Evolution of stratospheric chemistry in the Saturn storm beacon region

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\textbf{Abstract}

The giant northern-hemisphere storm that erupted on Saturn in December 2010 triggered significant changes in stratospheric temperatures and species abundances that persisted for more than a year after the original outburst. The stratospheric regions affected by the storm have been nicknamed “beacons” due to their prominent infrared-emission signatures (Fletcher, L.N. et al. [2011]. Science 332, 1413). The two beacon regions that were present initially merged in April 2011 to form a single, large, anticyclonic vortex (Fletcher, L.N. et al. [2012]. Icarus 221, 560). We model the expected photochemical evolution of the stratospheric constituents in the beacons from the initial storm onset through the merger and on out to March 2012. The results are compared with longitudinally resolved \textit{Cassini}/CIRS spectra from May 2011. If we ignore potential changes due to vertical winds within the beacon, we find that C\textsubscript{2}H\textsubscript{2}, C\textsubscript{2}H\textsubscript{6}, and C\textsubscript{3}H\textsubscript{8} remain unaffected by the increased stratospheric temperatures in the beacon, the abundance of the shorter-lived CH\textsubscript{3}C\textsubscript{2}H decreases, and the abundance of C\textsubscript{2}H\textsubscript{4} increases significantly due to the elevated temperatures, the latter most notably in a secondary mixing-ratio peak located near mbar pressures. The C\textsubscript{4}H\textsubscript{2} abundance in the model decreases by a factor of a few in the 0.01–10 mbar region but has a significant increase in the 10–30 mbar region due to evaporation of the previously condensed phase. The column abundances of C\textsubscript{6}H\textsubscript{6} and H\textsubscript{2}O above ~30 mbar also increase due to aerosol evaporation. Model-data comparisons show that models that consider temperature changes alone under-predict the abundance of C\textsubscript{2}H\textsubscript{x} species by a factor of 2–7 in the beacon core in May 2011, suggesting that other processes not considered by the models, such as downwelling winds in the vortex, are affecting the species profiles. Additional calculations indicate that downwelling winds of order ~10 cm s\textsuperscript{-1} near ~0.1 mbar need to be included in the photochemical models in order to explain the inferred C\textsubscript{2}H\textsubscript{2} abundances in the beacon core, indicating that both strong subsiding winds and chemistry at elevated temperatures are affecting the vertical profiles of atmospheric constituents in the beacon. We (i) discuss the general chemical behavior of stratospheric species in the beacon region, (ii) demonstrate how the evolving beacon environment affects the species vertical profiles and emission characteristics (both with and without the presence of vertical winds), (iii) make predictions with respect to compositional changes that can be tested against \textit{Cassini} and \textit{Herschel} data, and higher-spectral-resolution ground-based observations of the beacon region, and (iv) discuss future measurements and modeling that could further our understanding of the dynamical origin, evolution, and chemical processing within these unexpected stratospheric vortices that were generated after the 2010 convective event.

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1. Introduction

The pristine, hazy appearance of Saturn, with its muted atmospheric banding, is known to be disturbed on rare occasions by enormous convective storms dubbed “Great White Spots” (e.g., Sanchez Lavega, 1982; Sanchez Lavega and Battaner, 1987).

In December 2010, one such gigantic storm system erupted at northern mid-latitudes on Saturn (Sánchez-Lavega et al., 2011; Fischer et al., 2011; Fletcher et al., 2011). The “head” of the storm drifted westward with the prevailing zonal winds, leaving a turbulent wake of fresh cloud particles. Within a couple of months of the storm onset, the storm head had caught up with its wake “tail” to form a distinct planet-encircling band of clouds that persisted for more than a year after the storm’s initial appearance.

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(e.g., Sánchez-Lavega et al., 2012; Sayanagi et al., 2013). Although the convective disturbance originated in the troposphere and had a notable effect on the cloud structure, lightning activity, atmospheric dynamics, thermal structure, and distribution of molecular species within the troposphere (Fischer et al., 2011; Sánchez-Lavega et al., 2011, 2012; Fletcher et al., 2011, 2012; Hurley et al., 2012; Sanz-Requena et al., 2012; Janssen et al., 2013; Laraia et al., 2013; Sayanagi et al., 2013; Dyudina et al., 2013; Sromovsky et al., 2013; Achterberg et al., 2014; Trammell et al., 2014), the storm also had some profound and unexpected consequences for higher-altitude regions. In particular, temperatures in the stratosphere were found to be greatly elevated in latitude regions associated with the storm, perhaps as a result of momentum and energy redistribution from vertically-propagating atmospheric waves generated from tropospheric convective plume activity and/or from dynamical compression within the resulting vortex region (Sayanagi and Showman, 2007; Fletcher et al., 2011, 2012). In addition, the gas-phase abundances of ethylene and water were inferred to have increased by roughly two orders of magnitude in these high-temperature stratospheric regions in the months after the storm onset (Hesman et al., 2012; Cavalé et al., 2012).

The strong stratospheric temperature increase was initially confined to two broad air masses nicknamed “beacons” due to their distinctive bright signatures at infrared wavelengths (Fletcher et al., 2011). These two initial air masses, centered at different longitudes/lattitudes and associated with zonal winds of different relative velocities, encountered each other in April 2011, at which point the two beacons merged into a single, enormous, anticyclonic vortex (Fletcher et al., 2012). Temperatures within the initial two beacons rose rapidly in the months before the merger, intensified and reached a maximum in the combined beacon vortex after the merger, and then cooled slowly but steadily from May 2011 onward (Fletcher et al., 2012; see also Fletcher et al., 2011; Hesman et al., 2012).

The Cassini spacecraft was in a prime position to track the evolution of the storm and its associated beacon features. Fig. 1 shows the vertical temperature profiles derived by Fletcher et al. (2012) from spectra acquired with the Composite Infrared Spectrometer (CIRS) instrument aboard Cassini. These temperature retrievals were obtained from spectra coadded over broad areas of the beacons (i.e., within ±10° longitude, ±5° latitude of the beacon center)—temperatures within the hottest regions at the beacon centers were even higher. For example, on May 4, 2011, after the merger, 2-mbar temperatures at the central “core” of the beacon reached ~220 K, about 80 K greater than the pre-storm temperature (Fletcher et al., 2012; see also Hesman et al., 2012), whereas the broader-scale averages indicated temperatures of ~210 K at 2 mbar.

The higher temperatures resulted in increased infrared emission, making molecular bands from trace stratospheric constituents easier to identify. One such example is ethylene (C2H4), which was not identified in CIRS spectra before the storm at northern mid-latitudes, but which was detected by Hesman et al. (2012) in the post-storm beacon region in May 2011, from both Cassini CIRS data and ground-based infrared observations. Hesman et al. (2012) derived stratospheric temperatures at ~0.5–5 mbar using the v1 band of methane (CH4) in the 1250–1311 cm−1 wavenumber region, which then allowed them to retrieve the C2H4 abundance from the observed ethylene emission band near 950 cm−1. The retrievals of the ethylene abundance profile are complicated by the possibility that the C2H4 emission may not originate from the 0.5–5-mbar pressure levels where the temperatures are best constrained; however, the Hesman et al. (2012) analysis clearly indicates that the ethylene abundance in May 2011 was significantly increased in the beacon region at ~10−10−2 mbar in comparison with pre-storm observations and expectations (Fig. 2). In fact, Hesman et al. (2012) found that their pre-storm photochemical-model profile for C2H4 would need to be increased uniformly by almost two orders of magnitude in order to reproduce the observed ethylene emission from the beacon, whereas their photochemical models predicted only a factor of ~2 increase in the C2H4 mixing ratio due to the elevated temperatures in the beacon. Hesman et al. (2012) explored several ideas as to the mechanisms that could be the cause of the C2H4 enhancement, but they did not come up with a definitive conclusion. Fig. 2 shows...
their predicted pre-storm ethylene profile, in comparison with their post-storm ethylene profiles, in the mbar region.

Acetylene was also observed to increase in the beacon region after the merger (Fletcher et al., 2012; Hesman et al., 2014), albeit less dramatically than ethylene. In contrast, a preliminary analysis by Hesman et al. (2014) indicates that CH3C2H, C3H8, and C4H2 exhibit little or no enhancement in the beacon, and Fletcher et al. (2012) find that the beacon enhancement of C2H6 was at the level of retrieval uncertainty and therefore inferred to be smaller than that of C2H5.

In an attempt to explain these observations, Cavalié et al. (2015) used a photochemical model to track the expected evolution of hydrocarbon chemistry in the beacon region. Their models predicted a small fraction of C2H6 abundance at mbar pressures due to the elevated beacon temperatures—an amount that is insufficient to explain the observed ethylene emission reported by Hesman et al. (2012). The Cavalié et al. (2015) model also predicted very little change in the acetylene (C2H2) and ethane (C2H6) abundances at mbar pressures due to the elevated beacon temperatures, in contrast to the post-merger beacon observations of Fletcher et al. (2012) and Hesman et al. (2014).

In this paper, we further explore the theoretical chemical evolution of stratospheric hydrocarbon and oxygen species in the storm beacon region in an attempt to reconcile models and observations and to better understand the physical and chemical conditions within this unusual stratospheric anticyclonic vortex on Saturn.

2. Photochemical model

To calculate the vertical distribution of stratospheric species in the beacon region on Saturn, we use the Caltech/JPL KINETICS code developed by Allen et al. (1981) and Yung et al. (1984) to solve the coupled one-dimensional (1-D) continuity equations for each species i in the model:

$$\frac{\partial n_i}{\partial t} + \frac{\partial \Phi_i}{\partial z} = P_i - L_i,$$

where \(n_i\) is the number density (cm\(^{-3}\)), \(\Phi_i\) is the vertical flux (cm\(^{-2}\) s\(^{-1}\)), and \(P_i\) and \(L_i\) are, respectively, the chemical production and loss rates (cm\(^{-3}\) s\(^{-1}\)) of the \(i\)-th species, all of which are explicit functions of time \(t\) and altitude \(z\). The flux term is calculated for the vertical direction only and includes transport by molecular diffusion, eddy diffusion, and potential vertical winds:

$$\Phi_i = -n_i D_i \left( \frac{1}{n_i} \frac{\partial n_i}{\partial z} + \frac{1}{H_i} \frac{\partial H_i}{\partial z} \left( 1 + \gamma_i \frac{dT}{dz} \right) \right) - n_i K_{zz} \left( \frac{1}{n_i} \frac{\partial n_i}{\partial z} + \frac{1}{H_i} \frac{\partial H_i}{\partial z} + \frac{1}{T} \frac{\partial T}{\partial z} \right) + n_i w,$$

where \(D_i\) is the molecular diffusion coefficient (cm\(^2\) s\(^{-1}\)), \(H_i\) is the pressure scale height (cm) of the \(i\)-th constituent, \(K_{zz}\) is the vertical eddy diffusion coefficient (cm\(^2\) s\(^{-1}\)), and \(w\) is the vertical wind velocity (cm s\(^{-1}\)). Vertical winds are typically neglected in 1-D models, given that such models are most often intended to describe global, steady-state averages; however, we include vertical winds in some specific test cases to better describe the behavior in the beacon vortex.

The hydrocarbon chemical reaction mechanism in our model is taken from Model C of Moses et al. (2005), with updates to several association reactions (including radiative association) based on the recommendations of Vuitton et al. (2012), and updates to several reactions involving C5H4 species based on recommendations of Hébrard et al. (2013). The rate coefficients for reactions involving oxygen species are taken from Moses et al. (2000b). The model contains 70 hydrocarbon and oxygen species that interact via ~500 chemical reactions. Condensation and evaporation of water (H2O), diacetylene (C2H4), and benzene (C6H6) are considered in a manner described in Moses et al. (2000b). The expressions for the vapor pressures of H2O, C2H4, and C6H6 over their respective ices are taken from Marti and Mauersberger (1993), Orton et al. (2014), and Reid and Prausnitz (1987); see also Fray and Schmitt (2009). Model calculations are performed for 34° planetocentric latitude (~40° planetographic latitude), relevant to the beacon center after the merger, and we consider diurnally averaged fluxes, fixed seasonal parameters near equinox, and a low-to-average solar ultraviolet flux (see Moses et al., 2000a, for details). These choices are appropriate for the beacon situation in 2010–2011, and none have much influence on the time-variable results over the short time period of the beacon model.

