0.56 \ (1130 - 1700 \ {\rm \AA}); \\ 0.14 \times q_{neu}^{a} (< 1130 \ {\rm \bigstar}); 0.14 \ (1130 - 1700 \ {\rm \AA}); \\ 0.27 \times q_{neu}^{a} (< 1130 \ {\rm \circlearrowright}); 0.27 \ (1130 - 1700 \ {\rm \AA}); \\ 0.22 \times q_{neu}^{a} (< 1130 \ {\rm \circlearrowright}); 0.02 \ (1130 - 1700 \ {\rm \AA}); \\ 0.02 \times q_{neu}^{a} (< 1130 \ {\rm \circlearrowright}); 0.02 \ (1130 - 1700 \ {\rm \AA}); \\ 0.01 \times q_{neu}^{a} (< 1130 \ {\rm \bigstar}); 0.01 \ (1130 - 1700 \ {\rm \AA}); \\ 0.01 \times q_{neu}^{a} (< 1130 \ {\rm \bigstar}); 0.01 \ (1130 - 1700 \ {\rm \AA}); \\ 0.01 \times q_{neu}^{a} (< 1130 \ {\rm \bigstar}); 0.01 \ (1130 - 1700 \ {\rm \AA}); \\ 0.08 \ (Ly \ {\rm \circlearrowright}) \\ \text{see Quantum Yields; Tian and Vidal [1998b] \end{array}$	Au et al. [1993]; Mount and Moos [1978] at 200 K	<i>Adachi et al.</i> [1979] 60–1200 Å taken from electron impact quantum yields <i>Tian and Vidal</i> [1998b]; <i>Kameta et al.</i> [1977] <i>Mount et al.</i> [1977]

Table 4. (continued)			
Pathways	Branching Ratios	Cross Sections	Quantum Yields
[J10] $C_3H_3 + h\nu \rightarrow a$) $C_3H_2 + H$ b) $C_2H + H_2$	0.96 0.04	Fahr et al. [1997]	Jackson et al. [1991]
[J11] CH ₃ C ₂ H + $h_V \rightarrow 0$ C ₃ C ₃ C ₄ + H b) C-H + H ₂	$0.56 \times q_{neu}^{a}$ (<1200 Å); 0.56 (1200–2200 Å) 0.44 × c_{n}^{a} (<1200 Å); 0.44 (1200–2200 Å)	Ho et al. [1998]; Chen et al. [2000] at 200 K	Sun et al. [1999]; Ho et al. [1998]
[J12] CH ₂ CCH ₂ + $h\nu \rightarrow a$) C ₃ H ₃ + H	0.64	Rabalais et al. [1971];	Rabalais et al. [1971]
b) $C_{3H_2} + H_2$	0.36	<i>Chen et al.</i> [2000] at 200 K	
[J13] C ₃ H ₅ + $h\nu$ → a) C ₂ H ₂ + CH ₃ b) CH ₃ C ₅ H + H	0.79 0.16	Jenkan et al. [1993]	Gierczak et al. [1988]
c) $CH_2CCH_2 + H$	0.01		
[J14] $C_3H_6 + h\nu \rightarrow a$) $C_3H_5 + H$	0.0 (1000–1350 Å); 0.0 (1350–1550 Å); 0.565	Samson et al. [1962];	Collin [1988]
b) CH ₃ C ₅ H + H ₅	(1550–1750 A); 0.41 (1750–1950 A) 0.11 (1000–1350 Å): 0.11 (1350–1550 Å):	Fahr and Nayak [1996] at 223 K	
4	0.01 (1550–1750 Å); 0.01 (1750–1950 Å)		
c) $CH_2CCH_2 + H_2$	0.17 (1000–1350 A); 0.22 (1350–1550 A);		
H_{1}^{-} H_{1}^{-} H_{2}^{-} H_{2}^{-}	(A UCU (1200-1/10); A); U.U (1350-150); U.U (A UCU (1360-1350); A): A A (1350-1550); A		
u) V2114 - V112	0.02 (1550–1750 Å): 0.03 (1750–1950 Å)		
e) $C_2H_3 + CH_3$	0.21 (1000 - 1350 Å); 0.27 (1350 - 1550 Å);		
	0.335 ($1550 - 1750$ Å); 0.4 ($1750 - 1950$ Å)		
f) $C_2H_2 + CH_4$	0.05 (1000-1350 Å); 0.03 (1350-1550 Å);		
	0.05 (1550–1750 Å); 0.04 (1750–1950 Å)		
$[115]C_3H_8 + h\nu \rightarrow a) C_3H_6 + H_2$	0.19 (1000-1150 A); 0.34 (1150-1350 A); 0.66 (1350-1540 Å); 0.04 (1540-1630 Å)	Koch and Skibowski [1971]; Obaba and Bookar [1963]	Johnston et al. [1978]
h) $C_{S}H_{\epsilon} + {}^{1}CH_{s}$	$(x) \alpha \alpha \alpha 1 - \alpha + \alpha$	[COLI] ISANGA MIN SONNO	
	0.04 (1350–1540 Å); 0.00 (1540–1630 Å)		
c) $C_2H_5^+ + CH_3$	0.40 (1000-1150 Å); 0.35 (1150-1350 Å);		
	0.19 (1350–1540 A); 0.00 (1540–1630 A)		
d) $C_2H_4 + CH_4$	0.32 (1000–1150 Å); 0.22 (1150–1350 Å); 0.11 (1350–1540 Å); 0.06 (1540–1630 Å)		
[J16] $C_4H_2 + h\nu \rightarrow a$) $C_4H + H$	$0.20~(1200 - 1640~{ m \AA});~0.00~(1640 - 2640~{ m \AA})$	Okabe [1981]; Fahr and Nayak [1994]	Glicker and Okabe [1987]
b) 2C ₂ H	$0.03~(1200 - 1640~{ m \AA});~0.01~(1640 - 2640~{ m \AA})$	at 223 K; Smith et al. [1998] at 193 K.	
c) $C_2H_2^2 + C_2$	$0.10~(1200 - 1640~{ m \AA});~0.06~(1640 - 2640~{ m \AA})$		
d) $C_4H_2^2$	0.67 (1200 - 1640 A); 0.93 (1640 - 2640 A)		
$[J17] C_4H_4 + h\nu \rightarrow a) C_4H_2 + H_2$	0.8	Fahr and Nayak [1996] at 233 K.	Gladstone et al. [1996]
[J18] $C_4H_6 + h\nu \rightarrow a) C_4H_4 + H_2$	0.05	Samson et al. [1962]	Bergmann and Demtröder [1968]
b) $2C_2H_3$ c) $C_2H_4 + C_3H_3$	0.10 0.17		
d) $C_3H_3 + CH_3$	0.40		
e) $C_4H_5 + H$	0.28		

Pathways	Branching Ratios	Cross Sections	Quantum Yields
[J19] $C_4H_8 + h\nu \rightarrow a) C_4H_6 + 2H$	0.23 (1050 - 1350 Å); 0.14 (1350 - 1600 Å); 0.06 (1600 Å); 0.06	Samson et al. [1962]	Niedzielski et al. [1978, 1979]; Caliie and Wiscolomusti [1078]
b) $C_3H_5 + CH_3$	0.12 (1050-1350 & 0.39 (1350-1600 Å); 0.66 (1600-1350 Å);		COUNT RECEVENSAN [12/0]
c) $CH_3C_2H + CH_4$	0.03 (1050 - 1350 Å); 0.02 (1350 - 1600 Å); 0.02 (1350 - 1500 Å); 0.02 (1350 - 1600 Å)		
d) $CH_2CCH_2 + CH_4$	0.00 (1000 - 1350 A) 0.14 (1050 - 1350 Å); 0.10 (1350 - 1600 Å); 0.00 (1350 - 1080 Å);		
e) $C_2H_5 + C_2H_3$	0.25 (1050-1350 Å); 0.14 (1350-1600 Å);		
f) $2C_2H_4$	0.02 (1050 - 1350 Å); 0.04 (1350 Å); 0.05 (14500		
g) $C_2H_2 + 2CH_3$	0.02 (1050 - 1350 - 1300 - 1300) 0.02 (1050 - 1350) 0.04 (1500 - 1850)		
h) $C_{3}H_{6} + {}^{1}CH_{2}$	0.02 (1050 - 1350 - 1360 - 1360 - 1360); 0.02 (1050 - 1350); 0.02 (1350 - 1600);		
$[J20] C_4H_{10} + h\nu \rightarrow a) C_{4H_8} + H_2$	0.48 (1000 - 1550 Å); 0.31 (1350 - 1650 Å)	Koch and Skibowski [1971];	<i>Obi et al.</i> [1971]
$\begin{array}{c} 0 \ C_{2}H_{6} + C_{2}H_{4} \\ 0 \ C_{3}H_{6} + CH_{3} + H \\ 2 \ C \ H + C \ H + H \\ 2 \ C \ H + C \ H + H \end{array}$	0.28 (1000–1350 Å); 0.11 (1350–1650 Å) 0.28 (1000–1350 Å); 0.41 (1350–1650 Å) 0.00 (1000–1350 Å).	[covi] techer	
[J21] $C_6H_2 + h\nu \rightarrow a)$ $C_6H_3 + h\nu \rightarrow a)$	0.02 (1000-1530 Å), 0.11 (1530-1030 Å) 0.20 (1150-1640 Å); 0.00 (1640-3000 Å)	Kloster-Jensen et al. [1974]; Bénilan	Wilson and Atreya [2003]
[J22] $C_8H_2 + h\nu \rightarrow a$) $C_4H + C_2H$	0.13 (1150–1640 Å); 0.07 (1640–5000 Å) 0.20 (1150–1640 Å); 0.00 (1640–3000 Å)	et al. [1992] at 253 K Assumed same as $C_6H_2 + h\nu$	Wilson and Atreya [2003]
b) $C_{6}H + C_{2}H$ [J23] $C_{6}H_{6} + h\nu \rightarrow a) C_{6}H_{5} + H$	$0.13 (1150 - 1640 \text{ A}); 0.07 (1640 - 3000 \text{ A}) 0.8 \times q_{\text{neu}} < (1300 - 2200 \text{ A}); 0.8 (1300 - 2200 \text{ A});$	Rennie et al. [1998]; Pantos et al. [1978]	Yo
b) $C_6H_4 + H_2$	$0.16 \times q_{\text{heu}}^{-1}$ (c1300 Å); 0.16 (1300–2200 Å); 0.16 (1300–2200 Å); 0.16 (1300–2200 Å); 0.06 (22200 Å);		Kennte et al. [1998]
c) $C_{5}H_{3} + CH_{3}$	$0.04 \times q_{\text{neu}^{11}} (<1300 \times 0.00 \text{ k}) 0.04 \times q_{\text{neu}^{11}} (<1300 \times 0.01 \text{ k});$		
$ [124] N + h\nu \rightarrow N^+ + e^- $ $ [125] N_2 + h\nu \rightarrow a) N^{4s} + N^{2d} $ $ b) N_2^+ + e^- $	see Quantum Yields	Fennel Chan et al. [1993]	Fennelly and Torr [1992] Shaw et al. [1992]; Zipf and McLaughlin [1978];
$[J26] HCN + h\nu \rightarrow a) CN + N^{+s} + e^{-}$ $b) HCN + H$ $b) HCN + e^{-}$ $c) CN^{+} + H + e^{-}$ $d) U^{+} + a^{-} + d$	q _{neu} ^a see Quantum Yields; Kreile et al. [1982]	180–620 Å assumed equal to C ₂ H ₂ ; Nuth and Glicker [1982]; F. Raulin (personal communication, 1995)	Gallagher et al. [1988] Kreile et al. [1982]; Nuth and Glicker [1982]
$ [J27] HC_3N + h\nu \rightarrow a) C_3N + H b) CN + C_2H c) HC_3N^* $	0.09 0.05 0.24	Connors et al. [1974]; Clarke and Ferris [1996]; Andrieux et al. [1995];	Clarke and Ferris [1995]; Halpern et al. [1988]; see text
[J28] CH ₃ CN + $h\nu \rightarrow$ CN + CH ₃ [J29] C ₂ H ₃ CN + $h\nu \rightarrow$ a) CN + C ₂ H ₃ b) HCN + C ₂ H ₂ c) HCN + H ₂ c) HCAN + H ₂	1.0 0.01 0.15 0.59	Bénilan et al. [1994]; Bruston et al. [1989] Nuth and Glicker [1982]; Suto and Lee [1985] F. Raulin (personal communication, 1995), Derecskei-Kovacs and North [1999] Wilson and Atreya [2003]	Bruston et al. [1989] Nuth and Glicker [1982]; Suto and Lee [1985] mmunication, 1995), Derecskei-Kovacs and North [1999] 003]
$[130] C_2N_2 + h\nu \rightarrow 2CN$	67-0 0.1	Nuth and Glicker [1982]; F. Raulin (nercond) communication 1005)	
$[J31] C_4N_2 + h\nu \rightarrow C_3N + CN$	1.0	(personal communication, 1993) Comors et al. [1974	Commun. 1979) Commors et al. [1974]; Bénilan et al. [1996] at 233 K

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Onontrum Vialde	Wight et al. [1976]; Okabe [1978]	Okabe [1978]	Haddad and Samson [1986]; Stief et al. [1975]; Mordaunt et al. [1994]	•			Gitcker and Stief [19/1]; Moortgat et al. [1985]	Braun et al. [1970]; Rabalais et al. [1971]	Harich et al. [1999]	Shah et al. [1987]	Backx et al. [1976]	Orient and Srivastava [1987]	Zhona and Srivastava [1006]	[0661] namenate mi	Tian and Vidal [1998a]				Tian and Vidal [1998b]	1					Brook et al. [1978] Zipf et al. [1980]; Itikawa et al. [1986]			Rao et al. [1995]	
Cross Cartine	Wight et al. [1976];	Guttagner et al. [1966]; Ogawa [1971]; Nakata et al. [1965]; Ogawa [1971]; Lewis and Carver [1983] at 200 K	Haddad and Samson [1986]; Watanabe and Jursa [1964];	Watanabe et al. [1953]; Thomnson et al. [1963]			Gentteu and Mentall [19/0]; Suto et al. [1985]		Пал	Sh	Ba	Orient a	Zhona a	בווכנול מ	Tian				Tian					1	Bro Zipf et al. [19			R	
Breanching Dation	prarenting reation gneu ^a	1.0 (1000 - 1670 Å); 0.0 (1670 - 2150 Å) 0.0 (1000 - 1670 Å): 1.0 (1670 - 2150 Å)	$0.78 \times \dot{q}_{neu}^{n}$ (<900 Å); 0.78 (990–1450 Å); 1.0 (1450–1980 Å)	$0.11 \times q_{\text{heu}^{a}} (<990 \text{ Å}); 0.11 (990-1450 \text{ Å}); 0.0 (1450-1980 \text{ Å})$	$0.11 \times q_{heu}^{a}$ (<990 Å); 0.11 (990–1450 Å); 0.0 (1450–1980 Å)	see Quantum Yields; Haddad and Samson [1986]	C.0 2.0	1.0	co.u 71.0	1.0	see Quantum Yields	see Quantum Yields	see Annun Vields	SCC Quantum LICIUS	see Quantum Yields				see Quantum Yields						1.0 see Quantum Yields			see Quantum Yields	
Table 4. (continued)	$[J32] \text{ CO} + h\nu \rightarrow \text{C} + \text{O}^{3p}$	[J33] CO ₂ + $h\nu \rightarrow$ a) CO + O ^{3p} b) CO + O ^{1d}	[J34] $H_2O + h\nu \rightarrow a$) $OH + H$ b) $O^{1d} + H_2$	$\begin{array}{c} c) O^{3p} + H_2^{-} \\ d) H_2O^+ + e^- \end{array}$	e) $H^+ + e^- + \dots$		$[J32] H_2CO + hv \rightarrow a) CO + H_2$ b) CO + 2H	[J36] CH ₂ CO + $h\nu \rightarrow$ CO + $^{1}CH_{2}$	$[J_2 /] CH_3 OH + m \rightarrow a CH_3 O + H \rightarrow b H_2 CO + H \rightarrow b + b + b + b + b + b + b + b + b + b$	→ H ⁺ + 2	$\begin{bmatrix} E2 \end{bmatrix} H_2 + e^- \rightarrow a) H_2 + 2 e^- \\ b) H^+ + H + 2 e^- \\ b + H + 2 e^- \end{bmatrix}$	[E3] CH ₄ + $e^{-\rightarrow}$ a) CH ₄ + 2 $e^{-\rightarrow}$	b) CH ₃ + H + 2 e ⁻ FA1 C ₂ H ₂ + e^{-} a) C ₂ H ⁺ + 2 e ⁻	1	[E5] $C_2H_4 + e^- \rightarrow a) C_2H_4^+ + 2 e^-$	c) $C_2H_3 + H + 2$ e c) $C_2H_2^2 + 2$ e ⁻ +	d) $H^+ + 2 e^- + \dots$ e) $H^+_2 + 2 e^- + \dots$	$f_{11} + f_{22} = e^{-1} + \dots$	$[E6] C_{3}H_{6} + e^{-} \rightarrow a) C_{3}H_{6}^{+} + 2 e^{-}$	b) $C_2H_5^+ + H + 2 e^-$ c) $C_2H_7^+ + 2 e^- + -$	d) $C_{2H_{3}}^{-2H_{4}} + 2 e^{-+}$	e) $C_{2}H_{2} + 2 e^{-+}$ f) $H^{+} + 2 e^{-+}$	g) $H_2^+ + 2 e^- +$ h) $H_3^+ + 2 e^- +$	i	$ [E7] N + e \rightarrow N + 2 e [E8] N_2 + e^- \rightarrow a) N^{4s} + N^{4s} + e^- $	b) $N^{4s} + N^{2d} + e^{-1}$ c) $N^{2d} + N^{2d} + e^{-1}$	d) $N^+_{1+} + 2e^{1+}$	[E9] $H_2O + e^- \rightarrow a) H_2O^+ + 2 e^-$ b) $H_2O^+ + 2 e^-$	^a q _{neu} is the neutral quantum yield.

