COMPOSITIONAL DIVERSITY IN THE ATMOSPHERES OF HOT NEPTUNES, WITH APPLICATION TO GJ 436b

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ABSTRACT

Neptune-sized extrasolar planets that orbit relatively close to their host stars — often called “hot Neptunes” — are common within the known population of exoplanets and planetary candidates. Similar to our own Uranus and Neptune, inefficient accretion of nebular gas is expected produce hot Neptunes whose masses are dominated by elements heavier than hydrogen and helium. At high atmospheric metallicities of 10–10,000× solar, hot Neptunes will exhibit an interesting continuum of atmospheric compositions, ranging from more Neptune-like, H$_2$-dominated atmospheres to more Venus-like, CO$_2$-dominated atmospheres. We explore the predicted equilibrium and disequilibrium chemistry of generic hot Neptunes and find that the atmospheric composition varies strongly as a function of temperature and bulk atmospheric properties such as metallicity and the C/O ratio. Relatively exotic H$_2$O, CO, CO$_2$, and even O$_2$-dominated atmospheres are possible for hot Neptunes. We apply our models to the case of GJ 436b, where we find that a CO-rich, CH$_4$-poor atmosphere can be a natural consequence of a very high atmospheric metallicity. From comparisons of our results with Spitzer eclipse data for GJ 436b, we conclude that although the spectral fit from the high-metallicity forward models is not quite as good as the fit obtained from pure retrieval methods, the atmospheric composition predicted by these forward models is more physically and chemically plausible. High-metallicity atmospheres (orders of magnitude in excess of solar) should therefore be considered as a possibility for GJ 436b and other hot Neptunes.

Subject headings: planetary systems — planets and satellites: atmospheres — planets and satellites: composition — planets and satellites: individual (GJ 436b) — stars: individual (GJ 436)

1. INTRODUCTION

Observations from the Kepler spacecraft reveal that planets with radii between that of the Earth and Neptune constitute a dominant fraction of the known transiting exoplanet population [Borucki et al. 2011; Howard et al. 2012; Batalha et al. 2013; Fressin et al. 2013]. Most of these planets and planetary candidates orbit relatively close to their host stars and are therefore quite hot by Solar-System standards, with equilibrium temperatures $T_{eq} \gtrsim 400$ K. These so-called “hot Neptunes” and hot “Super Earths” represent terra incognita for planetary researchers. The atmospheric compositions of these intriguing worlds could be widely diverse, ranging from the more familiar hydrogen-dominated ice-giant atmospheres, like our own Uranus and Neptune, to relatively hydrogen-poor CO$_2$-, H$_2$O-, or N$_2$-dominated atmospheres...

We investigate the possible atmospheric diversity of hot Neptunes, i.e., close-in transiting exoplanets whose radii \( R_p \) are typically considered to fall in the 2\( R_J < R_p < 6 R_\oplus \) range (see Borucki et al. 2011; Howard et al. 2012), and whose atmospheres contain some H/He component. Our focus is on how atmospheric properties like temperature, metallicity, and bulk elemental ratios can affect the predicted equilibrium and disequilibrium composition of the atmospheres of generic hot-Neptune exoplanets, as well as specific hot Neptunes such as GJ 436b, for which eclipse observations suggest an unexpected and puzzling atmospheric composition (Stevenson et al. 2010, Madhusudhan & Seager 2011).

The discovery of GJ 436b by the radial-velocity technique (Butler et al. 2004; see also Maness et al. 2007), followed by its identification as a transiting planet (Gillon et al. 2007b), confirmed this intriguing object as the first Neptune-sized exoplanet ever detected. GJ 436b’s mass of 1.4\( M_{\text{Neptune}} \) (0.078\( M_\text{Jup} \), 25\( M_\oplus \)), radius of 1.1\( R_{\text{Neptune}} \) (0.37\( R_\text{Jup} \), 4.1\( R_\oplus \)), and density of 1.2\( \rho_{\text{Neptune}} \) (2.0 g cm\(^{-3} \)), according to von Braun et al. (2012), are all slightly larger than the corresponding values for Neptune. However, with an orbital semimajor axis of only 0.03 AU (Torres et al. 2008; Southworth 2010; von Braun et al. 2012), GJ 436b’s dayside atmosphere maintains an effective temperature of ~700–900 K (Deming et al. 2004; Demory et al. 2007; Stevenson et al. 2010; Madhusudhan & Seager 2011; Beaulieu et al. 2011; Knutson et al. 2011) as a result of the strong irradiation from its nearby M-dwarf host star, and thus the planet earns its “hot” designation. The planet’s relatively large radius in relation to its mass suggests that GJ 436b, like Neptune, cannot be a purely rocky body or ocean world but must contain a non-negligible component of light gases like hydrogen and helium in an outer atmospheric envelope (e.g., Fortney et al. 2007; Deming et al. 2007; Gillon et al. 2007; Demory et al. 2007; Stevenson et al. 2010; Knutson et al. 2011). Interior and evolution models for GJ 436b constrain this H/He component to be ~0.1-22% by mass, depending on other uncertain interior properties and the planet’s evolutionary history (Adams et al. 2008; Baraffe et al. 2008; Figueira et al. 2008; Rogers & Seager 2010a; Nettelmann et al. 2010; Miller & Fortney 2011).

The spectral and photometric behavior of GJ 436b’s atmosphere have been studied through secondary eclipse observations in the 3.6–24 \( \mu \)m range (Demory et al. 2007; Stevenson et al. 2010; Beaulieu et al. 2011; Knutson et al. 2011) and through transit observations in the ~0.5–8 \( \mu \)m range (Gillon et al. 2007a; Deming et al. 2007; Alonso et al. 2008; Bean et al. 2008; Coughlin et al. 2008; Ciceres et al. 2008; Pont et al. 2009; Shporer et al. 2009; Ballard et al. 2010; Beaulieu et al. 2011; Gibson et al. 2011; Knutson et al. 2011). Atmospheric models have been presented by Demory et al. (2007); Spiegel et al. (2010); Stevenson et al. (2010); Lewis et al. (2010); Madhusudhan & Seager (2011); Beaulieu et al. (2011); Shabram et al. (2011) and Line et al. (2011). The consensus from these models is that the large brightness temperatures derived from the Spitzer secondary eclipse data are indicative of inefficient heat redistribution from the dayside to the nightside of the planet and that the atmospheric metallicity may be greater than solar.

More contentious are the compositional inferences from the transit and eclipse data for GJ 436b — the transit depths and their implications, in particular (cf. Pont et al. 2009; Beaulieu et al. 2011; Shabram et al. 2011; Gibson et al. 2011; Knutson et al. 2011). From analyses of transit spectra from Hubble Space Telescope (HST) NICMOS instrument in the 1.1–1.9 \( \mu \)m range, Pont et al. (2009) and Gibson et al. (2011) both conclude that the GJ 436b transit spectrum at these wavelengths is relatively flat, with no evidence for strong molecular absorption features (including those from water); however, the smaller-scale wavelength dependence of the transit depths are notably different in the two investigations. Analyses of transit photometric data from the Spitzer IRAC instrument at 3.6, 4.5, and 8 \( \mu \)m have led to the conflicting conclusions of a methane-rich atmosphere (Beaulieu et al. 2011) and methane-poor atmosphere (Knutson et al. 2011). As is discussed extensively in Knutson et al. (2011), the GJ 436b Spitzer/IRAC transit depths appear to vary with time, which despite the relatively old and apparently quiet nature of the host star, could be due to the occultation of star spots or other regions of non-uniform brightness on the star’s surface as the planet transits across the disk. The eclipse data are less prone to such problems, although instrument systematics are still an issue, and stellar flares can complicate the data analyses (e.g., Stevenson et al. 2012). However, some disagreement still exists with respect to the inferred planetary flux at 3.6 and 4.5 \( \mu \)m and the associated error bars at these wavelengths in the eclipse data (cf. Stevenson et al. 2010; Beaulieu et al. 2011; Stevenson et al. 2012), although it is noteworthy that there is qualitative agreement in terms of the relative 3.6- to 4.5-\( \mu \)m flux ratio.

Resolving these discrepancies will be important (and/or obtaining new emission data for GJ 436b) because the Spitzer secondary-eclipse data at 3.6, 4.5, 5.8, 8.0, 16, and 24 \( \mu \)m suggest a very unexpected composition for GJ 436b’s dayside atmosphere (Stevenson et al. 2010). In particular, the very large flux in the 3.6-\( \mu \)m IRAC bandpass in combination with the negligible flux in the 4.5-\( \mu \)m bandpass suggest that CO and potentially CO\(_2\) are much more abundant than CH\(_4\) in the atmosphere of GJ 436b (Stevenson et al. 2010; Madhusudhan & Seager 2011; but see Beaulieu et al. 2011 for a contrary viewpoint), in contrast to theoretical models that predict that CH\(_4\) and H\(_2\)O will be the dominant carbon and oxygen constituents. If CH\(_4\) were the dominant carbon-bearing species under GJ 436b photospheric conditions, as thermochemical-equilibrium models for solar-like compositions suggest (e.g., Lodders & Fegley 2002), then absorption in the 3.6-\( \mu \)m channel would be stronger than is observed,
and the brightness temperature of the planet at those wavelengths would be much lower. Invoking a stratospheric temperature inversion in an attempt to explain the strong 3.6-µm emission does not improve the situation because the prominent \( \nu_1 \) band of CH\(_4\) would then produce a higher-than-observed flux at 8.0-µm (Stevenson et al. 2013; Madhusudhan & Seager 2011).

Through a systematic exploration of parameter space, Madhusudhan & Seager (2011) find that the best fit to all the Spitzer secondary-eclipse photometric data is obtained for atmospheres with very high CO mixing ratios, very low CH\(_4\) mixing ratios, and moderately low H\(_2\)O mixing ratios, with no thermal inversion. Both Stevenson et al. (2010) and Madhusudhan & Seager (2011) suggest that non-equilibrium chemical processes could be responsible for this unexpected atmospheric composition, with photochemistry destroying the methane in favor of complex hydrocarbons and carbon monoxide, and transport-induced quenching in combination with a high-metallicity atmosphere allowing a large quenched CO abundance. However, Line et al. (2011) demonstrate that for assumed atmospheric metallicities up to \( 50 \times \) solar, photochemistry does not effectively remove CH\(_4\) from the GJ 436b’s photosphere, and transport-induced quenching in combination with photochemistry cannot explain the large inferred CO abundance on GJ 436b. How then can the puzzling secondary-eclipse observations be explained?

