Evidence for Ammonia-bearing Species on the Uranian Satellite Ariel Supports Recent Geologic Activity

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Abstract

We investigated whether ammonia-rich constituents are present on the surface of the Uranian moon Ariel by analyzing 32 near-infrared reflectance spectra collected over a wide range of sub-observer longitudes and latitudes. We measured the band areas and depths of a 2.2 μm feature in these spectra, which has been attributed to ammonia-bearing species on other icy bodies. Ten spectra display prominent 2.2 μm features with band areas and depths >2σ. We determined the longitudinal distribution of the 2.2 μm band, finding no statistically meaningful differences between Ariel’s leading and trailing hemispheres, indicating that this band is distributed across Ariel’s surface. We compared the band centers and shapes of the five Ariel spectra displaying the strongest 2.2 μm bands to laboratory spectra of various ammonia-bearing and ammonium-bearing species, finding that the spectral signatures of the Ariel spectra are best matched by ammonia-hydrates and flash frozen ammonia-water solutions. Our analysis also revealed that four Ariel spectra display 2.24 μm bands (>2σ band areas and depths), with band centers and shapes that are best matched by ammonia ice. Because ammonia should be efficiently removed over short timescales by ultraviolet photons, cosmic rays, and charged particles trapped in Uranus’ magnetosphere, the possible presence of this constituent supports geologic activity in the recent past, such as emplacement of ammonia-rich cryolavas and exposure of ammonia-rich deposits by tectonism, impact events, and mass wasting.

Unified Astronomy Thesaurus concepts: Uranian satellites (1750); Surface ices (2117); Surface composition (2115); Planetary surfaces (2113); Surface processes (2116)

1. Introduction

Planetesimals rich in ammonia (NH₃) were likely incorporated into Ariel and the other proto-classical moons as they formed in the Uranian subnebula (e.g., Lewis 1972). NH₃ is an efficient antifreeze when mixed with liquid H₂O, which, if incorporated into Ariel’s interior as a primordial constituent, would have helped this moon retain liquid H₂O in its subsurface for a much longer period of time compared to “pure” liquid H₂O, at temperatures as low as ~176 K (e.g., Spohn & Schubert 2003). Analysis of crater densities suggests that some regions of Ariel’s surface may be relatively young (~1–2 Ga, Zahnle et al. 2003). These regions display numerous examples of landforms indicative of resurfacing driven by tectonism and cryovolcanism (e.g., Smith et al. 1986; Schenk 1991; Kargel 1992; Beddingfield & Cartwright 2020a). Morphological assessment of potential cryovolcanic features in these regions points to emplacement of material sourced from Ariel’s interior (Schenk 1991; Beddingfield & Cartwright 2020a). Furthermore, the estimated flow rheologies for potential cryovolcanic deposits on Ariel are consistent with emplacement of NH₃-rich cryolavas (e.g., Schenk 1991; Kargel 1992). The geologic evidence therefore suggests that NH₃-bearing deposits sourced from Ariel’s interior have played an important role in resurfacing this moon, in particular, in regions estimated to be fairly young.

Supporting the geologic evidence for NH₃-rich deposits on Ariel, recent ground-based, near-infrared (NIR) observations have revealed a subtle absorption band centered near 2.2 μm, consistent with the presence of NH₃-bearing species (Cartwright et al. 2018). Deposits rich in NH₃-hydrates and NH₃ ice, contained within the top meter of Ariel’s subsurface, are likely decomposed by magnetospheric charged particle bombardment over geologically short timescales (Moore et al. 2007), suggesting replenishment of NH₃ from Ariel’s interior in the recent past. However, the spatial distribution of the 2.2 μm band was not previously assessed, limiting our ability to investigate the origin of this feature. Furthermore, the species contributing to the 2.2 μm band have not been assessed, and both NH₃-hydrates and NH₂-rich salts could contribute to this feature. The presence of these constituents could have important implications for Ariel’s geologic history and the evolution of its surface composition. In this work, we investigated the longitudinal distribution and composition of the 2.2 μm band, conducting band parameter measurements on 32 NIR spectra, which confirm the presence of the 2.2 μm band on Ariel. We also investigated the possible presence of a 2.24 μm band, which could result from NH₃ ice.