Physical Quantity	Boundary Constraint
Lower Boundary Conditions	
Pressure	$p_0 = 1496 \text{ mb}$
Methane mole fraction	$\xi_{\rm CH_4} = 5.6\%$
Methane supersaturation	1.37
Molecular hydrogen mole fraction	$\begin{array}{l} \xi_{\rm H_2} = 1.1 \times 10^{-3} \\ \xi_{\rm CO} = 5.0 \times 10^{-5} \end{array}$
Carbon monoxide mole fraction	$\xi_{\rm CO} = 5.0 \times 10^{-5}$
Upper Boundary Conditions	
Atomic hydrogen escape velocity	$H_{vel}^{esc} = 2.7 \times 10^4 \text{ cm s}^{-1}$
Molecular hydrogen escape velocity	$H_{2_{vel}}^{esc} = 7 \times 10^3 \text{ cm s}^{-1}$ $\Phi_{H_2}O^a = 5 \times 10^6 \text{ cm}^{-2} \text{ s}^{-1}$
Water influx from micrometeorites	$\Phi_{\rm H_2} \dot{\rm O}^{\rm a} = 5 \times 10^6 \ {\rm cm}^{-2} \ {\rm s}^{-1}$

Table 5. Boundary Conditions Used in the Model

^aFlux referred to the surface.

mixtures of gases, but rather are composed of many constituents. For an atmosphere dominated by one constituent, binary coefficients involving that major constituent and a minor constituent are sufficient for calculating the molecular diffusion coefficient of that given minor constituent. However, an atmosphere with a minor constituent that makes up a significant portion of the bulk gas requires a multicomponent treatment. *Wilke* [1950] provides a convenient approximation for the diffusion of gas in a multicomponent mixture of J gases

$$D_i = \frac{1 - \xi_i}{\sum_{j=1, j \neq i}^J \frac{\xi_j}{D_{ij}}}$$
(17)

where D_{ij} are the binary diffusion coefficients and ξ_i is the mole fraction for species *i*. This formulation, which is good to about 10% for multicomponent systems where the individual diffusion coefficients are not sensitive to composition changes [*Wilke*, 1950], is useful for Titan where methane increases, due to diffusive separation, in the heterosphere, exceeding 20% above 1400 km [*Strobel et al.*, 1992]. Unfortunately, the measurements of binary coefficients have been conducted for only a select group of molecules. The diffusion coefficient follows the form of

$$D = \frac{AT^s}{n} \tag{18}$$

where T = temperature and n = bulk gas density. The binary gas mixtures pertinent to Titan that have been measured or empirically calculated are shown in Table 6. In the absence

of measurements, empirical correlations are often used, on the basis of the physical parameters of the given molecules.

[25] Thermal diffusion, which typically affects light gases, is given by the term in equation (2)

$$[1-\xi_i]\alpha_i\frac{n\xi_i}{T}\frac{\partial T}{\partial z},$$

where α_i is the thermal diffusion coefficient. In the model, $\alpha_H = \alpha_{H_2} = -0.38$ is adopted [*Banks and Kockarts*, 1973], while for all other species $\alpha = 0$.

3.5.2. Eddy Diffusion

[26] Owing to the complexity involved in relating microscopic turbulent processes to macroscopic transport, the eddy diffusion coefficient acts as a free parameter that must be estimated to match constituent observations. This task is relatively trivial for a completely inert species (e.g., argon), as the point where the constituent profile changes from well-mixed to diffusively separated marks the homopause where the eddy diffusion coefficient is equal to the molecular diffusion coefficient of the constituent. Unfortunately, there is no known vertical profile of such a species for Titan. Methane, however, is largely inert and is distributed largely through diffusive processes, as shown in Figure 5a. Thus the methane distribution can be used as a proxy for the determination of the homopause level. In the lower atmosphere, chemistry plays a major role for many species. However, HCN is largely formed in the upper atmosphere and transported to lower altitudes. In much of the lower regions of the atmosphere the eddy diffusion time constant is much smaller than the HCN chemical time constant (Figure 5b), and the millimeter observations of Tanguy et al. [1990] and Hidayat et al. [1997] provide a vertical

Table 6. Binary Molecular Diffusion Coefficients Used in the Model

Binary Mixture	A, $\mathrm{cm}^{-1} \mathrm{K}^{-\mathrm{s}}$	S	References
N_2-N_2	5.09×10^{16}	0.810	Massman [1998]
CH_4-N_2	7.34×10^{16}	0.750	Banks and Kockarts [1973]
H-N ₂	4.87×10^{17}	0.698	Banks and Kockarts [1973]
H_2-N_2	1.88×10^{17}	0.820	Mason and Marrero [1970]
$\tilde{N-N_2}$	9.69×10^{16}	0.774	Mason and Marrero [1970]
$O-N_2$	9.69×10^{16}	0.774	Banks and Kockarts [1973]
$CO-N_2$	5.15×10^{16}	0.810	Massman [1998]
$CO_2 - N_2$	4.08×10^{16}	0.810	Massman [1998]
$H_2 \tilde{O} - N_2$	6.26×10^{16}	0.810	Massman [1998]
$C_2H_6-N_2$	7.64×10^{16}	0.730	Wakeham and Slater [1973]
$C_3H_8-N_2$	6.54×10^{16}	0.660	Wakeham and Slater [1973]
C ₄ H ₁₀ -N ₂	7.34×10^{16}	0.610	Wakeham and Slater [1973]
CH ₄ -CH ₄	5.73×10^{16}	0.500	estimated from Lennard-Jones correlation; Reid et al. [1987]
H ₂ -CH ₄	2.30×10^{17}	0.765	Mason and Marrero [1970]

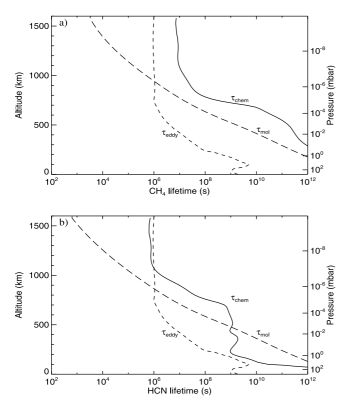


Figure 5. Local chemical, eddy, and molecular diffusive time constants for a) CH_4 and b) HCN.

profile for HCN in this region. These factors make HCN a good candidate for constraining the eddy diffusion profile in the lower atmosphere.

[27] Observations of CH₄ in the upper atmosphere were provided by Smith et al. [1982], who analyzed the Voyager ultraviolet spectrometer (UVS) observations and obtained mole fractions of 6 \pm 1% at 1000 km and 8 \pm 3% at 1130 km. However, the reanalysis of Voyager data by R. J. Vervack et al. (New perspectives on Titan's upper atmosphere from a reanalysis of the Voyager 1 UVS solar occultations, submitted to Icarus, 2003) (hereinafter referred to as Vervack et al., submitted manuscript, 2003) suggested a methane mixing ratio profile that might decrease with altitude, contrary to what would be expected from diffusive separation. Such a profile might be possible through a large chemical sink for CH₄, as suggested by Lara et al. [1999], but such a sink would most likely require much larger electron densities that what has been observed, as discussed in section 5.7. The authors argue that this profile may be the result of misassignment of CH₄ absorption to other species above 1050 km. With the large error bars and lack of coverage below the nanobar region, these results are unable to yield any firm conclusions about the homopause level, although the authors favor a high homopause level. Strobel et al. [1992], in their analysis of Voyager UVS solar occultation and airglow data, also infer a high homopause level around 1000 km. However, Steiner and Bauer [1990] in their study of diffusive processes on Titan, presented an analytical eddy profile corresponding to a homopause level of 660 km.

[28] The UVS reanalysis also points out a discrepancy with the *Smith et al.* [1982] C_2H_2 observations, retrieving

up to three orders of magnitude less acetylene in the upper atmosphere than was suggested by the *Smith et al.* [1982] observations. This finding is supported by photochemical models, which have not determined a mechanism to form upper atmospheric C_2H_2 at such high densities as suggested by *Smith et al.* [1982].

[29] The millimeter observations of *Hidavat et al.* [1997] and Tanguy et al. [1990] are in decent agreement with each other over the lower 200 km of the atmosphere. However, above this level they diverge, with Hidayat et al. [1997] observing an approximately uniform HCN mixing ratio profile while Tanguy et al. [1990] observed a profile increasing with altitude. The recent Marten et al. [2002] observations corroborate the finding of Tanguy et al. [1990] that HCN increases in mole fraction throughout the stratosphere. To determine the nominal eddy diffusion profile, the fitting of the Smith et al. [1982] CH₄ observations in the upper atmosphere and the HCN observations of Hidayat et al. [1997] and Tanguy et al. [1990] are used as a guideline. Considering this set of observations, 100 monotonically increasing, randomly generated eddy diffusion coefficients are used to determine the best fits for the CH₄ and HCN observations. These profiles are generated by obtaining diffusion coefficients at 12 altitude levels, determined by calculating a randomly generated positive slope from the previous altitude level within a predetermined range. The eddy diffusion coefficient at the lower boundary is randomly chosen between $100-10000 \text{ cm}^2 \text{ s}^{-1}$. The results of the bestfit profiles are then compared to determine which profile fits the remaining observational constraints, with slight adjustments made to achieve a better fit. The resulting nominal profile (NOM) along with the Steiner and Bauer [1990] profile (SB) is shown in Figure 6a, corresponding to a homopause level of 850 km. This process was repeated, replacing the Smith et al. [1982] results with those from the Vervack et al. (submitted manuscript, 2003) nominal CH₄ profiles up to 1050 km, to determine the best-fit high homopause eddy profile (HI), which is also displayed in Figure 6b.

[30] The nominal profile allows for more mixing in the middle atmosphere than previous models, with a larger eddy coefficient from 250–700 km (Figure 6c). The homopause level of 850 km is higher than the *Yung et al.* [1984], *Toublanc et al.* [1995], and *Lara et al.* [1996] profiles, although considerably lower than the *Lara et al.* [2002] model, which counteracts the result of a smaller methane upper atmosphere mole fraction from the higher homopause level by assuming a larger methane surface density.

4. Chemical Mechanisms

4.1. Hydrocarbons

[31] The formation of hydrocarbons begins with the photodissociation of CH_4 , which proceeds through the following channels:

$$\mathrm{J4a}) \qquad \qquad \mathrm{CH}_4 + h\nu \to \mathrm{CH}_3 + \mathrm{H}$$

$$(J4b) \longrightarrow {}^{1}CH_{2} + H_{2}$$

$$(J4c) \rightarrow {}^{1}CH_{2} + 2H$$

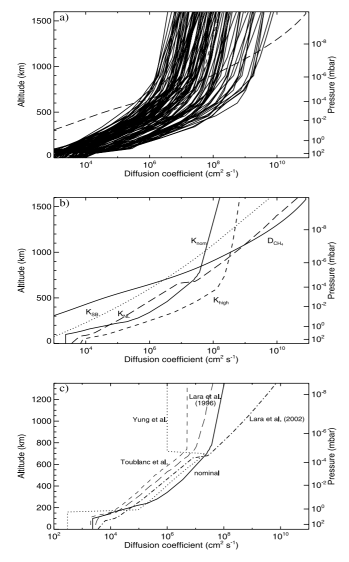


Figure 6. a) Randomly generated eddy diffusion profiles used to determine best fitting profile to constituent observations. b) Plots of tested eddy diffusion profiles - nominal profile (solid line), *Steiner and Bauer* [1990] profile (dotted line), the best-fit high homopause profile (short dashed line), and the *Lara et al.* [2002] profile (long dashed line). Also included in the methane molecular diffusion coefficient profile c) Eddy diffusion profiles from various photochemical models - nominal profile (solid line), *Yung et al.* [1984] profile (dotted line), *Lara et al.* [1996] profile (long dashed line), and *Lara et al.* [2002] profile (dotted line), and *Lara et al.* [2002] profile (dotted line).

$$(J4d) \rightarrow {}^{3}CH_{2} + 2H$$

$$(J4e) \rightarrow CH + H_2 + H.$$

The branching ratios for this process, particularly at Lyman α , which accounts for 75% of methane photodissociation above 700 km, are still unsettled. *Toublanc et al.* [1995] used the *Mordaunt et al.* [1993] model 1 scheme, which

places half of methane dissociation into (J4a) while splitting the other half between (J4b) and (J4d), and Lara et al. [1996] chose the Mordaunt et al. [1993] model 2 scheme, splitting methane dissociation into (J4a) and (J4e). The scheme adopted for this model is the *Romani* [1996] scheme where the quantum yields at Lyman α are as follows: q(J4a) = 0.41, q(J4b) = 0.28, q(J4c) = 0.0, q(J4d) = 0.21, q(J4e) = 0.10. At wavelengths other than Lyman α , methane is assumed to dissociate into the (J4a) channel. Wilson and Atreva [2000b] examined and compared these methane photolytic schemes along with that provided by Smith and Raulin [1999] and determined that there was little impact in the distribution of C2-hydrocarbons, while there was considerable difference in the profiles of the C₃H₄ isomers and C₃H₆, mainly arising from the differences in CH yield among the schemes. The progression of hydrocarbon chemistry follows with reactions involving the radicals produced from methane dissociation.

