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Ion Neutral Mass Spectrometer Results from the First Flyby of Titan

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The Cassini Ion Neutral Mass Spectrometer (INMS) has obtained the first in situ composition measurements of the neutral densities of molecular nitrogen, methane, molecular hydrogen, argon, and a host of stable carbon-nitrile compounds in Titan’s upper atmosphere. INMS in situ mass spectrometry has also provided evidence for atmospheric waves in the upper atmosphere and the first direct measurements of isotopes of nitrogen, carbon, and argon, which reveal interesting clues about the evolution of the atmosphere. The bulk composition and thermal structure of the moon’s upper atmosphere do not appear to have changed considerably since the Voyager 1 flyby.

This Report documents neutral composition measurements made with the INMS during the first low-altitude pass through Titan’s upper atmosphere by the Cassini-Huygens spacecraft on 26 October 2004. During the 15 min before and after its closest approach, INMS acquired a rich data set of information on Titan’s atmospheric composition and structure covering an altitude range from 3000 to 1174 km. The closest approach occurred at 38.774° latitude, 88.45° west longitude at a solar local time of 16.753 hours. The highest priority objective of the Cassini orbiter during the first flyby was to measure the densities of the major atmospheric constituents N₂, CH₄, and the inferred thermal structure between 1174 and 2000 km, because these data were important in planning the Huygens probe insertion and subsequent Cassini orbiter flybys. However, these measurements and subsequent analysis yielded substantial scientific results, including (i) the existence of large-amplitude (~10 K) and large-spatial scale (~180 km) atmospheric waves between 1100 and 1500 km, (ii) the identification of a suite of carbon-nitrile compounds at higher altitudes than anticipated (~1200 km), and (iii) the determination of the isotopic ratios of nitrogen in N₂, carbon in CH₄, and the abundance of ⁴⁰Ar as well as an upper limit for the mixing ratio of ³⁶Ar. We begin with a brief description of the instrument, then present the data set, and finish with a discussion of the major results enumerated above.

INMS is a dual–ion source quadrupole mass spectrometer covering the mass range of 0.5 to 8.5 and 11.5 to 99.5 daltons (1, 2). The dual-source design combines classic closed and open ionization source configurations designed to measure inert species and reactive species and ions, respectively. The primary data reported in this paper were obtained with the closed source, which is designed to measure nonreactive atmospheric gases. In the closed source, the neutral gas flows into a spherical antechamber where it thermally accommodates with the walls of the antechamber before flowing through a transfer tube to an electron ionization source; there, the gas is ionized by an electron impact at 70 eV. The high flyby velocity of the Cassini spacecraft with respect to Titan (~6 km s⁻¹) produces a dynamic pressure enhancement in the antechamber, which increases sensitivity (1, 2). Reactive species are not measured in this configuration.

Ionization of the primary constituents N₂ and CH₄ by the electron ionization source produced not only the primary ions N₂⁺ and CH₄⁺ but also dissociative fragments (over-
bars in Fig. 1). It also measured a host of ions and fragments due to minor atmospheric species (H₂, C₂H₂, C₂H₄, C₃H₆, C₄H₂, and Ar).

All of the measurable product channels were determined during the flight unit and engineering unit calibrations (with the exception of C₂N₂, C₂H₂, C₂H₄, HCN, and HC₃N, which were obtained from National Institute of Standards and Technology tabulations) and are subsequently used in the deconvolution of the spectra. The spectrum displayed in Fig. 1 (acquired every 4.6 s between the altitudes of 1230 and 1174 km and which can be co-added to enhance the signal-to-noise ratio) includes a mass scan covering the range from 1 to 99 daltons. This altitude range was chosen to be within approximately one atmospheric scale height of the Cassini closest approach (CA) altitude. The signal in each mass bin is a combination of the signals from the ionization or dissociative ionization of several species. Calibration data provide the sensitivity and the distribution of the dissociation products. Spacecraft velocity and altitude are used to compute the ram flow enhancement. From this data, a matrix is constructed relating instrumental response for various mass channels to the atmospheric composition. An inversion of this matrix with suitable numerical methods (J) yields abundances for a range of species. The measurements (and matrix elements) are weighted by the reciprocal measurement error.