The model atmospheric grid contains 198 pressure levels, ranging from 5.1 bar to 10\(^{-8}\) mbar. At the lower boundary, the helium and methane mole fractions are fixed at, respectively, 0.119 (Conrath and Gautier, 2000) and 4.7 \times 10\(^{-7}\) (Fletcher et al., 2009), and the carbon monoxide mole fraction is fixed at 1.0 \times 10\(^{-9}\), which is the upper limit for tropospheric CO derived by Cavalié et al. (2009). All other trace species are assumed to have a concentration gradient of zero at the lower boundary, which causes these species to flow through the lower boundary at a maximum possible velocity. The lower boundary is far removed from the stratospheric region of interest in this problem, and our choice of the lower boundary condition for the photochemically produced species has no effect on our results. Atomic H, some of which is produced photochemically in the high-altitude thermosphere and ionosphere above the top of our model, is assumed to have a downward flux of 1.0 \times 10\(^{6}\) cm\(^{-2}\) s\(^{-1}\) at the upper boundary of our model, whereas all other species are given zero flux boundary condition at the top of the model (cf. Moses et al., 2000a, 2005). Water, CO, and CO2 are assumed to be introduced to the atmosphere from external sources (Feuchtgruber et al., 1997, 1999; de Graauw et al., 1997; Moses et al., 2000b; Bergin et al., 2000; Cavalié et al., 2009, 2010). The ultimate origin of the external oxygen compounds is uncertain. Guerlet et al. (2010) demonstrate from back-of-the-envelope calculations that Enceladus could be the dominant source (see also Jurac and Richardson, 2007; Cassidy and Johnson, 2010; Hartogh et al., 2011; Flesham et al., 2012), while Cavalié et al. (2010) favor a relatively recent cometary impact within the past ~200–250 years. For simplicity, we assume that the external oxygen species are introduced to the atmosphere through ablation of small icy grains, with assumed influx rates of 8.5 \times 10\(^{7}\) H2O molecules cm\(^{-2}\) s\(^{-1}\), 4.1 \times 10\(^{10}\) CO molecules cm\(^{-2}\) s\(^{-1}\), and 1.2 \times 10\(^{10}\) CO2 molecules cm\(^{-2}\) s\(^{-1}\) (cf. Moses et al., 2000b).

These fluxes, in combination with our inferred pre-storm \(K_p\) profile, thermal structure, and chemical reaction mechanism, reproduce the observed global-average stratospheric abundances of H2O and CO2 from observations from the Infrared Space Observatory (ISO) (de Graauw et al., 1997; Feuchtgruber et al., 1997, 1999; Moses et al., 2000b).

The temperature–pressure profiles adopted in the model are shown in Fig. 1. The pre-storm temperature profile is taken from CIRS temperature retrievals averaged over 36–44° planetographic latitude from spectra acquired in May–August 2010 (see Section 3). The profiles adopted after the storm onset (hereafter called "post-storm") are the Fletcher et al. (2012) retrievals from coadded CIRS spectra acquired from within a 10° latitude and 20° longitude region centered over the initial ‘B1’ and merged ‘B0’ beacons from 18 separate dates ranging from January 2, 2011 (~1 month after storm onset) to March 14, 2012 (last available data from the Fletcher et al., 2012 study; see Fig. 1). The May 4, 2011 CIRS
observations are of particularly high quality (i.e., high signal-to-noise ratio), and we adopt the retrieved temperatures from the hottest longitude region at the beacon core (see the dotted line in Fig. 1) for some models. Although the CIRS temperature retrievals are most sensitive to the $\sim$0.5–230 mbar pressure range, Fletcher et al. (2012) present retrieved temperatures over a broader range from 10 bar to $10^{-3}$ mbar, and we adopt these values over that entire pressure range. At higher altitudes, we smoothly (and arbitrarily) connect the Fletcher et al. (2012) profiles to a thermospheric temperature profile derived from Voyager Ultraviolet Spectrometer (UVS) occultation observations (Vervack and Moses, 2015). Note that the full 198-level pressure range was used in the retrievals from the hot beacon core shown by the dotted line in Fig. 1, which is why the high-altitude profile for that curve differs from the others. We will show results assuming both of these May 4, 2011 profiles, with the hot-beacon core profile referred to as the “hot” nominal model, and the beacon-average profile as the “beacon-average” nominal model. We then determine the complete background atmospheric grid for these temperature profiles by assuming hydrostatic equilibrium. That is, the pressure grid is kept constant for the different dates, and the altitude and density profiles are calculated from the temperature–pressure profiles via solution of the hydrostatic-equilibrium equation.

Our modeling procedure is to first run the photochemical model for the fixed-season, pre-storm conditions at 40° planetographic latitude, allowing the solution to converge and reach a steady state. The eddy $K_{zz}$ profile, which is a free parameter in the model, is adjusted in this pre-storm model (see Fig. 3) until the C$_2$H$_6$ and C$_2$H$_2$ mixing ratios are consistent with the CIRS pre-storm emission at the relevant 40° latitude. The scaling factors—i.e., the uniform-with-altitude multiplicative factors—that the model mixing-ratio profiles need to be scaled by to reproduce the pre-storm (May–August 2010) CIRS zonal-mean nadir spectra are shown in Fig. 4 (see also the discussion of these observations in Section 3). The pre-storm model underestimates the acetylene abundance slightly, such that the C$_2$H$_2$ profile needs to be scaled by 1.07 to fit the CIRS spectra at 40° planetographic latitude, whereas the pre-storm model overestimates ethane slightly, such that the model C$_2$H$_6$ mixing ratios need to be scaled by $\sim$0.91 to explain the pre-storm ethane emission at that latitude. In fact, although the reaction mechanism used in this study provides a good representation of the global-average hydrocarbon abundances on Saturn (e.g., Moses et al., 2005), existing 1-D photochemical models for Saturn do not reproduce the CIRS observations for the meridional distribution—and in some cases the vertical distribution—of all observed hydrocarbon species at all locations on Saturn (Moses and Greathouse, 2005; Guerlet et al., 2009, 2010; Sinclair et al., 2013; Hue et al., 2015). Some of the model-data mismatch likely results from the lack of atmospheric circulation in the photochemical

Fig. 3. The eddy diffusion coefficient ($K_{zz}$) profile adopted in our models (solid line), along with the CH$_4$ molecular diffusion coefficient profile for the pre-storm thermal profile (dashed line) and the post-storm “hot” beacon core thermal profile from May 4, 2011 (dotted line). Note that because the $K_{zz}$ profile is defined as a function of pressure and because the temperature profiles are similar at high altitudes, the methane homopause pressure level (i.e., where $K_{zz}$ equals the CH$_4$ molecular diffusion coefficient) does not change between the pre-storm and post-storm models.

Fig. 4. Zonal mean retrievals of pre-storm conditions derived from CIRS nadir data from May–August 2010 as a function of planetographic latitude: temperatures (a), and scaling factors for acetylene (b), ethane (c). The scaling factors are uniform-with-height multiplicative factors that have been applied to the mixing-ratio profiles predicted by our pre-storm photochemical model, with several model iterations conducted to ensure that these scaling factors are close to unity at the location of the beacon core (approximately 40°N). Note that ethylene is not detected in these pre-storm observations, as is consistent with the predictions from the pre-storm model.
models, but the chemistry itself may also be incomplete or inaccurate. It is precisely for this reason that the elevated temperatures within the beacon region provide a useful “laboratory” test case to evaluate the viability of the chemical mechanism, or at least to provide insight into the key temperature-sensitive reactions involved in stratospheric chemistry on Saturn.

Once a pre-storm profile has been established, we use the converged pre-storm photochemical-model solution as our initial condition and run the time-variable model for 40° planetographic latitude, starting at December 5, 2010 with the pre-storm temperatures, and then let the temperature profiles (and atmospheric grid) vary as a function of time for the 15 months for which CIRS beacon data have been reported. The observational data are spaced unevenly in time, and we simply update the model temperature profiles at the halfway point between each of the observations. This choice of when to update the thermal structure is arbitrary and can affect the results for the shortest-lived molecules (including C2H4), but modifications to this assumption result in only small differences in the quantitative conclusions. The ending mole-fraction profiles for the results from one time segment at one temperature are passed on to the next run as initial conditions for the new temperature sequence. The stratospheric gas abundances thus evolve with time as the temperatures in the beacon regions change. For our initial set of models, we ignore any dynamical or eddy-diffusion changes within the beacon region, keeping Kzz fixed at pre-storm values, but we later explore how vertical winds affect the results.

3. Observations and spectral modeling

The observations discussed in this paper were obtained from the Cassini CIRS Fourier transform spectrometer (Flasar et al., 2004), using the mid-infrared focal planes (FP3, covering 600–1000 cm⁻¹; FP4, covering 1100–1500 cm⁻¹). We adopt without modification the Fletcher et al. (2012) retrievals of the average thermal structure within the beacons from CIRS observations acquired in 2010–2012 with a variety of observing strategies and spectral resolutions (see Table 1 of Fletcher et al., 2012, and associated discussion). We also present new analyses of CIRS spectra at 2.5 cm⁻¹ spectral resolution acquired during two epochs: (i) previously unpublished pre-storm observations obtained in mid-2010 covering the 25–55°N latitude range (130SA_MIRMAP001 on May 5, 2010; 134SA_MIRMAP001 on July 10, 2010; 135SA_MIRMAP001 on July 19, 2010; and 137SA_MIRMAP001 on August 28, 2010); and (ii) a reanalysis of previously published post-storm observations acquired on May 4, 2011, shortly after the beacon merger event (148SA_MIRMAP001) that have been shown to exhibit enhanced acetylene (Fletcher et al., 2012) and ethylene (Hesman et al., 2012) emission within the beacon. The May 2011 observations have been averaged in 10°-wide bins on a 5° longitude grid. All spectra use 4000 calibration reference spectra to improve the signal-to-noise ratio in the observed emission features. We simultaneously retrieve atmospheric temperatures and hydrocarbon scale factors, exploiting methane emission between 1250 and 1350 cm⁻¹. H₂–He collision induced absorption from 600 to 1000 cm⁻¹ is used as a reference to calibrate the hydrocarbon abundances. The vertical profiles of several hydrocarbons (as labeled) as they evolve with time in the beacon for our “beacon-average” nominal photochemical model that uses the beacon-average thermal profiles (see Fig. 1). The color coding corresponds to the dates listed in the top left figure. The shorter-lived species C2H4, CH3C2H, and C4H2 are affected by the changing temperatures, while the longer-lived species C2H6, C2H2, and C3H8 are not. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
Inversions of spectral data can be prone to extreme degeneracy, where a wide variety of potential solutions can reproduce the data equally well. This situation is particularly true at mid-infrared wavelengths, where the magnitude of hydrocarbon emission features is governed by both their mixing ratio profiles and the atmospheric temperature structure in the line-forming region. Spectral retrieval algorithms, such as the NEMESIS model employed here (Irwin et al., 2008), use \textit{a priori} profiles to constrain solutions, but the resulting retrieved thermal and chemical distributions can sometimes be biased towards this prior information. In this study, we combine the photochemical modeling with spectral inversion, taking advantage of the synergistic nature of the forward and reverse modeling, to better assess how well the CIRS spectra can constrain the photochemical model. Throughout the analysis, we adopt the spectral inversion techniques described by Fletcher et al. (2011, 2012), using identical sources of spectroscopic line data. In each case, we use the predicted mixing ratio profiles for all hydrocarbon species in the photochemical model as \textit{prior} information, and scale these profiles simultaneously with a temperature retrieval to reproduce the CIRS measurements. Our goal is to find a set of theoretically derived mixing-ratio profiles, based on chemistry and vertical motions, that require minimal scaling in order to reproduce the CIRS emission features.