[32] Ethylene is created in the upper atmosphere through two different addition/decomposition mechanisms - radical/ radical association

$$(R12) \qquad {}^{3}CH_{2} + CH_{3} \rightarrow C_{2}H_{4} + H,$$

and the CH-insertion-H-elimination process involving methane

$$(R5) \qquad \qquad CH + CH_4 \rightarrow C_2H_4 + H$$

to yield the net reaction $2CH_4 \rightarrow C_2H_4 + H_2 + 2H$, as well as through ion chemistry in the scheme

(S1)
$$\frac{N_{2} + h\nu \rightarrow N_{2}^{+} + e^{-}}{CH_{4} + N_{2}^{+} \rightarrow CH_{3}^{+} + N_{2} + H}{CH_{4} + CH_{3}^{+} \rightarrow C_{2}H_{5}^{+} + H_{2}}{HCN + C_{2}H_{5}^{+} \rightarrow H_{2}CN^{+} + C_{2}H_{4}} - \frac{e^{-} + H_{2}CN^{+} \rightarrow HCN + H}{2CH_{4} \rightarrow C_{2}H_{4} + H_{2} + 2H}$$

The ion contribution through (S1) controls ethylene production near the model-calculated ionospheric peak at 1060 km. However, the peak of ethylene production corresponds with the peak of methane dissociation near 800 km, controlled by processes (R12) and (R5) which accounts for 56% and 42%, respectively, of ethylene production above 600 km [Wilson and Atreya, 2000b]. Once ethylene is formed, it serves as the major source of acetylene above 500 km through photolysis (J7), responsible for 75% of the total acetylene column production rate above 500 km. This differs from the Toublanc et al. [1995] and Yung et al. [1984] models where recombination of methylene radicals plays a larger role due to the authors' choice of methane photolytic scheme and the larger ${}^{3}CH_{2}$ recombination rate coefficient. Most of this acetylene is diffused into the lower atmosphere where it is polymerized to form higher-order hydrocarbons (e.g., C₄H₂)

(S2)
$$\frac{C_2H_2 + h\nu \to C_2H + H}{C_2H + C_2H_2 \to C_4H_2 + H},$$

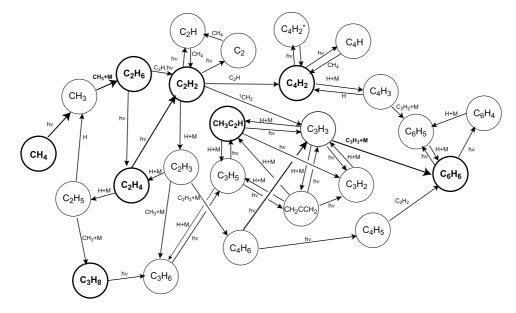


Figure 7. Schematic of Titan hydrocarbon chemistry. Stable species and primary reactions are shown in bold.

(S8

and undergoes reactions with hydrogen to reform ethylene:

(S3)
$$C_{2}H_{2} + H \xrightarrow{M} C_{2}H_{3}$$
$$C_{2}H_{3} + H \xrightarrow{M} C_{2}H_{4} \cdot \frac{1}{C_{2}H_{2} + 2H \rightarrow C_{2}H_{4}}$$

The progression of this lower atmosphere hydrocarbon chemistry is demonstrated in Figure 7.

[33] Methyl radicals are produced mainly through methane photolysis (J4a) above 700 km, while the catalytic dissociation of methane

(S4)
$$\frac{\begin{array}{c}C_{2}H_{2} + h\nu \rightarrow C_{2} + H_{2}\\C_{2} + CH_{4} \rightarrow C_{2}H + CH_{3}\\\underline{C_{2}H + CH_{4} \rightarrow C_{2}H_{2} + CH_{3}}\\CH_{4} \rightarrow CH_{3} + H\end{array}}{CH_{4} \rightarrow CH_{3} + H}$$

(S5)
$$\begin{array}{c} C_2H_2 + h\nu \rightarrow C_2H + H \\ C_2H + CH_4 \rightarrow C_2H_2 + CH_3 \\ \hline CH_4 \rightarrow CH_3 + H \end{array} .$$

accounts for 80% of methyl radical production or 5.6 \times 10⁹ CH₃ molecules cm⁻³ s⁻¹. The main destination of these CH₃ radicals is the formation of ethane through the addition of (S4), (S5), or (J4a) and methyl recombination (R21),

(S6)
$$\frac{\begin{array}{c} CH_4 + h\nu \to {}^{1}CH_2 + H_2 \\ CH_4 + {}^{1}CH_2 \to CH_3 + CH_3 \\ \hline CH_3 + CH_3 \xrightarrow{M} C_2H_6 \\ \hline 2CH_4 \to C_2H_6 + H_2 \end{array}}{CH_3 + CH_2}.$$
 (S9)

Methyl radicals also combine with C_2H_5 radicals, produced from H-attachment to ethylene (R48) or C_2H -insertion-H- abstraction from ethane (R31), to produce propane in the lower stratosphere

$$(S7) \qquad \begin{array}{c} 2CH_4 \to C_2H_4 + 2H + H_2 \\ CH_4 \to CH_3 + H \\ C_2H_4 + H \xrightarrow{M} C_2H_5 \\ \hline C_2H_5 + CH_3 \xrightarrow{M} C_3H_8 \\ \hline 3CH_4 \to C_3H_8 + 2H + H_2 \end{array}$$

with (S7) accounting for 61% of C_3H_8 production and (S8) producing 38% of the propane in the atmosphere.

[34] CH radical is mainly produced through (R10b) in the upper atmosphere, promoted by the formation of 3 CH₂ radicals through both methane dissociation (J4d)–43% of 3 CH₂ and 1 CH₂ collisional quenching–51% of 3 CH₂ above 600 km [*Wilson and Atreya*, 2000b]

$$CH_4 + h\nu \rightarrow {}^3CH_2 + 2H$$

and

$$\frac{\mathrm{CH}_{4}+h\nu\rightarrow\ ^{1}\mathrm{CH}_{2}+\mathrm{H}_{2}}{\mathrm{^{1}\mathrm{CH}}_{2}+\mathrm{N}_{2}\rightarrow\ ^{3}\mathrm{CH}_{2}+\mathrm{N}_{2}.}$$

$$\frac{\mathrm{CH}_{4}\rightarrow\ ^{3}\mathrm{CH}_{2}+\mathrm{H}_{2}}{\mathrm{CH}_{4}\rightarrow\ ^{3}\mathrm{CH}_{2}+\mathrm{H}_{2}}$$

Via (R10b) and (S5) CH radicals advance the production of methylacetylene and propylene. CH_3C_2H is produced in the

upper atmosphere from the direct CH-insertion-H-elimination process

(S10)
$$\begin{array}{c} CH_4 \to {}^{3}CH_2 + 2H \\ {}^{3}CH_2 + H \to CH + H_2 \\ \underline{CH + C_2H_4 \to CH_3C_2H + H} \\ \underline{CH_4 + C_2H_4 \to CH_3C_2H + H_2 + 2H} \end{array}$$

and indirectly through collisional isomerization of allene,

(S11)
$$CH_{4} \rightarrow {}^{3}CH_{2} + 2H$$
$${}^{3}CH_{2} + H \rightarrow CH + H_{2}$$
$$CH + C_{2}H_{4} \rightarrow CH_{2}CCH_{2} + H ,$$
$$CH_{2}CCH_{2} + H \rightarrow CH_{3}C_{2}H + H$$
$$CH_{4} + C_{2}H_{4} \rightarrow CH_{3}C_{2}H + H_{2} + 2H$$

while propylene is formed in the upper atmosphere through the mechanism

(S12)
$$\begin{array}{c} 2CH_4 \to C_2H_6 + H_2 \\ CH_4 \to {}^{3}CH_2 + 2H \\ {}^{3}CH_2 + H \to CH + H_2 \\ CH + C_2H_6 \to C_3H_6 + H \\ \hline {}^{3}CH_4 \to C_3H_6 + 2H_2 + 2H \end{array}$$

and more prominently in the lower atmosphere through

(S13)
$$\begin{aligned} & 2CH_4 \to C_2H_4 + 2H + H_2 \\ CH_4 \to CH_3 + H \\ C_2H_4 + h\nu \to C_2H_2 + H_2 \\ C_2H_2 + H \xrightarrow{M} C_2H_3 \\ \hline \\ & \frac{C_2H_3 + CH_3 \xrightarrow{M} C_3H_6}{3CH_4 \to C_3H_6 + 2H_2 + 2H} \end{aligned}$$

[35] Heavier hydrocarbons may also be formed in Titan's atmosphere - among the most prominent, benzene. *Wilson et al.* [2003] explored possible mechanisms for benzene formation, supported by the tentative detection of C_6H_6 by *Coustenis et al.* [2003]. Benzene in the lower stratosphere is primarily formed through the scheme

(S14)
$$\begin{array}{c} 4(C_{2}H_{2} + H \stackrel{M}{\rightarrow} C_{2}H_{3}) \\ 2(C_{2}H_{3} + C_{2}H_{3} \stackrel{M}{\rightarrow} C_{4}H_{6}) \\ 2(C_{4}H_{6} + h\nu \rightarrow C_{3}H_{3} + CH_{3}) \\ CH_{3} + CH_{3} \stackrel{M}{\rightarrow} C_{2}H_{6} \\ C_{3}H_{3} + C_{3}H_{3} \stackrel{M}{\rightarrow} n - C_{6}H_{6} \\ \underline{n - C_{6}H_{6} + H \rightarrow C_{6}H_{6} + H} \\ \hline 4C_{2}H_{2} + 4H \rightarrow C_{6}H_{6} + C_{2}H_{6} \end{array}$$

with H-addition onto phenyl radical

(S15)

$$\begin{array}{c}
C_{2}H_{2} + h\nu \to C_{2}H + H \\
C_{2}H_{2} + C_{2}H \to C_{4}H_{2} + H \\
C_{4}H_{2} + H \xrightarrow{M} C_{4}H_{3} \\
C_{4}H_{3} + C_{2}H_{2} \xrightarrow{M} C_{6}H_{5} \\
\underline{C_{6}H_{5} + H \xrightarrow{M} C_{6}H_{6}} \\
\hline
3C_{2}H_{2} \to C_{6}H_{6}
\end{array}$$

providing a minor pathway.

4.2. Nitriles

[36] Nitrogen, the predominant molecule in Titan's atmosphere, is largely chemically inert. The primary manner in

which nitrogen is involved in chemistry is through dissociation via both photons and electrons. Nitrogen is photodissociated mainly due to radiation in the 800–1000 Å region. *Zipf and McLaughlin* [1978] and *Zipf et al.* [1980] have determined the yields of nitrogen atoms from this process to be approximately 50% for ground-state N^{4s} and 30-40% and 10-20% for excited states N^{2d} and N^{2p}, respectively. However, *Zipf et al.* [1980] found that N^{2p} atoms are rapidly de-excited to the N^{2d} state. Therefore N^{4s} and N^{2d} are assumed to adopt a quantum yield of 0.5 each from N₂ photolysis. Nitrogen also undergoes electronimpact dissociation, with quantum yields determined by *Zipf et al.* [1980] and *Itikawa et al.* [1986].

[37] Nitrogen atoms combine with hydrocarbons to form an assortment of nitrile neutrals and ions in the upper atmosphere, as shown in Figure 8. The flux of N^{4s} atoms combines with methane photolytic product, CH_3 , to form the basis of nitrile chemistry–HCN. HCN is formed through photodissociation

(S16)

$$0.5(N_2 + h\nu \rightarrow N^{4s} + N^{4s})$$

$$CH_4 + h\nu \rightarrow CH_3 + H$$

$$N^{4s} + CH_3 \rightarrow H_2CN + H$$

$$\frac{H_2CN + H \rightarrow HCN + H_2}{0.5N_2 + CH_4 \rightarrow HCN + H_2 + H}$$

electron impact processes

(S17)
$$\begin{array}{c} N_{2}+e^{-} \rightarrow N^{+}+N^{4s,2d}+2e^{-} \\ CH_{4}+N^{+} \rightarrow H_{2}CN^{+}+H_{2} \\ e^{-}+H_{2}CN^{+} \rightarrow HCN+H \\ \overline{N_{2}+CH_{4}} \rightarrow HCN+N+H_{2}+H \end{array} ,$$

and photoionization

$$\begin{split} N_2 + h\nu \rightarrow N^+ + N^{2d} + e^- \\ CH_4 + N^+ \rightarrow H_2 CN^+ + H_2 \\ N^{2d} + C_2 H_4 \rightarrow CH_3 CN + H \\ \underline{CH_3 CN + H_2 CN^+ \rightarrow CH_3 CNH^+ + HCN} \\ N_2 + C\overline{H_4} + \overline{C_2 H_4} \rightarrow HCN + H_2 + H + \overline{CH_3 CNH^+} + e^- \end{split}$$
(S18)

with (S16) accounting for 60% of non-recycled HCN in the atmosphere. HCN is then diffused down into the lower atmosphere, providing the major source of HCN in this region, evident by an HCN flux of 3.4×10^8 cm⁻² s⁻¹ at 600 km, where it undergoes recycling mechanisms, commenced through photolysis and H-addition

м

(S19)
$$\begin{array}{c} HCN + h\nu \rightarrow CN + H \\ CN + CH_4 \rightarrow HCN + CH_3 \\ \hline CH_4 \rightarrow CH_3 + H \end{array}$$

(S20)
$$\frac{\text{HCN} + \text{H} \stackrel{\text{M}}{\to} \text{H}_2\text{CN}}{\text{H}_2\text{CN} + \text{H} \rightarrow \text{HCN} + \text{H}_2} \cdot \frac{\text{HCN} + \text{H}_2}{\text{H} + \text{H} \rightarrow \text{H}_2} \cdot \frac{\text{HCN} + \text{H}_2}{\text{H}_2} \cdot \frac{\text{HCN} + \frac{\text{HCN} + \text{H}_2}{\text{H}_2} \cdot \frac{\text{HCN} + \frac{H}_2}{\text{H}_2} \cdot \frac{\frac{H}_2}{\text{H}_2} \cdot$$

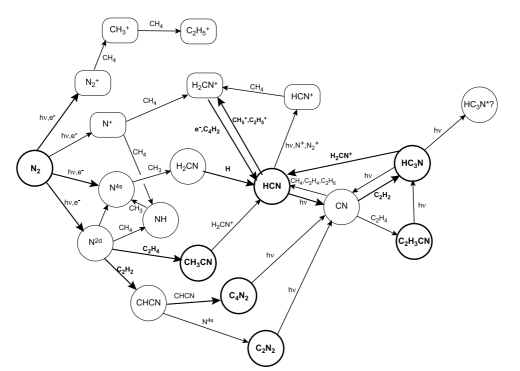


Figure 8. Schematic of Titan nitrile chemistry. Stable species and primary reactions are shown in bold, with ions shown in rounded rectangles.