2. Data and Methods

2.1. Observations and Data Reduction

The nine new reflectance spectra reported here were collected between 2017 and 2019 using the NIR SpeX spectrograph/imager at NASA’s Infrared Telescope Facility (IRTF), operating in low resolution PRISM mode and moderate...
resolution short cross-dispersed (SXD) mode (e.g., Rayner et al. 2003). These IRTF/Spex observations were made using “AB” nodding, where the target is observed in two different positions along the 15° slit, separated by 7°.5. The resulting “A” and “B” exposures are separated into sequential pairs, and the “B” exposures are subtracted from the “A” exposures to perform first-order sky emission correction. Calibration frames (flat fields and arc lamps) were generated using quartz and argon lamps with an internal integrating sphere. Data extraction and calibration were conducted using custom software and the Spextool data reduction suite (Cushing et al. 2004). Extracted spectra from each night were co-added during reduction to boost signal-to-noise. Ariel spectra were divided by solar analog spectra, observed multiple times on the same nights, to remove the solar spectrum and provide additional atmospheric correction. All star spectra were collected within ±0.1 airmass of the Ariel spectra. The analog stars we observed between 2017 and 2019 were: HD 3628, HD 11532, HD 12124, and BD+09 213. The 23 previously reported Ariel spectra were also collected using IRTF/Spex between 2000 and 2016 (Grundy et al. 2003, 2006; Cartwright et al. 2015, 2018). We refer the reader to these papers for detailed descriptions of the data reduction routines utilized by each team. Observation details for all 32 Ariel spectra are summarized in Table 1. Each team used different slit widths (0°3, 0°5, and 0°8) with Spex to achieve a variety of observing goals. The 2.2 μm band can be detected in both PRISM and SXD mode using all slit widths reported here.

2.2. Band Parameter Analyses

The central wavelength of the 2.2 μm band can vary between ~2.20 and 2.22 μm in spectra of Ariel (Figure 1). Furthermore, the wavelength range covered by the 2.2 μm band varies between 2.18 and 2.23 μm, with bandwidths of ~0.03–0.05 μm. To assess the spatial distribution and spectral signature of the 2.2 μm band, we measured the band area and band depth of this feature in each of the 32 Ariel spectra, utilizing a custom data processing program that our team has used previously to conduct band parameter analyses of other icy constituents detected on the Uranian moons (Cartwright et al. 2015, 2018). The program ingested individual spectra and fit their continua between 2.17 and 2.24 μm, spanning the entire wavelength range covered by the 2.2 μm band. To simulate the continuum of each spectrum (i.e., continua without 2.2 μm or 2.24 μm absorption bands), we generated synthetic spectra using Hapke-based radiative transfer models (e.g., Hapke 2012, Appendix A.1). These synthetic spectra were generated using laboratory measured optical constants for the primary constituents that have been detected previously on Ariel: H2O ice (Mastrapa et al. 2008), amorphous carbon (Rouleau & Martin 1991), and CO2 ice (Hansen 1997). The program then divided each Ariel spectrum by its modeled continuum between 2.17 and 2.24 μm and measured the areas of the resulting continuum-divided bands using the trapezoidal rule. To estimate uncertainties, the program utilized Monte Carlo simulations that resample the 1σ errors of each spectral channel within the wavelength range of each band (iterated 20,000 times).

To measure the 2.2 μm feature depths, we first assigned the wavelength at the deepest part of each continuum-divided feature as the band center. The program then averaged the reflectance for all spectral channels within ±0.004 μm of these band centers, thereby calculating a mean reflectance for each band center. To calculate the 2.2 μm band depths, the program subtracted these mean reflectances from 1. To estimate the uncertainties of each band depth measurement, the program added the 1σ errors of all spectral channels included in the mean reflectance measurement, in quadrature, and divided by the number of channels (n), thereby calculating the mean uncertainty (σ). The program then calculated the standard deviation of the mean (σ = σ/√n) to estimate the point-to-point variation for each band depth measurement and calculated the final band depth error by summing σ and σ in quadrature. We show an illustrative example of our band area and depth measurement procedure in Figure A1.

We also conducted band parameter measurements of the 2.24 μm band, using the same procedures described above for the 2.2 μm band. The 2.24 μm band spans ~2.22–2.26 μm (Figure 1), slightly overlapping the wavelength range of the 2.2 μm band, with band centers between ~2.24 and 2.245 μm (continua spanning 2.21–2.27 μm).