Using two independent methods, we found the derived volume-mixing ratio of methane to be between 2.2 and 2.7% (for the first and second method, respectively) in the upper atmosphere. For the first method, we calculated the average ratio from the mass deconvolution of the average spectra between 1230 and 1180 km in the ingress data set as described above; for the second method, we used a diffusion model containing CH₄ and H₂ with an altitude-independent eddy diffusion coefficient and temperature as free parameters to fit the data in the altitude range of 1174 to 1500 km. The temperature structure between 1174 and 1500 km was assumed to be isothermal with a temperature (±SD) of 149 ± 3 K for the combined ingress and egress N₂ and CH₄ data sets (Fig. 2). A weak horizontal temperature gradient (~2 K/1000 km) was seen between ingress (145.8 K) and egress (151.8 K). From Fig. 2 we also note the close agreement of the present observations with the Voyager ultraviolet spectrometer data [reanalysis by Verhacck (J)]. The extension of the INMS density profile (solid line) to lower altitudes in this case is with the assumption of a 149 K isothermal profile down to 900 km.

Some other important atmospheric quantities can also be derived from the density measurements. An eddy coefficient of $K = 4.2 \times 10^6$ cm$^2$ s$^{-1}$ was determined by fitting the combined set of ingress and egress data ($K = 1.6 \times 10^6$ cm$^2$ s$^{-1}$ was found for ingress data alone and $K = 8.3 \times 10^6$ cm$^2$ s$^{-1}$ for egress data). Adequate fits were obtained assuming constant $K$ with altitude. These values imply a homopause altitude defining the boundary between the well-mixed and diffusively separated portions of the atmosphere; the altitude was calculated by finding the intersection between $K$ and the N₂-CH₄ binary diffusion coefficient profiles. We found the homopause at 1195 ± 65 km. Finally, the exobase above which the mean free path of a particle is equal to the atmospheric scale height was calculated from the measured densities to be at an altitude of 1429 ± 5 km.

We observed strong temperature perturbations superimposed on the isothermal profile derived from altitude analysis of the combined ingress and egress N₂ and CH₄ data sets. These perturbations along the spacecraft track can be mapped either horizontally or vertically and appear to represent either a vertically propagating oscillation or an intrinsic harmonic of the atmosphere. Only the vertical mapping case is presented in Fig. 3, suggesting a wave amplitude of 10 K and a vertical wavelength of 180 km. The origin of these waves in Titan’s upper atmosphere could be an energy or momentum source located at lower altitudes in the

**Fig. 1.** The average mass spectrum from 1 to 99 daltons measured by INMS on the 26 October flyby ($T_s$) between the altitudes of 1174 and 1230 km. The average spectrum includes only inbound data where the INMS was pointed along the spacecraft velocity vector. The solid black line represents the measured, background-corrected spectra and the red circles represent the reconstructed spectra using the matrix with suitable numerical methods for the first and second method, respectively.

**Table 1.** Minor species determined from the mass spectral deconvolution with 1σ error. C₃H₆ may represent a contaminant. Data are expressed in values of the mole fraction. C₃H₆ value depends on the value adopted for HCN. The following species were identified at less than 5 ppm: C₃H₄, C₄H₂, HCN, HC₃N, C₅H₆, and C₆N₂.