[38] HC₃N is produced primarily through the addition of the photolytic product CN and acetylene

(S21)
$$\begin{array}{c} \mathrm{HCN} + h\nu \to \mathrm{CN} + \mathrm{H} \\ \mathrm{CN} + \mathrm{C}_{2}\mathrm{H}_{2} \to \mathrm{HC}_{3}\mathrm{N} + \mathrm{H} \\ \mathrm{HCN} + \mathrm{C}_{2}\mathrm{H}_{2} \to \mathrm{HC}_{3}\mathrm{N} + \mathrm{H} \end{array} .$$

[39] C_4N_2 can also be formed through CN-insertion, as studied by *Halpern et al.* [1989]

$$(R149) \qquad \qquad CN + HC_3N \to C_4N_2 + H.$$

Yung [1987] proposed another mechanism for the formation of C_4N_2 as well as C_2N_2 through the action of the CHCN radical in the reactions

$$(R161) \qquad CHCN + CHCN \rightarrow C_4N_2 + H_2$$

$$(R160) CHCN + N^{4s} \rightarrow C_2N_2 + H.$$

Considering estimations by *Yung* [1987] for (R161), the main source of dicyanoacetylene is the scheme

(S22)
$$\frac{N_2 + h\nu \to 2N^{2d}}{2[N^{2d} + C_2H_2 \to CHCN + H]} \frac{2[N^{2d} + C_2H_2 \to CHCN + H]}{N_2 + 2C_2H_2 \to C_4N_2 + 2H + H_2},$$

responsible for 98% of C_4N_2 production throughout Titan's atmosphere. Likewise, C_2N_2 is produced through the scheme

(S23)
$$\frac{\begin{array}{c} N_{2} + h\nu \to N^{4s} + N^{2d} \\ N^{2d} + C_{2}H_{2} \to CHCN + H \\ N^{4s} + CHCN \to C_{2}N_{2} + H_{2} \\ \hline N_{2} + C_{2}H_{2} \to C_{2}N_{2} + 2H \end{array},$$

responsible for virtually all of C_2N_2 production. Reactions with CH_3 and H_2CN , to produce HCN, severely limit the availability of N^{4s} atoms, giving a preference for CHCN to produce C_4N_2 in the upper atmosphere.

[40] In the upper atmosphere, C_2H_3CN production follows along the same procedure as (S19),

(S24)
$$\begin{array}{c} HCN + h\nu \rightarrow CN + H \\ CN + C_2H_4 \rightarrow C_2H_3CN + H \\ HCN + C_2H_4 \rightarrow C_2H_3CN + 2H \end{array}$$

while in the lower atmosphere, C_2H_3CN is produced through

(S25)
$$\begin{array}{c} C_2H_2 + H \stackrel{M}{\longrightarrow} C_2H_3 \\ HCN + C_2H_3 \rightarrow C_2H_3CN + H \\ HCN + C_2H_2 \rightarrow C_2H_3CN + H \end{array}$$

4.3. Oxygen Species

[41] Stratospheric measurements of CO [e.g., *Gurwell* and Muhleman, 2000], CO₂ [Coustenis et al., 1989], and H₂O [Coustenis et al., 1998] have indicated the presence of oxygen chemistry in the atmosphere of Titan. A likely contributor to this oxygen inventory is an influx of micrometeorites into the atmosphere, providing a source of H₂O molecules. Primarily, H₂O molecules undergo UV photolysis. The formation of hydroxyl radicals through H₂O photodissociation comprises 60% of H₂O destruction in Titan's atmosphere. Some of the OH radicals serve to recycle H₂O. In fact, 54% of H₂O destruction is recycled back to H₂O, mainly through charged-particle catalysis,

(S26)

$$N_{2} + h\nu \rightarrow N_{2}^{+} + e^{-}$$

$$N_{2}^{+} + H_{2}O \rightarrow H_{2}O^{+} + N_{2}$$

$$H_{2}O^{+} + CH_{4} \rightarrow H_{3}O^{+} + CH_{3}$$

$$\frac{H_{3}O^{+} + HCN \rightarrow H_{2}CN^{+} + H_{2}O}{CH_{4} + HCN \rightarrow H_{2}CN^{+} + CH_{3} + e^{-}}$$

(S27)
$$\begin{array}{c} N_{2} + h\nu \rightarrow N_{2}^{+} + e^{-} \\ N_{2}^{+} + C_{2}H_{2} \rightarrow N_{2}H^{+} + C_{2}H \\ N_{2}H^{+} + H_{2}O \rightarrow H_{3}O^{+} + N_{2} \\ H_{3}O^{+} + HCN \rightarrow H_{2}CN^{+} + H_{2}O \\ \hline CH_{4} + HCN \rightarrow H_{2}CN^{+} + CH_{3} + e^{-} \end{array}$$

through the reversibility of methylene-water addition

$$(R214a) \qquad {}^{1}CH_{2} + H_{2}O \rightarrow CH_{3} + OH$$

$$(R208a) \qquad \qquad CH_3 + OH \rightarrow H_2O + {}^1CH_2$$

and through

(S28)
$$\frac{H_2O + h\nu \rightarrow OH + H}{OH + CH_4 \rightarrow H_2O + CH_3}.$$

The main destiny of OH radicals that are not recycled back to H_2O is the formation of CO_2 through reaction with primordial CO,

(R219)
$$CO + OH \rightarrow CO_2 + H.$$

[42] CO is engaged in chemistry mainly in the lower atmosphere, where it produces oxygenated compounds through pressure-dependent reactions

(R217)
$$CO + {}^{3}CH_{2} \xrightarrow{M} CH_{2}CO$$

and

(S29)
$$\frac{2(CO + H \xrightarrow{M} HCO)}{HCO + HCO \rightarrow H_2CO + CO} \cdot \frac{HCO + HCO \rightarrow H_2CO + CO}{CO + 2H \rightarrow H_2CO} \cdot \frac{1}{CO}$$

However, photolysis of these compounds simply leads to CO recycling, the dominant mechanism in Titan oxygen chemistry.

4.4. Ion Chemistry

[43] Ion chemistry in Titan's atmosphere commences with the ionization of molecular nitrogen. Nitrogen is ionized by solar photons below 796 Å and by electron impact. These sources provide N_2^+ and N^+ ions that readily react with neutrals to form larger ions. Likewise, the abundance of CH_4 in the upper atmosphere makes its ionization an

important early process in the formation of the ionosphere. Methane is ionized by photons and electrons producing a number of hydrocarbon and hydrogen ions.

[44] The formation of most of the ions in Titan's ionosphere begin with the mechanism

(S30)
$$\begin{array}{c} N_2 + h\nu \to N_2^+ + e^- \\ N_2^+ + CH_4 \to CH_3^+ + N_2 + H \\ \underline{CH_3^+ + CH_4 \to C_2H_5^+ + H_2} \\ \hline 2CH_4 \to C_2H_5^+ + H_2 + H + e^- \end{array}$$

which serves to produce the abundant ion $C_2H_5^+$. H_2CN^+ , found to be the major ion in the *Keller et al.* [1992] and *Banaszkiewicz et al.* [2000] models, is produced through

(S31)
$$\begin{array}{c} 2CH_4 \to C_2H_5^+ + H_2 + H + e^- \\ \frac{C_2H_5^+ + HCN \to H_2CN^+ + C_2H_4}{2CH_4 + HCN \to H_2CN^+ + C_2H_4 + H_2 + H + e^-} \end{array}$$

and

(S32)
$$\begin{array}{c} N_{2} + h\nu \rightarrow N_{2}^{+} + e^{-} \\ N_{2}^{+} + H_{2} \rightarrow N_{2}H^{+} + H \\ N_{2}H^{+} + CH_{4} \rightarrow CH_{5}^{+} + N_{2} \\ \underline{CH_{5}^{+} + HCN \rightarrow H_{2}CN^{+} + CH_{4}} \\ \overline{HCN + H_{2} \rightarrow H_{2}CN^{+} + H + e^{-}} \end{array}$$

where (S31) is responsible for 78% of H_2CN^+ production and (S32) accounts for 10%. As indicated previously, the main function of H_2CN^+ ions is the formation of HCN through electron recombination (R349) and through reactions (R347) and scheme (S18), while $C_2H_5^+$ serves to convert nitriles to hydrocarbons through (S31).

5. Results and Discussion

5.1. Sensitivity to Eddy Diffusion and Methane Abundance

[45] A test of the accuracy of a photochemical model is comparison with observations of constituent abundances. However, exactly what is being considered when dealing with observations must be taken into account. The large error bars associated with the Vervack et al. (submitted manuscript, 2003) UVS reanalysis and the questions about the CH₄ and C₂H₂ observations that affect the retrieval of other species, as well, have been pointed out in section 3.5.2. However, the absence of other observations of constituent densities in Titan's upper atmosphere, with the exception of CH₄ and C₂H₂, and the difficulty in reconciling Titan chemistry with the Smith et al. [1982] >825 km C_2H_2 observation suggests the use of the Vervack non-CH₄ retrievals as a preliminary guide to how Titan's constituents might be distributed in the upper atmosphere. Furthermore, many of the existing observations in the stratosphere (e.g., IRIS) are determined assuming a uniform mole fraction profile, which is not realistic for chemically active constituents, affecting the abundances derived. For instance, Coustenis et al. [1989] derived a CO2 mole fraction of $1.4^{+0.3}_{-0.5} \times 10^{-8}$ at 105 km, the altitude where the contribution function of CO2 peaks. However, as dictated by the contribution function, the IRIS instrument received signal

contribution from layers above and below this layer. Consequently, an uncertainty of +55, -30 km is included with this result, associated with the region where the contribution function is greater than half of the peak value. The authors were able to fit the obtained spectra with a CO₂ mole fraction profile that reaches 1.4×10^{-8} at ~180 km and decreases to 5×10^{-9} at 105 km, outside the range of values indicated at the contribution peak assuming a uniform mole fraction profile. A similar case is found for H₂O as the stratospheric value of 4×10^{-10} is 20 times smaller than the value of the best-fitting H₂O mole fraction profile, provided by *Lara et al.* [1996], at the peak of the contribution function for the observation [*Coustenis et al.*, 1998]. Thus the assumption of a uniform profile must be kept in mind when comparing model results to observations.

[46] To demonstrate the effect of the eddy diffusion coefficient on the distribution of constituents, mole fraction profiles of key constituents for the eddy diffusion profiles demonstrated in Figure 6 are displayed in Figures 9 and 10. Figure 9 shows the results for CH_4 and HCN, key constituents used as guides in obtaining the eddy coefficient along with the observations referred to in section 3.5.2. The Vervack et al. (submitted manuscript, 2003) and *Vervack* [1997] UVS reanalysis observations displayed represent the extremes of both ingress and egress results, converted to mole fractions on the basis of the nominal total number density profile of this model.

[47] The nominal CH₄ profile fits the *Smith et al.* [1982] 1125 km observation along with the *Strobel et al.* [1992] 1400 km observation and the Vervack et al. (submitted manuscript, 2003) observations up to \sim 1000 km. SB approximates the upper limits of *Smith et al.* [1982] while HI matches Vervack et al. (submitted manuscript, 2003) up to higher levels but does not approach the *Strobel et al.* [1992] observation.

[48] For HCN the nominal profile fits the *Tanguy et al.* [1990] observations at 100 km and 200 km, as well as provides a good fit to the *Coustenis et al.* [1989] IRIS observations. Above 300 km, the nominal profile splits the divergent observations of *Tanguy et al.* [1990] and *Hidayat et al.* [1997] HI was generated to fit the *Hidayat et al.* [1997] HCN observations below 300 km as closely as possible, while simultaneously matching the CH_4 Vervack et al. (submitted manuscript, 2003) observations up to 1000 km, but is unable to completely match the HCN observations below 200 km. On the other hand, SB clearly overpredicts the HCN abundance in the lower stratosphere.

[49] In the upper atmosphere, the nominal profile provides a reasonable fit to the *Vervack* [1997] reanalysis, below 800 km, in all cases with the exception of C_2H_2 . The C_2H_2 nominal profile overpredicts acetylene abundance in relation to the *Vervack* [1997] results, consistent with the other photochemical models but on a lesser scale than the *Yung et al.* [1984] or *Toublanc et al.* [1995] models. Vervack et al. (submitted manuscript, 2003) provide reanalysis profiles assuming C_2H_2 to be the only absorbing species that exceeds the nominal profile. Furthermore, it is difficult to imagine a C_2H_2 profile with a shape that fits both the stratospheric and nominal upper atmosphere observations, simultaneously, absent a significant localized sink around to $10^{-1}-10^{-2}$ millibar region. This raises the possibility that more absorption should be ascribed to

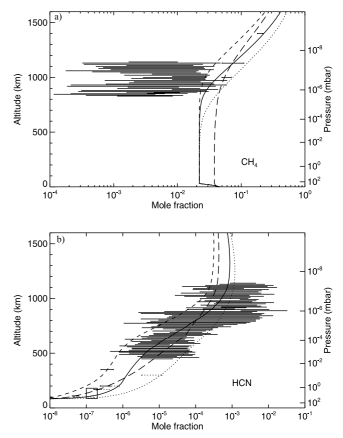


Figure 9. a) CH₄ and b) HCN profiles using the nominal eddy diffusion profile (NOM, solid line), *Steiner and Bauer* [1990] profile (SB, dotted line), the high homopause profile (HI, short dashed), and the *Lara et al.* [2002] profile (NL, long dashed line). The group of horizontal lines represent the *Vervack* [1997] reanalysis of UVS observations. In a) the horizontal bars at 1000 km and 1130 km represent the *Smith et al.* [1982] observations, while the horizontal bar at 1400 km indicates the *Strobel et al.* [1992] Voyager UVS observations. In b) the solid horizontal lines in the lower portion of the plot shows the *Hidayat et al.* [1997] observations with the dotted lines representing the *Tanguy et al.* [1990] results, and the box represents the stratospheric Voyager IRIS observations including error bars in abundance and altitude.

 C_2H_2 in the UVS reanalysis than their nominal observations suggest. Above 800 km, NOM falls off with respect to the *Vervack* [1997] observations for C_4H_2 and HC₃N. However, these constituents have short lifetimes in this region of the atmosphere compared with a solar cycle and thus are sensitive to changes in the solar radiation output, as discussed in the next section, and assuming solar maximum conditions provides a better fit to this data. Furthermore, as *Vervack* [1997] points out, the uniformity in number density with respect to altitude for the C_4H_2 and HC₃N observations suggest that these species may not have been well-retrieved. Misassignment of CH₄ absorption may have understated CH₄ densities at high altitudes while overstating these species.

[50] The high homopause eddy diffusion profile limits the amount of methane in the upper atmosphere, resulting in

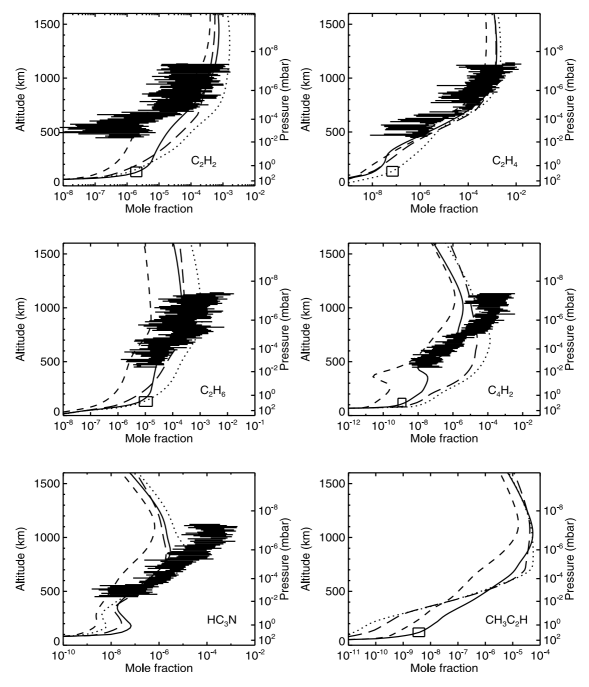


Figure 10. Constituent profiles using the nominal eddy diffusion profile (NOM, solid line), *Steiner and Bauer* [1990] profile (SB, dotted line), the high homopause profile (HI, short dashed), and the *Lara et al.* [2002] profile (NL, long dashed line). The boxes represent the stratospheric Voyager IRIS observations including error bars in abundance and altitude, while the horizontal lines represent the *Vervack* [1997] reanalysis of UVS observations.

less conversion to higher order hydrocarbons and, consequently, smaller hydrocarbon abundances. HI matches the C_2H_2 and C_2H_4 upper atmosphere observations well, but underpredicts for C_2H_6 , C_4H_2 and HC_3N . SB, on the other hand, provides reasonable predictions only for HC_3N and C_2H_6 above 700 km, according to the Vervack et al. (submitted manuscript, 2003) observations. The HI profile significantly underpredicts constituent abundances in the stratosphere, while SB matches IRIS observations only for the C_2 -hydrocarbons. [51] Recently, *Lara et al.* [2002] reanalyzed this question of the eddy diffusion coefficient in relation to the methane density in the upper atmosphere. They proceeded by recalculating the methane density with an assumption of CH_4 stratospheric mole fraction at 3.8%, compared to the *Lara et al.* [1996] assumption of 1.7%. This value assumes very high methane supersaturation in the troposphere, which according to *Courtin et al.* [1995], can be as high as 230%. In contrast, the nominal model uses the results of *Samuelson et al.* [1997], which places the level of maxi-

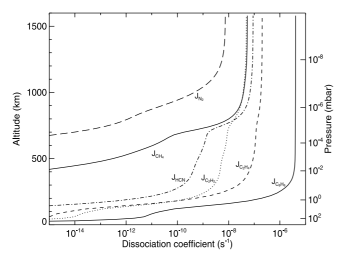


Figure 11. Photodissociation coefficients of key species.

mum methane supersaturation at 1.48 ± 0.11 , using a methodology that, according to the authors, avoids some of the sources of systematic error inherent in the Courtin et al. [1995] study. A methane supersaturation value of 1.37 yields of CH₄ stratospheric mole fraction of 2.2%. Nevertheless, with the large methane abundance, Lara et al. [2002] adopt an eddy diffusion profile (NL) that increases rapidly in the stratosphere and middle atmosphere, resulting in a high homopause level around 1000 km (Figure 6b). This profile with a CH₄ stratospheric mole fraction of 3.8% was tested along with the nominal case. As Figure 9 shows, NL provides good agreement with the HCN infrared and microwave observations, which eliminates the necessity for a large sink for HCN to the haze due to polymerization, as promoted by Lara et al. [1999]. However, demonstrated by Figure 10, the NL case underpredicts hydrocarbons in the lower stratosphere, with the exception of C₄H₂. C₃-hydrocarbons are especially diminished in the NL case, with a CH₃C₂H profile that is two orders of magnitude smaller and a C_3H_8 profile (not shown) that is an order of magnitude smaller than IRIS observations. In the upper atmosphere, the nominal case provides profiles more consistent with the UVS reanalysis than the NL case, as well.