3. Results and Analysis

3.1. IRTF/Spex Spectra of Ariel

We report nine new disk-integrated spectra of Ariel, collected with IRTF/Spex operating in PRISM and SXD mode (Figure A2 in Appendix A.2). All nine of these spectra display the 1.52 μm and 2.02 μm H2O ice bands detected in previously collected NIR spectra of this moon (e.g., Cruikshank & Brown 1981; Grundy et al. 2003, 2006; Cartwright et al. 2015, 2018). Furthermore, the five new spectra collected in SXD mode display clear evidence for the narrow CO2 ice bands detected previously between 1.9 and 2.1 μm (Grundy et al. 2003, 2006; Cartwright et al. 2015, 2018). Analysis of these H2O and CO2 ice absorption features is beyond the scope of this paper and will be included in future work.

3.2. Band Parameter Analyses

We assessed the presence of the 2.2 μm band by conducting band area and depth measurements, which we report in Table 2 (along with their 1σ uncertainties). Our analysis determined that 10 of the 32 Ariel spectra have 2.2 μm bands with area and depth measurements that are both >2σ. The band centers for these 10 spectra cluster around three wavelength intervals: 2.198–2.203 μm, 2.209 μm, and 2.214 μm (Figure 1). Evidence for the 2.2 μm band is weaker in the other 22 spectra, making robust assignment of their band centers challenging, and consequently, we do not report band centers for these data. Along with the 2.2 μm band, four spectra of Ariel also display 2.24 μm bands with >2σ band areas and depths (Table A1 in Appendix A.3). In total, 12 different Ariel spectra display 2.2 and/or 2.24 μm bands (Figure 1).