We suggest that other bulk properties of the atmosphere, such as a very high metallicity or non-solar elemental compositions, could potentially resolve the current discrepancies between models and the secondary-eclipse observations of GJ 436b. As the atmospheric metallicity is increased in a planet with GJ 436b’s effective temperature, the overall hydrogen mole fraction is decreased, and species like CO and CO\(_2\) that do not contain hydrogen become progressively favored over hydrogen-containing species like H\(_2\)O and CH\(_4\). Similarly, as the C/O ratio is decreased, CH\(_4\) becomes progressively less important in relation to CO and CO\(_2\) as a major carbon-bearing constituent (Madhusudhan et al. 2012; Moses et al. 2013). We note that high-metallicity atmospheres are not unexpected for Neptune-mass planets (see section 3.1 below). In fact, in our own solar system, Neptune’s atmosphere is observed to have a C/H ratio of 40-120× solar (Baines et al. 1993; Karkoschka & Tomasko 2011) and is estimated to have an O/H ratio greater than 400× solar (Lodders & Fegley 1994; Luszcz-Cook & de Pater 2013).

We use both chemical-equilibrium models and thermochemical-photocatalytic gas and transport models (e.g., Moses et al. 2011; Visscher & Moses 2011; Moses et al. 2013) to investigate ways in which CO could be enriched at the expense of CH\(_4\) in the atmosphere of GJ 436b. We also explore how bulk properties like atmospheric temperature, metallicity, and the C/O ratio can affect the predicted composition of more generic hot Neptunes, leading to potentially widely diverse spectral properties for such planets.

Two chemical models are used in this study. The first is a chemical-equilibrium model using the NASA CEA code of Gordon & McBride (1994), and the second is a one-dimensional (1-D) thermochemical and photochemical kinetics and transport model based on the Caltech/JPL KINETICS code of Allen et al. (1981). The kinetics/transport model is described more fully in Moses et al. (2011), Visscher & Moses (2011), and Moses et al. (2013). For the equilibrium calculations, we consider \( \sim 500 \) gas-phase species and condensates containing the elements H, He, C, N, O, Ne, Na, Mg, Al, Si, P, S, Cl, Ar, K, Ca, Ti, Cr, Mn, Fe, and Ni. For the kinetics/transport calculations, we solve the continuity equations for 92 atmospheric species via \( \sim 1600 \) forward and reverse chemical-reaction pairs. Only species containing the elements H, He, C, N, and O are considered in the kinetics/transport model due to a lack of key rate-coefficient data for species containing the other elements. Note that the included elements are the dominant ones that will not be sequestered within condensates in the atmosphere of GJ 436b, and thus the GJ 436b results for the major gas-phase species are not expected to change significantly with the inclusion of additional elements.

The thermodynamic principle of microscopic reversibility, which is expected to be accurate even for complex multiple-potential-well chemical reactions (Miller et al. 2009), is assumed in the kinetics/transport model. Chemical equilibrium is achieved kinetically in the deepest, hottest regions of these models, but chemical reactions tend to be slower in the upper, cooler regions. When transport time scales drop below kinetic conversion time scales — at a pressure called the “quench pressure” or “quench level” — transport begins to dominate over chemical kinetics in controlling the species’ vertical profiles (e.g., Prim & Barshay 1977; Lewis & Fegley 1984; Fegley & Lodders 1994; Moses et al. 2010; Visscher & Moses 2011). When this situation occurs, the species can be “quenched” at mole fractions that remain constant with altitude above the quench level (and thus diverge from chemical-equilibrium predictions), as long as transport times scales remain shorter than chemical-kinetics time scales. Different assumptions about the rate coefficients for some key reactions can lead to differences in the predicted abundances of the quenched species (Visscher & Moses 2013; Venot et al. 2012); some of the current assumptions in published exoplanet models are reviewed by Moses (2013). The abundances of the carbon- and nitrogen-bearing species are particularly affected when kinetic interconversion between the dominant carbon constituents, CO and CH\(_4\), and dominant nitrogen constituents, N\(_2\) and NH\(_3\), ceases to be effective.

Changes to the Moses et al. (2011) chemical mechanism, other than what is discussed in Moses et al. (2013), include the addition of O\(_3\) and related reactions, which could potentially become important as the metallicity is increased. Ions and ion chemistry are not included in the models, and our solutions at the highest altitudes in the model will be unrealistic due to our neglect of ion chemistry, high-temperature thermospheres, and possible hydrodynamic escape (e.g., García Muñoz 2007). The models contain 198 vertical levels separated uniformly in log pressure, with the hydrostatic equilibrium equation being used to solve for the background atmospheric param-
eters along the vertical grid. The chemical-equilibrium abundance profiles from the CEA code are adopted as initial conditions in the kinetics and transport models, with zero flux boundary conditions being assumed for all species at the top and bottom of the model. Our assumed solar composition is taken from the protosolar abundances listed in Table 10 of Lodders et al. (2003). Multiple Rayleigh scattering of incoming stellar radiation by gases is considered in the kinetics/transport models, but we assume that aerosols are not present.

We assume that vertical transport occurs through molecular and "eddy" diffusion, with the eddy diffusion coefficients $K_{zz}$ being free parameters in the model. In the deep, convective portion of the atmosphere, free-convection and mixing-length theories (e.g., Stone[1977]) predict relatively large eddy diffusion coefficients and short mixing time scales (e.g., $K_{zz} \approx 10^9 \text{ cm}^2 \text{s}^{-1}$ for the convective regions in GJ 436b), but $K_{zz}$ values tend to be much smaller in the radiative regions in the upper troposphere and lower stratosphere. Turbulence due to atmospheric tides and upward-propagating gravity waves (non-breaking as well as breaking waves) is expected to cause effective $K_{zz}$ values to increase roughly with the inverse square root of atmospheric pressure in planetary stratospheres (e.g., Lindzen[1981] and references therein) — a scaling that appears consistent with inferred vertical mixing in exoplanet general circulation models (GCMs) (Parmentier et al. 2013; cf. also the $K_{zz}$ profiles inferred from Showman et al. 2009, as shown in Moses et al. 2011). In our kinetics and transport models, the $K_{zz}$ profile influences the quench behavior of molecules like CO, CH$_4$, and NH$_3$, and the observations themselves may ultimately provide the best means for defining both the $K_{zz}$ values at the quench levels and the pressures at which those quench levels occur (e.g., Fegley & Lodders 1994; Bézard et al. 2002; Visscher et al. 2003; Moses et al. 2010; Visscher & Moses 2011). However, the broadband photometric eclipse observations obtained to date for GJ 436b are not sufficient to constrain either the compositional or thermal profile accurately enough to derive $K_{zz}$ values in such a manner at this time. We therefore treat $K_{zz}$ as a free parameter and will explicitly specify our assumptions for each model presented in section 3.3.

2.2. Thermal Models

Both the chemistry and the predicted spectrum of our hot-Neptune exoplanets will depend strongly on the adopted thermal structure. We do not self-consistently calculate temperatures within the kinetics/transport models. For our generic hot Neptunes, we explore a wide range of temperature-pressure conditions. For GJ 436b, we consider a variety of theoretically-derived thermal profiles, including the dayside-average profiles at conditions of secondary eclipse from the 1× and 50× solar metallicity GCMs of Lewis et al. (2010), as shown in Fig. 1. For higher-metallicity GJ 436b scenarios, we use the PHOENIX atmospheric model (Hauschildt et al. 1999; Allard et al. 2001) in the presence of an external radiation field, as described in Barman et al. (2001) and Barman et al. (2003), to compute 1-D temperature-pressure profiles under the assumption of inefficient daytime heat redistribution and efficient gravitational settling of condensates (T. S. Barman, personal communication, 2012). The resulting profile for the case of 1000× solar metallicity is shown in Fig. 1. For all the non-solar-metallicity thermal models, solar ratios of elements other than hydrogen and helium are scaled by a constant factor. In certain instances, we also adopt thermal profiles from Madhusudhan & Seager (2011) that provide a good fit to the Stevenson et al. (2010) GJ 436b secondary-eclipse data, or we adopt profiles derived from new retrievals such as are described in Line et al. (2012, 2013) (see also section 3.2). We typically extend the thermal profiles upward in altitude nearly isothermally to a pressure of $10^{-11}$ bar, where all major UV absorbers are optically thin, and downward assuming an adiabat to a lower boundary that reaches at least 2400 K, to ensure that the N$_2$–NH$_3$ quench levels are contained within the models (i.e., to encompass the pressure at which kinetic interconversion between N$_2$ and NH$_3$ slows down enough to prevent equilibrium from being maintained, see Moses et al. 2010, 2011). The adopted thermal profiles will be clearly described when we discuss the different GJ 436b models.