Several iterations between the photochemical model priors and the spectral fitting were required to (a) converge on a pre-storm model that reproduced the emission at 40°N planetographic latitude (Section 2); (b) determine that photochemistry at elevated temperatures alone is insufficient to explain the enhanced emissions (Section 4; see also Hesman et al., 2012; Cavalié et al., 2015); and (c) converge on a solution with subsiding winds that required minimal scaling of the model hydrocarbon profiles (Section 4.5). Examples of the spectral fits are shown in Section 5.

Throughout the paper, the errors in the retrieved species mixing ratios that we quote are the formal uncertainties from the optimal estimation procedure used by NEMESIS. These formal errors take into account measurement uncertainties, temperature uncertainties due to the degeneracy between abundance and temperature when fitting the observed emission, and a fractional error accounting for uncertainties in the spectral line database and other spectral modeling assumptions. The quoted uncertainties do not account for systematic errors. More importantly, they do not account for errors due to the uncertain shape of the species vertical profiles used to define the priors—the vertical model profiles are simply scaled uniformly at all altitudes until a best fit is obtained. As such, the formal errors will underestimate the true uncertainties, especially for pressure regions far removed from the peak of the contribution functions.

4. Results and discussion

For our nominal beacon model, we keep the \( K_{12} \) profile fixed at pre-storm values (Fig. 3), and we neglect vertical winds. Because time-variable dynamics are not being considered in the nominal model, changes in the mixing-ratio profiles of the hydrocarbons are caused solely by temperature-dependent reactions. Fig. 5 shows the predicted time variation for several important species from January 2, 2011 through March 14, 2012 from our “beacon-average” nominal model that assumes the beacon-average temperature profiles for all dates, including May 4, 2011. Note that \( \text{C}_2\text{H}_4 \) experiences a strong increase at mbar pressures due to the increased temperatures, while the \( \text{C}_2\text{H}_4 \) isomer methylacetylene \((\text{CH}_3\text{C}_2\text{H})\) and diacetylene \((\text{C}_2\text{H}_2)\) decrease in the \( 10^{-10} \) to \( 10^{-12} \) mbar region, and \( \text{C}_2\text{H}_2, \text{C}_2\text{H}_6, \) and \( \text{C}_3\text{H}_8 \) (propane) are unaffected by the temperature increase. Species that condense under pre-storm stratospheric conditions on Saturn, such as \( \text{H}_2\text{O}, \text{C}_2\text{H}_2, \) and \( \text{C}_2\text{H}_6 \) exhibit strong increases in abundance in the lower-stratosphere due to evaporation of the aerosols at the elevated temperatures in the beacon.

In Fig. 6, the pre-storm and post-storm (for May 4, 2011) \( \text{C}_2\text{H}_6 \) model profiles are compared with derived abundances from various global and local observations. The lighter green squares in Fig. 6 show the \( \text{C}_2\text{H}_2 \) and \( \text{C}_2\text{H}_6 \) mole fractions from our analysis of the May–August 2010 pre-storm CIRS nadir spectra, the darker green circles show the 2005–2006 pre-storm mole fractions from the CIRS limb data analysis of Guerlet et al. (2009, 2010) for 40° planetographic latitude, and the red squares show our retrieved mole fractions from the hot beacon-core region on May 4, 2011. Note that the \( \text{C}_2\text{H}_2 \) and \( \text{C}_2\text{H}_6 \) mole fractions are actually observed to increase in the hot beacon-core region in comparison with the pre-storm retrievals, whereas our models predict virtually no change in these species in the beacon over that time period. These nominal models without a temporally variable dynamical component underpredict the \( \text{C}_2\text{H}_4 \) abundances in the beacon by factors of \( \approx 2-7 \). If the observed beacon increases in \( \text{C}_2\text{H}_4 \) abundance were caused by temperature-dependent chemistry alone, the additional carbon would have to come from methane, as methane is the only sufficiently large source of local carbon. We are unable to identify any temperature-sensitive chemical reactions that efficiently convert methane to \( \text{C}_2\text{H}_4 \) species at mbar levels on Saturn on the short time scales involved, suggesting that dynamics may be contributing to the observed increase in \( \text{C}_2\text{H}_4 \) species in the beacon.

Fig. 6 also illustrates that the “hot” nominal model that uses the beacon-core temperatures from May 4, 2011 produces slightly more \( \text{C}_2\text{H}_6 \) at mbar pressures than the “beacon-average” nominal model that uses beacon-average temperatures from that date. In particular, the \( \sim 10^{-10} \) K temperature difference between the “hot”
and “beacon-average” model at ~2 mbar leads to 16% higher C2H2 mole fraction in the hot model.

Fig. 7 shows the scaling factors that need to be applied to our hot nominal beacon-core model profiles in order to fit the molecular emission from the longitudinally resolved spectra across the beacon. The resulting retrieved mole fractions at 1.8 mbar at the beacon central core at 294.8° System III longitude on May 4, 2011 are 1.08 (±0.02) × 10^{-6} for C2H2, 2.0 (±0.3) × 10^{-9} for C2H4, and 9.4 (±0.1) × 10^{-6} for C2H6, with the formal error bars neglecting both systematic uncertainties and uncertainties in the hydrocarbon vertical profiles (see Section 3). From Fig. 7, it is obvious that the beacon photochemical model underpredicts the C2H2 hydrocarbon abundances in the beacon vortex by factors of ≥ 2–7. These scale factors are in relation to the hot beacon-core model only, and are not necessarily directly comparable to the pre-storm model scaling factors shown in Fig. 4. However, recall that the C2H2 and C2H6 vertical profiles in the hot beacon-core model have not changed much in comparison with the pre-storm model. Fig. 7 therefore indicates that the C2H2 abundance has experienced a general factor of ~2 increase outside of the beacon vortex across the entire beacon latitude region observed on the May 4, 2011 date, with larger increases found within with the beacon vortex itself, and especially in the central core region at 294.8° System III longitude and a secondary peak at C2H2 peak at 273.3° longitude (the latter associated with a local high-altitude increase in temperature). Ethane also exhibits longitudinal structure, with the largest abundances located within the vortex itself, again at 273.3° and 294.8° longitude. At longitudes away from the vortex itself, the C2H6 scaling factors trend back to the pre-storm case on the east side, but the “hot” nominal model profile apparently overestimates the C2H6 abundance on the west side of the vortex, where temperatures are not as large. Note from Fig. 7a that the stratospheric temperatures themselves have also increased compared to the pre-storm case (cf. Fig. 4a) throughout the observed region, but especially within the vortex itself.

The large-scale longitudinal perturbations in temperatures and abundances throughout the storm latitude are not too surprising, as neither the tropospheric storm head nor the stratospheric beacon vortex were stationary with respect to System III longitude. The storm head drifted westward and encountered the southern branch of its wake tail within about 50 days of the storm’s appearance, forming a planet encircling band of active tropospheric clouds that continued to be influenced by each ~120-day circum-navigational pass of the storm head through the region (Sánchez-Lavega et al., 2011, 2012; Sayanagi et al., 2013). Similarly, the beacon vortices also drifted longitudinally at their own rates, with the final merged beacon vortex circling the planet once every 130 days or so (Fletcher et al., 2012). The fact that stratospheric warming was observed throughout the entire latitude band in which the beacon resides (Fletcher et al., 2012, see also Fig. 7) indicates that large regions of the stratosphere were perturbed by the storm, rather than the effects just being limited to the merged vortex itself. That is especially true at higher altitudes (see Fig. 11a of Fletcher et al., 2012), where a large swath of the northern hemisphere was observed to have enhanced temperatures in comparison with pre-storm values. If the tropospheric convective plumes provided a source of upward-propagating planetary and gravity waves that transported energy and momentum to the stratosphere (e.g., Fletcher et al., 2011, 2012), both stratospheric dynamics and temperatures could have been affected over broad regions, leading to different thermal and vertical abundance profiles across the latitude band. Indeed, the different zonal profiles for the different species suggest different chemical–dynamical–thermal coupling as a function of longitude across the beacon. Ethylene is more sensitive to the thermal structure than either acetylene or ethane, but the difference in morphology of all the

![Fig. 7.](image-url) Retrievals of temperatures and hydrocarbon distributions as a function of longitude through the beacon core on May 4, 2011. Spectra were averaged over the 36–43°N latitude range, and model-predicted mixing-ratio profiles from our nominal hot beacon-core photochemical model were uniformly scaled to reproduce the emission observed by CIRS. In all cases, the required scaling factors still exceed unity within the beacon, implying that the distributions must be affected by processes in addition to photochemistry.
C$_2$H$_6$ scaling factors as a function of longitude across the beacon suggests complicated longitudinally and vertically variable wind fields and a different chemical/dynamical response of the different species to the winds and temperatures at different altitudes.

The results regarding the chemical response of the individual species to the elevated beacon temperatures are discussed in detail below, while the potential effects of vertical winds are discussed in Section 4.5.

4.1. Nominal model results: ethane, acetylene, and propane

The lack of significant temporal evolution of C$_2$H$_6$, C$_2$H$_2$, and C$_3$H$_8$ in our nominal beacon model (Fig. 5) is due to the long chemical lifetime of these species, even at the higher temperatures experienced in the beacon. If we define the net chemical lifetime $\tau_{chem}$ of a species $i$ as $n_i/|P_i - L_i|$, where $n_i$ is the concentration (cm$^{-3}$) and $P_i$ and $L_i$ are the chemical production and loss rates (cm$^{-3}$ s$^{-1}$) at any particular altitude, we find that $\tau_{chem}$ at the temperature maximum at ~2 mbar in the hot beacon core is 580 (Earth) yrs for C$_2$H$_6$, 1.4 yrs for C$_2$H$_2$, and 6.9 yrs for C$_3$H$_8$. In contrast, the highest beacon temperatures were observed in May of 2011, only 5 months after the storm onset. The C$_2$H$_6$, C$_2$H$_2$, and C$_3$H$_8$ abundances in the model have not had enough time to respond chemically to the elevated temperatures in a notable way by the May 2011 observations. Considering the fact that the beacon has been cooling slowly but steadily since May 2011, we do not expect continued evolution of C$_2$H$_6$ and C$_2$H$_2$ due to chemistry alone, but Fig. 5 does illustrate some slight expected changes in the C$_2$H$_6$ abundance over the ~1.3-year span of the published CIRS observations.

Chemical loss exceeds production of C$_2$H$_6$ over much of the pressure region in which temperatures are elevated within the beacon. The rate coefficient for the reaction number R130 in our list (H + C$_2$H$_6$ + M → C$_2$H$_3$ + M, with M representing any third-body molecule or atom) becomes much larger at the higher beacon temperatures, but the background density and H mole fraction drop in the elevated-temperature region (the latter largely due to reaction R130), leading to an overall ~60% increase in the loss rate of C$_2$H$_6$ at 2 mbar in the beacon on May 4, 2011; however, there is sufficient C$_2$H$_6$ at these pressures that the perturbation to the C$_2$H$_6$ abundance is minor over the relatively short time period involved. On the other hand, this increased loss of C$_2$H$_6$ in the beacon contributes to the increased production rate of less-abundant species such as C$_2$H$_4$ and ultimately C$_2$H$_2$.

The small loss of C$_2$H$_2$ in the beacon contributes to a ~70% increase in the production rate of C$_3$H$_8$ at 2 mbar in the hot beacon core on May 4, 2011, through the dominant pathway H + C$_2$H$_2$ + M → C$_2$H$_3$ + M, C$_2$H$_3$ + H → C$_2$H$_4$ + H, and H + C$_2$H$_4$ + M → C$_2$H$_2$ + M, followed by reaction of atomic H with C$_2$H$_2$ to either form C$_2$H$_4$ directly, or to form two CH radicals, which recombine to form C$_2$H$_2$. Given the already large C$_2$H$_4$ mole fraction at the relevant pressure levels, this increased production has an insignificant effect on the C$_2$H$_4$ mole fraction in the beacon over the time scales involved, however.