3.3. Spatial Distribution of the 2.2 μm Band

Previous work has demonstrated that Ariel displays clear longitudinal trends in its surface composition and photometric properties, with significantly stronger (>3σ) H2O ice bands on its leading hemisphere (1°–180° longitude) and stronger CO2 ice bands and higher NIR geometric albedos (>3σ) on its trailing hemisphere (181°–360° longitude) (e.g., Grundy et al. 2006; Cartwright et al. 2020a). Consequently, we searched for similar longitudinal trends in the distribution of the 2.2 μm band on Ariel by calculating the mean 2.2 μm band areas for Ariel’s leading and trailing hemispheres and comparing them: 3.13 ± 0.84 μm² and 1.44 ± 0.48 μm², respectively. We also calculated and compared
| Sub-observer Long. (°) | Sub-observer Lat. (°) | UT Date (mid-expos) | UT Time (minutes) | Integration Time (minutes) | SpeX Observing Mode | Slit Width (″) | Average Resolving Power (λ/Δλ) | References |
|------------------------|-----------------------|--------------------|-------------------|--------------------------|---------------------|--------------|--------------------------------|------------|
| 9.2                    | 31.8                  | 9/18/15            | 10:05             | 7.5                      | PRISM               | 0.8          | 93.8                           | Cartwright et al. (2018) |
| 15.3                   | 27.8                  | 9/15/14            | 11:35             | 88                       | SXD                 | 0.8          | 750                            | Cartwright et al. (2018) |
| 38.8                   | 35.8                  | 9/20/16            | 14:15             | 7.5                      | PRISM               | 0.8          | 93.8                           | Cartwright et al. (2018) |
| 53.6                   | -16.0                 | 8/9/03             | 12:15             | 156                      | SXD                 | 0.5          | 2000                           | Grundy et al. (2006)    |
| 79.8                   | -19.4                 | 7/17/02            | 13:25             | 108                      | SXD                 | 0.8          | 750                            | Cartwright et al. (2015) |
| 87.8                   | 24.0                  | 9/5/13             | 11:10             | 92                       | SXD                 | 0.3          | 2000                           | Grundy et al. (2006)    |
| 93.5                   | -18.1                 | 10/4/03            | 5:45              | 108                      | SXD                 | 0.8          | 750                            | Cartwright et al. (2018) |
| 110.1                  | 32.0                  | 9/11/15            | 13:30             | 44                       | SXD                 | 0.8          | 750                            | Cartwright et al. (2018) |
| 132.2                  | 28.5                  | 8/24/14            | 14:05             | 40                       | SXD                 | 0.3          | 2000                           | Grundy et al. (2006)    |
| 137.6                  | 34.6                  | 10/21/16           | 12:40             | 7.5                      | PRISM               | 0.8          | 93.8                           | Cartwright et al. (2018) |
| 144.8                  | 43.2                  | 10/12/18           | 9:30              | 73                       | SXD                 | 0.5          | 1200                           | This work              |
| 159.9                  | -11.1                 | 7/15/04            | 12:00             | 112                      | SXD                 | 0.3          | 2000                           | Grundy et al. (2006)    |
| 200.0                  | -15.9                 | 8/5/03             | 12:00             | 84                       | SXD                 | 0.3          | 2000                           | Grundy et al. (2006)    |
| 201.3                  | 33.2                  | 1/23/17            | 5:45              | 10.5                     | PRISM               | 0.8          | 93.8                           | Cartwright et al. (2018) |
| 205.5                  | 46.1                  | 11/7/19            | 11:20             | 60                       | SXD                 | 0.5          | 1200                           | This work              |
| 219.8                  | -17.2                 | 9/7/03             | 9:35              | 90                       | SXD                 | 0.3          | 2000                           | Grundy et al. (2006)    |
| 224.8                  | 31.8                  | 9/17/15            | 9:40              | 10                       | PRISM               | 0.8          | 93.8                           | Cartwright et al. (2018) |
| 233.6                  | 32.0                  | 9/12/15            | 10:15             | 8                        | PRISM               | 0.8          | 93.8                           | Cartwright et al. (2018) |
| 233.8                  | -23.1                 | 7/5/01             | 14:10             | 50                       | SXD                 | 0.5          | 1200                           | Grundy et al. (2005)    |
| 242.0                  | 42.2                  | 11/7/18            | 6:50              | 10.5                     | PRISM               | 0.8          | 93.8                           | This work              |
| 244.8                  | 47.1                  | 10/18/19           | 14:05             | 7.5                      | PRISM               | 0.8          | 93.8                           | This work              |
| 253.9                  | 29.2                  | 12/2/15            | 5:25              | 9                        | PRISM               | 0.8          | 93.8                           | Cartwright et al. (2018) |
| 257.6                  | -29.5                 | 9/6/00             | 7:35              | 76                       | SXD                 | 0.8          | 750                            | Cartwright et al. (2015) |
| 268.3                  | 39.0                  | 10/15/17           | 8:00              | 40                       | SXD                 | 0.5          | 1200                           | This work              |
| 273.2                  | 42.2                  | 11/7/18            | 12:00             | 42                       | SXD                 | 0.5          | 1200                           | This work              |
| 278.3                  | 24.8                  | 8/7/13             | 13:20             | 44                       | SXD                 | 0.8          | 750                            | Cartwright et al. (2015) |
| 294.8                  | -19.3                 | 7/16/02            | 13:10             | 140                      | SXD                 | 0.5          | 1200                           | Grundy et al. (2003)    |
| 297.0                  | 39.2                  | 10/10/17           | 11:50             | 12.5                     | PRISM               | 0.8          | 93.8                           | This work              |
| 304.8                  | -23.2                 | 7/8/01             | 14:40             | 48                       | SXD                 | 0.5          | 1200                           | Grundy et al. (2003)    |
| 316.6                  | -18.2                 | 10/8/03            | 7:55              | 132                      | SXD                 | 0.3          | 2000                           | Grundy et al. (2006)    |
| 334.4                  | 39.7                  | 9/25/17            | 15:10             | 7.5                      | PRISM               | 0.8          | 93.8                           | This work              |
the mean 2.2 μm band depths for Ariel’s leading and trailing hemispheres: 0.015 ± 0.004 μm and 0.010 ± 0.003 μm, respectively. Comparison of these values demonstrates that Ariel’s mean 2.2 μm band areas are slightly stronger on its leading hemisphere but not at a statistically meaningful level (<2σ). Similarly, Ariel’s mean 2.2 μm band depths display essentially no difference (<1σ) between its leading and trailing hemispheres.

As another test, we plotted the 32 individual 2.2 μm band area and depth measurements as a function of longitude, and fit them with a mean and sinusoidal model to determine whether the 2.2 μm band displays leading/trailing trends in its distribution (Figure 2). The mean model represents a surface displaying no longitudinal trends in the distribution of the 2.2 μm band. Conversely, the sinusoidal model represents a surface with significant differences in the longitudinal distribution of the 2.2 μm band (i.e., stronger 2.2 μm bands on Ariel’s leading or trailing hemisphere). We compared these model fits using an F-test (e.g., Lomax & Hahs-Vaughn 2013), with a null hypothesis that there is no meaningful difference between these two model fits. The results of the F-test demonstrate that there is no statistically significant difference between the mean and sinusoidal models for either the band area or depth measurements (Table A2 in Appendix A4).