Although the thermal profiles derived from the 3-D GCM and the 1-D PHOENIX models have some notable differences for any given metallicity (not shown in Fig. 1), both models predict a general upward vertical shift in the thermal profile as the metallicity is increased above solar. An increased metallicity results in increased mole fractions of opacity sources like water and other key molecular absorbers at lower pressures, which moves the optical-depth unity level (and "photosphere" in general) upwards to higher altitudes. However, because the overall atmospheric number density decreases with increasing altitude, there is a limit to how high the photosphere can be shifted upwards in altitude from this increased metallicity (see also Lewis et al. 2010; Miller-Ricci & Fortney 2010) because optical thickness...
cannot be approached at very high altitudes. As an example, the PHOENIX-based models show virtually no differences in the thermal profiles between 1000× and 10,000× solar metalicities. The chemical equilibrium

\[
\text{CH}_4 = \text{CO} \quad \text{equilibrium abundance of CH}_4 \quad \text{at 1000× solar metallicity}
\]

crosses H-like CH4 quench point (i.e., where \( T_{\text{int}} \approx 60 \, \text{K} \) for GJ 436b (comparable to that of Neptune with its \( T_{\text{int}} \approx 50 \, \text{K} \)), defined in terms of an intrinsic heat source would make higher CO/CH4 ratios more likely. In contrast, the thermal structure of the 1000× solar model shown in Fig. 4 lies completely within the CO-dominated regime, down to at least the 30-bar level. Since the CO=CH4 quench point (i.e., where transport processes dominate over the chemical-kinetic interconversion of CO and CH4) is likely to be within the ≈0.1-30 bar region on GJ 436b (Moses et al. 2011; Visscher & Moss 2011; Line et al. 2011), the quenching will occur in the CO-dominated regime for the 1000× solar metallicity model, and carbon monoxide will dominate over methane.

In general, the hotter the photosphere of GJ 436b, the more abundant CO is likely to be. Aside from the effect of increased metallicities, hotter photospheres can result from the addition of tidal heating or a large residual internal heat source. GJ 436 is a relatively quiet star and slow rotator (Saffe et al. 2005; Demory et al. 2007; Jenkins et al. 2009; Sanz-Forcada et al. 2010; Knutson et al. 2010, 2011), suggesting that the system is relatively old (e.g., \( 6_{-5}^{+4} \) Gyr, according to Torres et al. 2008). For an older planet of GJ 436b’s mass, the interior would be expected to have cooled significantly such that \( T_{\text{int}} \approx 60 \, \text{K} \) for GJ 436b (comparable to that of Neptune with its \( T_{\text{int}} \approx 50 \, \text{K} \)), defined in terms of an intrinsic heat source would make higher CO/CH4 ratios more likely. In contrast, the thermal structure of the 1000× solar model shown in Fig. 4 lies completely within the CO-dominated regime, down to at least the 30-bar level. Since the CO=CH4 quench point (i.e., where transport processes dominate over the chemical-kinetic interconversion of CO and CH4) is likely to be within the ≈0.1-30 bar region on GJ 436b (Moses et al. 2011; Visscher & Moss 2011; Line et al. 2011), the quenching will occur in the CO-dominated regime for the 1000× solar metallicity model, and carbon monoxide will dominate over methane.
downloadable file are photons cm$^{-2}$ s$^{-1}$ per wavelength bin, which then reproduces the correct integrated x-ray and EUV luminosity from Table 6 of Sanz-Forcada et al. (2011). At wavelengths between Lyman beta and the Lyman alpha line at 1215.7 Å, we use the solar spectrum of Woods & Rottman (2002) at solar-cycle minimum, scaled downward by a factor of 6.3 to transition smoothly to the longer-wavelength flux. For the Lyman alpha line itself, we use the reconstructed GJ 436 flux from Ehrenreich et al. (2011). For wavelengths between 1215.7 Å and 1800 Å, we use International Ultraviolet Explorer (IUE) data for GL 15B (a M3.5V star) from the MAST archive (http://archive.stsci.edu), and for wavelengths longer than 1800 Å, we use the spectrum of GL 15B from the Next Generation Spectral Library (Heap & Lindler 2010). For our dayside atmospheric models, we scale this normalized 1-AU spectrum to the 0.027 AU orbital distance relevant to the GJ 436b secondary eclipse. The key spectral region as far as the neutral atmospheric chemistry is concerned is the wavelength region from Lyman alpha out to ∼2400 Å. We use a fixed solar zenith angle $\theta = 48^\circ$ to simulate the eclipse conditions in the calculations (see Moses et al. 2011). A directly measured ultraviolet spectrum for GJ 436 has recently been made available by France et al. (2013), and we emphasize that such studies are of great utility to theoretical photochemical models for exoplanets.

2.4. Spectral Models

To calculate the emergent planetary spectrum for GJ 436b from the assumed thermal structure and derived abundance profiles from the chemical models, we use a plane-parallel radiative-transfer code as described in Line et al. (2013). Opacity from H$_2$, He, H$_2$O, CO, CO$_2$, CH$_4$, and NH$_3$ is included in the modeling, although NH$_3$ itself is not included in the retrievals. The source line parameters are described in Line et al. (2013). Local thermodynamic equilibrium is assumed throughout, and scattering is ignored. In a manner similar to Sharp & Burrows (2007), absorption cross sections are precomputed using a line-by-line code at high spectral resolution and tabulated on a grid with 1 cm$^{-1}$ wavenumber resolution and twenty evenly spaced temperature points between 500–3000 K and log(temperature) points between 50–10$^{-6}$ bar (see http://www.atm.ox.ac.uk/RFM/). The cross sections from this pre-tabulated grid can then be interpolated for the pressure-temperature-abundance conditions relevant to the model atmosphere, which has 90 grid layers equally spaced between 50–10$^{-6}$ bar. As is typical, we plot the model emission spectrum in terms of the flux of the planet divided by the flux of the star. The flux emergent from the entire disk of the planet is calculated using a four-point gaussian quadrature. The stellar flux is derived from a PHOENIX model (Hauschildt et al. 1999; Allard et al. 2001) assuming $T_{eff} = 3350$ K (Maness et al. 2007), although we should note that the latest determinations of the stellar effective temperature from von Braun et al. (2012) are somewhat hotter at $T_{eff} = 3416\pm54$ K (see also Bean et al. 2006; Nettelmann et al. 2010; Southworth 2010).

3. RESULTS

Results from our chemical equilibrium and kinetics/transport models are presented below. We first investigate how the predicted equilibrium composition of generic hot Neptunes changes as a function of atmospheric temperature, metallicity, and C/O ratio. We discuss various interesting atmospheric compositional regimes that are not representative of planets in our own solar system, and we identify conditions for which CO rather than CH$_4$ is likely to be the dominant carbon component. We then focus on GJ 436b, determining whether the retrieval technique of Line et al. (2013) can shed any new light on the atmospheric composition of the planet, and we calculate disequilibrium chemical abundance profiles for several possible metallicities and thermal profiles. Finally, synthetic spectra from these disequilibrium models are compared with Spitzer transit and eclipse data, and observational consequences are discussed.

3.1. Chemical Equilibrium for Generic Hot Neptunes

With the core-accretion model of giant-planet formation (Mizuno et al. 1978; Bodenheimer & Pollack 1986; Pollack et al. 1996), a rocky or rock-ice protoplanetary core initially forms and grows from the accretion of solid planetesimals within a protoplanetary disk, with gaseous envelopes forming around these emerging protoplanets through core outgassing, direct accretion of nebular gas, and ablation of incoming solid planetesimals within the gaseous envelope. Rapid accretion of the surrounding largely H$_2$ and He nebular gas occurs when the protoplanet reaches a certain critical mass (∼10$^{15}$ M$_{\oplus}$ in these traditional core-accretion models). The standard explanation for the “ice giants” Uranus and Neptune being so much more enriched in heavy elements than Jupiter and Saturn is that the accretion rate of solids was slow enough for the proto-Uranus and -Neptune that they did not completely reach the runaway gas-accretion phase before the nebular gas was dispersed from the disk (e.g., Lissauer & Stevenson 2007). Elements heavier than hydrogen and helium therefore make up 80–85% of Uranus and Neptune by mass (Podolak et al. 1995; Hubbard et al. 1994; Fortney & Nettelmann 2010). Depending on the degree of mixing between the core and atmosphere, the outermost gaseous envelope could have a heavy-element enrichment by mass ($Z_{env}$) different from that of the bulk planet (e.g., Nettelmann et al. 2013); however, observations of CH$_4$ on the giant planets support the picture of a greater atmospheric $Z_{env}$ for the ice giants Uranus and Neptune in comparison with the gas giants Jupiter and Saturn (e.g., Moses et al. 2004; Fouquet et al. 2009; Fegley et al. 1991; Gautier et al. 1995; Karkoschka & Tomasko 2009, 2011).

A metal-enriched atmospheric envelope might also be true for hot-Neptune exoplanets, despite potentially widely different evolutionary and migration histories. Although the hot Neptunes for which we have good constraints on both mass and radius show a large scatter in the mass-versus-radius relation, indicating likely different compositions, Weiss et al. (2013) demonstrate that there is a general trend toward increasing density with decreasing mass for planetary masses below ∼150$M_{\oplus}$ (see also Miller & Fortney 2011). This trend suggests that smaller planets become progressively more enriched
in heavy elements, similar to the situation in our own solar system. Planet-formation and population-synthesis models (e.g., Alibert et al. 2005; Figueira et al. 2009; Mordasini et al. 2012a,b) also reflect this trend, with Fortney et al. (2013) predicting a significant increase in metallicity in these calculations then becomes a proxy for the evolution of smaller Neptune-mass planets as a function of temperature, bulk metallicity, and bulk elemental ratios in the atmosphere. The increase in metallicity in these calculations then becomes a proxy for the evolution of smaller Neptune-mass planets, which are more likely to have high Z_{env} due to either inefficient gas accretion or more efficient atmospheric escape.

Figure 4 illustrates how the atmospheric composition is expected to change as a function of either the atmospheric metallicity or the atmospheric C/O ratio for a few different temperatures at a pressure of 100 mbar. The 100-mbar pressure was selected here because it represents a typical infrared photospheric emission region in transiting-planet atmospheres. For these calculations, we use the protosolar abundances of Lodders et al. (2009) and assume an increase in metallicity occurs uniformly for all species except H, He, and Ne; moreover, we de-

C/O ratio, a pressure of 100 mbar, and a temperature of 500 K. The middle row is similar to the top row, except the assumed temperature is 1200 K. The bottom row shows variations as a function of the C/O ratio for an assumed metallicity of 300 times solar (i.e., protosolar abundances of all species except H, He, Ne, and O are multiplied by 300, with O being defined through the C/O ratio), a pressure of 100 mbar, and a temperature of 800 K. Note the very large variation in composition for different bulk atmospheric properties. A color version of this figure is available in the online journal.