Like ethane, the production rate of propane exceeds its loss rate over much of the elevated temperature region of the beacon. The increased acetylene loss rate in the beacon leads to increased amounts of C$_2$H$_4$ and therefore C$_2$H$_2$ through the reaction R232: CH$_3$ + C$_2$H$_5$ + M → C$_2$H$_4$ + M. Again, however, the perturbation in the net production rate has little effect on the C$_2$H$_4$ mole fraction in the beacon over the relevant time scales.

We therefore do not expect much change in the abundance of ethane, acetylene, and propane in the beacon due to chemistry alone. Any observed changes in these species (e.g., Figs. 6 and 7) are likely caused by changes in dynamics within the beacon region. Cavalié et al. (2015) arrived at a similar conclusion for these species, although Cavalié et al. did predict small changes to the abundances of C$_2$H$_2$ and C$_2$H$_4$ at high stratospheric altitudes within the beacon. The differences at high altitudes between our model and that of Cavalié et al. (2015) are caused by transport and the prescription of high-altitude temperatures, which are not constrained by CIRS. In the Cavalié et al. (2015) model, the high-altitude temperatures are assumed to remain isothermal above 10$^{-3}$ mbar; the different dates then have different high-altitude temperatures, which leads to different high-altitude density structures and different pressure levels for the methane homopause at the different dates within the Cavalié et al. (2015) model. As is shown in Figs. 1 and 3, we assume in our models here that the upper-atmospheric temperatures are unperturbed by the beacon, so the thermal profiles from all dates converge at high altitudes, and the methane homopause pressure level does not change significantly with time in our model. High-altitude diffusion therefore does not have much effect on the evolution of the profiles in our nominal models. The elevated temperatures within the beacon region do expand the atmosphere in terms of the altitude scaling at the relevant pressures within the beacon, but the diffusion time scales at these pressures are longer than the total observational period after the storm, so changes due to diffusion are minor in our nominal model.

4.2. Nominal model results: ethylene

Because ethylene has one of the shortest chemical time scales of all the stable species (e.g., 40 days at the 2-mbar temperature maximum in the hot beacon-core model), it has one of the most pronounced responses to the elevated temperatures in the beacon (Fig. 5). From a column-integrated standpoint, the dominant reactions producing C$_2$H$_4$ in Saturn’s unperturbed (pre-storm) stratosphere are reaction R184 (CH + CH$_3$ → C$_2$H$_4$ + H), reaction R132 (H + C$_2$H$_4$ + M → C$_2$H$_3$ + M), and reaction R267 (C$_2$H$_4$ + H$_2$ → C$_2$H$_2$ + H), with lesser contributions from C$_2$H$_4$ photolysis and from R136 (H + C$_2$H$_3$ → C$_2$H$_4$ + H$_2$) (see also Moses et al., 2000a, 2005). The dominant loss processes are reaction R134 (H + C$_2$H$_4$ + M → C$_2$H$_2$ + M) and photolysis. The reaction C$_2$H$_4$ + H$_2$ → C$_2$H$_2$ + H (R267) contributes only 21% to the stratospheric column-integrated production rate of C$_2$H$_4$ in the pre-storm model. However, when the beacon temperature increase dramatically over pre-storm values, the highly temperature-sensitive reaction R267 overwhelmingly dominates the production of ethylene (see also Cavalié et al. (2015) and is correspondingly responsible for the major increase in the C$_2$H$_4$ abundance at ~mbar pressures in our nominal beacon model (see Fig. 5). Although the rate coefficient for R267 is relatively modest at room temperature and below (Callear and Smith, 1986; Tsang and Hampson, 1986; Weissman and Benson, 1988; Fahr et al., 1995; Mebel et al., 1995; Litwinowicz et al., 1995; Knazyev et al., 1996; Li et al., 2004; Laufer and Fahr, 2004; Tautermann et al., 2006; Agarwal et al., 2011), Saturn’s atmosphere contains enough background H$_2$ to make this reaction important.

The rate coefficient for this temperature-sensitive abstraction reaction R267 at low temperatures is poorly known, and extrapolations of the various published literature expressions to the lower temperatures relevant to Saturn can differ by many orders of magnitude (see Fig. 8). For our nominal model, we adopt one of the largest available published rate coefficients at low temperatures (i.e., the expression of Weissman and Benson, 1988), which results in a large predicted post-storm spike in the C$_2$H$_4$ abundance. Note, however, that the transition-state theory estimation method used by Weissman and Benson (1988) to derive the rate-coefficient expression of $k_{267} = 5.25 \times 10^{-15} T^{0.7}$ exp($-2574 k/T$) cm$^3$ s$^{-1}$ for this reaction is outdated in comparison to more modern techniques. Many recent theoretical
The sparse available experimental data do not help resolve the situation, with Fahr et al. (1995) deriving rate coefficients at room temperature that are significantly smaller than those favored by Tsang and Hampson (1986) (see dotted purple line in Fig. 8), while most other methods even smaller values at low temperature: \( k_{267} = 1.0738 \times 10^{-38} \exp(-3145.4 \, \text{K}/\text{T}) \, \text{cm}^3\,\text{s}^{-1} \) (see Fig. 8). Mebel et al. (1995) and Li et al. (2004) use various \textit{ab initio} theoretical techniques to predict the rate coefficients for this reaction; Li et al.’s calculations that use improved canonical variational transition-state theory with small-curvature tunneling corrections result in rate constants as large as those derived by Weissman and Benson (1988) (see dotted purple line in Fig. 8), while most other methods result in much slower rates at low temperature. Moreover, most of the theoretical expressions were developed for high-temperature combustion studies and were not designed to be extrapolated to the \( \sim 100–220 \, \text{K} \) temperatures relevant to Saturn’s stratosphere. The sparse available experimental data do not help resolve the situation, with Fahr et al. (1995) deriving rate coefficients at room temperature that are significantly smaller than those favored by Callear and Smith (1986) and Knyazev et al. (1996). The most recent study of the reaction energetics of \( \text{C}_2\text{H}_3 + \text{H}_2 \rightarrow \text{C}_2\text{H}_4 + \text{H} \) by Agarwal et al. (2011) provides further evidence that the rate coefficient for this abstraction reaction has yet to be well resolved.

The fact that the \( \text{C}_2\text{H}_4 \) abundance was observed to increase so significantly with increasing temperatures in the beacon region on Saturn (Hesman et al., 2012) makes it tempting to rule out the lowest estimates or determinations of the rate coefficient for this reaction (such as that of Tsang and Hampson, 1986 or Fahr et al., 1995), because such low rate coefficients prevent the \( \text{C}_2\text{H}_3 + \text{H}_2 \rightarrow \text{C}_2\text{H}_4 + \text{H} \) reaction from being significant for ethylene production on Saturn, even at the elevated 220-K beacon-core maximum temperature. The observed beacon behavior itself suggests that some reaction with a strong temperature dependence dominates ethylene production in the beacon, and the \( \text{C}_2\text{H}_3 + \text{H}_2 \rightarrow \text{C}_2\text{H}_4 + \text{H} \) reaction fills that role nicely. However, the modeled behavior of \( \text{C}_2\text{H}_4 \) depends on a complex coupling of many reactions, not all of which are well constrained from experimental or theoretical data, so it is possible that other yet-to-be-identified reactions are contributing to the observed ethylene increase, or that dynamical changes in the beacon are responsible for the observed increase.

In any case, Fig. 8 shows the sensitivity of the model results to the adopted expression for the rate coefficient for \( \text{C}_2\text{H}_3 + \text{H}_2 \rightarrow \text{C}_2\text{H}_4 + \text{H} \). A distinct peak in the \( \text{C}_2\text{H}_4 \) is formed at \( \sim \text{mbars} \) pressures for most of the cases, with the larger rate-coefficient assumptions for this reaction leading to larger predicted \( \text{C}_2\text{H}_4 \) abundances in the beacon. However, even with the adoption of the comparatively fast Weissman and Benson (1988) reaction rate coefficient, our “hot” nominal model underpredicts the emission in the \( \text{C}_2\text{H}_4 \) bands in the beacon core on May 4, 2011 by a factor of \( \sim 3.4 \) (see Fig. 7); that is, the photochemical model profile for \( \text{C}_2\text{H}_4 \) using the Weissman and Benson (1988) rate coefficient for reaction R267 would need to be multiplied uniformly by a factor of \( \sim 3.4 \) in order to reproduce the observed CIRS emission.

Fig. 8 also shows that our derived vertical profile for \( \text{C}_2\text{H}_4 \) at the beacon center differs from that of Hesman et al. (2012), despite the fact that both analyses use the NEMESIS retrieval program and both consider the same beacon-center CIRS data set from May 4, 2011. This difference is due to different analysis strategies with NEMESIS. The thermal structure and \( \text{C}_2\text{H}_4 \) vertical profile in the beacon region are not known \textit{a priori}. Hesman et al. (2012) proceeded by first determining the thermal structure from the CIRS data over a broad wavelength range, using the constrained linear inversion algorithm described in Conrath et al. (1998) and Achterberg et al. (2008). Then, Hesman et al. kept that temperature structure fixed and allowed NEMESIS to freely adjust the \( \text{C}_2\text{H}_4 \) vertical profile within a certain pressure range to provide a best fit to the \( \text{C}_2\text{H}_4 \) emission. We, on the other hand, simultaneously fit both the thermal structure and the scale factor for the hydrocarbon vertical profiles with NEMESIS, letting the temperature be adjusted freely but retaining the overall shape of the hydrocarbon vertical profiles from the photochemical models and requiring NEMESIS to scale these profiles uniformly to provide the best fit to the emission from all the hydrocarbons (see Section 3). Both procedures have their strengths and weaknesses. The photochemical model profiles provide a welcome connection to physical reality, but when the models do not adequately reproduce the observations—as is the case with these beacon models where vertical winds are not included—those constraints may not be meaningful. At the 2.5 cm\(^{-1}\) spectral resolution of these nadir observations, the CIRS data provide little concrete information about the vertical profile of \( \text{C}_2\text{H}_4 \). Instead, the retrievals provide \( \text{C}_2\text{H}_4 \) abundance information that is most reliable in the pressure region where the emission contribution function peaks, which is near the \( \sim 2 \, \text{mbar} \) region for the \( \text{C}_2\text{H}_4 \) emission bands observed here (see the black square with the pink outline in Fig. 8). Therefore, it is interesting to note that
the vertical profiles retrieved from both our technique and that of Hesman et al. (2012) converge on a similar C₂H₄ abundance in this ~2 mbar region. Although the two techniques lead to vastly different C₂H₄ abundances at pressures less than 1 mbar, those high-altitude regions have less influence on the C₂H₄ emission seen by CIRS than the deeper ~2 mbar region. Hereafter, we plot a single observational data point at the peak of the contribution function for C₂H₄ and the other hydrocarbons rather than the full retrieved vertical profile, but we also note that the location at which the contribution function peaks depends on the vertical profiles of both the temperature and the hydrocarbon in question, so that data point will be located at different pressures for different prior model profiles.

Although uncertainties in the C₆H₆ + H₂ → C₆H₆ + H production reaction cause the most dramatic changes in the C₆H₆ profile in the photochemical model, uncertainties in the dominant loss reaction R134, H + C₂H₄ + M → C₂H₅ + M, can also affect the predicted ethylene abundance. This reaction is important as an intermediate in the conversion of C₂H₂ to C₂H₅ on the giant planets (e.g., Allen et al., 1992; Romani, 1996; Moses et al., 2000a, 2005)—a slow rate will short-circuit this conversion, leading not only to more C₂H₅ but to increased abundances of C₂H₄ and most other hydrocarbon photochemical products, as a result of C₂H₅ being a key “parent” molecule for many species. As discussed in the review of Baulch et al. (2005), the high-pressure limiting rate coefficient for the H + C₂H₅ + M → C₂H₄ + M reaction has been measured in the 400–600-K temperature range (e.g., Lee et al., 1978; Sugawara et al., 1981; Lightfoot and Pilling, 1987; Michael et al., 2005) and is fairly well established. Experimental measurements at low pressures and in the intermediate fall-off pressure regime are also available, but only at room temperature and higher (e.g., Braun and Lenzi, 1967; Kurylo et al., 1970; Brouard et al., 1986; Lightfoot and Pilling, 1987; Hanning-Lee et al., 1993; Sillesen et al., 1993; Clarke et al., 2000). The low-pressure limiting rate coefficient at the low temperatures relevant to Saturn’s atmosphere is not well established, nor is the influence of tunneling on the high-pressure rate coefficient at low temperatures.