<table>
<thead>
<tr>
<th>Species</th>
<th>INMS-derived values</th>
<th>CIRS values ($\sigma$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CH₄</td>
<td>$2.19 \times 10^{-2} \pm 0.002 \times 10^{-2}$</td>
<td>$1.6 \times 10^{-2} \pm 0.5 \times 10^{-2}$</td>
</tr>
<tr>
<td>H₂</td>
<td>$4.05 \times 10^{-3} \pm 0.03 \times 10^{-3}$</td>
<td>Not measured</td>
</tr>
<tr>
<td>C₂H₂</td>
<td>$1.89 \times 10^{-4} \pm 0.05 \times 10^{-4}$</td>
<td>$3.3 \times 10^{-6} \pm 0.3 \times 10^{-6}$</td>
</tr>
<tr>
<td>C₂H₄</td>
<td>$2.59 \times 10^{-4} \pm 0.70 \times 10^{-4}$</td>
<td>$1.6 \times 10^{-7} \pm 0.7 \times 10^{-7}$</td>
</tr>
<tr>
<td>C₃H₆</td>
<td>$5.26 \times 10^{-4} \pm 0.08 \times 10^{-4}$</td>
<td></td>
</tr>
<tr>
<td>C₄H₄</td>
<td>$1.21 \times 10^{-4} \pm 0.06 \times 10^{-4}$</td>
<td>$2.3 \times 10^{-5} \pm 0.4 \times 10^{-5}$</td>
</tr>
<tr>
<td>C₅H₆</td>
<td>$3.86 \times 10^{-6} \pm 0.22 \times 10^{-6}$</td>
<td>$1.2 \times 10^{-8} \pm 0.3 \times 10^{-8}$</td>
</tr>
</tbody>
</table>
Fig. 2. The INMS-derived densities of methane (mass channel 16) and molecular nitrogen (mass channels 14 and 28) in red (ingress) and blue (egress) from $T_\nu$ compared with the Vervack-derived (4) values from Voyager ultraviolet spectrometer in pink (ingress) and green (egress). The solid black lines represent the best-fit densities, which correspond to a 149 K isothermal temperature profile. The methane mixing ratio from this modeling fit to the data is 2.7% at 1174 km.

Fig. 3. Vertical profiles of temperature derived from observations of $N_2$ and $CH_4$ by INMS during $T_\nu$. Red data refer to ingress results and blue to egress.

Composite Infrared Spectrometer (CIRS) measurements (7) is shown in Table 1. Modeling and further analysis of the new atmospheric data will lead to explanations of the large abundance of complex hydrocarbons and nitriles at high altitudes.

The measured isotopic ratios have important implications for the evolution of the Titan atmosphere and interior over geological time. The isotopic ratios for nitrogen ($^{14}N/^{15}N$) and carbon ($^{12}C/^{13}C$) can be used to constrain the role of atmospheric escape and other physical and chemical processes in the Titan system. The $^{14}N/^{15}N$ ratio of 172 to 215 (Table 2) is between that of Mars and Earth, indicating either substantial escape of the atmospheric reservoir of nitrogen since capture or release from the interior or an initial source of nitrogen with an isotopic ratio closer to the solar wind value of 200 ± 55 (8, 9). Comparison of these results with model results of Lunine et al. (10) is informative. If one assumes that the initial nitrogen isotopic ratios were similar to those of Earth [$^{14}N/^{15}N = 272.0 ± 0.3$ (11)], that the exospheric temperature has remained near 150 K, and that the relative position of the homopause and exobase has remained unchanged over the course of atmospheric evolution, then the formulation of Lunine et al. combined with INMS measurements suggests a loss of atmospheric nitrogen over the history of the atmosphere by a factor of 1.7 ± 0.05 to 10 ± 5 from its original levels, depending on the efficiency (0 to 0.6) for dissociative fractionation of the isotopes. This also assumes that the upper end of the INMS measurement range for this ratio (i.e., 215) is representative of the present-day isotopic ratio. Correspondingly, if we use the lower end of the INMS-measured range (172), the range of atmospheric loss factor becomes 2.8 ± 0.2 to 100 ± 75. Clearly, it is hard to constrain the amount of atmospheric loss over geological time with the present data set. The quoted uncertainty is largely due to our imprecise knowledge of the position of the homopause and exobase. However, large uncertainties about the initial Titan isotopic ratio (8, 9) and the role of dis-

Atmosphere, or they could represent an external source from Saturn’s magnetosphere, or both.

Analysis of an average spectra for the altitude range of 1230 to 1174 km near CA allowed us to determine a preliminary list of carbon-nitrite compounds and their abundances. The spectral fit from our deconvolution, discussed above, is shown in Fig. 1, and the resulting list of compounds is provided in Table 1. At this preliminary stage in the analysis, the abundances of molecular species are not well-established because we have not completed a thorough analysis of systematic uncertainties, most notably the interaction of potentially reactive species (such as HCN) with the wall of the antechamber for an impact velocity of $6 \text{ km s}^{-1}$. However, we are confident in the detection of all species listed in the table.