Fig. 4.—Pie charts illustrating equilibrium gas-phase compositions on generic hot Neptunes for different assumptions about atmospheric properties. The top row shows variations as a function of metallicity (10-10,000 times solar, as labeled) for an atmosphere with a solar C/O ratio, a pressure of 100 mbar, and a temperature of 500 K. The middle row is similar to the top row, except the assumed temperature is 1200 K. The bottom row shows variations as a function of the C/O ratio for an assumed metallicity of 300 times solar (i.e., protosolar abundances of all species except H, He, Ne, and O are multiplied by 300, with O being defined through the C/O ratio), a pressure of 100 mbar, and a temperature of 800 K. Note the very large variation in composition for different bulk atmospheric properties. A color version of this figure is available in the online journal.
Fig. 5.— Equilibrium mole fractions for different gas-phase species as a function of temperature and metallicity for a solar ratio of elements (except H, He, and Ne remain solar at all metallicities), at a pressure of 100 mbar. When we present a metallicity \([X/H]\) in square brackets here and in subsequent figures, we use the standard logarithmic definition; e.g., a metallicity of \([\text{Fe/H}] = 3\) corresponds to a heavy element enrichment of 1000× solar.

fine the oxygen abundance through the C/O ratio, so the “metallicity” in this context refers to the C/H ratio (or X/H ratio, where X is any species except H, He, Ne, or O). This “C/H metallicity” should not be confused with bulk atmospheric metallicity, as the overall heavy-element enrichment will change with the C/O ratio — at very low C/O ratios, for example, the overall atmospheric metallicity, as defined by the abundance of C + O + all heavy elements in relation to that of the Sun, can be quite a bit higher than the C/H metallicity due to the exaggerated importance of the enhanced O. Condensates are not assumed to rain out in this equilibrium model. Given such simplifications and given that solar ratios of elements may not be preserved as a planetary atmosphere forms and evolves, the relevance of these calculations to real planets is questionable, but the exercise does effectively demonstrate that the atmospheric composition of hot Neptunes could be highly variable with atmospheric properties. Exo-Neptunes with solar-like elemental ratios and moderately low metallicities could have hydrogen-dominated atmospheres very reminiscent of our own Neptune, but water and methane will make up an increasing fraction of the atmosphere with increasing metallicity, until \(\text{H}_2\) itself becomes less important. At high-enough metallicities, exo-Neptunes can have \(\text{CO}_2\)-dominated atmospheres, qualitatively reminiscent of Venus. Carbon monoxide becomes an increasingly important constituent at higher temperatures and higher metallicities, and even \(\text{O}_2\) can become dominant in equilibrium at very high metallicities and very low C/O ratios, further emphasizing that \(\text{O}_2\) by itself is not necessarily a good indicator of biological activity on exoplanets (e.g., see Selsis et al. 2002, Segura et al. 2007, Schaefer et al. 2012, Hu et al. 2012).
These scenarios are borne out more clearly in Fig. 6, which shows how the equilibrium abundances of several atmospheric species vary as a function of temperature and metallicity for an assumed pressure of 100 mbar and an assumed C/O ratio that is maintained at the protosolar value (C/O = 0.46) with an increase in metallicity (i.e., all elements except H, He, and Ne are increased by a constant factor). Because we do not have a complete set of condensates in these equilibrium calculations (and in particular, we are missing several Ca-Ti-Al silicates, see Lodders 2010), these gas-phase mixing ratios are not completely accurate, but the general trends with temperature and metallicity hold true. For example, Fig. 6 demonstrates that H$_2$ becomes relatively less abundant at all temperatures as the metallicity is increased, whereas the water (H$_2$O) mole fraction increases roughly linearly with increasing metallicity at most temperatures, until H$_2$O starts to decrease at very high metallicities (e.g., $\geq 1000 \times$ solar) due to the overall decrease in the bulk H fraction. Methane (CH$_4$) is more stable at low temperatures, where its abundance also increases with increasing metallicity until the bulk hydrogen abundance drops and makes less H available to form methane. Carbon monoxide (CO) becomes more stable at high temperatures and high metallicities, where it is a significant atmospheric component under these conditions. Carbon dioxide (CO$_2$) is not very abundant at low metallicities, especially at low temperatures, but CO$_2$ increases significantly with increasing metallicity, becoming the dominant constituent at very high metallicities for all temperatures. Carbonyl sulfide (OCS) has a behavior similar to CO$_2$, although it is never as abundant. Solid graphite is stable in the lower-right corner of these plots in Fig. 6 (high metallicities, low temperatures) and sequesters a notable fraction of the available carbon, thereby lowering the gas-phase C/O ratio in
that metallicities is particularly worth mentioning. We should also note that atmospheric metallicities of 10, 100, 1000, and 10,000 times solar correspond to $Z_{\text{env}} = 0.13$, $0.61$, $0.94$, and $0.99$, respectively, for solar ratios of the full suite of elements considered in the models (see section 2.1). This range of $Z_{\text{env}}$ variation is expected for Neptune-sized planets in the population-synthesis formation and evolution models of Fortney et al. (2013), with $Z_{\text{env}} = 0.6-0.9$ being the most common for such planets. We might therefore expect the whole range of possible phase space in Fig. 5 to be represented in the hot-Neptune exoplanet population. Although Uranus and Neptune are good examples of the cool, moderate-metallicity, H$_2$-dominated end members, many hot-Neptune exoplanets would have atmospheric compositions that are not found in our own solar system, with various H$_2$O-, CO-, or CO$_2$-rich possibilities being particularly worth mentioning. We should also note that metallicities $\gtrsim 500-600 \times$ solar are not possible if the “metals” are brought in predominantly via water or very water-rich volatiles (and/or through H-rich species such as CH$_4$, NH$_3$, and H$_2$S) — assuming no further atmospheric evolution due to hydrogen escape or other fractionation processes — because such volatiles would also deliver large amounts of hydrogen to the atmosphere (Nettelmann et al. 2011).

Aside from bulk metallicity, the atmospheric composition is also dependent on the C/O ratio (Seager et al. 2005; Kuchner & Seager 2005; Lodders 2010; Line et al. 2010; Madhusudhan et al. 2011a; Madhusudhan 2012; Kopparapu et al. 2012; Moses et al. 2013). In Fig. 6 we show how the equilibrium abundances of several atmospheric constituents vary as a function of temperature and C/O ratio for an assumed 100-mbar pressure and a moderately-high assumed C/H metallicity of 300$\times$ solar (i.e., where all elements other than H, He, Ne, and O are assumed to be 300 times the protosolar abundances of Lodders et al. 2009). Recall here that we define the oxygen abundance through the C/O ratio, such that the very low C/O ratios at the left edge of these plots also correspond to large overall atmospheric metallicities, in addition to O being a dominant element. For example, our nominal solar C/H ratio is $2.78 \times 10^{-4}$ and O/H ratio is $6.06 \times 10^{-4}$. For a uniform enrichment of all elements of 300$\times$ solar, the corresponding C/H and O/H ratios would be $8.33 \times 10^{-2}$ and $1.82 \times 10^{-1}$, respectively. For a C/H ratio of 300$\times$ solar but oxygen defined via the C/O ratio from O/H = (C/H)/(C/O), a C/O ratio of 0.08 would correspond to an O/H ratio of 1.04, which is $\sim 1700 \times$ the solar O/H ratio. Under such very low C/O ratio conditions (i.e., C/O $\lesssim 0.08$) in this otherwise 300$\times$ solar case, hydrogen is no longer the dominant element, and the H$_2$ abundance drops precipitously, as seen in Fig. 6. Molecular oxygen then becomes the dominant gas at C/O $\lesssim 0.04$ in this scenario — if such conditions have any relevance to real atmospheres — but H$_2$O quickly takes over with increasing C/O ratio to dominate at 0.05 $\lesssim$ C/O $\lesssim$ 0.19 (or at even greater C/O ratios for lower temperatures), whereas H$_2$ dominates at moderate-to-high C/O ratios (C/O $\gtrsim 0.2$). Carbon monoxide becomes an important constituent at moderate-to-high temperatures and moderate-to-high C/O ratios. CH$_4$ and NH$_3$ are important at low temperatures for all but the lowest C/O ratios, and CO$_2$ is important under most conditions except for the lowest temperatures considered at high C/O ratios. Condensed graphite is stable above $\sim 750$ K for the highest C/O ratios, and HCN increases in significance at high temperatures and high C/O ratios. Molecular nitrogen (not shown in Fig. 6) is relatively unaffected by the atmospheric C/O ratio.

The phase boundaries and stability regimes for the dominant carbon-bearing gases under the conditions plotted in Figs. 5 & 6 are further illustrated in Fig. 7. The conditions under which graphite is stable are also shown. Carbon dioxide dominates at very high metal-
licity and/or very low C/O ratios. At more moderate metallicities and C/O ratios, CH₄ dominates at low temperatures and CO at high temperatures.

It is unclear what C/O ratio to expect for hot Neptunes, as that will depend greatly on the planet’s original formation location (especially in relation to specific condensation fronts in the protoplanetary disk), as well as to its evolutionary history (e.g., Moses et al. 2013, and references therein). For high initial bulk C/O ratios, graphite condensation followed by gravitational settling of the condensates (i.e., precipitation) is expected to maintain the gas-phase C/O ratio \( \lesssim 1 \) above any graphite clouds in exoplanet atmospheres (Lodders & Fegley 1997; Seager et al. 2003; Lodders 2010); therefore, we do not consider C/O ratios > 1 in these models. However, C/O ratios close to 1 are possible (e.g., Lodders 2004; Madhusudhan et al. 2011a; Oberg et al. 2011; Moussis et al. 2012; Moses et al. 2013) if the planet accreted carbon-rich solids inward of the water-ice line in the disk; if the planet accreted CO-rich, H₂O-poor gas from a region between the H₂O and CO ice lines, from a region in the innermost disk, or from within a heterogeneous disk or water-poor feeding zones; or if the disk was carbon-rich to begin with. Similarly, C/O ratios < 0.1 can occur if the disk or feeding zones were oxygen-rich or if planetesimals composed largely of water ice or clathrate hydrates dominated the delivery of heavy elements to the protoplanetary envelope.