Theoretical calculations could potentially help bridge the gap, but such studies seldom extend to the low pressures and temperatures required for Saturn (e.g., Miller and Klippenstein, 2004; Michael et al., 2005). The recent ab initio transition-state theory based master-equation calculations presented by Vuitton et al. (2012) are an exception, as expressions are provided that are valid at low pressures in the 50–300-K range. The Vuitton et al. (2012) expressions, which result in very efficient C₂H₅ adduct formation under Saturn stratospheric conditions, are adopted in our nominal model. In contrast, Li et al. (2014) suggest that the underestimation of the C₂H₅ abundance in many photochemical models of the giant planets and Titan could be the result of an overestimation of the rate coefficient of R134 at low temperatures, and they suggest adopting an expression that leads to rate coefficients much smaller than our adopted ones under the relevant conditions. Such low rate coefficients would imply that tunneling is very inefficient for this reaction, in conflict with existing theoretical calculations (Miller and Klippenstein, 2004; Michael et al., 2005; Vuitton et al., 2012), but the use of this expression does provide a better fit to Titan observations (Li et al., 2014).

Fig. 9 shows how the models results are affected by variations in the rate coefficient for R134 (H + C₂H₅ + M → C₂H₄ + M). Our adoption of the relatively efficient Vuitton et al. (2012) expression for R134 results in a notably smaller C₂H₅ abundance at mbar levels than is predicted from using the Li et al. (2014) expression. Our nominal model profiles using the Vuitton et al. (2012) expression for R134 are consistent with the non-detection of C₂H₂ before the storm, but the model underpredicts the beacon-core C₂H₄ abundance after the storm. On the other hand, the Li et al. (2014) expression leads to a better fit to the post-storm C₂H₄ abundance, but results in too much C₂H₄ pre-storm, as well as excessive amounts of C₂H₂, C₃H₄, C₄H₂, C₄H₆, C₆H₆, and most other hydrocarbons in comparison with pre-storm observations. We therefore favor the Vuitton et al. (2012) expression for R134, which has a strong theoretical basis, and we seek other non-chemical explanations for the underestimation of the beacon C₂H₄ abundances from our nominal model (see Section 4.5).

More information about the rate coefficients for the reaction C₂H₂ + H₂ → C₂H₄ + H at low temperatures (50 ≤ T ≤ 250 K) and experimental confirmation of the theoretical reaction rate coefficients for H + C₂H₅ + M → C₂H₄ + M at low pressures and temperatures (P ≤ 1 mbar, T ≤ 200 K) are needed before we can feel confident about the quantitative predictions for the C₂H₄ abundance in photochemical models of the giant planets and Titan. However, the highly temperature-sensitive reaction C₂H₅ + H₂ → C₂H₄ + H is the most likely culprit of the enormous observed increase in the abundance of C₂H₄ detected by Hesman et al. (2012) in Saturn’s beacon region.

4.3. Nominal model results: methylacetylene and diacetylene

Both CH₂C₂H and C₂H₆ have relatively short chemical lifetimes at mbar regions in the beacon models—just 26 days for CH₂C₂H and 28 days for C₂H₆ at the 2-mbar temperature maximum in the beacon. Fig. 5 demonstrates that the stratospheric mole fraction of CH₂C₂H is expected to decrease with time in the beacon region. As discussed by Cavalié et al. (2015), the depletion is due in large part to the increased loss rate of CH₂C₂H due to the reaction H + CH₂C₂H + M → C₂H₆ + M, which has a moderately large energy barrier and is more effective at elevated temperatures. The results for C₂H₆ are more complicated and interesting. Fig. 10 shows the model results for the hot beacon-core model on May 4, 2011, in comparison with the unperturbed, pre-storm model profile. In the ~10–17 mbar pressure region, C₂H₆ becomes depleted due to the elevated temperatures in the beacon. As discussed by Cavalié et al. (2015), the decrease of C₂H₆ at these pressures in the beacon is due to the decrease in C₂H₄ photolysis and, more importantly, to the increase in the rate coefficient for the temperature-sensitive reaction R252, C₂H₂ + H₂ → C₂H₆ + H, which both result in a decreased concentration of C₂H₆ at these pressures. The reduction in C₂H reduces the effectiveness of the primary non-recycling C₂H₆ production mechanism, C₂H...
+ C\textsubscript{2}H\textsubscript{2} → C\textsubscript{4}H\textsubscript{2} + H, resulting in less C\textsubscript{4}H\textsubscript{2} in the beacon at these pressures. Although the chemical mechanism we are adopting underestimates the pre-storm C\textsubscript{4}H\textsubscript{2} abundance (see Fig. 10), the rate coefficients for the reactions C\textsubscript{2}H + H → C\textsubscript{4}H\textsubscript{2} + H and C\textsubscript{4}H\textsubscript{2} + H → C\textsubscript{4}H\textsubscript{2} + H have been well studied experimentally (see the review of Laufer and Fahr, 2004), and the prediction regarding the more rapid depletion of C\textsubscript{4}H\textsubscript{2} at increased temperatures is robust. Condensation is a major loss process for C\textsubscript{4}H\textsubscript{2} in Saturn’s lower stratosphere over much of the planet, and as the beacon temperatures increase, our models predict a large increase in the C\textsubscript{4}H\textsubscript{2} vapor abundance at pressures greater than ~10 mbar due to evaporation of solid C\textsubscript{4}H\textsubscript{2} aerosols. The very large magnitude of the post-storm spike in the C\textsubscript{4}H\textsubscript{2} abundance shown in Fig. 10 is partially an artifact of our model in that we only include condensation and evaporation and neglect other aerosol–microphysical processes like gravitational settling; in particular, we allow the condensates to diffuse through the atmosphere as a just another heavy gas, which enables more of the condensed phase to persist in the lower stratosphere than it would in the real atmosphere. However, some sort of large evaporation “spike” is expected in the beacon, even in the real atmosphere, because in situ production of C\textsubscript{4}H\textsubscript{2} occurs readily within the C\textsubscript{4}H\textsubscript{2} condensation region as a result of C\textsubscript{2}H\textsubscript{2} photolysis, followed by C\textsubscript{2}H + C\textsubscript{2}H\textsubscript{2} → C\textsubscript{4}H\textsubscript{2} + H, and because gravitational settling times for the aerosols are relatively long (e.g., Roman et al., 2013). The condensation of C\textsubscript{4}H\textsubscript{2} shuts off the local recycling back to C\textsubscript{2}H\textsubscript{2}, so acetylene photolysis continues to produce a steady, irreversible leak of carbon into condensed C\textsubscript{4}H\textsubscript{2}, which then can become a major aerosol component in Saturn’s stratosphere (see also Moses et al., 2000a,b). This evaporation spike in the beacon is too deep to be detectable by infrared instruments like CIRS, but it would be worthwhile to search for signatures of stratospheric aerosol thinning or clearing with error bars are from ISO (Moses et al., 2000a). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

4.4. Nominal model results: benzene and water

Benzene and water can also condense in Saturn’s lower stratosphere, but because there is less in situ production of these species within their condensation regions, the models do not predict that same kind of evaporation spike as was predicted for C\textsubscript{4}H\textsubscript{2}. Instead, the condensable vapor flows into the condensation region from higher altitudes. Evaporation of the aerosols then causes a smoother local increase in the vapor abundance (see Fig. 11) and an increase in the overall stratospheric column abundance of H\textsubscript{2}O and C\textsubscript{4}H\textsubscript{2}. Again, this increased vapor abundance in the beacon is at deep-enough pressures that it might be difficult to detect, but we note that Cavalié et al. (2012) reported a factor of 30–100 increase in the column abundance of H\textsubscript{2}O in the beacon with the PACS instrument on the Herschel Space Observatory. Our hot beacon-core model for May 4, 2011, predicts a similar factor of 30–100 increase in the water mole fraction in the ~2–3 mbar region compared to pre-storm values, but only a factor of 3 increase in the integrated water column abundance in the beacon compared to pre-storm levels. Direct comparison of synthetic model spectra...
with the Herschel/PACS observations would be needed before we can determine whether our predictions from the beacon models are consistent with the observations. In any case, we know of no chemical mechanisms that would increase the water abundance so severely in the beacon, so the Herschel/PACS observations most likely result from evaporation of water ice, with the water originating from outside the planet, and/or an increase due to vertical winds in the beacon (see Section 4.5). The benzene column abundance in our May 4, 2011, model increases by a factor of 4 in comparison with the pre-storm model as a result of evaporation.

4.5. The potential influence of transport in the beacon

As discussed in Section 4, the fact that our beacon models (and those of Cavalie et al., 2015) underpredict the abundances of all the C$_2$H$_i$ hydrocarbons in the hot beacon core on May 4, 2011 suggests that dynamics may play a large role in controlling the observed abundance increases within the beacon region. The beacon vortex is inherently a 3-D atmospheric structure that is difficult to capture accurately in a 1-D model, even if the 1-D model equations were appropriately solved. However, if horizontal advection of species is small in comparison to vertical advection, then some useful estimates can still be made, so we proceed with the examination of the 1-D behavior in the presence of vertical winds. For the situation with winds included, it is convenient to reformulate Eq. (1) in terms of mixing ratios rather than number densities. In the well mixed region of the atmosphere below the homopause, assuming hydrostatic equilibrium is maintained, and accounting for the continuity of the total atmospheric density $n_s$, Eq. (1) can be rewritten as

$$\frac{\partial q_i}{\partial t} + \frac{\partial q_i}{\partial z} = \frac{\partial}{\partial z} \left( n_s K_{zz} \frac{\partial q_i}{\partial z} \right) = \frac{Q_i}{n_s},$$

where $q_i$ is the mole fraction of species $i$, $w$ is the vertical wind velocity, $z$ is the altitude, $K_{zz}$ is the vertical eddy diffusion coefficient, and $Q_i$ is the net chemical source/sink term for species $i$.

If we further assume that chemistry and turbulent eddy transport have a minor influence in comparison with large-scale vertical winds, then Eq. (3) reduces to

$$\frac{\partial q_i}{\partial t} = -w \frac{\partial q_i}{\partial z}.$$  

Eq. (4) demonstrates that for subsidence (i.e., a downward wind, $w < 0$), the local mixing ratio will increase with time if the mixing-ratio gradient of the species is positive ($dq_i/dz > 0$), and it will decrease with time if the mixing-ratio gradient of the species is negative ($dq_i/dz < 0$). Both C$_2$H$_4$ and C$_2$H$_6$ have mixing-ratio gradients that are positive throughout the stratospheric beacon region, implying that subsidence is needed to increase the mixing ratios of these species locally within the beacon (see also the discussion of subsidence on the mixing ratios in the unperturbed atmosphere; Flasar et al., 2005; Fletcher et al., 2008, 2015; Fouchet et al., 2009; Guerlet et al., 2009, 2010; Sinclair et al., 2013, 2014). The C$_2$H$_4$ profile is more complicated, and chemical production and loss of C$_2$H$_4$ cannot be ignored, so we focus on C$_2$H$_2$ and C$_2$H$_6$ for the moment. The CH$_4$ mixing ratio actually decreases with height, especially in the upper stratosphere, and a downwelling wind would reduce the CH$_4$ mixing ratio locally in the beacon, which may in turn affect derived temperatures in the spectral analysis.