5.2. Sensitivity to Solar Flux and Aerosol Structure

[52] The stable constituents in Titan's atmosphere are dissociated and ionized by solar photons and electrons that are transferred through the atmosphere. A comparison of Figure 11 and absorption cross sections will yield the deposition regions of different wavelength regions. N_2 , which is dissociated by EUV photons, undergoes attenuation in dissociation in the upper atmosphere, with $J_{N_{a}}$ having been reduced by an order of magnitude at 1000 km. Below 900 km, electron impact dominates N_2 absorption as J_{N_2} drops off at the base of the ionosphere at 720 km. Abundant species that undergo significant absorption of photons at 1000 Å $\leq \lambda \leq$ 1450 Å experience dissociation attenuation around 800 km, as these photons are deposited through methane absorption. Deposition of UV radiation longer than 1450 A occurs lower in the atmosphere, at 200-300 km, as a result of absorption by molecules such as HCN ($\lambda \leq 1800$ Å) and C₂H₂ ($\lambda \leq 2100$ Å) as well as by aerosols ($\lambda > 1800$ A).

[53] During much of the upcoming Cassini-Huygens encounter with Titan, the Sun will undergo moderate solar activity, likely resulting in a reduced solar flux compared with what took place during the Voyager flybys. In Titan's atmosphere, constituents above 500 km are typically destroyed chemically within one solar cycle, while for certain stable constituents, like C₄H₂, their relatively short lifetimes may allow changes in solar output to affect their lower stratospheric abundances, as well. With this in mind, it is constructive to analyze the effect of variations in solar flux on the distribution of various key constituents in Titan's atmosphere. Figure 12 provides such a comparison, assuming solar fluxes in the FUV and NUV obtained by SOLSTICE for average solar maximum, solar moderate, and solar minimum conditions, with the corresponding EUV calculated fluxes for (F10.7, F10.7A) = (233,211.9), (130, 130), and (70, 70), respectively. The C_2 hydrocarbons, which are more directly affected by methane and its dependence in the upper atmosphere on transport rather than chemistry, show little dependence on solar activity. However, nitriles like HCN and HC₃N show much more sensitivity in the upper atmosphere to solar output as HCN photolysis plays a considerably larger role with respect to transport than CH₄ photolysis.

[54] In comparing the nominal dissociation coefficients with the photodissociation coefficients of Toublanc et al. [1995] and Lara et al. [1996] the most glaring differences reside in the lower 300 km. The nominal dissociation coefficients fall off much more rapidly than Lara et al. [1996] most likely due to differences in the treatment of aerosols. Solar radiation at wavelengths 1800 Å $\leq \lambda \leq$ 2100 Å is relatively unattenuated in the Lara et al. [1996] model, reaching to lower regions, while those photons are deposited above 200 km in the nominal model due to C₂H₂ absorption and aerosol opacity. The assertion of significant opacity at these wavelengths is corroborated by the analysis of UV spectra by McGrath et al. [1998]. The Toublanc et al. [1995] photodissociation rates fall off more significantly than Lara et al. [1996] which used the Yung et al. [1984] parameterization of aerosol absorption, but less than the nominal case, probably a result of consideration of Mie haze particles. For instance, $J_{C_2}H_2(100 \text{ km})/J_{C_2}H_2(500 \text{ km}) =$ 1.0×10^{-5} in the nominal model while *Toublanc et al.* [1995] and Lara et al. [1996] yield 4.8 \times 10⁻² and 8.2 \times 10^{-3} , respectively. Fractal particles are more opaque than Mie particles at short wavelengths [Rannou et al., 1995], limiting the penetration of radiation at deeper levels. However, aerosol opacities amount to a significant uncertainty affecting photodissociation coefficients, as Titan aerosol densities and optical constants are still not well understood. The effect of the type of aerosol particle considered is shown in Figure 13, which compares constituent mole fraction profiles from the nominal case with those assuming Mie opacities derived from Rannou et al. [1995]. In the Mie case more radiation is allowed to penetrate into the lower atmosphere, promoting greater formation of CH₃ radical through catalytic dissociation of methane (S5) and increasing the levels of ethane and propane in the atmosphere. Furthermore, the smaller opacity provided by Mie particles longward of 1800 A results in much larger dissociation of HC₃N and C₂H₃CN molecules which significantly absorb in that wavelength region. The IRIS HC₃N upper limit of

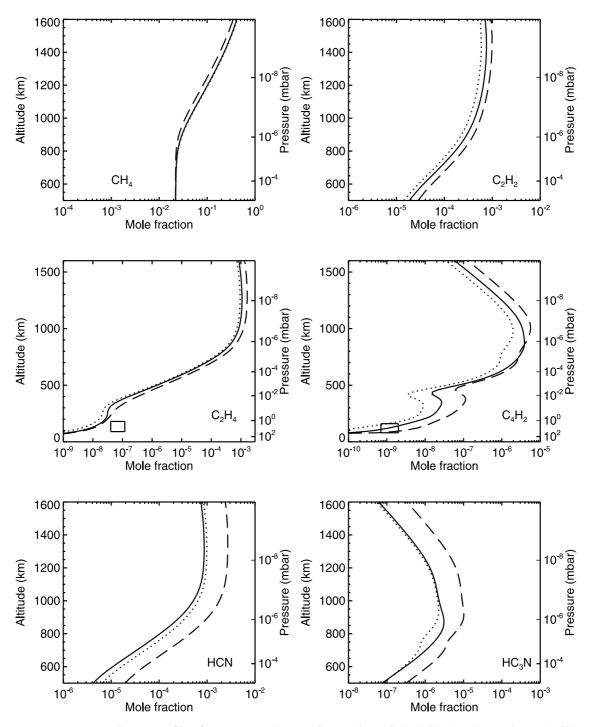


Figure 12. Constituent profiles for average solar maximum (long dashed line), solar moderate (solid line), and solar minimum conditions (dotted line).

 1.5×10^{-9} and the lack of observations of C₂H₃CN suggest that more absorption of these molecules takes place in the lower atmosphere than what is determined by the nominal profile. These factors along with the C₃H₈ Mie profile hint at possibly less aerosol opacity in Titan's stratosphere than what is considered in the nominal case.

5.3. Hydrocarbons

[55] Figure 10 shows that the model slightly underpredicts ethylene density in Titan's stratosphere. This issue is a factor in the *Toublanc et al.* [1995] model, as well. *Lara et al.* [1996] assumed a boundary condition of 1.5×10^{-7} to match IRIS observations, necessitating a flux of 7.0 $\times 10^7$ cm⁻² s⁻¹ from the surface, resulting from some surficial process (e.g., outgassing). The nominal model calculates a C₂H₄ net loss of 8.1×10^7 cm⁻² s⁻¹ (Table 7), requiring a corresponding flux from the surface in order to maintain steady state equilibrium. Such a flux, which would not be cold-trapped as C₂H₄ does not condense at those abundances, is a viable mechanism to account for the IRIS obser-

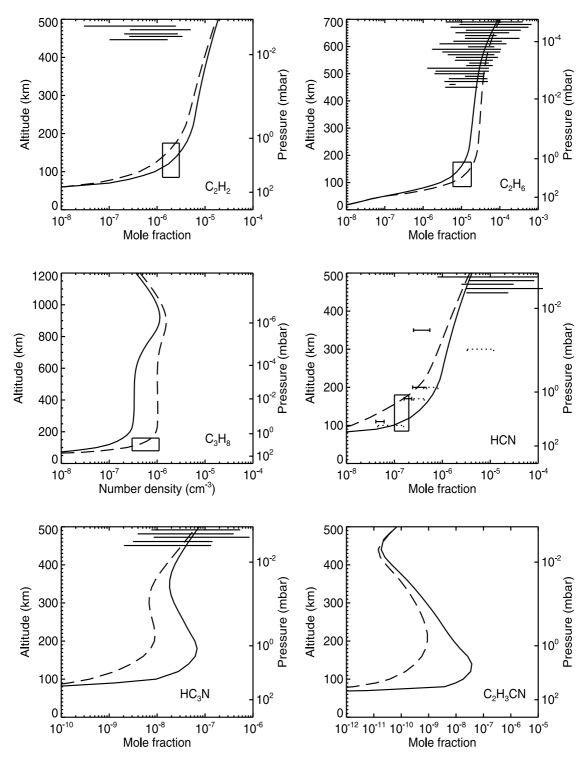


Figure 13. Constituent profiles assuming fractal haze particles (solid line) and Mie haze particles (long dashed line). The boxes represent the stratospheric Voyager IRIS observations including error bars in abundance and altitude, while the horizontal lines represent the *Vervack* [1997] reanalysis of UVS observations. For HCN, the solid horizontal bars show the *Hidayat et al.* [1997] observations with the dotted bars representing the *Tanguy et al.* [1990] observations.

vations. Possible irradiation of hydrocarbon condensates on Titan's surface may also provide a source of C_2H_4 in the stratosphere.

[56] Canosa et al. [1997] measured the $CH + CH_4$ rate coefficient in a temperature range of 23–295 K, but could

not positively identify products. Thermochemical and kinetic analysis indicate, however, that H-atom elimination is the most favored process [*Canosa et al.*, 1997], yielding the C_3H_4 complex. With potential products CH_3C_2H and CH_2CCH_2 similarly exothermic, these isomers are assumed

Species ^a	$\begin{array}{c} \text{Production} (P), \\ \text{cm}^{-2} \text{ s}^{-1} \end{array}$	$\begin{array}{c} \text{Loss (L),} \\ \text{cm}^{-2} \text{ s}^{-1} \end{array}$	$P - L, cm^{-2} s^{-1}$	Chemical Lifetime at 300 km, s
N ₂	1.1×10^{7}	2.9×10^{8}	-2.8×10^{8}	8.4×10^{15}
H ₂	3.2×10^{9}	3.9×10^7	3.2×10^9	8.7×10^{12}
CH ₄	1.1×10^{8}	4.9×10^{9}	-4.8×10^{9}	8.4×10^{11}
C_2H_2	3.3×10^{9}	2.9×10^{9}	3.7×10^{8}	3.1×10^{8}
C_2H_2 C_2H_4	6.7×10^{8}	7.6×10^{8}	-9.3×10^{7}	3.7×10^{7}
C_2H_6	1.7×10^{9}	1.3×10^8	1.6×10^{9}	2.3×10^{10}
CH ₃ C ₂ H	1.9×10^8	2.0×10^8	-7.6×10^{6}	2.8×10^{7}
CH ₂ CCH ₂	1.5×10^{8}	1.5×10^8	-1.1×10^{6}	2.5×10^{6}
$C_3 \tilde{H_6}$	1.1×10^{8}	1.1×10^8	6.6×10^{5}	3.9×10^{6}
C ₃ H ₈	3.7×10^7	4.1×10^{6}	3.3×10^7	1.0×10^{10}
C_4H_2	1.8×10^{9}	1.8×10^{9}	-2.2×10^{6}	1.4×10^{6}
C_4H_6	9.8×10^{7}	9.8×10^{7}	6.5×10^{4}	4.4×10^{4}
C_4H_8	2.7×10^{6}	2.6×10^{6}	9.2×10^{4}	3.9×10^{6}
C_4H_{10}	3.6×10^{7}	6.2×10^{6}	3.0×10^{7}	5.5×10^{9}
C_6H_2	3.7×10^{6}	3.7×10^{6}	2.5×10^{4}	3.4×10^{6}
C_6H_6	2.0×10^{8}	2.0×10^{8}	4.6×10^{5}	4.8×10^{5}
C_8H_2	5.7×10^{4}	5.8×10^{4}	-1.0×10^{3}	3.4×10^{6}
CO	1.8×10^{7}	2.2×10^{7}	-4.1×10^{6}	7.8×10^{11}
CO_2	3.3×10^{6}	2.0×10^{5}	3.1×10^{6}	2.2×10^{10}
H_2O	3.9×10^{6}	7.3×10^{6}	-3.4×10^{6}	1.3×10^{8}
H_2CO	1.3×10^{6}	3.7×10^{5}	9.1×10^{5}	7.7×10^{7}
CH ₃ OH	4.3×10^{3}	4.6×10^{3}	-2.6×10^{2}	1.0×10^{8}
CH ₂ CO	4.6×10^{6}	4.6×10^{6}	9.3×10^{3}	1.9×10^{6}
HCN	4.1×10^{8}	4.0×10^{8}	2.1×10^{6}	1.4×10^{9}
CH ₃ CN	8.5×10^{6}	8.5×10^{6}	-3.4×10^{4}	1.3×10^{9}
C_2N_2	2.4×10^{6}	3.5×10^{5}	2.0×10^{6}	8.0×10^{6}
HC_3N	2.0×10^{8}	1.9×10^{8}	1.3×10^{7}	2.5×10^{7}
C_2H_3CN	2.5×10^{8}	2.3×10^{8}	1.6×10^{7}	1.6×10^{6}
C_4N_2	7.3×10^{6}	6.4×10^{6}	9.7×10^{5}	3.7×10^{6}
N ⁺	4.7×10^{7}	4.7×10^{7}		
N_2^+	1.6×10^{8}	1.6×10^{8}		
CH_3^+	1.5×10^{8}	1.5×10^{8}		
CH_{4}^{+}	1.0×10^{7}	1.0×10^{7}		
$C_2H_5^+$	9.3×10^{7}	9.3×10^{7}		
H_2CN^+	1.5×10^{8}	1.5×10^{8}		
<u>e</u>	1.8×10^8	1.8×10^8		

 Table 7. Production and Loss Rates and Chemical Lifetimes of Various Stable Species

^aSpecies that have long lifetimes compared to a solar cycle are italicized.

in this model to have a branching ratio of 0.5 for this reaction. This mechanism for forming methylacetylene and allene is not considered in other Titan photochemical models. With this assumption, the CH_3C_2H profile shown in Figure 10 agrees well with observations as opposed to *Toublanc et al.* [1995], which is a factor of six too large or *Lara et al.* [1996], which is two orders of magnitude too small (Table 8).