The chemical-equilibrium results are also sensitive to pressure, as is shown in Lodders & Fegley (2002) and Visscher et al. (2006), for example. However, given that transport-induced quenching will affect the predicted abundances as a function of pressure in real atmospheres, we have simply chosen a single representative photospheric pressure for the above figures. The quench level may occur at higher pressures in hot-Neptune atmospheres, thereby affecting the predicted compositions, and the infrared photosphere of high-metallicity exoplanets may reside at lower pressures than our nominal choice of 100 mbar. In fact, we emphasize that these simple equilibrium models are largely phenomenological and are not designed to represent all the complex processes that have gone into shaping the atmospheric composition of actual exoplanet atmospheres. These equilibrium models do serve a useful purpose, though, in illustrating the possible diversity of hot Neptunes and in highlighting specific trends such as the increasing dominance of CO₂ at very high metallicities, the importance of H₂O at moderate-to-high metallicities for a variety of other conditions, and the change in the relative importance of methane and CO with increasing temperature. These general trends can be useful for considerations of the likely bulk atmospheric properties of specific hot Neptunes such as GJ 436b, based on the compositional clues provided by transit and eclipse data.

3.2. CHIMERA retrieval methods applied to GJ436b

The secondary-eclipse data for GJ 436b (Stevenson et al. 2010; see also Beaulieu et al. 2011; Knutson et al. 2011; Stevenson et al. 2012) provide important constraints for compositional models. Madhusudhan & Seager (2011) use a Markov-chain Monte-Carlo (MCMC) method to help identify the range of parameter space allowed for GJ 436b from comparisons of synthetic emission spectra with the Spitzer photometric data. Consistent with the earlier conclusions of Stevenson et al. (2010), Madhusudhan & Seager (2011) find that plausible GJ 436b models require a low methane abundance (e.g., mole fractions of \( 10^{-6} \) to \( 10^{-7} \)) and a large CO abundance (mole fraction \( \geq 10^{-3} \)), along with a H₂O mole fraction \( \leq 10^{-4} \) and a CO₂ mole fraction in the range \( 10^{-6} \) to \( 10^{-4} \) in order to adequately reproduce the Stevenson et al. (2010) eclipse data. As is noted by Stevenson et al. (2010) and Madhusudhan & Seager (2011), such compositions are inconsistent with low-metallicity equilibrium models. Line et al. (2011) further demonstrate that disequilibrium chemical processes like photochemistry and transport-induced quenching do not help resolve this problem.

The relatively sparse spectral coverage, low signal-to-noise ratio, and systematic uncertainties of the Spitzer secondary-eclipse data from GJ 436b and other exoplanets make retrieving atmospheric information difficult, as many solutions are statistically valid (Madhusudhan & Seager 2009, 2011; Lee et al. 2012; Line et al. 2012, 2013; Benneke & Seager 2012; Barstow et al. 2013). As a check on the derived best-fit abundances and thermal profile for GJ 436b, we apply the CHIMERA retrieval code of Line et al. (2013) to the Stevenson et al. (2010) secondary-eclipse data. Briefly, CHIMERA employs a suite of Bayesian retrieval algorithms — optimal estimation, bootstrap Monte Carlo, and differential-evolution MCMC — to determine the allowed range of temperatures and gas-phase mixing ratios for GJ 436b. Here we discuss the results of the differential-evolution MCMC approach. With this tech-
unique, the posterior probability distribution for each of the parameters that controls the temperature structure (based on the analytic parameterizations of Guillo 2010) and assumed constant-with-altitude mole fractions for H₂O, CH₄, CO, and CO₂ is characterized using a genetic algorithm that generates ~10⁶ models (see Line et al. 2013, for further details). The statistics from the retrieval are shown in Figs. 8-10.

First, Fig. 8 shows both the retrieved thermal structure and the corresponding spectra for the ensemble of models for GJ 436b generated from the differential-evolution MCMC approach. Rather than plotting the many thousands of spectra and thermal profiles generated from this retrieval approach, we instead plot the median of the spectra (or temperatures in the plot insert) in black, along with the 1-sigma and 2-sigma spread in the spectra (or temperatures) in dark gray and light gray, respectively. In essence, these spreads reflect the fact that if we were to draw a random set of parameters from the posterior probability distributions, there would be a 95% chance that the flux at any one wavelength corresponding to the Spitzer bandpasses would fall within the 2-sigma spread, and so on. Note that although the synthetic spectra qualitatively reproduce the 3.6-to-4.5 µm flux ratio from the Spitzer photometric data, the non-detection of the eclipse in the 4.5-µm bandpass is particularly difficult to reproduce (see also Stevenson et al. 2010; Madhusudhan & Seager 2011).

Next, Fig. 9 shows histograms of the H₂O, CH₄, CO, and CO₂ mole fractions for GJ 436b from the marginalized posterior probability distribution, as derived from the differential-evolution MCMC approach. These distributions show that H₂O and CO are relatively well constrained from the GJ 436b secondary-eclipse spectra, with the most probable H₂O mole fraction residing within the range of a few ×10⁻⁷ to a few ×10⁻⁴, and the most probable CO mole fraction restricted to ≥ 10⁻⁴, although the solutions for both H₂O and CO contain an extended, highly unconstrained tail at lower mixing ratios. The methane distribution also shows that the CH₄ mole fraction is restricted to values less than 10⁻⁶. These results are consistent with the analysis of Madhusudhan & Seager (2011). The posterior CO₂ distribution has an interesting double-peaked structure that makes the precise value for the CO₂ mole fraction less well constrained, but viable solutions are found for mole fractions between 10⁻⁸ and 10⁻². This result for CO₂ is also consistent with the χ²/N栏 ≤ 2 results of Madhusudhan & Seager (2011), where N栏 is the number of available photometric data points. Our results confirm the picture of a GJ 436b atmosphere that possesses a large CO abundance, a very low methane abundance, and a moderately low water abundance.

Finally, Fig. 10 shows the correlations in the retrieved abundances amongst the different gases. The CH₄ abundance remains low, regardless of the abundance of the other species, and thus the CH₄ mole fraction appears uncorrelated with either CO, CO₂, or H₂O. An apparent correlation between CO and CO₂ results from the very low flux (non-detection) in the 4.5-µm Spitzer channel where both CO and CO₂ absorb: if the CO mole fraction is large enough, the CO₂ mole fraction is relatively unconstrained and can be low, but if the CO mole fraction is small, the CO₂ mole fraction must be large in order to explain the very low 4.5-µm flux. Note also that solutions with low CO and high CO₂ do not fit the data as well as models with relatively high CO, perhaps because the high-CO₂ solutions end up with too much absorption in the 16-µm Spitzer channel. Figure 10 also illustrates a correlation between CO₂ and H₂O. If water is fairly abundant, the CO₂ abundance is not well constrained, but if the water abundance is low, the CO₂ abundance is more tightly constrained to fall within a mole fraction range of ~10⁻⁸ to 10⁻⁶. At the highest water abundances, there is also a positive correlation between H₂O and CO₂ in that very high water abundances require very high CO₂ abundances. This may reflect a strong correlation between H₂O (as the gas that has the largest overall influence on the spectrum) and temperatures, in that a very large water abundance can only be accommodated through high temperatures, at which point more CO₂ is needed to keep the low flux at 4.5 µm. Given that more solutions are found for low-to-moderate water abundances, the low-abundance peak for CO₂ in Fig. 10 dominates over the secondary high-CO₂ abundance peak.

These new retrievals do not help resolve the apparent puzzles with respect to atmospheric composition on GJ 436b. Recalling Figs. 5 and 6, a very low methane abundance can occur in equilibrium at very high temperatures at low metallicities (i.e., T ∼ 1200 K in the photosphere, hotter than is likely for GJ 436b) or at more moderate temperatures when the C/O ratio is low or when the atmospheric metallicity is very high. However, under those conditions, the H₂O abundance is likely to be significantly larger than is indicated by all the retrievals (above, and Madhusudhan & Seager 2011). Similarly, the preference for high CO mole fractions in the retrievals suggests a high metallicity for GJ 436b, which again implies unacceptably large water (and potentially CO₂) abundances. In fact, looking at Figs. 5 and 6 we need to emphasize that there are no equilibrium chemistry solutions — or even, as we will show below, no disequilibrium chemistry solutions — that fall within the favored temperature-abundance ranges indicated by the retrievals. This result illustrates one potential drawback of retrievals from sparse, noisy, systematics-prone data: the solutions may reproduce the data well but not make any physical sense.

Going to higher metallicities does help in general in that the photosphere is both hotter and more likely to reside in the CO-dominated regime (see Fig. 1), but a high water abundance is a necessary consequence of high metallicity, unless the metallicity is so large (Z ∼ 10 solar) that the atmosphere contains very little H. That solution, which corresponds to Z绿v = 0.994, (i.e., 0.6% H/He by mass), is unfortunately ruled out by GJ 436b’s mass-radius combination, according to most interior models (cf. Adams et al. 2008; Baraffe et al. 2008; Figueira et al. 2009; Rogers & Seager 2010a; Nettelmann et al. 2010; Miller & Fortney 2011).

Figure 11 illustrates the possible C/O-ratio vs. metallicity phase space that can accommodate a low methane abundance on GJ 436b for the assumption of a moderately hot, high-altitude photosphere (e.g., assuming 900 K at 10 mbar), which may be relevant for a high-metallicity GJ 436b atmosphere. The shaded regions in this plot are consistent with the retrieved CH₄ mole fractions of < 1 ppm, which can occur at low atmospheric
Fig. 9.— Gas mole-fraction histograms of the marginalized posterior probability distribution derived from the differential-evolution Markov-chain Monte-Carlo retrieval approach (Line et al. 2013), as applied to the GJ 436b Spitzer eclipse data of Stevenson et al. (2010). The horizontal dot-dashed blue curves represent the priors, which are assumed to be flat (uninformative). The vertical red lines represent the mole fractions that correspond to the best-fit solution. A color version of this figure is available in the online journal.