Subsidence within the beacon is also consistent with the increased beacon temperatures (i.e., due to adiabatic compression of the atmosphere resulting from the subsidence). Although energy deposition from atmospheric waves may have contributed to the stratospheric heating (Fletcher et al., 2011, 2012), especially initially or outside of the vortices themselves (i.e., to the extent of 10–20 K), we can make a first-order estimate of the magnitude of the subsiding winds in the final merged beacon by assuming that the observed temperature increase in the beacon is entirely due to adiabatic heating, and then by solving for the wind speeds needed to produce this temperature increase. In other words, we assume that the adiabatic heating (e.g., Holton, 1979, p. 52) is balanced by radiative relaxation:

$$w \left( \frac{dT}{dz} + \frac{g}{c_p} \right) = T - T_b,$$  

where $w$ is the vertical velocity, $T$ is the unperturbed atmospheric temperature, $g$ is the gravitational acceleration, $c_p$ is the specific heat at constant pressure, $T_b$ is the beacon temperature, and $T_{rad}$ is the radiative time constant, all of which are functions of altitude $z$. Given the observed $T_b$ and $T$, we can solve for $w$. The only difficult term to handle in Eq. (5) is $T_{rad}$. Conrath et al. (1990) determine that the globally and annually averaged $T_{rad}$ is approximately $3 \times 10^4$ s (almost 10 yrs) in the mbar region of Saturn. However, actual temperature-evolution observations and more recent models (Fletcher et al., 2007, 2010; Fouchet et al., 2008; Greathouse et al., 2008; Guerlet et al., 2009, 2010, 2014; Friedson and Moses, 2012; Sinclair et al., 2013, 2014) suggest that stratospheric cooling times may be shorter than this value, and $T_{rad}$ likely decreases more rapidly with height than is described in Fig. 2 of Conrath et al. (1990) because the mixing ratios of major coolants like C$_2$H$_2$ and C$_2$H$_6$ are increasing with height, whereas Conrath et al. (1990) assumed profiles that are constant with height. From the Friedson and Moses (2012) general circulation model (GCM), we note that a roughly $0.027$ cm s$^{-1}$ wind at 1 mbar at 25°N latitude (Fig. 9 of Friedson and Moses, 2012) results in a temperature increase from $\sim 137$ K to $\sim 142$ K (Fig. 5 of Friedson and Moses, 2012). Plugging this information back into Eq. (5) and solving for $T_{rad}$ gives us $\sim 3 \times 10^5$ s. However, $T_{rad}$ will strongly depend on the emission temperature: $T_{rad} = T_I/(dT/dt)$, and $dT/dt \propto T^4$, so $T_{rad} \propto 1/T^3$. Using the results from the Friedson and Moses (2012) GCM situation described above, we can estimate $T_{rad} = 3 \times 10^5(142/K/T_b)^3$ s. At the elevated temperatures of the beacon, $T_{rad}$ is considerably shorter than the nominal radiative time constant of the unperturbed atmosphere. For example, at the beacon maximum temperature of $\sim 220$ K, the radiative time constant is just 3 months.

When we solve Eq. (5) for $w$, using the CIRS-derived thermal structure and the temperature- and altitude-dependent $T_{rad}$, we derive the vertical wind profile shown in Fig. 12. The resulting winds exhibit upwelling in the troposphere (see also Fletcher et al., 2011) and strong, but narrowly focused, downwelling in the stratosphere, with a peak magnitude of roughly $1$ cm s$^{-1}$ centered at $\sim 2$ mbar. While a $1$ cm s$^{-1}$ vertical wind in the stratosphere is very strong by terrestrial standards, we will show that such a wind velocity is not sufficient to transport the necessary amount of C$_2$H$_2$ and C$_2$H$_6$ from higher altitudes ($p \leq 0.1$ mbar) to the mbar region, where these species are observed to be enhanced in the beacon on May 4, 2011. Moreover, the overall profile is not consistent with the steady 1-D continuity equation for the total atmospheric density, $d/dz(n_s w) = 0$, which would require the vertical wind to be proportional to $1/n_s$, such that the magnitude of the downwelling wind exponentially increases with height. The stratospheric region below the wind peak in Fig. 12 roughly exhibits this behavior, while the implied winds above the $\sim 2$ mbar peak do not, indicating that horizontal winds are important in conserving mass in the beacon at higher altitudes. Although the relatively short radiative time scale $T_{rad} \approx 3$ months at the maximum 2-mbar beacon temperature in May 2011 (five months after the storm onset, less than 1 month after the merger) could be contributing to the situation such that some of the excess energy from potentially higher temperatures at high altitudes could have radi-
are proportional to $1/a_b$ abundances in the beacon. The inclusion of vertical winds that the possible effects of strong downwelling winds on the species winds being present at lower altitudes.

could be converging toward the beacon center at high altitudes, do not extend indefinitely in altitude. Instead, horizontal winds followed by vertical descent through the mbar region, with diverging winds being present at lower altitudes.

In any case, we can use the above concepts to help us explore the possible effects of strong downwelling winds on the species abundances in the beacon. The inclusion of vertical winds that are proportional to $1/n_a$, which is required for the continuity of the total density in 1-D, causes major instability problems with KINETICS. We therefore solve Eq. (4) for chemically long-lived species outside of KINETICS, using finite-difference techniques with the Lax method (e.g., Press et al., 1986). Fig. 13 shows the solution for a wind profile that varies as $w(z) = w(0)n_a(0)/n_a(z)$, where $w(0) = -1.0 \text{ cm s}^{-1}$, $n_a(0) = 6.8 \times 10^{16} \text{ cm}^{-3}$ at $p(0) = 2 \text{ mbar}$. The wind is assumed to be constant with time and is applied for 25 days (for consistency with the KINETICS beacon model for the May 4, 2011 date), with boundary conditions of $dq_i/dz = 0$ (i.e., constant flux in this 1-D description, allowing the species to flow through the boundaries) and initial species profiles given by the pre-storm KINETICS results shown in Fig. 6. Although the results are sensitive to the boundary conditions, the assumed vertical extent of the atmosphere, and the time scale over which the winds are applied, Fig. 13 shows that a subsiding wind acts to redistribute the species from high altitudes to lower altitudes. Because the mixing-ratio profiles for C$_2$H$_2$ and C$_2$H$_6$ are positive in this region, the mixing ratios of C$_2$H$_2$ and C$_2$H$_6$ increase at mbar pressures within the beacon, while the CH$_4$ mixing ratio decreases. From exploring various wind profiles that are proportional to $1/n_a$, we find that the beacon-core observations from May 4, 2011 are best reproduced when winds are of order $-10 \text{ cm s}^{-1}$ in the 0.1–0.3 mbar region.

However, the wind profile inferred from the adiabatic heating (e.g., Fig. 12) implies that the downwelling wind speeds do not increase exponentially with height indefinitely within the upper stratosphere. There is a limit to the vertical extent of the beacon, and horizontal winds must dominate at some point in the upper stratosphere. There is a limit to the vertical extent of the beacon, while the CH$_4$ mixing ratio decreases. Downwelling winds of order $-10 \text{ cm s}^{-1}$ at $0.1\text{–}0.3 \text{ mbar}$ are again required to transport sufficient C$_2$H$_2$ and C$_2$H$_6$ to the $\sim 0.2 \text{ mbar}$ region to explain the elevated beacon-core abundances from May 4, 2011; that is, other wind profiles that fulfill this criterion provide similar results. The main advantage of the Gaussian wind profile is that it mitigates the severe instability problems that plague the inverse-density wind profile within the KINETICS code, so we can use KINETICS to study how the downwelling winds affect all species, including...
those with short chemical lifetimes. The main disadvantage is that the Gaussian profile violates the 1-D continuity equations for total density, and horizontal winds must be present to maintain mass conservation. The only way we can justify the use of the Gaussian vertical wind profile in the 1-D photochemical model is to demonstrate that the vertical advection term dominates over the horizontal advection term in the continuity equation.

To pursue this justification, we examine the density continuity equation assuming the beacon exhibits cylindrical symmetry, such that the continuity equation for the total atmospheric density becomes:

$$\frac{1}{r} \frac{\partial}{\partial r} (r \rho v) + \frac{\partial \rho}{\partial z} + w \frac{\partial n_a}{\partial z} = 0,$$

where $r$ is the radial distance, $v$ is the velocity in the radial direction, $w$ is the vertical velocity, $n_a$ is the total atmospheric density, and $z$ is the altitude. Right at the Gaussian wind maximum, $\partial v/\partial z = 0$, so $v \approx w_{\text{max}} r / 2 H_a$, where $H_a$ is the density scale height and $w_{\text{max}}$ is the maximum wind speed. If we go back to Eq. (3) in the cylindrical coordinate system now and assume eddy diffusion and chemistry have a negligible influence, then Eq. (3) becomes

$$\frac{\partial q_i}{\partial t} + v_0 \frac{\partial q_i}{\partial r} + w_{\text{max}} \frac{\partial q_i}{\partial z} = 0.$$

We can compare the magnitude of the radial horizontal advection term [$v_0 \partial q_i / \partial r$] with the vertical advection term [$w_{\text{max}} \partial q_i / \partial z$]. Using the longitudinally resolved species scaling factors to determine $\partial q_i / \partial r$ near the beacon center, considering a distance $r$ as the average of the nearest longitude points from the beacon center for which we have retrievals (e.g., see Fig. 7), and considering the pressure at which the Gaussian wind has its maximum, we find that the vertical advection term is indeed much stronger than the horizontal advection term at the beacon center:

$$v_0 \frac{\partial q_i}{\partial r} \ll \left( \frac{w_{\text{max}} r}{2 H_a} \right) \frac{\partial q_i}{\partial z},$$

for both C$_2$H$_2$ and C$_2$H$_6$ near the beacon center, so the vertical advection dominates. In fact, using Eq. (7) to define $v_0$ at any altitude near the beacon center, we can show that $w_{\text{max}} \partial q_i / \partial z$ dominates over horizontal advection up to 0.01 mbar and beyond for our assumed Gaussian profile, so we are justified in considering this vertical wind profile in our 1-D calculations.

We therefore proceed with considering Gaussian wind profiles within our 1-D photochemical model. We continue to solve the continuity equations from Eq. (1) with KINETICS, but now include the vertical wind term in the flux equation (Eq. (2)). The vertical winds are introduced only at the point when the model switches to the thermal structure for the May 4, 2011 date, which happens on April 9, 2011, roughly coincident with the beacon merger. When we include the weaker, roughly Gaussian, wind profile calculated from the adiabatic heating (see Fig. 12 and Eq. (5)), the resulting C$_2$H$_2$ and C$_2$H$_6$ mole fractions are only slightly increased in the mbar region, suggesting, as expected, that winds that are about an order of magnitude stronger are needed to explain the chemical abundances in the beacon.

Fig. 15 shows the results for the C$_2$H$_2$ species when we include the stronger Gaussian winds considered in the finite-difference calculations described earlier; i.e., a Gaussian downward wind in log($P$) space with a peak magnitude of $-10$ cm s$^{-1}$ centered at log$_{10}(P)$ mbar $= -0.5$, with a standard deviation of log$_{10}(P)$ mbar $= 0.8$. The eddy diffusion coefficient is not changed from the nominal model, but the transport time scale due to these stronger winds is much shorter than the diffusion time scale in the mbar region, so the downwelling winds very effectively transport the C$_2$H$_2$ from high altitudes to lower altitudes, increasing the local mole fractions in the mbar region. Note that because the unperturbed C$_2$H$_2$ mixing-ratio gradient in the mbar region is greater than that for C$_2$H$_6$ (see Fig. 15), the local C$_2$H$_6$ abundance increases more significantly than that of C$_2$H$_2$ when the subsiding winds are included (recall Eq. (4)). In contrast, the main effect of the winds on C$_2$H$_4$ is to "push" the local mixing-ratio peak downward, making it deeper but narrower. Note that because CH$_4$ has a negative mixing-ratio gradient, the subsiding winds actually deplete the local CH$_4$ mixing ratio in the mbar region, which in turn affects the retrieved temperatures. The results shown in Fig. 15 have been through one additional iteration in which we re-retrieved the thermal structure in the hot-beacon core using the initial wind-model species profiles as priors, and then we adopted the resulting retrieved thermal structure for a final beacon model with the winds imposed. Additional tests indicated that further iterations were unnecessary because the retrieved temperatures and hydrocarbon abundances converged on the same solution. The new retrieved “wind-derived” thermal structure for the hot beacon core is cooler by $\sim 7$ K at 1 mbar and warmer by $\sim 7$ K at 5 mbar; in essence, the winds shown in Fig. 15 caused the peak temperatures to migrate downward in altitude.