[57] Notwithstanding the effect of aerosol opacity, the model-generated propane profile has a similar uniform shape in the homosphere as the Yung et al. [1984], Lara et al. [1996], and Lebonnois et al. [2001]. The Toublanc et al. [1995] C_3H_8 profile shape differs with propane peaking at 500 km, mainly a product of their chemical scheme which results in a C₃H₈ net production rate 15 times larger than the Lara et al. [1996] or our nominal rate. In our model, the $C_{3}H_{8}$ abundance is governed by the synthesis of ethyl and methyl radicals (R51b) above 140 km, and the self-reaction of C₃H₇ radicals (R81) below 140 km, which facilitates the recycling of propane in the stratosphere, via H₂-elimination in propane photolysis (J15a) and subsequent hydrogen attachment (R72). The omission of this recycling mechanism in the Lara et al. [1996] scheme is responsible for their C₃H₈ fall-off above 100 km. 89% of propane created is lost to condensation below 50 km.

[58] The diacetylene profile compares well with the IRIS observations while both *Toublanc et al.* [1995] and *Lara et al.* [1996] overpredict C_4H_2 abundance, by a factor of three and two, respectively, due to smaller eddy mixing in the stratosphere. The C_4H_2 profile shape compares similarly to the *Lebonnois et al.* [2001] C_4H_2 equatorial profile shape, although the nominal profile exhibits larger diacetylene densities in the stratosphere as a result of larger acetylene mixing ratios, compared to the *Lebonnois et al.* [2001] C_2H_2 equatorial profile. Diacetylene reacts with the photo-

Table 8. A Comparison of Model-Generated Species Abundances Along With Available Observations^a

	-		1		0		
Species	Altitude, km	Observational Limits	Y84	T95	La96/B00	Le01/Le02	Nominal With Fractal/Mie Haze
C_2H_2	125	1.3-2.9(-6) (IRIS)	4.3(-5)	2.2(-6)	$3.0(-6)^{b}$	1.9(-6)	1.9(-6)/1.1(-6)
C_2H_4	125	0.4 - 1.2(-7) (IRIS)	3.1(-7)	3.2(-9)	8.3(-8)	2.1(-8)	9.4(-9)/1.5(-8)
C_2H_6	125	0.6 - 1.8(-5) (IRIS)	2.0(-4)	1.2(-5)	8.7(-6)	2.7(-6)	5.8(-6)/1.2(-5)
CH_3C_2H	105	2.3-6.1(-9) (IRIS)	9.5(-7)	1.4(-8)	2.3(-11)/<1.0(-11)	9.8(-10)	1.8(-9)/6.6(-10)
C_3H_8	105	0.3 - 1.1(-6) (IRIS)	4.2(-6)	2.8(-7)	$1.0(-7)^{c}$	2.4(-7)	6.3(-8)/2.8(-7)
C_4H_2	105	0.7-2.0(-9) (IRIS)	1.6(-10)	6.8(-9)	$4.7(-9)^{d}$	3.9(-9)	6.2(-10)/1.9(-9)
C_6H_6	110	1.0-7.0(-10) (ISO)	_	_	-	<1.0(-13)	6.1(-10)/5.8(-11)
CO_2	105	0.9 - 1.7(-8) (IRIS)	5.7(-9)	4.6(-13)	5.5(-9)	-	6.2(-9)/5.8(-9)
H_2O	400	0.4 - 1.4(-8) (ISO)	1.2(-9)	3.1(-9)	1.9(-8)	-	1.1(-8)/1.0(-8)
HCN	110	0.5 - 1.6(-7) (IRAM)	3.8(-6)	1.6(-7)	1.2(-7)/4.0(-8)	1.3(-7)	1.4(-7)/3.2(-7)
	300	0.03 - 1.2(-5) (IRAM)	9.7(-6)	2.2(-6)	6.4(-6)/2.3(-6)	7.9(-7)	1.3(-6)/2.0(-6)
CH ₃ CN	450	2.2-6.2(-8) (IRAM)	_	1.2(-7)	$7.3(-9)^{\rm e}$	7.8(-7)	9.6(-9)/8.0(-9)
HC_3N	105	1.5 - 8.5(-10) (ISO)	3.3(-7)	<1.0(-12)	2.4(-8)/1.4(-8)	3.2(-8)	1.2(-8)/1.2(-10)
	450	2.2-6.2(-8) (IRAM)	3.1(-6)	4.1(-6)	3.9(-6)/3.1(-6)	1.3(-6)	4.0(-8)/3.7(-8)
C_2N_2	105	$\leq 1.0(-9)$ (IRIS)	1.1(-7)	<1.0(-12)	4.5(-12)/1.6(-11)	1.5(-9)	1.1(-9)/1.6(-13)
$C_2H_3CN^{f}$	105	_	_	_	-	6.6(-9)	2.1(-8)/8.8(-12)
$C_4 N_2^{t}$	105	_	_	_	1.2(-11)/1.7(-12)	-	1.4(-9)/6.2(-12)
H ₂ CO ^f	120	—	<1.0(-11)	9.1(-12)	-	_	1.0(-9)/4.1(-9)
CH ₃ OH ^f	120	—	_	1.6(-9)	-	_	4.5(-13)/1.0(-14)
CH_2CO^{f}	120	-	<1.0(-11)	3.2(-9)	-	_	2.4(-11)/2.2(-11)

^aRead 1.0(-9) as 1.0×10^{-9} . Y84 = Yung et al. [1984]; T95 = Toublanc et al. [1995]; La96 = Lara et al. [1996]; B00 = Banaszkiewicz et al. [2000]; Le01/Le02 = Lebonnois et al. [2001] (equator), Lebonnois et al. [2002].

^bMixing ratio at 130 km.

^cMixing ratio at 110 km.

^dMixing ratio at 125 km.

^eMixing ratio at 400 km.

^fThis species has not yet been observed in Titan's atmosphere.

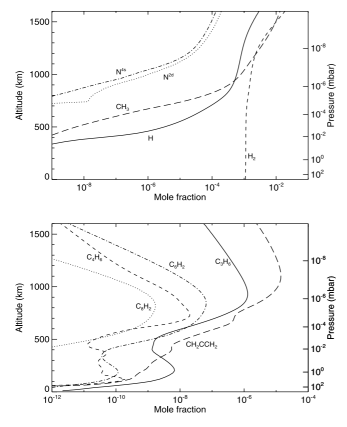


Figure 14. Nominal constituent profiles for various hydrocarbon species along with atomic nitrogen, atomic hydrogen, and molecular hydrogen.

lytic products of C_2H_2 and C_4H_2 to form C_6H_2 and C_8H_2 , whose profiles are shown in Figure 14. These processes may have implications in the formation of Titan haze between 500–800 km.

[59] Wilson et al. [2003] suggested the possibility of benzene formation in Titan's atmosphere, corroborated by the tentative detection of benzene by the Infrared Space Observatory (ISO) [Coustenis et al., 2003]. The observation fit by a uniform mole fraction profile of $4 \pm 3 \times 10^{-10}$ was matched by a vertical profile of benzene from Wilson et al. [2003], multiplied by 3.0 ± 0.5 , an upper-limit profile calculated assuming reaction rates for aromatic compounds that correspond to a temperature of 300 K. However, aerosol opacity was not considered in that calculation. Haze particles act to shield benzene from photodissociation, which extends as far as 2700 Å [Pantos et al., 1978]. Figure 15 compares the nominal benzene profile from Wilson et al. [2003] with the present nominal profiles including aerosols and calculated with the nominal temperature profile, taken from Yelle et al. [1997], which ranges from 71 K at the tropopause to 175 K in the thermosphere. The present profile matches the ISO observation with a column density of 2.4×10^{15} cm⁻² s⁻¹ above 30 mbar, with a benzene mole fraction of 6.1×10^{-10} at 110 km. The only other modeling study that covers benzene abundance on Titan is Lebonnois et al. [2002], which calculates a benzene mole fraction at least three orders of magnitude smaller at the observation altitude. Figure 15 also demonstrates the large dependence on solar flux for benzene

abundance in the upper atmosphere. *Wilson and Atreya* [2003] explored possible mechanisms for haze formation and suggested that aromatic compounds like benzene could be a source of Titan haze. Assuming this nominal benzene profile and that the mechanism for haze formation begins with the reactions

(R115)
$$C_6H_6 + C_2H \rightarrow haze$$

(R112) $C_6H_5 + C_2H_2 \rightarrow haze,$

the aromatic pathway provides a haze production rate of 9.5×10^6 cm⁻² s⁻¹ or 3.2×10^{-14} g cm⁻² s⁻¹, assuming a nucleation mass of 2000 amu [*Richter and Howard*, 2000], compared with $0.5-2 \times 10^{-14}$ g cm⁻² s⁻¹ derived by microphysical models [*McKay et al.*, 2001].

[60] An important uncertainty in modeling chemistry in outer planetary atmospheres is the methyl recombination rate. Methyl recombination serves as the primary loss mechanism for the chemically important methyl radical as well as the source of ethane production. However, until recently, rate measurements have only been taken as low as room temperature, with extrapolations down to lower temperatures yielding widely varying results [Atreva et al., 1999]. ISO observations of CH₃ seem to indicate that the more widely used Slagle et al. [1988] rate expression is too low, perhaps by an order of magnitude. For this reason, we use a methyl recombination rate that is ten times that obtained with the Slagle et al. [1988] expression. Studies comparing hydrocarbon mixing ratio profiles using both the Slagle et al. [1988] rate and our rate indicate little difference in the ethane stratospheric profile and a factor a two increase in ethane density in the upper atmosphere with our rate [Wilson and Atreya, 2000a]. Recently, Cody et al. [2003] have measured methyl recombination at lower temperatures, obtaining results at as low as 155 K. These results find

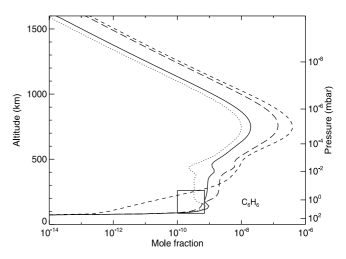


Figure 15. Benzene profiles for average solar maximum (long dashed line), solar moderate (solid line), and solar minimum conditions (dotted line), along with the *Wilson et al.* [2003] nominal benzene profile (short dashed line). The box represents the ISO benzene observations including error bars in abundance and altitude.

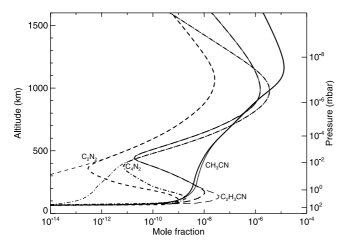


Figure 16. Constituent profiles of various nitrile species including cosmic ray deposition (thick line) and no cosmic ray deposition (thin line).

methyl recombination to proceed at roughly 1.6 times the *Slagle et al.* [1988] rate at high pressures. However, these experiments were conducted with He as a bath gas, and the reaction may proceed faster with N₂ as the background atmosphere as exhibited by the hydrogen atom recombination reaction, which proceeds 60% faster at 150 K with the *Tsang and Hampson* [1986] rate which uses an N₂ bath gas, as opposed to the *Baulch et al.* [1992] rate which uses an H₂ bath gas. Using the *Cody et al.* [2003] rate, the nominal ethane density in the upper atmosphere is reduced by 55% at 1100 km and reduced by 6% at 200 km, while the CH₃ density is increased by 145% and 35%, respectively, at those altitudes.

5.4. Nitriles

[61] Cosmic rays can play a substantial role in the formation of nitriles. Cosmic rays penetrate the atmosphere and dissociate nitrogen molecules (Figure 4b), producing N atoms that serve as a source for stratospheric nitriles, in particular, nitriles that are formed directly from N-atom addition. In addition, Molina-Cuberos et al. [2002] demonstrated that the possible formation of ion clusters in this region through high pressure reactions may be a source of stratospheric nitriles. The kinetics and identification of products of these reactions, however, are still to be determined. Figure 16 shows the profiles of several nitrile compounds with and without the inclusion of galactic cosmic rays. In particular, the stratospheric abundances of C₂N₂ and C₄N₂ are highly dependent on cosmic ray dissociation of nitrogen, through the schemes (S20) and (S21). Treatment of cosmic ray interaction through particle cascade, illustrated by Capone et al. [1983], may increase C₂N₂ and C₄N₂ densities in the lower atmosphere further.

[62] Lara et al. [1996] in order to explain the CH₃CN observations announced by *Bézard et al.* [1993] propose a source provided by CH₄ + CN and C₂H₆ + CN. Previous studies [*Hess et al.*, 1989; *Yang et al.*, 1992b; *Sims et al.*, 1993], however, give no indication of an acetonitrile channel, while *Balla et al.* [1991] point out that though the CH₃CN channel is thermodynamically possible, their mea-

surements only indicate traces of CH₃CN. Nevertheless, Lara et al. assume a quantum yield for the production of 0.05 for CH₄ + CN and 0.035 for C₂H₆ + CN. The reaction N^{2d} + C₂H₄ was taken to proceed at a rate of 1.2×10^{-12} cm⁻³ s⁻¹ by Lellouch et al. [1994] in their Neptune model, on the basis of N^{2d} quenching coefficients by Black et al. [1969]. They predicted the reaction to proceed in an insertion/abstraction manner, producing products NH and C₂H₃. This reaction was included in the Lara et al. model, while Toublanc et al. [1995] and Yung [1987] did not include this reaction. However, Sato et al. [1999] measured N^{2d} + C₂H₄ down to 230 K and obtained a rate of 2.6 × 10^{-11} cm⁻³ s⁻¹, a rate over 20 times that assumed by Lellouch et al. [1994] and Lara et al. [1996]. Furthermore, crossed-beam experiments conducted by Balucani et al. [2000] indicate CH₃CN to be the likely product

$$(R127) N^{2d} + C_2H_4 \rightarrow CH_3CN + H_4$$

Including (R127) with the *Sato et al.* [1999] rate produces a profile (Figure 16) that matches observations reasonably well, producing acetonitrile primarily in the upper atmosphere through the scheme

0.1

(S33)
$$\begin{array}{c} 0.5(N_2 + h\nu \to 2N^{2d}) \\ \frac{N^{2d} + C_2H_4 \to CH_3CN + H}{0.5N_2 + C_2H_4 \to CH_3CN + H} \end{array} .$$

[63] Notwithstanding the nominal C2H3CN profile (Figure 16), which presents a stratospheric mixing ratio as high as 3.8×10^{-8} , observations suggest the lack of significant abundance of acrylonitrile in this region of the atmosphere. C₂H₃CN has not been identified in the atmosphere of Titan by IRIS or ISO observations. Furthermore, Voyager infrared spectra analysis do not seem to advocate the presence of acrylonitrile condensates in larger abundances than say C_2N_2 or C_4N_2 . In the nominal model, acrylonitrile stratospheric production, despite the low rate coefficient for (R156), is clearly dominated by (S23) over (S22). Monks et al. [1993] assume an addition-decomposition process as the favored mechanism of (R156) through analogy with (R155) and (R29). However, a definitive yield for acrylonitrile production was not given as only trace amounts of C₂H₃CN were identified in the experiment. Thus an excessively large reaction rate for acrylonitrile production through (R156), as well as excessive aerosol opacity in the NUV as discussed in section 5.2, are possible explanations for the large stratospheric abundance of C₂H₃CN calculated by the model.