Fig. 10.— Contours of the gas-abundance constraints for GJ 436b derived from the differential-evolution MCMC posterior probability distributions, illustrating the correlations between the different gases. The dark gray regions represent the 1-sigma confidence interval and light gray regions represent the 2-sigma confidence interval. The red dot in each plot is the best-fit (maximum-likelihood) solution from the ensemble of $10^5$ fits. A color version of this figure is available in the online journal.
Fig. 11.— Equilibrium mole fractions for different species as a function of C/H metallicity (see text) and C/O ratio at 900 K and 10 mbar (i.e., a relatively hot, high-altitude photosphere, which may be relevant to a high-metallicity GJ 436b). The shaded region in each plot is where methane has a mole fraction below $10^{-6}$, as is indicated by retrievals based on the Stevenson et al. (2010) Spitzer GJ 436b secondary-eclipse data (our work, and that of Madhusudhan & Seager 2011).
C/O ratios and/or high metallicities. Again, there are no equilibrium solutions within the shaded regions of Fig. 11 that are entirely consistent with the retrievals, with the main problem being too much H$_2$O and too much CO$_2$. However, the atmosphere is not likely to be in equilibrium [Line et al. 2010; Moses 2013], and we use these constraints as a guide for our disequilibrium kinetics and transport models, in an attempt to find forward models with more physically meaningful atmospheric properties that can provide a reasonable fit to the GJ 436b Spitzer secondary-eclipse data.

3.3. Disequilibrium chemistry modeling of GJ 436b

Photochemistry and transport-induced quenching can affect the predicted photospheric abundances on extrasolar giant planets, resulting in mixing ratios that are often orders of magnitude different from chemical-equilibrium expectations [Liang et al. 2003, 2004; Zaline et al. 2009, Line et al. 2010, 2011; Moses et al. 2011, 2013; Miller-Ricci Kempton et al. 2012, Kopparapu et al. 2012, Venot et al. 2012, Hu et al. 2012]. Most disequilibrium models to date have focused either on giant planets with a near-solar-like complement of elements or on hydrogen-poor terrestrial exoplanets or super-Earths. We now test various scenarios with moderate and high metallicities (but still containing a non-negligible hydrogen mass fraction) for GJ 436b using disequilibrium kinetics/transport models to see how photochemistry and transport-induced quenching can alter the predicted transit and eclipse spectra. We start with the first-principles-based temperature profiles shown in Fig. 1 and adopt a solar C/O ratio and a constant eddy diffusion coefficient of $K_{zz} = 10^9$ cm$^2$ s$^{-1}$ for these initial models. Fig. 12 shows how the model results for CH$_4$, H$_2$O, CO, and CO$_2$ compare with the mixing ratios retrieved from the Spitzer secondary-eclipse data. This figure demonstrates that even when photochemistry and transport-induced quenching are considered, none of these models can reproduce the abundances derived from the retrievals.

The mixing-ratio profiles for other potentially interesting species in these disequilibrium models are shown in Fig. 13. The species profiles for the 1× solar model are qualitatively similar to those of Line et al. (2011): behind H$_2$ and He, the next most abundant gases are H$_2$O, CH$_4$, and NH$_3$. Water survives at a near-equilibrium abundance until very high altitudes (~1 microbar), at which point photolysis and other destruction mechanisms start to irreversibly convert the H$_2$O to CO and other oxygen-bearing species. Although photolysis and other chemical mechanisms operate to destroy H$_2$O at lower altitudes, water is efficiently recycled in the background H$_2$ atmosphere, and H$_2$O remains the dominant infrared opacity source in these 1× solar models. Methane is even less stable than water at high altitudes, due primarily to the large H abundance released by H$_2$O photolysis and subsequent catalytic destruction of H$_2$ (e.g., Liang et al. 2003; Moses et al. 2011). The carbon that was in high-altitude CH$_4$ gets photochemically converted to CO, C$_2$H$_2$, HCN, and atomic carbon, primarily (see Moses et al. 2011 for details). Ammonia is photolyzed by longer-wavelength UV radiation that can penetrate a bit deeper in the atmosphere, and the nitrogen liberated by the NH$_3$ photodestruction largely ends up as HCN and atomic N. There are some quantitative differences between our 1× solar model and that of Line et al. (2011) due to different adopted reaction rate coefficients, $K_{zz}$ assumptions, stellar ultraviolet flux, and atmospheric temperatures, but the results from both Line et al. (2011) and our own models indicate that CH$_4$ survives photochemical destruction throughout the bulk of the ~0.0001–1 bar photosphere on the 1× solar GJ 436b. As a result, the 1× solar model has more methane than is indicated by the retrieval analyses of the Spitzer secondary-eclipse data (see Fig. 12 section 3.2 and Madhusudhan & Seager 2011).

The resulting spectrum for the 1× solar model does not compare well with the Spitzer secondary-eclipse data (Fig. 13). One obvious problem is a reversed 3.6-to-4.5-μm flux ratio in comparison with observations. The CO-CH$_4$ quench point in the 1× solar model occurs within the CH$_4$-dominated regime, so that the quenched CO abundance is relatively small, and the methane abundance is large. Since the equilibrium abundance of CO is everywhere lower than that of CH$_4$ with the 1× solar thermal profile (see Fig. 1), there is no eddy $K_{zz}$ value we could adopt that would change this conclusion. Some CO is produced at high altitudes from CH$_4$ and H$_2$O photochemistry, but the CO column abundance is never large enough to influence the relative 3.6-to-4.5-μm flux ratio. The large methane abundance therefore results in a lower flux (more absorption) at 3.6 μm in comparison with 4.5 μm. Another obvious problem is the overall lower brightness temperature at most wavelengths in comparison with the data. The thermal profile from Lewis et al. (2010) adopted in the 1× solar model is not significantly colder than the profiles derived from the thermal retrievals shown in Madhusudhan & Seager (2011) or in section 4.2 but the 1× solar model does have a water abundance at the upper end (or greater than) the retrievals indicate (see Fig. 12). Water accounts for most of the absorption in the 1× solar model, and the relatively large water abundance leads to more absorption at most wavelengths than can be accommodated by the observations – hence the requirement of a low water abundance from the retrievals in the first place. Since photochemistry does not permanently destroy the photospheric water and methane in our 1× solar models, such models cannot explain the Spitzer secondary-eclipse data.

The quench point for the 50× solar model is closer to the equilibrium CO-CH$_4$ equal-abundance boundary due to an overall hotter atmosphere (see Fig. 1), and both CO and CH$_4$ end up being major atmospheric constituents (see Fig. 13). Carbon dioxide and molecular nitrogen also become more important constituents at the higher 50× solar metallicity, while ammonia becomes relatively less important due to the N$_2$-NH$_3$ quench point falling within the N$_2$-dominated regime. Because NH$_3$ is more photochemically active than N$_2$, the photochemical production of HCN and complex nitriles does not increase significantly at high altitudes in the higher-metallicity model because the overall NH$_3$ mixing ratio has not changed much between 1× and 50× solar metallicities. However, the column abundance of HCN actually in the atmosphere, and the relatively large water abundance leads to more absorption at most wavelengths than can be accommodated by the observations – hence the requirement of a low water abundance from the retrievals in the first place. Since photochemistry does not permanently destroy the photospheric water and methane in our 1× solar models, such models cannot explain the Spitzer secondary-eclipse data.
Fig. 12.— Mixing-ratio profiles for CH$_4$ (top left), H$_2$O (top right), CO (bottom left), and CO$_2$ (bottom right) from our kinetics/transport models for GJ 436b, for assumed atmospheric metallicities of 1× solar (blue solid lines), 50× solar (green solid lines), 1000× solar (orange solid lines), and 10,000× solar (purple solid lines), as described more fully in Fig. 13. The corresponding equilibrium solutions for the 10,000× solar model are shown as dashed purple lines. The red horizontal bar illustrates the most-probable solutions derived from the differential-evolution MCMC approach discussed in section 3.2, with the star representing the best-fit solution. The black horizontal bar represents the model solutions from Madhusudhan & Seager (2011) that fit the Spitzer data to within $\chi^2/N_{\text{obs}} \leq 1$. Note that none of the disequilibrium models for the different metallicities have mole fractions that fall within the retrieval constraints for all four species. A color version of this figure is presented in the online journal.

in the photosphere maintains a pseudo-equilibrium with the quenched NH$_3$ and CH$_4$; see Moses et al. 2011, 2013. Carbon dioxide is produced effectively at high altitudes from the photochemistry of CO and H$_2$O, but equilibrium through the net reaction CO + H$_2$O $\rightleftharpoons$ CO$_2$ + H$_2$ is maintained kinetically through much of the photosphere. Although the CO mole fraction in the 50× solar model now falls within the range required by the retrievals discussed in section 3.2, both H$_2$O and CH$_4$ are much more abundant than is allowed by the retrievals, and the 50× solar model does not fit the Spitzer secondary-eclipse data well.

At metallicities of 1000× solar, hydrogen and helium now make up only $\sim$6% of the atmosphere by mass. The methane abundance has dropped significantly in the 1000× solar metallicity model because the CH$_4$-CO quench point lies within the CO-dominated regime. The resulting CO/CH$_4$ ratio is finally in the right direction to explain the observed relative 3.6- to 4.5-μm flux ratio from the Spitzer eclipse data. Carbon monoxide begins to rival H$_2$ as the dominant constituent, and H$_2$O, CO$_2$, and N$_2$ are all very abundant. The atmosphere as a whole is more oxidized, and species like O$_2$ and NO that are produced from photochemistry at high altitudes are able to survive more readily than they do in a more reducing atmosphere. The fact that elements other than O, C, N, and H have been neglected from these disequilibrium models (particularly Cl and S) will have an impact on the resulting abundances in higher-metallicity models due to omitted catalytic cycles (e.g., Yung & DeMott 1999), and atmospheric escape and other evolutionary processes will likely alter this simple picture of steadily increasing metallicity, but the general supplantation of H$_2$ and the increasing dominance of CO$_2$ with increasing metallicity is a robust conclusion. At atmospheric metallicities of 10,000× solar, for instance, hydrogen and helium make up only $\sim$0.6% of the atmosphere by mass, and Fig. 13 shows that CO$_2$ has solidly replaced H$_2$ as the dominant constituent, and even CO, N$_2$, and H$_2$O are more abundant than H$_2$.