The spectral analysis procedure also leads to a determination of isotopic ratios for nitrogen ($^{14}N/^{15}N$) and carbon ($^{12}C/^{13}C$), a determination of the $^{40}Ar$ mixing ratio, and an upper limit on the $^{38}Ar$ mixing ratio in the upper atmosphere. Because the lowest altitudes sampled by INMS on this Titan flyby were evidently below the homopause, these volumemixing ratio values can be used to infer mixing ratios for the well-mixed atmosphere at lower altitudes. The statistical error associated with determining the volume-mixing ratios by this method is on the order of 1%, but the uncertainties associated with the calibration data lead to an overall uncertainty of 20% (Table 2). However, the calibration uncertainties cancel for the isotopic ratios reported so that the only error sources are statistical or systematic errors introduced by the analysis method. Another method was also used to determine the isotopic abundances in which integrated sums of measured abundances (and errors) below 1230 km are used. The results of the two approaches are given in Table 2. There is good agreement between the methods for the carbon ratio, leading us to conclude that systematic errors are small and that statistical errors define the uncertainty. In the case of the nitrogen ratio, the saturation in the mass-28 channel required scaling of the low-sensitivity counter to replace data from the saturated high-sensitivity counter. At this time, the scale factor is not well-determined, resulting in systematic errors that are differential between the two methods. Thus we quote a range of values that define our uncertainty in this regard and we have verified the reported isotopic ratio value at higher altitudes where the $N_2$ data in channel 28 are not yet saturated.

INMS measurements obtained during the first pass of the Cassini orbiter through Titan’s upper atmosphere lead to some important conclusions. A key finding is the apparent stability of the atmosphere over the 25 years above, is shown in Fig. 1, and the resulting list of compounds is provided in Table 1. At this preliminary stage in the analysis, the abundances of molecular species are not well-established because we have not completed a thorough analysis of systematic uncertainties, most notably the interaction of potentially reactive species (such as HCN) with the wall of the antechamber for an impact velocity of $6 \text{ km s}^{-1}$. However, we are confident in the detection of all species listed in the table.

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INMS measurements obtained during the first pass of the Cassini orbiter through Titan’s upper atmosphere lead to some important conclusions. A key finding is the apparent stability of the atmosphere over the 25 years since the Voyager flyby, as indicated by the comparison shown in Fig. 2. We suggest that this may be due to the thermostatic control exercised on the upper atmosphere by the infrared active species HCN. Another key result is the location of the homopause at ~1195 km, which is much higher than was expected [table 1 in (1)] (5). This result indicates that turbulent mixing extends to very high altitudes, and is consistent with the discovery by INMS of large scale–size and large-amplitude atmospheric waves in the upper atmosphere (Fig. 3). The large-scale size of the atmospheric waves was anticipated and is a consequence of atmospheric filtering by wind shears and thermal gradients at lower altitudes, and the large wave amplitude is within the limits of model predictions (6).

The existence of heavy hydrocarbon species at altitudes above 1200 km is another possible consequence of the well-mixed atmosphere. An alternative explanation is that previous determinations of the production rates for these chemical species at high altitudes are too small. Most of the observed compounds were identified earlier in the lower atmosphere [table 1 in (1)] (5). A comparison of INMS-derived high-altitude values with those determined in the lower atmosphere from Cassini...
Sociative fractionation limit us to conclude only that the Titan atmosphere was at least 50% denser in the past.

A perplexing aspect of the nitrogen isotopic ratios comes from comparing the INMS-measured molecular nitrogen isotopic ratio value to earlier ground-based determinations of $^{14}$N/$^{15}$N in HCN measured in Titan’s stratosphere (12, 13). The HCN-derived values show enrichment of the heavier isotope of nitrogen that is more than a factor of 2 larger than the INMS-measured values for the molecular nitrogen isotope ratio. Chemical fractionation is the most likely source of the difference, but estimates so far do not come close to explaining the difference (14, 15).