Thus, comparison of the mean 2.2 μm band areas and depths, and F-test analysis of the individual band parameter measurements, indicates that there are no meaningful trends in the longitudinal distribution of the 2.2 μm band across the surface of Ariel, unlike the longitudinal distribution of H$_2$O and CO$_2$ ice on this moon. The small number of Ariel spectra displaying a 2.24 μm band (four) prevents similar quantitative analysis of the longitudinal distribution of this feature. Nevertheless, we note that three of the four detected 2.24 μm bands are clustered between longitudes 305° to 335° on Ariel’s trailing hemisphere, which is dominated by large chasmas, as well as other landforms suggestive of geologic activity in the recent past. The fourth 2.24 μm band (137°6 longitude) is proximal to two of the strongest 2.2 μm bands detected on Ariel (Spectra 5 and 6 in Figure 1).

### 3.4. Comparing Ariel Spectra to Laboratory Spectra of NH$_3$- and NH$_4$-bearing Species

To investigate the composition of the constituents contributing to the detected 2.2 and 2.24 μm bands, we compared the Ariel data to reflectance spectra of different NH$_3$-bearing species measured in the laboratory, including mixtures of multiple NH$_3$-hydrates (e.g., Brown et al. 1988), flash frozen NH$_3$-H$_2$O solutions (T. Nordheim 2020, private communication), and ammonium carbonate ((NH$_4$)$_2$CO$_3$) (Berg et al. 2016). Although other NH$_4$ salts display similar 2.2 μm bands to (NH$_4$)$_2$CO$_3$, we selected this particular constituent for comparison because of the large abundance of CO$_2$ ice on Ariel, which could chemically interact with NH$_3$ and other constituents due to magnetospheric charged particle bombardment of Ariel’s surface. We refer the reader to Berg et al. (2016) for more information on a wide variety of NH$_4$-bearing constituents. We also compared the Ariel data to

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**Figure 1.** Left: the 12 IRTF/SpeX spectra of Ariel displaying 2.2 and/or 2.24 μm bands with >2σ band areas and depths (Table 2), offset vertically for clarity and numbered 1 through 12. The 1σ uncertainties for each spectrum are shown in light gray. These spectra have been lightly smoothed using a binning routine with a 3 to 10 pixel-wide window. The central wavelengths of the detected 2.2 μm bands are located between 2.198 and 2.205 μm (blue markers), at 2.209 μm (green markers), and at 2.214 μm (yellow markers). The central wavelength of the 2.24 μm band is between 2.237 and 2.244 μm (red markers). Right: Voyager 2/Imaging Science System image mosaic of Ariel (courtesy of NASA/JPL/Caltech/USGS, http://maps.jpl.nasa.gov/Uranus.html), with night-side sections illuminated by “Uranus shine” (Stryk & Stooke 2008). The mid-observation longitudes and latitudes for all 32 Ariel spectra are indicated with dots that represent the center of the target disk at the time of each observation (each collected spectrum averages over an entire hemisphere). The 12 spectra shown on the left are indicated by color-filled dots (numbered 1 through 12). Spectra that do not display 2.2 μm bands are shown as black-filled dots (i.e., spectra with band areas and depths <2σ). For continuum-divided examples of the 2.2 and 2.24 μm bands, please see Figure 3. NH$_3$- and NH$_4$-rich constituents that could be contributing to these bands are investigated in Section 3.4.
Hapke-based synthetic spectra (Appendix A.1), generated using optical constants for NH₃ ice (Sill et al. 1980) and amorphous NH₃ (Roser et al. 2018), mixed with H₂O ice, CO₂ ice, and amorphous C.