Figure 14 shows that the 1000× solar metallicity model fares better in reproducing the Spitzer secondary-eclipse data than the 1× solar metallicity model did. The flux in the 3.6 μm channel is predicted to be greater than that in...
Fig. 13.—Mixing-ratio profiles for several species of interest (as labeled) in our kinetics/transport models for GJ 436b, for assumed atmospheric metallicities of $1 \times$ solar (top left), $50 \times$ solar (top right), $1000 \times$ solar (bottom left), and $10,000 \times$ solar (bottom right). The thermal profiles adopted in these models are described in section 2.2 (see Fig. 1); the profile for the $10,000 \times$ solar model (not shown in Fig. 1) was derived from the PHOENIX model (Hauschildt et al. 1999; Allard et al. 2001; Barman et al. 2005) and is very similar to the $1000 \times$ solar profile. The C/O ratio is assumed to be the protosolar value of Lodders et al. (2009) with 21% of the oxygen unavailable due to being bound up in condensates at deep atmospheric levels (Visscher et al. 2010a), and the eddy diffusion coefficient $K_{zz}$ is assumed to be constant at $10^9$ cm$^2$ s$^{-1}$. A color version of this figure is presented in the online journal.

the 4.5 $\mu$m model now with the $1000 \times$ solar model, due to the CH$_4$ abundance being much less than the CO abundance. The $1000 \times$ solar model still provides too much absorption at 3.6 $\mu$m to be consistent with the eclipse depth as determined by Stevenson et al. (2010), indicating too much methane in the model; however, we should note that Beanleu et al. (2011) derive both a lower 3.6-$\mu$m flux and a larger uncertainty in the eclipse depth at this wavelength (and at 4.5 $\mu$m), and the $1000 \times$ solar model more readily satisfies those constraints. More intriguing is the excellent fit in the 5.8 and 8.0 $\mu$m channels where water in this model is providing the dominant opacity source. This good fit with a water abundance that exceeds a 10% mixing ratio — in contrast to the much lower abundances derived from the retrievals — reflects the influence of the higher-temperature photosphere that results from the increased atmospheric opacity at high metallicities. The water abundance and photospheric temperatures are closely linked, and a hotter photosphere requires more water to remain consistent with the 5.8 and 8.0 $\mu$m eclipse depths, whereas, conversely, a colder photosphere requires less water. Still, this $1000 \times$ solar model provides a relatively poor fit to the eclipse data of Stevenson et al. (2010, 2012), providing a $\chi^2/N_{obs} \sim 11.5$, with the main problem being a 3.6-$\mu$m flux that is much lower than observed, but also a 4.5-$\mu$m flux that is in excess of the observational upper limit and a 16-$\mu$m flux that is much lower than observed (due to CO$_2$ absorption).

Although the $1000 \times$ solar metallicity model itself does not provide a good fit to the Stevenson et al. (2010, 2012) secondary-eclipse data, and none of the models in Fig. 12 fall within the constraints imposed by the retrievals for all the species, the model-data comparisons do suggest further avenues to explore with higher-metallicity models. The larger observed brightness temperature at 3.6 $\mu$m in relation to that at 5.8 and 8 $\mu$m in the Spitzer data suggests that the atmosphere has a sharper thermal gradient than the one derived from the 1-D PHOENIX model, given that the contribution functions at the three wavelengths are not hugely separated in terms of the pressures at which they peak in the high-metallicity models. We note that both the dayside-average profiles from the 3-D GCMs of Lewis et al. (2010) and the thermal profiles derived from the retrievals exhibit a larger thermal gradient in the relevant photospheric region than the
The red and gold circles represent, respectively, the fluxes from the 1× and 1000× solar models, averaged over the Spitzer bandpasses. The black dotted lines represent the planetary emission (smoothed) assuming GJ 436b radiates as a blackbody at a temperature of 500 K (lower curve), 800 K (middle curve), or 1100 K (upper curve). See Fig. 14 for further details. A color version of this figure is available in the online journal.

For the model presented in this figure, the atmospheric C/H and N/H metallicity are assumed to be 300× solar, the C/O ratio above the silicate clouds is assumed to be 0.6 (which actually requires a slightly sub-solar initial bulk C/O ratio if ~21% of the oxygen is sequestered in silicates and other condensates at deeper atmospheric levels, as expected by Lodders et al. 2010 and Visscher et al. 2010a), and the eddy diffusion coefficient follows the relation $K_{zz} = 10^7 \text{ cm}^2 \text{ s}^{-1}$ at pressures $P \geq 10^{-4}$ bar, with $K_{zz}$ increasing with the inverse square root of atmospheric pressure at $P < 10^{-4}$ bar. The thermal profile as shown in the insert of Fig. 4 of Madhusudhan & Seager (2011) has been shifted upward uniformly by $-1.2$ in $\log_{10}(P)$ (i.e., pressures have been multiplied by $10^{-1.2}$) to account for the higher-altitude photosphere that will result from the higher metallicity. The 300× solar atmospheric metallicity in this model is consistent with the interior models for GJ 436b (Adams et al. 2008, Baraffe et al. 2008, Figueira et al. 2009, Rogers & Seager 2010a, Nettelmann et al. 2010, Miller & Fortney 2011), and the lower value of $K_{zz}$ in the lower atmosphere allows CH$_4$ to remain in equilibrium.

1000× solar 1-D PHOENIX models do. A larger thermal gradient could also result in a hotter atmosphere at the CH$_4$-CO quench point, if the quench point occurs at a high-enough altitude, pushing the atmosphere towards greater CO dominance at the expense of CH$_4$. That would improve the relative 3.6- to 4.5-µm flux ratio in comparison with observations. To investigate such a scenario, we simply scale one of the thermal profiles derived from the Madhusudhan & Seager (2011) MCMC analysis upward uniformly in altitude to lower pressures and run different disequilibrium chemistry models, assuming different bulk atmospheric properties. The results from one such model are shown in Fig. 15.
until ~0.1 bar, at which point the methane quenches at the desired < 1 ppm abundance. Judging from the PHOENIX models, the thermal profile at photospheric pressures may be hotter in this model than is expected on average in the dayside hemisphere of GJ 436b at this metallicity, so the likelihood of this scenario will need to be investigated by further 3-D circulation modeling, but given that the observed emission will derive preferentially from the hottest regions on the observed disk, the profile does not appear to be unreasonably hot.

Figure 15 shows that H₂ is still the dominant atmospheric constituent (by number) in the atmosphere in this 300× solar metallicity model, although both CO and H₂O at mole fractions of ~10% each are encroaching on the H₂ dominance. Kinetic production of CO₂ from H₂O and CO occurs in the middle and upper photosphere in this model, allowing the CO₂ abundance to exceed that of water at high altitudes. Quenching of ammonia at non-negligible values occurs and is responsible for maintaining HCN at greater than ppm levels. The CH₄ abundance remains low, and the emission spectrum (Fig. 16) is dominated by opacity from H₂O, CO₂, and CO. This model finally has a low-enough methane abundance that the 3.6-μm flux from the Spitzer secondary-eclipse analysis of Stevenson et al. (2010) is better reproduced. The resulting 3.6-to-4.5-μm flux ratio is also more in line with observations, although this model, like all others including the retrievals, still produces too much flux in the 4.5-μm band. This result is more a function of the atmosphere being too hot where the contribution function peaks at 4.5 μm, rather than being due to insufficient CO and CO₂, as can be readily seen in Figs. 12 & 15.

In fact, the very large CO₂ abundance in this 300× solar metallicity model produces too much absorption in the 16-μm Spitzer channel, where the contribution function peaks at cooler, higher altitudes due to the large CO₂ column abundance. The non-detection of the GJ 436b secondary eclipse at 4.5 μm is difficult to explain by any known physically reasonable scenario without adversely affecting the fit at other wavelengths. In terms of the statistical fit to all the Stevenson et al. (2010, 2012) secondary eclipse depths, this model fares better, at χ²/N₁obs = 2.8. The primary problems are insufficient flux at 3.6, 16, and 24 μm, and too much flux at 4.5 μm.

This particular forward model is not unique in terms of fitting the data, and there are numerous models with atmospheric metallicities in the couple-hundred to couple-thousand times solar range that fit the Spitzer data to within χ²/N₁obs ≤ 3 (see, for example the 2000× solar metallicity model we presented in Richardson et al. 2013), but we have not found any disequilibrium chemistry models that fit to within χ²/N₁obs ≤ 2. All of these high-metallicity models produce too much CO₂ to be consistent with the observed flux in the 16-μm channel. Models with high metallicities in the 100–10,000× solar range also end up with a large water abundance, which then requires a hot photosphere to explain the 5.8, 8.0, and 24-μm fluxes, at which point there is too much emission at 4.5 μm. Slightly cooler models than the one shown in Figs. 15 & 16 tend to fit better at 4.5 μm, but then the fit tends to be worse at 3.6 and 24 μm. Hotter models fit the 24-μm flux better, but then the fit at 4.5, 5.8, and 8.0 μm is worse. Going to a higher C/O ratio will help remove excess CO₂ to improve the fit at 16 μm, but then CH₄ becomes too abundant to explain the 3.6-μm flux. Lower C/O ratios help keep the methane abundance low, but then the problems with excess H₂O and CO₂ are exacerbated. Therefore, while these high-metallicity models are promising in their ability to qualitatively reproduce several aspects of the observed secondary-eclipse observations, several quantitative problems remain. The solutions derived from the retrieval methods (section 3.2 and Madhusudhan & Seager 2011) provide a better overall fit to the data, but problems with the fit still remain at 4.5 and/or 16 μm; more importantly, these solutions have problems with plausibility in that it is difficult to theoretically explain how such abundances could be obtained chemically or physically in an atmosphere with any reasonable bulk properties. Therefore, despite the less-than-perfect fit to the existing GJ 436b secondary-eclipse data, such high-metallicity models should not be dismissed out of hand.