5.5. Will Ammonia Be Detected?

[64] Ammonia has not been detected in Titan's atmosphere but does serve prominently in Saturn's atmosphere and may have played a significant role in the formation of Titan's present nitrogen inventory [*Atreya et al.*, 1978; *Atreya and Wilson*, 2001]. *Bernard et al.* [2003] detected ammonia in their electron discharge simulations of Titan's atmosphere. They correctly point out that the source of ammonia molecules would likely arise through chargedparticle chemistry. Apart from a supply of primordial NH₃ from the interior, any ammonia in Titan's strato-

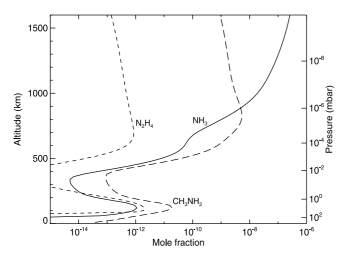


Figure 17. Constituent profiles for various amine species.

sphere will be the result of cosmic-ray ionization of N_2 , through the mechanism

(S34)

$$N_{2} + GCR \rightarrow N^{+} + N + e^{-}$$

$$CH_{4} + N^{+} \rightarrow HCN^{+} + H_{2} + H$$

$$CH_{4} + HCN^{+} \rightarrow C_{2}H_{3}^{+} + NH_{2}$$

$$\frac{NH_{2} + H \xrightarrow{M} NH_{3}}{NH_{2} + 2CH_{4} \rightarrow NH_{3} + N + H_{2} + C_{2}H_{3}^{+} + e^{-}$$

[65] However, this mechanism results in an ammonia mole fraction of only 9.6×10^{-13} at 120 km (Figure 17). The profile of NH₃ indicates a decrease in mole fraction above this level as a result of smaller cosmic ray precipitation, followed by an increasing profile with altitude at higher levels due to the greater influence of precipitating electrons in the upper atmosphere. In this region, electron recombination of ammonium ions, as suggested by *Atreya* [1986], is the most likely ammonia source. The larger estimate for NH₃ by *Bernard et al.* [2003] is most likely a result of the use of electron discharge as the primary energy source.

[66] The amino radical (NH₂), produced through (R336a), is responsible for the formation of other amine species, such as $N_2H_4(R174b)$ and $CH_3NH_2(R175)$. Figure 17 shows methylamine (CH₃NH₂) as the most abundant of the amine species in the stratosphere. However, barring a significant source of primordial ammonia, the low densitites of ammonia may preclude detection by Cassini-Huygens.

5.6. Oxygen Species

[67] Buoyed by the recent detection of H₂O on the giant planets, *Feuchtgruber et al.* [1997] estimate a water influx of $3-50 \times 10^5$ cm⁻² s⁻¹. With the recent detection of H₂O in Titan's atmosphere, it is evident that this applies to Titan as well. CO is the most abundant oxygen-bearing molecule in Titan's atmosphere. Sensitive millimeter observations have found a uniformly mixed carbon monoxide profile at ~5 × 10^{-5} [*Muhleman et al.*, 1984; *Gurwell and Muhleman*, 1995], although further millimeter observations [*Hidayat et al.*, 1998] and near infrared observations [*Noll et al.*, 1996] have suggested lower abundances with stratospheric deple-

tion. Using a surface CO mole fraction of 5×10^{-5} from the *Gurwell and Muhleman* [1995] finding of $5 \pm 1 \times 10^{-5}$ and an external H₂O influx of 5×10^{6} cm⁻² s⁻¹ [*Feuchtgruber et al.*, 1997], abundances are calculated for oxygen-bearing species, shown in Figure 18.

[68] ISO confirmed the existence of water vapor in Titan's atmosphere as *Coustenis et al.* [1998] found a uniform H₂O profile at 4×10^{-10} consistent with the obtained H₂O spectrum. However, scaling the *Lara et al.* [1996] H₂O model profile by a factor of 0.4 yielded a profile with an H₂O mole fraction of $8^{+6}_{-4} \times 10^{-9}$ at 400 km to fit the ISO data. The nominal model finds agreement with this observation as shown in Figure 18.

[69] With (R219) as the primary source of CO₂, the IRIS carbon dioxide observations argue for an external source of water. *Coustenis et al.* [1989] found a uniform mole fraction profile of $1.4^{+0.3}_{-0.5} \times 10^{-8}$ to fit IRIS observations, although a linear scaling of a altitude-dependent CO₂ profile of *Samuelson et al.* [1983], yielding a CO₂ mole fraction of 1.4×10^{-8} at 1.5 mbar where the value of the CO₂ contribution function was about 35% of the peak value at 4 mbar, provided a decent fit. The nominal profile of CO₂ is consistent with the IRIS observations, rendering a CO₂ mole fraction of 1.4×10^{-8} at 4 mbar.

[70] The nominal model predicts much lower abundances for CH₂CO and CH₃OH in the stratosphere than *Toublanc et al.* [1995] as a result of including photolysis which provides

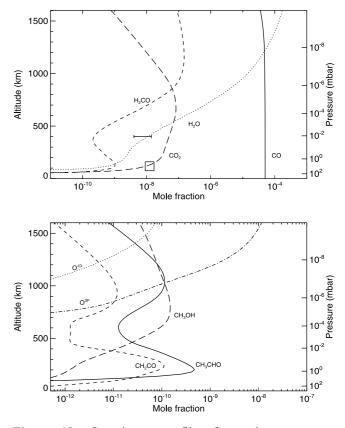


Figure 18. Constituent profiles for various oxygenbearing species with the horizontal bar representing the ISO H_2O observations based on the *Lara et al.* [1996] profile, and the box representing the Voyager IRIS CO_2 observations.

the main sink for these compounds. *Toublanc et al.* [1995] refer the absorption cross section of ketene to that of H₂O, although the absorption cross section for CH₂CO extends past 3000 Å [*Rabalais et al.*, 1971], much further than that of H₂O (2000 Å). Furthermore, the nominal model includes the measurement of the CH₃OH cross section [*Wodtke and Lee*, 1987], not included in previous models.

[71] However, formaldehyde is predicted to be in greater abundance than in previous models, perhaps near the levels of detection in the stratosphere. H₂CO is formed through self-reaction of formic acid (HCO), which is produced via pressure-dependent H-attachment to CO, demonstrated in (S29). The calculated H₂CO mole fraction of 1.0×10^{-9} at 4 mbar and the corresponding condensation at the tropopause indicate the possibility of detection of H₂CO gas and condensate by the GCMS, CIRS, and ACP instruments during the upcoming Cassini-Huygens exploration.

[72] The significant abundance of formaldehyde also suggests the possibility of detection of ethylene oxide (C_2H_4O) in Titan's stratosphere. Ethylene oxide was detected in the Titan simulations of Bernard et al. [2003], although they did not detect formaldehyde, presumably a result of the lack of water in their experiments. The isomer of ethylene oxide, acetaldehyde (CH₃CHO) is largely produced through the synthesis of methyl radicals and formic acid at higher pressures (R223) and it is redistributed in the oxygen family via reaction with N atoms (R283), released by cosmic rays, and through photolysis. This balance results in a calculated peak CH_3CHO mole fraction of 4.7×10^{-10} at 1 mbar, suggesting possible identification by Cassini-Huygens. Ethylene oxide, on the other hand, is formed less efficiently in Titan's atmosphere, produced by the synthesis of ethylene and oxygen atoms [Gaedtke et al., 1973]. Considering this synthesis and the photodissociation of C₂H₄O [Fleming et al., 1959], ethylene oxide is expected to be present in Titan's atmosphere in abundances of 1-2 orders of magnitude less than acetaldehyde.

[73] A major question involving the distribution of these oxygen-bearing molecules is what the source of carbon monoxide is. Samuelson et al. [1983] suggested a source from the water influx through the mechanism $OH + CH_3 \rightarrow$ $CO + 2H_2$. However, this was an overall reaction mechanism that was measured by Fenimore [1969]. No laboratory studies [e.g., De Avillez Pereira et al., 1997; Fagerström et al., 1993] have detected CO as a product of this reaction, and no mechanism involving the products of CH₃ + OH forming CO is evident. Lara et al. [1996] calculated an upward flux from the surface of 1.6 imes 10⁶ cm⁻² s⁻¹ necessary to maintain an mole fraction of 5 \times 10⁻⁵ in equilibrium, which they deem to be unlikely over the course of Titan evolution. Consequently, they have suggested that CO may be provided directly from the influx of micrometeorites, although a typical cometary inventory of CO does not provide enough influx to achieve equilibrium.

[74] Primordial CO remains the most likely source. *Bézard et al.* [2002] report the likelihood that CO is primordial on Jupiter, on the basis of the CO abundance in the troposphere. The Titan nominal model calculates an upward CO flux of 3.9×10^6 cm⁻² s⁻¹ necessary to maintain photochemical equilibrium. Oxygen is lost from the atmosphere through condensation of CO₂ and H₂CO, which combines for an upper limit condensation flux of 4.0×10^6 cm⁻² s⁻¹ as determined by the net production rate shown in Table 7. Surficial processes such as outgassing from the interior or irradiation of CO₂ and H₂CO condensates may balance this oxygen loss to the surface, providing a source for CO. Another possibility is that CO is not in equilibrium and was more abundant in Titan's past. *Wong et al.* [2002] postulate that CO may have been as much as 14 times more abundant after the initial escape stage in Titan's early evolution.

5.7. Charged Particles

[75] As shown in section 4.4, ions produced through N_2 and CH₄ ionization, in particular N_2^+ , N^+ , CH_4^+ , and CH_3^+ begin the chemical processes that furnish Titan's ionosphere. Figure 19 shows the relative importance of photoionization and photoelectron-impact ionization in the formation of these ions. Electron impact becomes important only below 1000 km, providing an important source in the lower part of this ionospheric region. Five times more N_2^+ than N⁺ is produced at the peak of ion production through N₂ ionization, facilitating the formation of abundant ions $C_2H_5^+$ and H_2CN^+ through (S28) and (S29). Comparing these production rates with previous Titan ionosphere models show that the Keller et al. [1992] model provides an N_2^+ production rate peak due to photoionization about three times larger than the nominal profile and about twice as large for CH₄⁺, while *Banaszkiewicz et al.* [2000] exceeds the nominal rates by 3.5 and 3 times, respectively. The reason for this discrepancy is not evident, but the fact that Banaszkiewicz et al. [2000] consider photoionization at a solar zenith angle of 30° would certainly be a factor in enhancing their photoionization rates with respect to the nominal model.

[76] The *Banaszkiewicz et al.* [2000] model shows a significant decrease in the CH₄ ionospheric density due to the inclusion of ion chemistry. However, our results indicate that loss of CH₄ through chemistry, which is larger than determined by *Lara et al.* [1999], peaking at 13 cm⁻³ s⁻¹ at 1040 km for solar maximum conditions, is more than replenished by the transport of methane from lower altitudes. An increase in the electron flux by a factor of 10 would be necessary to deplete methane whereby the mole fraction decreases with altitude in this region (Figure 20a), but such an increase would result in a peak electron density of 9740 cm⁻³ (Figure 20b), 2–4 times larger than observations.

[77] With the influence of EUV radiation on the formation of electrons, the electron density has a strong dependence on solar conditions, as exhibited in Figure 21a. For solar maximum conditions, consistent with those which took place during the Voyager flyby, the nominal model calculates an electron density profile which peaks at a density of 4200 cm⁻³ at 1060 km, 20% larger than the upper limit of the *Bird et al.* [1997] radio occultation observations. Considering the differences in production rates, it is not surprising that electron densities of *Keller et al.* [1992] and *Banaszkiewicz et al.* [2000] are larger by about 40% than the nominal model. Assuming electronimpact processes to be scaled for solar minimum and solar moderate conditions, Figure 21a shows a 24% decrease in the peak electron concentration from solar maximum

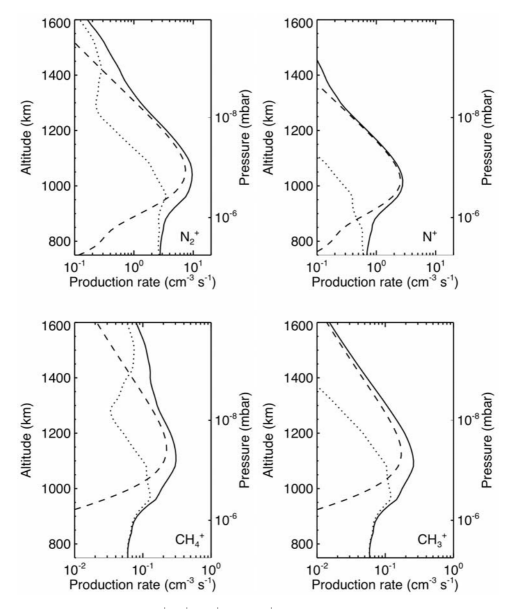


Figure 19. Production rates of N_2^+ , N^+ , CH_4^+ , and CH_3^+ ions from photoionization (dashed line), electron impact (dotted line), and the sum of the two processes (solid line).

conditions for moderate solar activity, which should have implications in the INMS observations from the upcoming Cassini-Huygens mission.

[78] Nominal ion densities, calculated for solar maximum conditions, are shown in Figures 21b and 21c. In agreement with *Banaszkiewicz et al.* [2000] and *Keller et al.* [1992] the nominal model finds H_2CN^+ as the major ion above 1000 km. Below this level, a pseudoion representing the collection of larger ions not considered separately in the model is found to dominate, necessitating the kinetic study and modeling of higher order ions.

[79] The most abundant oxidized ion is H_3O^+ , formed by

(S35)
$$\begin{array}{c} 2CH_4 \to C_2H_5^+ + H_2 + H + e^- \\ \frac{C_2H_5^+ + H_2O \to H_3O^+ + C_2H_4}{2CH_4 + H_2O \to H_3O^+ + C_2H_4 + H_2 + H + e^-}. \end{array}$$

However, the primary function of H_3O^+ is recycling back to H_2O , through

(R494)
$$H_3O^+ + HCN \rightarrow H_2CN^+ + H_2O$$
,

as shown in (S23).

[80] In the stratosphere, an ionosphere develops as well, as a result of cosmic ray ionization of nitrogen. Methane cosmic ray destruction, which is shielded by nitrogen, occurs at a much lesser extent. This cosmic ray deposition, which plays a significant role in nitrile chemistry in the stratosphere as shown previously, results in an electron peak of 1410 cm⁻³ at 120 km for maximum solar conditions. This value increases to 2010 cm⁻³ (Figure 4b) with a moderate sun, as cosmic rays diffuse more efficiently during periods of reduced solar activity [*Jokipii and Kopriva*, 1979]. As products of reactions with larger stable molecules

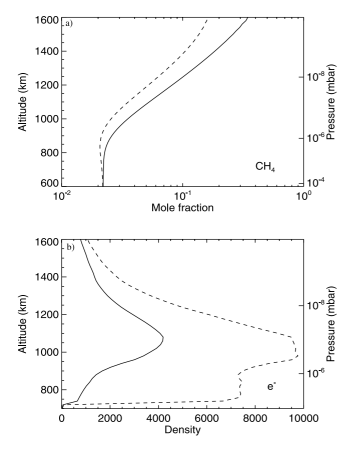


Figure 20. a) CH₄ mole fraction and b) electron density profiles for maximum solar conditions with nominal electron fluxes (solid line) and $10 \times$ nominal electron fluxes (dashed line).

like ethane and acetonitrile, ions like $C_4H_7^+(R413b)$, $C_2H_7^+(R376b)$, $HCO^+(R379)$, $CH_3CNH^+(R347)$, and $C_5H_5^+(R430)$, serve as the gateway to the formation of the large ions which populate Titan's lower ionosphere, as demonstrated by *Molina-Cuberos et al.* [1999]. Ion clusters, as stated before, may also play a significant role in the lower ionosphere.