Because the detected 2.2 μm features are fairly weak and display morphological differences between the different Ariel spectra (i.e., variable band centers, areas, and depths), we focused our comparison on the five Ariel spectra displaying the strongest 2.2 μm bands (>3σ band areas and depths; Figure 3). This comparison demonstrates that the 2.2 μm bands in the spectra collected at mid-observation longitudes 79°8, 144°7, and 159°7 (Spectra 3, 5, and 6, respectively, in Figures 1 and 3) have band shapes and central wavelengths that are remarkably consistent with the laboratory spectra of NH₃–H₂O solutions (2.210 μm) and NH₃-hydrates (2.215 μm). The 2.2 μm bands in the spectra collected at mid-observation longitudes 53°6 and 316°6 (Spectra 2 and 11, respectively, in Figures 1 and 3) have band centers shifted to shorter wavelengths, possibly consistent with (NH₃)₂CO (2.181 μm), or perhaps other NH₂-bearing salts. Albeit, the shape of the 2.2 μm band in these two Ariel spectra is not a close match to the shape of the spectral continuum of (NH₃)₂CO. Three of the four spectra displaying 2.24 μm bands have band centers that are close matches to NH₃ ice (2.238 μm), while the fourth spectrum (Spectrum 12 in Figures 1 and 3) has a 2.24 μm band shifted to slightly longer wavelengths. Amorphous NH₃ does not provide a good match to either the 2.2 μm or 2.24 μm band. This mismatch is perhaps unsurprising given that amorphous NH₃ transitions to poly-crystalline ice at 65 K (Dawes et al. 2007), and is therefore unstable at Ariel’s estimated peak surface temperatures (~80–90 K near the subsolar point, Sori et al. 2017). Thus, laboratory reflectance spectra of NH₃–H₂O solutions and NH₃-hydrates, and synthetic spectra including NH₃ ice, represent the best matches to the 2.2 and 2.24 μm bands detected in the Ariel spectra.

4. Discussion and Conclusions

Our analysis demonstrates that a 2.2 μm band is present at the >2σ level in 10 spectra collected over a wide range of sub-observer longitudes and latitudes on Ariel. We find no leading/trailing hemispherical trends in the distribution of the 2.2 μm band, unlike the distribution of CO₂ and H₂O ice on this moon (e.g., Grundy et al. 2006). The longitudinal distributions of CO₂ and H₂O are likely controlled by exogenic processes that modify the surface chemistry of Ariel, in particular charged particle radiolysis and dust impacts (Grundy et al. 2003, 2006; Cartwright et al. 2015, 2018). The lack of spatial trends in the distribution of the 2.2 μm band supports an origin from geologic landforms distributed across Ariel’s surface. Impact craters, tectonic faults and fractures, potential cryovolcanic constructs, and mass wasting features have been identified across Ariel, and these landforms could represent suitable
Although other constituents like CH$_4$ ice (e.g., Gerakines et al. 2005), complex organics (e.g., Cruikshank et al. 1991), and hydrated silicates (e.g., Clark et al. 1990) have prominent absorption features spanning the wavelength region of the 2.2 and 2.24 $\mu$m bands, these species are less likely to be the dominant contributors to these two bands. It is unlikely that CH$_4$ ice is currently present on Ariel’s surface because it is highly volatile and should sublimate rapidly at Ariel’s estimated peak surface temperatures ($\sim$80–90 K). Complex organics and hydrated silicates are much less volatile and could be present in the spectrally red material observed on the Uranian moons (Cartwright et al. 2018). This red material likely originated as dust particles liberated from the surfaces of Uranus’ retrograde irregular satellites, which migrated inward due to Poynting–Robertson drag and mantled the leading hemispheres of the classical moons, in particular the outer moons Titania and Oberon (e.g., Tamayo et al. 2013). However, the 2.2 and 2.24 $\mu$m bands are stronger on Ariel than on Titania and Oberon, and red material is relatively scarce on Ariel compared to the outer moons (Cartwright et al. 2018). Furthermore, the 2.2 and 2.24 $\mu$m bands are present over a wide range of longitudes on Ariel, spanning its leading and trailing hemispheres (Figure 1), unlike the distribution of red material, which is concentrated on the leading hemispheres of the Uranian moons. Therefore, NH$_3$-bearing constituents sourced from Ariel’s interior are better candidate species to explain the presence of the 2.2 and 2.24 $\mu$m bands compared to complex organics or hydrated silicates delivered by dust impacts.