The higher the atmospheric metallicity, the higher the mean molecular weight, and the smaller the atmospheric scale height for any given temperature and gravity profile. High-metallicity atmospheres will then have flatter transmission spectra (e.g., Fortney et al. 2013). Examples of transmission spectra from some of our GJ 436b models are shown in Fig. 17 in comparison with observations. The relatively flat spectrum observed
in HST/NICMOS data (Pont et al. 2009; Gibson et al. 2011) seems more consistent with the high-metallicity models, and such models also seem to compare well to the results from the Knutson et al. (2011) Spitzer analysis (see the open squares at 3.6, 4.5, and 8 µm in Fig. [17]). However, the high-metallicity models do not compare well with the Beaulieu et al. (2011) Spitzer transit analysis (see the open diamonds in Fig. [17] or the K-band transit data of Caceres et al. [2009]), and in fact none of the models provides a reasonable fit to all the available transit data for GJ 436b. Given the observed variability in the apparent transit depths with time from different observations (e.g., Knutson et al. [2011] and the lack of agreement between transit results analyzed with different procedures (e.g., Pont et al. [2009]; Gibson et al. [2011]; Knutson et al. [2011]; Beaulieu et al. [2011]), the poor fits of these models to the transit data are not particularly surprising. Obtaining simultaneous spectral observations at multiple wavelengths should help resolve some of these issues and help distinguish between competing models.

Note that we have not adjusted for potential differences in temperatures or chemical abundances between the terminator and dayside atmospheres when performing these calculations. The dayside atmosphere observed at secondary eclipse is likely hotter than the limb atmosphere at the terminators probed in the planetary transit (see Lewis et al. [2010]). The non-negligible eccentricity of GJ 436b will also lead to variations in temperature over the orbit, which is captured in the GCMs of Lewis et al. [2010]. It is likely that horizontal quenching will help homogenize the atmospheric abundances with respect to longitude due to strong zonal winds or rapid thermal changes over the orbit (Cooper & Showman 2009; Iro & Deming 2010; Visscher 2012; Agundez et al. 2012). But without more detailed multi-dimensional and temporal modeling, it is difficult to predict the exact terminator composition. Regardless of the exact molecular composition, however, the high-metallicity models will have a higher atmospheric mean molecular mass, and a flatter overall transit spectrum.

4. GJ 436b COMPOSITIONAL CONSTRAINTS FROM STRUCTURE AND EVOLUTION MODELS

As mentioned earlier, a relatively high metallicity for GJ 436b is also indicated by the planet's observed radius in relation to its mass. Figure [13] shows some of the compositional constraints imposed by models of the planet's interior structure, formation, and evolution (e.g., Adams et al. 2008; Baraffe et al. 2008; Figueira et al. 2009; Rogers & Seager 2010a; Nettelmann et al. 2010; Miller & Fortney 2011). The amount of H/He required to explain the mass and radius of GJ 436b depends on several other assumptions in the models, such as the relative abundance of ices (i.e., volatile heavy elements) versus rocks (i.e., refractory heavy elements), the age and evolutionary history of the planet, the presence/absence of a central core, the degree of mixing of elements within the planet, and the thermal structure within the planet. Not much H/He is needed to explain the observed radius if the planet is very hot (i.e., young, high intrinsic interior $T_{\text{int}}$ or tidal heating, early onset of convective interior, high-temperature isothermal radiative region), if the heavy elements in the planet have low atomic weight (i.e., ices, not rocks), and/or if the heavy elements are confined to deep levels and are not mixed throughout the atmospheric envelope. Conversely, more H/He is needed to explain the radius if the planet is old and cool, contains more rocky than icy elements, and has heavy elements mixed throughout the atmospheric envelope. Parameters such as the ice-to-rock ratio of GJ 436b are not well constrained from structure and evolution models alone, and the planet's age, evolutionary history, and atmospheric and interior thermal structure are also uncertain. As is clear from Fig. [13] the formation models of Figueira et al. (2009) provide the tightest constraints on the bulk composition. When the uncertainties in GJ 436b's mass, radius, and age are taken into account (e.g., Miller & Fortney 2011), the final uncertainty in the possible H/He mass fraction for GJ 436b is significantly increased. The assumed temperature of the isothermal radiative portion of the atmosphere somewhat influences the inferred H/He mass fraction (Nettelmann et al. 2010).

Accounting for the various uncertainties and adopting reasonable assumptions about the atmosphere and interior, most of these models predict H/He mass fractions of $\sim 3$–22% for GJ 436b, corresponding to metallicities of $\sim 230$–2000 $\times$ solar. Nettelmann et al. (2010) demonstrate that even lower H/He mass fractions (higher metallicities) are possible if the planet contains few rocky elements (only ices), if the ices are confined to the core, and if the pure H/He outer envelope is hot as a result of a thin radiative region. A hot isothermal radiative region is possible for GJ 436b (see Spiegel et al. 2010 and our Fig. 4 above); however, the other assumptions are less secure, and given that the secondary eclipse data show evidence for molecular absorbers rather than a blackbody atmosphere, it seems likely that at least some heavy elements are mixed into the outer atmospheric envelope. Therefore, the $\sim 230$–2000 $\times$ solar models seem most probable for GJ 436b, and our high-metallicity disequilibrium models discussed in section 3.3 appear reasonable in terms of the interior and evolution models for GJ 436b.

5. CONCLUSIONS

Moderate-to-high metallicity models ($\sim 230$–2000 $\times$ solar) for GJ 436b are appealing in that they provide a natural explanation for the apparent CO-rich, CH$_4$-poor nature of GJ 436b’s atmosphere (see Stevenson et al. 2010; Madhusudhan & Seager 2011) and are consistent with the heavy-element enrichment as inferred from the planet’s mass and radius from interior modeling (Adams et al. 2008; Baraffe et al. 2008; Figueira et al. 2009; Rogers & Seager 2010a; Nettelmann et al. 2010; Miller & Fortney 2011). Although such high-metallicity models are not without problems in terms of explaining all the transit and secondary-eclipse data (see section 3.3), they do have one significant advantage over the solutions obtained to date from retrieval methods (e.g., Madhusudhan & Seager 2011 and section 3.3 above) — the advantage of physical and chemical plausibility. The relative abundances of CO, H$_2$O, CH$_4$, and CO$_2$ derived for GJ 436b from the retrieval methods do not appear to be achievable from either equilibrium or disequilibrium chemistry in a GJ 436b atmosphere with any plausible imagined bulk properties (i.e., metallicity, C/O ratio, temperature). Either our theoretical understanding is incomplete, and we are
Fig. 18.— Constraints on the bulk composition of GJ 436b as determined from models of its structure, evolution, and formation. Possible compositions are shown for six different model assumptions and/or imposed constraints. For each, the sum over the displayed mass fractions of H/He (green diamonds), water (blue squares), and “rock” (includes silicates plus iron; orange triangles) equals 1. All calculations take into account the planet’s mass \( M_p \) and radius \( R_p \) as one set of constraints; the two leftmost models consider uncertainties in \( M_p \) and \( R_p \); while the four on the right adopt specific \( M_p \) and \( R_p \) values from Torres et al. (2008). From left to right, the derived constraints are based on (i) the thermal and structural evolution models of Miller & Fortney (2011) that assume an ice:rock ratio of 1:1 and that impose constraints based on the age of the GJ 436 system; (ii) the formation, disk-planet evolution (including migration), and structural models of Figuera et al (2009) that compute various formation paths consistent with \( M_p \) and \( R_p \); (iii) the structure models of Adams et al. (2008) that consider precise values for \( M_p \), \( R_p \), and atmospheric temperatures; (iv) the structural models of Nettelmann et al. (2010) that assume that water is mixed into the outermost H/He envelope; (v) structural models based on Nettelmann et al. (2010) that assume that a pure H/He layer resides above a water layer with a warm (1300 K) deep isothermal radiative region; (vi) a model similar to the one on the immediate left, except the isothermal radiative region is cool (700 K) and the planet is old (> 10 Gyr). Note that the tightest constraints on the composition are imposed by the formation models, that including the uncertainties in \( M_p \) and \( R_p \) enhance the uncertainty in the H/He mass fraction by a factor of ~2, and that assumptions about atmospheric temperatures affect the derived H/He mass fraction. A color version of this figure is available in the online journal.
Regardless of whether GJ 436b itself has a high-metallicity atmosphere, radial-velocity and transit surveys show that Neptune-class exoplanets are exceedingly common in the galaxy (Howard et al. 2010, 2012; Mayor et al. 2011; Borucki et al. 2011; Batalha et al. 2013), and based on our current views of exoplanet formation (see section 3.1), we would expect many of these planets to have atmospheric metallicities that are orders of magnitude in excess of solar. There is some evidence for that trend in the existing population of exoplanets for which we have derivations of both mass and radius (e.g., Miller & Fortney 2011; Weiss et al. 2013). As is shown in Figs. 4, 5, & 6, hot Neptunes could have very diverse atmospheric compositions depending on atmospheric temperatures and bulk atmospheric properties such as metallicity and C/O ratio. The higher-metallicity atmospheres described in this paper have no analogs from within our own solar system, but one can imagine a continuum of hot Neptunes that should exist in the exoplanet population, with atmospheres ranging from moderate-to-low metallicity, hydrogen-rich, Neptune-like compositions to high-metallicity, hydrogen-poor, super-Venus-like compositions, along with more exotic CO-dominated planets and H2O-dominated “water worlds”. As the statistics regarding exoplanet properties continues to grow, so does our amazement at the diversity of these worlds beyond the narrow confines of our solar system.

We thank Travis Barman for calculating and providing 1-D thermal profiles for high-metallicity GJ 436b atmospheres, Nikole Lewis for providing her dayside-average GCM thermal profiles for GJ 436b, and Heather Knutson for useful communications. This work was supported by the NASA Planetary Atmospheres Program grant number NNX11AD64G.

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