The heavy isotope enrichment of nitrogen stands in contrast to the carbon isotopic ratio ($^{13}$C/$^{12}$C = 95 ± 1). The ratio is well-determined and slightly above the terrestrial value of 89.0 ± 0.38, but certainly within the present constraints of determinations for Jupiter (12), Saturn (89 ± 25) (10). This suggests that the methane in the atmosphere is being replenished by an interior source at a rate fast enough that escape is not the dominant effect on the isotopic ratio. This is consistent with our understanding of photodissociation rates for methane and the associated deposition of complex hydrocarbons on the surface. This dissociation process acts as the dominant sink for the atmospheric portion of the carbon cycle on Titan. INMS measurements of H$_2$ outflow can help quantify this process. H$_2$ is a byproduct of the methane photodissociation, and the measured H$_2$ escape rate of 6.1 × 10$^{19}$ × 0.2 × 10$^9$ cm$^{-2}$ s$^{-1}$ represents a photodissociation rate of 5 × 10$^{27}$ methane molecules per second and an atmospheric carbon residence time of 5 × 10$^7$ years. However, this does not explain the elevated isotopic ratio value indicating an increase of the light $^{12}$C isotope relative to $^{13}$C isotope when compared with the terrestrial value of 89.0 ± 0.38 (11). Such a light isotope enhancement on Earth of up to ~10% is associated with biology (16); on Titan a more likely process involves vapor pressure isotope effects at the surface (17).

However, using a diffusion model and our present estimate of thermal structure and atmospheric diffusion to propagate the upper atmospheric value to the surface (Table 2), we obtained a surface value of 81, which is below the terrestrial value. Clearly, combining the INMS data with information obtained by the Huygens Gas Chromatograph Mass Spectrometer is necessary to reach closure on this issue.

The volume-mixing ratios of the isotopes of argon can be used to study both interior and accretion processes. $^{40}$Ar is likely produced by radioactive decay of $^{40}$K, as it is on Earth, and therefore the mixing ratio of $^{40}$Ar can be compared to estimates of the radiogenic production and release of potassium in the mantle rocks. The measured $^{40}$Ar is 7.081 × 10$^{-6}$ ± 0.1 × 10$^{-6}$ as compared with an upper limit of 0.05 to 0.07% set by McKinnon’s estimate (18). This suggests either that the outgassing of the interior has an efficiency of ~1% or (less likely) that the $^{40}$Ar was lost to space at an early phase of atmospheric evolution. On the other hand, the nondetection of $^{36}$Ar and $^{38}$Ar may suggest that N$_2$ was not trapped in nebula ices in the form of clathrate hydrates (19–22). The upper limit of less than 6 × 10$^{-7}$ by volume set by INMS for the $^{40}$Ar abundance ratio leads to the conclusion that the nitrogen in the moon’s atmosphere was not brought to Titan as N$_2$ trapped in clathrates but was likely derived initially from ammonia ice or ammonia hydrate formed in the local Saturn nebula (23) or in the solar nebula (24) and subsequently converted to N$_2$ by photochemical processes (25) and/or from high-temperature shock formation of N$_2$ (26).

The first measurements of Titan’s neutral composition in the upper atmosphere lead to a number of notable conclusions about the current atmospheric structure and about the evolution of Titan’s atmosphere and interior. The atmospheric likely formed from the outgassing after planetesimals composed of silicates, water ice, clathrates of methane, and ammonia hydrates coalesced to form the moon. Subsequent photochemistry and/or shock-induced chemistry likely converted the atmospheric nitrogen into molecular nitrogen. The early atmosphere was 1.6 to 100 times more substantial and was lost through escape over the intervening 4.5 billion years due to the reduced gravity associated with the relatively small mass of Titan. Carbon in the form of methane has likely continued to outgas over time from the interior, given that much of its subsequent photolysis products have been deposited in the form of complex hydrocarbons on the surface (~5 × 10$^{27}$ s$^{-1}$ as estimated from the H$_2$ escape rate of 6.1 × 10$^{19}$ ± 0.2 × 10$^9$ cm$^{-2}$ s$^{-1}$ measured by INMS) and some of it has also been lost to space [2 × 10$^{25}$ s$^{-1}$ as estimated from modeling of the exospheric loss of carbon (27)]. The carbon isotopes in the upper atmosphere appear to be isotopically light due to atmospheric diffusion effects or vapor phase isotope effects at the surface resulting from condensation, evaporation, and sublimation of surface methane.