Bombardment by charged particles trapped in Uranus’ magnetosphere should efficiently decompose NH$_4$-rich deposits exposed on Ariel’s surface in only $\sim$10$^6$ yr (Moore et al. 2007). Albeit, accurate models of moon–magnetosphere interactions at Uranus are lacking, and it is possible that charged particle weathering of NH$_4$-rich species is more efficient in some locations (i.e., on Ariel’s trailing hemisphere) compared to others (i.e., on Ariel’s leading hemisphere). Energetic protons and ultraviolet (UV) solar photons are absorbed within the top 10 $\mu$m of H$_2$O icerich surfaces (e.g., Delitsky & Lane1998). The surface of Ariel and the other Uranian moons could be mantled by a thin layer of small H$_2$O ice grains ($\sim$10 $\mu$m thick), with other constituents like CO$_2$ ice retained beneath this topmost layer (e.g., Cartwright et al. 2020a). The average NIR photon penetration depth into Ariel’s regolith at 2.2 $\mu$m and 2.24 $\mu$m is $\sim$1.2 mm and $\sim$1.6 mm, respectively (e.g., Cartwright et al. 2020a). Therefore, NH$_3$-bearing species retained beneath the top $\sim$10 $\mu$m of Ariel’s surface might not interact with UV photons or energetic protons. Energetic electrons ($\sim$1 MeV), however, can penetrate up to centimeter-scale depths into icy satellite regoliths (e.g., Nordheim et al. 2017) and should readily interact with NH$_3$-bearing species and other constituents retained at depth. Furthermore, the flux of $\sim$1 MeV electrons is relatively high at the Uranian moons, comparable to the Jovian system (e.g., Mauk & Fox 2010). Nevertheless, an overlying veneer of H$_2$O ice might reduce the destruction rate of NH$_4$-rich deposits, and thereby increase their retention timescales on Ariel.

NH$_3$ molecules will either recombine back into NH$_3$ (e.g., Cruikshank et al. 2019) or form NH$_4^+$ ions (e.g., Moore et al. 2007). These NH$_4^+$ ions could then interact with other constituents, including CO$_2$ ice, to form less volatile salts like (NH$_4$)$_2$CO$_3$, which would contribute to the 2.2 $\mu$m band. Charged particle bombardment of NH$_3$, H$_2$O ice, and CO$_2$ ice...
could spur production of N-rich complex organics as well (e.g., Allamandola et al. 1988), which would also contribute to the 2.2 μm band. Therefore, NH₃-bearing deposits retained at depth, and/or converted into other N-bearing species, could persist for longer periods of time than the estimated destruction rate for NH₃ exposed on Ariel’s surface.

Alternatively, NH₃-rich geologic landforms could be younger than the regional scale crater density age estimates (~1–2 Ga), and thus, NH₃ might have been exposed more recently. A similar scenario exists on Pluto’s moon Charon where young and fresh craters with bright ejecta deposits, like Organa, Nasreddin, Skywalker, and Candide, exhibit notably higher concentrations of NH₃ compared to the surrounding terrain (e.g., Grundy et al. 2016; Protopapa et al. 2020). NH₃ diffuses fairly rapidly through H₂O ice (4.0 × 10⁻¹⁰ cm² s⁻¹ at 140 K, Livingston et al. 2002), and geologic processes that increase near-surface porosity and fracturing could increase this diffusion rate further. Consequently, geologic features formed by cryovolcanism, tectonism, impact events, and mass wasting might represent ideal conduits for the diffusion of NH₃ through H₂O ice-rich regoliths, thereby increasing the timescales over which the spectral signature of NH₃-bearing species persist in these landforms. However, the low estimated surface temperatures of Ariel and the other Uranian moons (30–90 K; Sori et al. 2017) could substantially reduce the diffusion rate of NH₃ through H₂O ice, complicating this scenario. To more thoroughly investigate the possible connection between NH₃ and geologic landforms on Ariel, updated estimates of the retention timescales for NH₃ and surface age estimates, using local scale crater densities, are needed. Nevertheless, the spectral evidence presented here is consistent with Ariel experiencing geologic activity in the recent past, including possible emplacement of NH₃-rich cryolavas sourced from its interior. These results support the interpretation that Ariel is a possible ocean world, which has, or had, a global or regional subsurface liquid H₂O layer that communicated with its surface (Hendrix et al. 2019).