The multiplicative factors by which the wind-model species vertical profiles need to be scaled in order to reproduce the emission in the hot beacon core on May 4, 2011 are shown in Fig. 16.
comparison with our nominal hot-core model without winds, the wind model does a much better job of reproducing the amount of C$_2$H$_2$, C$_2$H$_4$, and C$_2$H$_6$ that is needed to explain the observed CIRS emission—scaling factors at the beacon core at 294.8° System III longitude are all 1.0 ± 0.3 now for these species (cf. Figs. 7 and 16). The resulting beacon-center volume mixing ratios from the CIRS spectral retrievals using the wind model profiles as priors are (9.17 ± 0.11) × 10^{-7} for C$_2$H$_2$, (1.86 ± 0.23) × 10^{-8} for C$_2$H$_4$, and (8.19 ± 0.08) × 10^{-6} for C$_2$H$_6$ at 2.4 mbar. The wind model actually predicts slightly too much C$_2$H$_6$ at the beacon core, and slightly too much C$_2$H$_2$, but not quite enough C$_2$H$_4$.

Although not shown in Fig. 15, the middle-stratospheric mixing-ratio peaks for CH$_3$C$_2$H and C$_3$H$_8$ are simply pushed downward in the wind model, with the overall column abundance above 10 mbar increasing only slightly. For C$_2$H$_2$, the middle-stratospheric peak also migrates downward, but given that the mixing-ratio gradient in the mbar region is large for C$_2$H$_2$ in the unperturbed model, the downwelling winds cause a comparatively large factor of 4.5 increase in the C$_2$H$_2$ column abundance above 10 mbar in comparison with the nominal hot-beacon model without winds (or an overall increase of a factor of 2.2 in comparison with the cooler pre-storm column abundance). The subsiding winds also have an interesting effect on the oxygen species, whose source is external material deposited at high altitudes. In comparison with the pre-storm column abundances, the post-storm column abundance above 30 mbar has increased by a factor of 3.2 for CO$_2$ and a factor of 8.5 for H$_2$O (which includes the contribution from evaporation).

Judging from Fig. 16, however, it appears that the strong downwelling winds are not uniform across the beacon vortex, as the scale factors have apparent structure as a function of longitude (see also Fig. 7). Our wind model tends to notably overestimate the species' abundances at longitudes away from the beacon center, except for an additional strong enhancement in C$_2$H$_6$ and C$_2$H$_2$ at 273° System III longitude. This feature at 273° longitude appears to be associated with a higher-altitude temperature increase (see Fig. 16a), suggesting that the strongest vertical winds were located both at 273° and at the core 295° longitude at the time of the observations, but that the winds at 273° longitude may have been confined to higher altitudes than at the beacon core. Interestingly, the C$_2$H$_4$ longitudinal cross section across the beacon does not show this same 273° feature, perhaps because the C$_2$H$_4$ abundance is not predicted to increase much due to subsiding winds in the ~0.1 mbar region, whereas C$_2$H$_2$ and C$_2$H$_6$ are (see Fig. 15). The narrower C$_2$H$_4$ central enhancement as a function of longitude better tracks the temperature structure across the beacon at ~1–5 mbar.

In fact, further analysis of the retrievals of temperature profiles and species abundances a function of latitude and longitude across the beacon could potentially illuminate the details of the 3-D dynamics within the beacon vortex, which may in turn reveal the complex dynamical coupling that was responsible for the stratospheric response in the first place. The prevailing theory for the stratospheric beacon formation is that the tropospheric convective plumes in the stratosphere served as a source of upward-propagating planetary waves and/or gravity waves that deposited energy and momentum in the stratosphere (Fletcher et al., 2012); upwelling and divergence of air on a rotating planet will naturally cause an anticyclonic vorticity, although these waves may also have interacted with the mean flow in the stratosphere to form the observed anticyclonic beacon vortices. Because we expect C$_2$H$_6$ (and to a lesser extent C$_2$H$_2$) to be chemically stable in the beacon region, the C$_2$H$_2$ abundance remains high, which is consistent with the strong vertical winds over the core of the beacon vortex. However, we expect C$_2$H$_4$ to be chemically unstable in the beacon region, which is consistent with the strong vertical winds over the core of the beacon vortex.
even at elevated temperatures, C$_2$H$_6$ and potentially C$_2$H$_2$ could serve as tracers of atmospheric vertical motions within the beacon(s), whereas C$_2$H$_4$ is also strongly sensitive to temperatures. Assuming chemical stability, the whole system could be potentially modeled with a 3-D mesoscale circulation model, which could provide insights into the wind fields before and after the storm event. Very little is currently known about stratospheric dynamics on Saturn, and the response to the storm could be a fertile source of information for the burgeoning stratospheric GCMs that are now being developed (e.g., Friedson and Moses, 2012; Guerlet et al., 2014).

An astute reader may notice that the original tendency of the model to overestimate the pre-storm ethane mole fraction and to underestimate the pre-storm acetylene mole fraction in comparison with the Cassini CIRS pre-storm data (see Figs. 4 and 6) is also still present (and even magnified) in the post-storm models that include vertical winds, in comparison with the post-storm CIRS data (see Figs. 13–16). Uncertainties in the chemical mechanism are likely at fault for some part of this model-data discrepancy, because the overall C$_2$H$_6$/C$_2$H$_4$ ratio is controlled in large part by chemistry (Moses et al., 2005). The effect of seasonal variations in solar insolation and/or dynamics due to large-scale stratospheric circulation may also play a role (e.g., Moses and Greathouse, 2005; Hue et al., 2015; Friedson and Moses, 2012)—neither of these effects were considered in the model presented here. The fact that the Cassini CIRS limb observations of Guerlet et al. (2009, 2010) indicate that our pre-storm model underestimates the C$_2$H$_6$ abundance at high altitudes suggests that we may be underestimating the vertical winds needed to carry sufficient amounts of C$_2$H$_6$ down to the ~mbar region to explain the May 2011 beacon observations. However, given the likely complicated 3-D nature of the problem and uncertainties in the time scales over which the winds are operating, we do not further pursue 1-D models to attempt to narrow down the necessary vertical wind magnitudes for an assumed initial C$_2$H$_6$ vertical profile that follows the Guerlet et al. (2009, 2010) retrieved profile.

5. Comparison with CIRS spectra

In Fig. 17, we demonstrate how synthetic spectra generated from the photochemical model results compare with the CIRS beacon observations from May 4, 2011. The green curves show the results from the pre-storm photochemical model. Although this model compares well with CIRS spectra acquired 4–5 months before the 2010 storm event, the predicted molecular emission features clearly fall far short of the observed intensities in the beacon—illustrating why the high-temperature air masses were nicknamed “beacons” in the first place. The purple curves show the synthetic emission assuming that the molecular abundances remain at these pre-storm model values, but assuming that the temperature profile follows that from the CIRS retrievals from the May 4, 2011 beacon core. From a comparison of the purple-curve emission intensity with that of the observations, one can see that the increased temperatures alone cannot explain the

![Fig. 17. CIRS spectra (dots with error bars) from May 4, 2011, averaged over ±5° longitude surrounding the beacon core at 294.8°W, and between 36 and 43°N latitude, compared to a series of synthetic spectra. The red line shows emission based on our nominal hot beacon-core model with no winds, and with no scaling of hydrocarbon profiles (see Fig. 6). The black line is our best-fitting model for the beacon core (in this case from scaling the hydrocarbon profiles with the beacon-center scaling factors shown in Fig. 16, associated with the predictions of the photochemical model that considers the Gaussian downwelling wind profile from Fig. 15); the blue line shows emission based on this same wind model, but with no scaling of hydrocarbon profiles. The green line is the best fitting pre-storm model for latitude 40°N. The purple line shows emission based on this same pre-storm model, but with temperatures matching those retrieved from the beacon, to demonstrate that (i) temperature variations alone cannot provide a good fit the hydrocarbon emission features and that (ii) ethylene experiences the largest differences in emission based on temperature-dependent chemistry alone (i.e., difference between the red and purple lines). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)](image-url)
observed emission intensities in the $\text{C}_2\text{H}_x$ molecular bands. The beacon is clearly characterized by both increased temperatures and increased $\text{C}_2\text{H}_x$ abundances. The red curve shows the predicted emission from our hot-beacon core model with no vertical winds (e.g., see Fig. 6). The temperature-dependent chemistry in the beacon region has led to an increase in the $\text{C}_2\text{H}_4$ abundance in this model, but the emission in the $\text{C}_2\text{H}_4$ band near 950 cm$^{-1}$ is still clearly underpredicted. Similarly, because the photochemical model predicts little change in the $\text{C}_2\text{H}_2$ and $\text{C}_2\text{H}_6$ abundances due to high-temperature chemistry in the beacon, there is little difference between the red and purple synthetic emission curves for these species.

The blue curves in Fig. 17 represent the synthetic spectra predicted from our hot-beacon core model with the Gaussian-profile downwelling winds described in Fig. 15. This wind-aided model clearly provides a much better fit to the data, although the model slightly underpredicts the $\text{C}_2\text{H}_2$ emission and slightly overpredicts the $\text{C}_2\text{H}_4$ and $\text{C}_2\text{H}_6$ emission. The best fit (black curves) occurs when we allow NEMESIS to scale the hydrocarbon profiles by the observed emission intensities in the $\text{C}_2\text{H}_x$ welling winds are needed to reproduce the $\text{C}_2\text{H}_2$ and $\text{C}_2\text{H}_6$ emission. The best fit (black curves) occurs when we allow NEMESIS to scale the hydrocarbon profiles by the scale factors at the beacon-center longitude, shown in Fig. 16. Note that our favored wind profile is by no means unique; other wind profiles that consider winds of roughly $-10$ cm s$^{-1}$ in the $0.1-0.3$ mbar region produce similar results. However, from a comparison of the red and blue curves with the observational data it is clear here that both high-temperature chemistry and strong downwelling winds are needed to reproduce the $\text{C}_2\text{H}_x$ emission features observed in the beacon in May 2011.

6. Conclusions

Although Saturn’s gigantic northern-hemisphere storm of 2010–2011 generated obvious changes in tropospheric cloud structure and dynamics in the weeks and months following the outburst (Sánchez-Lavega et al., 2011, 2012; Sayanagi et al., 2013), the more unexpected consequence has been a radical and long-lived change in the dynamics, energetics, and chemistry of the stratosphere. Detailed analyses of these stratospheric changes can shed new light on the complex coupling of physical and chemical processes throughout the atmosphere.