6. Conclusions

[81] The results of a one-dimensional photochemical model of Titan's neutral constituents and charged particles have been reported. This model contains updated chemistry and an extensive treatment of dissociation processes from solar photons at 50-3000 Å and electrons at 15-1000 eV, as well as parameterization of processes including galactic cosmic rays, magnetospheric electrons, and opacity provided by fractal haze particles.

[82] A test of various eddy diffusion profiles has revealed a profile with a homopause level of 850 km to provide the best fit to IRIS and ISO stratospheric observations as well as UVS observations in the upper atmosphere. With such a profile, fitted with the assumption of methane supersaturation near the tropopause, as analyzed by the *Samuelson et al.* [1997] study, the nominal model provides a good fit for the bulk of Titan stratospheric observations and provides a reasonable fit for most of the Voyager UVS reanalysis observations, an improvement over previous models. In this model, loss of HCN due to polymerization to Titan haze is not required to match observations. The profile of Lara et al. [2002], derived with a more extreme scenario of tropospheric methane supersaturation suggested by Courtin et al. [1995], was also tested. This profile, although providing a good fit of CH₄ and HCN observations, significantly underestimates stratospheric CH₃C₂H and overestimates C₄H₂ in the lower and middle atmosphere. The nominal profile, assuming opacity provided by fractal haze particles from Lebonnois et al. [2001], underpredicts the C₃H₈ stratospheric mole fraction by about a factor of two, while rendering profiles for HC₃N and C₂H₃CN that are not consistent with the lack of firm detection by the IRIS equatorial observations. Assuming Mie haze opacities, these profiles fall much more in line with what has and has not been observed, suggesting that the opacities for the fractal case used in the model may be too large.

[83] Dynamics is certain to play an important role in the distribution of Titan's constituents and the effect of dynamical processes should certainly be explored. However,

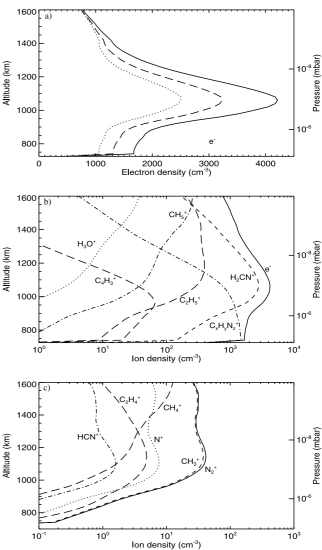


Figure 21. Density distribution of ions in Titan's upper ionosphere.

considering the limitations of existing observations, the uncertainties in other parameters such as aerosol opacity, and the results of the model, it is not possible to rule out the possibility that the globally averaged distribution of Titan's constituents can be accurately and simultaneously described with a single eddy-diffusion profile.

[84] Considering the differences in the expected solar flux during the upcoming Cassini-Huygens mission and the more enhanced solar flux during the Voyager flyby of Titan, sensitivity to variations in solar flux during the course of solar cycle has been explored. With the large chemical lifetimes of most of the stable constituents in Titan's atmosphere, only C_4H_2 demonstrated significant sensitivity among likely observed species, with stratospheric densities for moderate solar conditions reduced by about 60% from solar maximum conditions and a variation of a factor of three from solar maximum to solar minimum.

[85] The profile of C_6H_6 is improved on the basis of ISO observations [*Coustenis et al.*, 2003], compared with previous studies by the authors [*Wilson et al.*, 2003]. The inclusion of haze opacity sharply reduces the amount of benzene photolysis, the primary sink for C_6H_6 in the stratosphere. With this inclusion, the profile of C_6H_6 under nominal Titan temperature dependency matches the ISO observations and suggests a mechanism for haze production peaking at 180 km.

[86] A mechanism for the formation of CH_3C_2H is proposed with the aid of the kinetic measurements of Canosa et al. [1997]. Assuming a branching ratio of 0.5 for methylacetylene production through $CH + C_2H_4$, the profile of CH₃C₂H finds good agreement with observations, over the results of previous models. In agreement with previous models exploring Titan's ionosphere, the dominant ion at the electron peak is found to be H₂CN⁺, with larger ions making up the bulk of Titan's ionosphere below 1000 km. The peak of electron density is found to be 4200 cm^{-3} , a 20% enhancement over the observations of Bird et al. [1997]. The electron peak density is expected to be reduced by about 24% for moderate solar conditions. Cassini-Huygens will encounter variations in the magnetospheric input into the ionosphere. Any major enhancement of magnetospheric electrons over what was found by Voyager should affect the electron density above 1150 km, with no major effect on atmospheric neutrals.

[87] The Cassini-Huygens mission will seek to answer many of the questions explored in this study, among many others. Results from this model indicate the possible detection of H₂CO and C₂H₃CN by the Composite Infrared Spectrometer (CIRS) and the Gas Chromatograph Mass Spectrometer (GCMS), although acrylonitrile abundances may be overpredicted due to chemistry or opacity assumptions. Benzene, already tentatively detected by ISO [Coustenis et al., 2003] is predicted to be observed by these instruments, and the analysis of aerosols by the Aerosol Collector Pyrolyser (ACP) will enhance what we know about the composition and formation processes of Titan's aerosols. Furthermore, observations of constituents like C_2H_2 and CH_3C_2H by the Ultraviolet Imaging Spectrograph (UVIS) and the Ion and Neutral Mass Spectrometer (INMS) will further increase our understanding of the chemical and diffusive processes in the upper atmosphere. This model in conjunction with data

can be used as a helpful tool in furthering the understanding of Titan's atmosphere.

Appendix A: Finite Differencing and Matrix Solver

[88] The photochemical model solves the continuitydiffusion equation in spherical coordinates, accounting for condensation,

$$P_i - L_i - \gamma_i = \frac{1}{r^2} \frac{\partial (r^2 \Phi_i)}{\partial r}$$

where Φ_i is taken to be the flux of species *i* in the radial direction. Multiplying both sides by r^2 and integrating over the interval $[\tilde{r}_1, \tilde{r}_2]$ yields

$$\int_{\tilde{r}_2}^{\tilde{r}_1} \frac{\partial (r^2 \Phi_i)}{\partial r} dr = \int_{\tilde{r}_2}^{\tilde{r}_1} r^2 [P_i - L_i - \gamma_i] dr$$

or

$$\begin{split} \tilde{r}_1^2 \Phi_i(\tilde{r}_1) - \tilde{r}_2^2 \Phi_i(\tilde{r}_2) &= \frac{1}{3} \tilde{r}_1^3 [P_i(r) - L_i(r) - \gamma_i(r)] \\ &- \frac{1}{3} \tilde{r}_2^3 [P_i(r) - L_i(r) - \gamma_i(r)] \end{split}$$

assuming $P_i - L_i - \gamma_i$ constant over the interval $[\tilde{r}_1, \tilde{r}_2]$. Equations (2) and (4) indicate that $n = n(\xi_1^A, \ldots, \xi_S^A)$ where the ξ_1^A, \ldots, ξ_S^A are the S abundant species which affect the mean molecular weight, defined as species with a mole fraction greater than 10^{-3} for a particular level *r*. So, $\Phi_i = \Phi_i(n(\xi_1^A, \ldots, \xi_S^A), \xi_i)$ and the P_i , L_i , and γ_i are functions of $n(\xi_1^A, \ldots, \xi_S^A)$ and the ξ_1, \ldots, ξ_N for M species and N atmospheric levels. Charged particles are assumed to be governed solely by chemistry, with $\Phi_i = 0$, and the assumption of charge neutrality

$$n_e = \sum_i n_i$$

where n_e is the electron density and n_i is the density of each ion *i*.

[89] This system of equations is set up through finite differencing on a grid where the fluxes are expressed in terms of its values at $\tilde{r}_1 = r_{i+1/2}$ and $\tilde{r}_2 = r_{i-1/2}$, assuming that they are linear in the intervals [i - 1, i] and [i, i + 1], while the chemical terms are expressed at $r = r_i$. Thus for a quantity Q,

$$Q(\tilde{r}_1) = \frac{Q(r_{i+1}) + Q(r_i)}{2}, Q(\tilde{r}_2) = \frac{Q(r_i) + Q(r_{i-1})}{2}$$

while

$$\frac{\partial Q(\tilde{r}_1)}{\partial z} = \frac{Q(r_{i+1}) - Q(r_i)}{r_{i+1} - r_i}, \frac{\partial Q(\tilde{r}_2)}{\partial z} = \frac{Q(r_i) - Q(r_{i-1})}{r_i - r_{i-1}}$$

At the lower boundary, the mixing ratios ξ_1, \ldots, ξ_N and number density *n* are set to zero or their lower boundary

conditions, shown in Table 5, where n is described by equation (5). At the upper boundary, the flux equations are finite differenced where, using equation (2),

$$\begin{split} \Phi_i(\tilde{r}_N) &= -\left[D_i(\tilde{r}_N) + K(\tilde{r}_N)\right] n(\tilde{r}_N) \frac{\partial \xi_i(\tilde{r}_N)}{\partial z} \\ &- D_i(\tilde{r}_N)\xi(\tilde{r}_N) \left[\frac{\partial n(\tilde{r}_N)}{\partial z} + n(\tilde{r}_N) \left(\frac{1}{T(\tilde{r}_N)} \frac{\partial T(\tilde{r}_N)}{\partial z} + \frac{1}{H_i(\tilde{r}_N)}\right)\right] \\ &= \beta_i^{\Phi} \end{split}$$

where for quantity Q,

$$Q(\tilde{r}_{N}) = \frac{Q(r_{N}) + Q(r_{N-1})}{2}, \frac{\partial Q(\tilde{r}_{N})}{\partial z} = \frac{Q(r_{N}) - Q(r_{N-1})}{r_{N} - r_{N-1}}$$

and β_i^{Φ} = flux boundary condition for species *i*. For species H and H₂, the velocity equations at the boundary from equation (15) are expressed as

$$\begin{split} w_i(\tilde{r}_N) &= -\left[D_i(\tilde{r}_N) + K(\tilde{r}_N)\right] \frac{\partial \ln(\xi_i(\tilde{r}_N))}{\partial z} \\ &- D_i(\tilde{r}_N) \left[\frac{\partial \ln(n(\tilde{r}_N))}{\partial z} + \left((1 + \alpha_i(\tilde{r}_N)) \frac{1}{T(\tilde{r}_N)} \frac{\partial T(\tilde{r}_N)}{\partial z} \right. \\ &+ \frac{1}{H_i(\tilde{r}_N)} \right) \right] \\ &= \beta_i^w \end{split}$$

where for quantity Q,

$$\frac{\partial \ln(Q_i(\tilde{r}_N))}{\partial z} = \frac{1}{Q_i(\tilde{r}_N)} \frac{\partial Q_i(\tilde{r}_N)}{\partial z} = \frac{\ln(Q_i(r_N)) - \ln(Q_i(r_{N-1}))}{r_N - r_{N-1}}$$

and β_i^w = velocity boundary condition for species *i*.

[90] Meanwhile, equations (3) and (4) can be combined with expression (1) and expressed in terms of r_{i-1} , r_i , and r_{i+1} to calculate the total density

$$r^2 \nabla_r p = -r^2 nmg$$

$$\int_{\tilde{r}_2}^{\tilde{r}_1} \frac{\partial}{\partial r} (r^2 p) dr = -GM \int_{\tilde{r}_2}^{\tilde{r}_1} \frac{r^2 nm}{r^2} dr$$

$$\tilde{r}_1^2 p(\tilde{r}_1) - \tilde{r}_2^2 p(\tilde{r}_2) = -GMn(r)m(r)[\tilde{r}_1 - \tilde{r}_2]$$
(A1)

or, taking condensation into account,

$$\tilde{r}_{1}^{2}p(\tilde{r}_{1}) - \tilde{r}_{2}^{2}p(\tilde{r}_{2}) = -GM[n(r)m(r) - \gamma^{c}][\tilde{r}_{1} - \tilde{r}_{2}]$$

where

$$p(\tilde{r}_1) = \frac{n(\tilde{r}_1)kT(\tilde{r}_1)}{f(\tilde{r}_1)}, p(\tilde{r}_2) = \frac{n(\tilde{r}_2)kT(\tilde{r}_2)}{f(\tilde{r}_2)},$$

$$\gamma^{c} = \begin{cases} [nm(r) - nm^{sat}(r)]^{p}, m(r)m^{sat}(r) \\ \\ -[nm^{sat}(r) - nm(r)]^{p}, m^{sat}(r)m(r) \end{cases},$$

and

$$m(r) = \sum_{j=1}^{S} \xi_{j}^{A}(r) m\left(\xi_{j}^{A}(r)\right).$$

At the boundary $(r = r_N)$, since $\tilde{r}_1 = R_0 + z - \frac{1}{2}\Delta z$ and $\tilde{r}_2 = R_0 + z + \frac{1}{2}\Delta z$, one can write in equation (A1) $\tilde{r}_2 = 2(R_0 + z) - \tilde{r}_1$ or $\tilde{r}_2 = 2r_N - \tilde{r}_1$, where $\tilde{r}_1 = \frac{r_N + r_{N-1}}{2}$, and as follows, $n(\tilde{r}_2) = 2n(r_N) - n(\tilde{r}_1)$, $T(\tilde{r}_2) = 2T(r_N) - \tilde{T}(\tilde{r}_1)$, and $f(\tilde{r}_2) = 2f(r_N) - 2f(r_N) - f(\tilde{r}_1)$.

[91] To solve this set of $T = (N) \times (M + 1)$ nonlinear equations, the set can be linearized by expressing the equations as

$$f_i(\xi_1,\ldots,\xi_M,n)=0,$$

expanding f_i in a Taylor series expansion, and dropping the higher order terms,

$$f_i(\xi_1 + \Delta\xi_1, \dots, \xi_M + \Delta\xi_M, \dots, n + \Delta n) = 0$$

= $f_i(\xi_1, \dots, \xi_M, n)$
+ $\sum_j^M \frac{\partial f(\xi_1, \dots, \xi_M, n)}{\partial \xi_j} \Delta \xi_j + \frac{\partial f(\xi_1, \dots, \xi_M, n)}{\partial n} \Delta n$

This results in the formation of a Jacobian matrix

$$\mathbf{J} = \begin{bmatrix} \frac{\partial f_1}{\partial \xi_1} & \cdots & \frac{\partial f_1}{\partial \xi_M} & \frac{\partial f_1}{\partial n} \\ \vdots & \vdots & \ddots & \vdots \\ \frac{\partial f_T}{\partial \xi_1} & \cdots & \frac{\partial f_T}{\partial \xi_M} & \frac{\partial f_T}{\partial n} \end{bmatrix}, \Delta \mathbf{x} = \begin{bmatrix} \Delta \xi_1 \\ \vdots \\ \vdots \\ \Delta \xi_M \\ \Delta n \end{bmatrix}$$

such that

$$\mathbf{J} \bullet \Delta \mathbf{x} = -\mathbf{f}.$$

The Jacobian can then be inverted to solve for the $\Delta \mathbf{x} = (\Delta \xi_1, ..., \Delta \xi_M, \Delta n)$, which is then added to the $\mathbf{x} = (\xi_1, ..., \xi_M, n)$ and the following equation is solved iteratively

$$\mathbf{x}^{(k+1)} = \mathbf{x}^{(k)} - \left[\mathbf{J}^{(k)}\right]^{-1} \mathbf{f}^{(k)}$$

until $\Delta \mathbf{x} = \mathbf{x}^{(k+1)} - \mathbf{x}^{(k)}$ falls below the tolerance level. The Jacobian matrix is solved by the Crout-LU decomposition method with scaled partial pivoting [*Yakowitz and Szidarovszky*, 1989], optimized for banded matrices.

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