Similar to Ariel, the four other classical Uranian moons display geologic landforms that suggest surface-interior communication via tectonism and cryovolcanism, in particular Miranda (e.g., Smith et al. 1986; Schenk 1991; Bedingfield et al. 2015; Bedingfield & Cartwright 2020b). NIR reflectance spectra of these four moons also display 2.2 μm bands (Bauer et al. 2002; Cartwright et al. 2018). Therefore, the presence of
NH₃-rich constituents might have contributed to geologic activity on the other classical Uranian moons as well. Future telescope observations, made with available and proposed facilities, are needed to further investigate the spectral signature and spatial distribution of the 2.2 µm band on the Uranian moons (Cartwright et al. 2019). Data collected by an orbiting spacecraft in the Uranian system would revolutionize our understanding of these moons, providing spatially resolved data sets that would allow for unprecedented investigation of whether they are, or were, ocean worlds (Cartwright et al. 2020b).

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Appendix

A.1. Methods: Hapke–Mie Spectral Modeling Procedure

The radiative transfer models utilized in this study were generated using a hybrid Hapke–Mie approach (example shown in Figure A1), which has been applied previously to IRTF/Spex data sets of these moons (Cartwright et al. 2015, 2018, 2020a). This hybrid approach calculates the single scattering albedo ($\bar{\omega}_0$) for each constituent using Mie theory (e.g., Bohren & Huffman 1983). These $\bar{\omega}_0$ values are then utilized by Hapke equations to model the scattering properties of the observed regolith (e.g., Hapke 2012). Mie theory models scattering and absorption by spherical particles that are distributed at random distances from one another. Mie theory can be used to model particles of any grain size. Consequently, $\bar{\omega}_0$ values calculated using Mie theory represent a valuable technique for generating synthetic spectra because many scattering models, including “pure” Hapke models, cannot generate robust results when considering constituents with grain diameters comparable to, or smaller than, the wavelength of incident light (e.g., Emery et al. 2006). To remove low amplitude resonance artifacts that can occur in scattering models that utilize Mie theory, our modeling software uses a small range of grain diameters (~10% spread) that are subsequently averaged together into the specified grain size for each constituent.

A.2. Results: IRTF/Spex Spectra

In this appendix, we present nine new near-infrared spectra of Ariel in Figure A2.

A.3. Results: 2.24 µm Band Parameter Measurements

In this appendix, we report band parameter measurements for the 2.24-micron feature in Table A1.

A.4. Results: Spatial Distribution of the 2.2 µm Band

In this appendix, we report the results of our F-test analysis in Table A2.
Figure A2. Nine new IRTF/Spex spectra of Ariel and their 1σ uncertainties, collected between 2017 and 2019. The mid-observation, sub-observer longitude for each spectrum is included in the bottom left-hand corner of each plot (see Table 1 for observation details). All spectra have been normalized to 1 between 2.24 and 2.25 μm.

Table A1
Measurements of the 2.24 μm Band Areas and Depths

| Sub-observer Long. (°) | Sub-observer Lat. (°) | Wavelength Range (μm) | Band Area (10^-4 μm) | Band Depth (μm) | >2σ Band Area and Depth Measurement? | Band Center (μm) |
|------------------------|------------------------|------------------------|------------------------|-----------------|--------------------------------------|------------------|
| 137.6                  | 34.6                   | 2.218–2.253            | 3.71 ± 1.22            | 0.023 ± 0.011   | Yes                                  | 2.237            |
| 304.8                  | –23.2                  | 2.225–2.255            | 4.37 ± 1.85            | 0.033 ± 0.016   | Yes                                  | 2.238            |
| 316.6                  | –18.2                  | 2.220–2.260            | 6.22 ± 1.55            | 0.037 ± 0.013   | Yes                                  | 2.238            |
| 334.4                  | 39.7                   | 2.227–2.262            | 5.09 ± 0.86            | 0.028 ± 0.006   | Yes                                  | 2.244            |

Table A2
F-test Analysis of the Longitudinal Distribution of the 2.2 μm Band

| Analyzed Measurement | One Tailed F-test Ratio | Sample Size (n) | Mean Model Degree of Freedom (n – 1) | Sinusoidal Model Degree of Freedom (n – 3) | Probability (p) | Reject Null Hypothesis? |
|----------------------|-------------------------|-----------------|--------------------------------------|--------------------------------------------|----------------|------------------------|
| 2.2 μm Band Areas    | 1.659                   | 32              | 31                                   | 29                                         | 0.087          | No                     |
| 2.2 μm Band Depths   | 1.603                   | 32              | 31                                   | 29                                         | 0.102          | No                     |
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