We have used a photochemical model to track the expected evolution of the stratospheric hydrocarbon and oxygen species in the anticyclonic vortex “beacons” that formed in Saturn’s northern-hemisphere stratosphere after the eruption of this massive storm system. We start from a fully converged 1-D stratospheric photochemical model for the appropriate northern mid-latitude region, and then allow the temperature and density structure in the model to change with time as described by the Fletcher et al. (2012) Cassini/CIRS observational retrievals of the thermal structure within the initial beacon “B1” and the final single beacon “B0” after the two initial beacons merged sometime in April 2011. From our photochemical models that consider the increased temperatures in the beacon but no corresponding changes in the dynamics of the region, we obtain the following results:

- The beacon models predict a large factor of 7 increase in the $\text{C}_2\text{H}_4$ mole fraction in a localized mid-stratospheric peak centered at $-0.4$ mbar within 5 months of the storm onset, resulting solely from the increased temperatures within the beacon regions (see Fig. 6).
- The large predicted increase in $\text{C}_2\text{H}_4$ in our models is caused by the strong temperature dependence of the reaction $\text{C}_2\text{H}_3 + \text{H}_2 \rightarrow \text{C}_2\text{H}_4 + \text{H}$. Although laboratory measurements of the rate coefficient for this reaction are challenging at room temperature and below, more definitive information at low temperatures is needed before we can make quantitative predictions regarding the expected $\text{C}_2\text{H}_4$ abundance in the beacon, and before we can fully understand $\text{C}_2\text{H}_4$ chemistry both within Saturn’s beacon regions and at ambient conditions on all the giant planets.
- In contrast to $\text{C}_2\text{H}_4$, our photochemical models predict little or no change in the stratospheric mixing ratios of the longer-lived hydrocarbons $\text{C}_2\text{H}_2$, $\text{C}_2\text{H}_4$, and $\text{C}_2\text{H}_6$ due to the beacon-temperature increase alone, whereas the mixing ratios of the less chemically stable species $\text{CH}_3\text{C}_2\text{H}$ and $\text{C}_2\text{H}_2$ decrease with time in the $10^{-10}$–$2$ mbar region as a result of the elevated temperatures (see also Cavalié et al., 2015).
- Constituents that condense in Saturn’s lower stratosphere, such as $\text{C}_2\text{H}_4$, $\text{C}_2\text{H}_6$, and $\text{H}_2\text{O}$ (the latter from external sources) exhibit strong increases in abundance in the beacons at pressures greater than a few mbar due to the evaporation of icy aerosols.
- The evaporation of $\text{C}_2\text{H}_2$-bearing ices is a particularly interesting result and could lead to a large local “spike” in the gas-phase $\text{C}_2\text{H}_2$ abundance at depth because the $\text{in situ}$ production of $\text{C}_2\text{H}_2$ has made condensed $\text{C}_2\text{H}_2$ a major local sink of carbon in the pre-storm model, and conversion back to $\text{C}_2\text{H}_2$ and other hydrocarbons once the $\text{C}_2\text{H}_2$ evaporates is not instantaneous. The predicted clearing of the lower-stratospheric hazes (which are optically thin under normal undisturbed conditions) may be observable when the beacon features are near the planetary limb.
- The increased temperatures alone in the beacon cannot explain the $\text{C}_2\text{H}_2$ band emission intensities observed by Cassini/CIRS in May 2011, just after the beacon merger. Our beacon model with the temperature increase alone (and no winds) underestimates the $\text{C}_2\text{H}_4$ abundance in the hot central core of the beacon on May 4, 2011 by a factor of $\sim 3.5$, and underestimates the abundance of $\text{C}_2\text{H}_2$ and $\text{C}_2\text{H}_6$ by factors of $\sim 7.5$ and $\sim 2$, respectively (see Fig. 7).
- If the inferred beacon increases in the abundance of $\text{C}_2\text{H}_2$, $\text{C}_2\text{H}_4$, and $\text{C}_2\text{H}_6$ (see also Hesman et al., 2012, 2014) were due to chemistry alone, the carbon would have had to have originated in $\text{CH}_4$ as methane is the only local source of carbon large enough to explain the observed increases (e.g., Figs. 6 and 7). We could identify no temperature-dependent loss reaction for $\text{CH}_4$ in the middle stratosphere that is effective enough on the time scales involved to produce the observed increase in $\text{C}_2\text{H}_4$, hydrocarbon abundances. We therefore conclude that vertical winds are contributing to the observed hydrocarbon increases in Saturn’s beacon regions.

Our results considering the effects of temperature changes alone are qualitatively consistent with those of Cavalié et al. (2015), except that we predict a larger increase in the $\text{C}_2\text{H}_4$ abundance (our factor of 7 versus their factor of $\sim 3$), and Cavalié et al. do not mention effects due to evaporation of lower-stratospheric aerosols.

In Section 4.5, we demonstrate that the observed increase in $\text{C}_2\text{H}_2$, $\text{C}_2\text{H}_4$, and $\text{C}_2\text{H}_6$ in the beacon is best explained both by altered chemistry due to the increased beacon temperatures and by strong descent of air in the middle stratosphere within the beacon. We also discuss the magnitude of the subsiding winds that are implied by the observations. The main results from our photochemical models that include vertical winds in the merged beacon are the following:

- Downwelling winds of order $-10$ cm s$^{-1}$ in the $0.1-0.3$ mbar region are needed to carry the necessary amounts of $\text{C}_2\text{H}_2$ and $\text{C}_2\text{H}_4$ from higher altitudes, where the primary chemical production regions reside and where the $\text{C}_2\text{H}_2$ and $\text{C}_2\text{H}_6$ mixing ratios are larger, to the $\sim 1-5$ mbar pressure region, where the $\text{C}_2\text{H}_4$ mixing ratios were observed to increase.
• When we include downwelling winds of the appropriate magnitude in our 1-D photochemical model, the resulting C$_2$H$_2$, C$_2$H$_4$, and C$_2$H$_6$ profiles are within 30% of the abundances needed to reproduce the observed CIRS emission within the core of the merged beacon on May 4, 2011 (see Figs. 15 and 16).

• The resulting volume mixing ratios derived from CIRS spectral retrievals from the observations of the beacon center on May 4, 2011 are $(9.17 \pm 0.11) \times 10^{-7}$ for C$_2$H$_2$, $(1.86 \pm 0.23) \times 10^{-8}$ for C$_2$H$_4$, and $(8.19 \pm 0.08) \times 10^{-6}$ for C$_2$H$_6$ at 2.4 mbar.

• The corresponding inferred abundance increases above pre-storm values are a factor of 7.6 for C$_2$H$_2$ and 2.2 for C$_2$H$_4$ based on CIRS observations alone, and a factor of ~25 for C$_2$H$_4$ based on the pre-storm photochemical model in comparison with CIRS beacon observations (i.e., C$_2$H$_4$ was not detected in the CIRS observations before the storm).

• Because the unperturbed CH$_4$ mixing-ratio gradient decreases with height due to molecular diffusion in the upper stratosphere, the downwelling winds cause a depletion of CH$_4$ within the beacon model (see Fig. 15). This tendency complicates retrievals of the thermal structure within the beacon, given that using the observed emission within the v$_4$ band of methane is a typical way of deriving the stratospheric temperatures—one can no longer assume that the CH$_4$ mixing-ratio profile is well known within the beacon, and model-data iterations are needed to ensure a consistent solution in terms of both the temperature and hydrocarbon profiles.

• The subsiding wind can also affect oxygen species like CO$_3$ and H$_2$O, whose source is presumed to be external to the planet (Feuchtgruber et al., 1997, 1999; Moses et al., 2000b; Bergin et al., 2000; Hartogh et al., 2011; Fleshman et al., 2012). Our model that includes downwelling winds predicts a factor 3.2 and 8.5 increase, respectively, in the column abundance of CO$_3$ and H$_2$O above 30 mbar in comparison with the pre-storm model abundances. These predictions are testable with further analysis of CIRS observations of hydrocarbons and CO$_2$ within the beacon region (e.g., Hesman et al., 2014) and of H$_2$O in the beacon from longer wavelength Herschel and Cassini observations (e.g., Cavalié et al., 2012; Bjoraker et al., 2014).

• Our photochemical model with winds included also predicts a factor of 2.2 increase in the column abundance of C$_2$H$_6$ above 10 mbar, but only a minor increase in the column abundances of CH$_4$ and C$_2$H$_2$.

• As with the photochemical model without winds, our wind model predicts that evaporation of icy C$_2$H$_6$, H$_2$O, and C$_6$H$_6$ aerosols in the lower stratosphere at the elevated beacon temperatures should cause a localized clearing or thinning of the stratospheric haze. The predicted increased gas-phase abundances in the lower stratosphere due to evaporation of these aerosols are likely too deep to affect the CIRS emission spectra, but the clearing of the stratospheric aerosol layer may affect the scattering behavior in Cassini visible and ultraviolet images (and at near-infrared wavelengths in regions where methane strongly absorbs), particularly at high phase angles, and the increased C$_2$H$_6$ abundance at depth may affect the ultraviolet spectra within the beacon region, in comparison with regions outside the beacon. These potential effects are worth further investigation.

Our model results were compared only with the Cassini CIRS observations. Our resulting inferred vertical profile for C$_2$H$_4$ from our wind-aided model differs from that derived by Hesman et al. (2012) for the May 2011 beacon from observations both from Cassini/CIRS (see Fig. 8) and from ground-based observations with the Celeste instrument. However, both analyses predict a similar C$_2$H$_4$ mole fraction at the ~2 mbar region, where the contribution function for the C$_2$H$_4$ emission band has a maximum. The C$_2$H$_4$ emission from the 2.5 cm$^{-1}$ resolution CIRS data are less sensitive to the higher-altitude ~0.5 mbar region where Hesman et al. (2012) derive a large localized maximum in the C$_2$H$_4$ mixing-ratio. Our solution does not require such a large peak; however, it remains to be seen whether our preferred C$_2$H$_4$ profile is consistent with the higher-spectral-resolution Celeste observations of Hesman et al. (2012). Future investigations comparing our model predictions with the Celeste data and with other ground-based high-spectral-resolution observations, such as those obtained with the TEXES instrument (e.g., Fouchet et al., 2013), would provide useful tests of the models. Note that the very large peak abundance (volume mixing ratio of nearly $10^{-9}$) at ~0.1 mbar from the Hesman et al. (2012) Celeste retrieval (see their Fig. 5) would be particularly difficult to explain photochemically, as C$_2$H$_4$ photochemical production in that pressure region is not very strong, and subsiding winds cannot explain such an abundance because the high-altitude C$_2$H$_4$ mixing ratios never reach such large values.

Our model provides a solution consistent with known physical and chemical principles, within the limitations of model parameter uncertainties. Our conclusion that the air within the beacon is subsiding is reasonable for anticyclonic vortices, but the magnitude of the necessary downwelling is larger than is expected based on the observed increased temperatures if adiabatic compression were responsible for the heating (see Section 4.5 and Eq. (7)). A $10^{-10}$ cm$^{-1}$ downwelling wind is also much larger than typical stratospheric subsiding winds on Earth or those predicted for the giant planets due to residual (diabatic) circulation (e.g., Andrews et al., 1987; Conrath et al., 1990; Friedson and Moses, 2012). However, some apparently similar extreme downwellings do occur in the middle atmospheres of terrestrial planets. For example, from temperature measurements from solar and stellar occultations with the SPIRAC/ESO spectrometers on Venus Express, Bertaux et al. (2007) inferred a downwelling wind of $43$ cm$^{-1}$ on the night side of Venus at 90–100 km as a result of the subsolar-tosub-solar flow in the Venus upper atmosphere. Another example is the so-called “stratospheric sudden warming” (SSW) events on Earth (e.g., Matsuno, 1971; Andrews et al., 1987), where descent rates as large as $1$ to $5$ cm$^{-1}$ have been inferred in the middle atmosphere (e.g., Holt et al., 2013; Bailey et al., 2014). These SSW events are marked by very rapid temperature increases in the high-latitude winter hemisphere caused by enhanced downwelling. The downwelling in turn is triggered by the dissipation of large-amplitude, planetary-scale waves that are generated in the troposphere and propagate upwards to the stratosphere, where they interact with the mean zonal flow, causing deceleration and even reversal of the polar-night jet that surrounds the polar vortex. Although the Saturn beacons are not high-latitude features, they too may have been caused by upward-propagating storm-generated waves interacting with the mean stratospheric circulation (e.g., Fletcher et al., 2012).

Future work should include 3-D dynamical modeling of the beacon systems. The Saturn storm beacons are inherently 3-D dynamical phenomena that would be best studied with mesoscale circulation models, particularly in terms of investigating how the anticyclonic stratospheric vortices formed and evolved in the months following the 2010 tropospheric convective outburst. Given the chemical stability of C$_2$H$_6$, and to a lesser extent C$_2$H$_2$, these species could act as useful tracers to diagnose winds within the beacon—their chemistry over the lifetime of the beacon can be ignored to first order.

The stratospheric beacons were an intriguing and unexpected consequence of the gigantic 2010–2011 convective outburst on Saturn. Studying the underlying factors controlling the evolution of the beacon temperatures and chemical-constituent abundances
can further our knowledge of dynamical coupling between the troposphere and stratosphere, the mean wind fields within the unperturbed and perturbed stratosphere, the energetics and long-term energy balance of the atmosphere, and the dominant chemical processes both within the unperturbed and beacon stratospheric environments.

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