Table 2. Key isotopic ratios measured by INMS (uncertainties are quoted as a 95% confidence interval).

<table>
<thead>
<tr>
<th>Isotopic ratio</th>
<th>Integration method</th>
<th>Deconvolution method</th>
<th>Combined results</th>
<th>Estimated value at surface</th>
</tr>
</thead>
<tbody>
<tr>
<td>$^{14}$N/$^{15}$N</td>
<td>188 ± 14</td>
<td>214 ± 1</td>
<td>172 &lt; $^{14}$N/$^{15}$N &lt; 215</td>
<td>168 &lt; $^{14}$N/$^{15}$N &lt; 211</td>
</tr>
<tr>
<td>$^{13}$C/$^{12}$C</td>
<td>95.6 ± 0.1</td>
<td>93.8 ± 1.9</td>
<td>95.6 ± 0.1</td>
<td>Not applicable</td>
</tr>
<tr>
<td>$^{36}$Ar mixing ratio</td>
<td>&lt;4 × 10$^{-7}$ ± 1 × 10$^{-7}$</td>
<td>&lt;6 × 10$^{-7}$</td>
<td>&lt;5 × 10$^{-7}$ ± 2 × 10$^{-7}$</td>
<td>7 Not applicable</td>
</tr>
<tr>
<td>$^{40}$Ar mixing ratio</td>
<td>7.1 × 10$^{-6}$ ± 0.1 × 10$^{-6}$</td>
<td>7.3 × 10$^{-6}$ ± 2.1 × 10$^{-6}$</td>
<td>7.1 × 10$^{-6}$ ± 0.1 × 10$^{-6}$</td>
<td>Not applicable</td>
</tr>
</tbody>
</table>

References and Notes
7. Lower atmospheric values of the hydrocarbon mixing ratio at a pressure of 1.5 to 3.0 mbar and a latitude of 39° were taken from Fraser et al. (28). The methane value comes from the same reference but at a pressure of 3 to 20 mbar.
9. If instead of the solar wind value, we choose initial values close to the newly determined protoneutron value (430), determined by Owen et al. (29), the requirement for escape is much larger than our assumption of a terrestrial initial value of 272.
14. One likely mechanism for the enrichment of HCN/N$_2$ is the photodissociated isotopic fractionation effect (PHIF) proposed by Yung and Miller (30). However, a preliminary estimate by Yung (31) shows that the magnitude of PHIF for HCN is not sufficient to account for the observed enrichment.
15. Another possibility is ion-neutral reactions as studied in the case of the interstellar medium by R. Terzieva and E. Herbst (32), but such effects have not been examined at Titan temperatures (150 K).
17. Because methane is near the triple point near the Titan surface, one would expect a heavy isotope depletion of the gas as compared with the liquid or solid due to vapor pressure isotope effects similar to those observed for H$_2$O on Earth (33).
18. W. B. McKinnon, personal communication. McKinnon assumes an anhydrous rock mass fraction for Titan of 0.55 ± 0.05 and a CI carbonaceous [K = 550 parts per million (ppm)] or H chondrite (780 ppm) composition for the rock (34). Of a primordial $^{40}$K/$^{40}$Ar = 1.47 × 10$^{-3}$, 9.6% has decayed to $^{40}$Ar. Perhaps we shouldn’t be surprised by the very low degassing efficiency, if we assume that Titan has a rock core and that core was volcanically active, those magmas would still be erupting at great depths, and at pressures around 2 GPa. There is no thermodynamic incentive for volcanics to degas at such pressures.
Cassini Measurements of Cold Plasma in the Ionosphere of Titan


The Cassini Radio and Plasma Wave Science (RPWS) Langmuir probe (LP) sensor observed the cold plasma environment around Titan during the first two flybys. The data show that conditions in Saturn’s magnetosphere affect the structure and dynamics deep in the ionosphere of Titan. The maximum measured ionospheric electron number density reached 3800 per cubic centimeter near closest approach, and a complex chemistry was indicated. The electron temperature profiles are consistent with electron heat conduction from the hotter Titan wake. The ionospheric escape flux was estimated to be $10^{25}$ ions per second.

The giant planet Saturn and its magnetosphere rotate with a period of $\sim 11$ hours, which can be inferred from radio measurements. At the distance of the large moon Titan [20 Saturn radii ($R_S$)], this corotation causes a magnetospheric plasma flow of several hundred kilometers per second that affects the upper ionized part of the thick atmosphere of Titan. Atoms and molecules in the upper atmosphere are ionized by solar ultraviolet (UV) radiation and by impacts of energetic particles that originate mainly from the magnetosphere. The aeronomic and electrodynamic processes involved in the interaction produce a complex organic chemistry within the nitrogen- and methane-rich atmosphere of Titan as well as a loss of atmospheric constituents, providing in turn a source of plasma for the magnetosphere of Saturn (2).

The two first flybys of Titan (TA and TB) on 26 October and 13 December 2004 were very similar. Both occurred at 10.5 Saturn local time (LT) near the front side of Saturn’s magnetosphere, and both approached inbound from the sunlit side and through the wake of Titan (Fig. 1). The outbound passes partly traversed a shaded region caused by Titan’s thick atmosphere at a latitude of 30° to 40°N.

The structure and thermal state of the ionosphere of Titan were affected by the saturnian magnetosphere all the way down to closest approach (1176 km) during both flybys (Figs. 2 and 3). The general shape of the two main number density maxima near closest approach in Fig. 2 is broadly consistent with photoionization by UV light from the Sun and impact ionization by magnetospheric electrons (3, 4). However, the plasma density was otherwise very structured and could be related to similar structures in the magnetic field data. The density data in Figs. 2 and 3 were collected at a high time resolution (20 samples/s) and are derived from the probe current at a constant bias voltage ($\sim +10$ V), corrected for electron temperature and adjusted to fit the potential sweep data points for density. The LP sensor (5) samples the total electron number density surrounding the spacecraft, which includes the naturally occurring plasma electrons as well as a nearly constant level of photoelectrons around the spacecraft. The spacecraft-generated photoelectrons dominated the magnetospheric part of the flyby. According to measurements by the Cassini Plasma Spectrometer (CAPS), the magnetospheric electron density near Titan was just below 0.1 cm$^{-3}$ during TA (6).

The calculated magnetic pressure ($B^2/2\mu_o$) of the magnetosphere and the electron thermal pressure ($n_i k_B T_i$) of the ionosphere (where $B$ is magnetic field strength, $\mu_o$ is magnetic permeability, $n_i$ is electron number density, $k_B$ is the Boltzmann constant, and $T_i$ is electron temperature) were largely of the same order during the TA flyby (Fig. 2B). Both flybys traversed the wake of Titan, and the effect of the magnetospheric dynamic pressure ($n_i m_i v_i^2/2$), where $n_i$ is ion number density, $m_i$ is average ion mass, and $v_i$ is ion ram velocity) should therefore be small near closest approach. The ionopause, the region where the ionospheric thermal pressure balances the magnetospheric pressure, covered an extended region on both the inbound and outbound trajectories (yellow area in Fig. 2). The close relationship between cold plasma density signatures and magnetic field fluctuations and comparable values of ionospheric electron thermal and magnetospheric magnetic field pressures confirms the view that the saturnian magnetosphere is important for the structure and dynamics in Titan’s ionosphere.

The Titan wake “neutral” sheet, where the magnetic field sharply rotates and relates to an electrical current, was identified by the Dual-Technique Magnetometer (MAG) near 15:35 UT (7). At this time, the RPWS LP detected a sharp peak in both electron number density (850 cm$^{-3}$) and electron thermal pressure (900 eV/cm$^3$). The altitude thickness of the compressed plasma sheet was 60 km, which is on the order of the ion gyroradius.

The LP provides estimates of $n_i$ and $T_i$, but under some circumstances, values can also be obtained for $m_i$, $v_i$, $n_i$ ion temperature ($T_i$), solar UV intensity, and spacecraft potential. An estimate of the effective ion temperature ($T_{i\text{eff}}$), which is the sum of the thermal random ion temperature and the ram ion directed flow energy [$T_{i\text{eff}} = T_i + (m_i v_i^2/2e)$], expressed...