Observable Signatures of Wind-driven Chemistry with a Fully Consistent Three-dimensional Radiative Hydrodynamics Model of HD 209458b

B. Drummond\(^1\), N. J. Mayne\(^1\), J. Manners\(^1,2\), A. L. Carter\(^1\), I. A. Boutle\(^1,2\), I. Baraffe\(^1,3\), É. Hébrard\(^1\), P. Tremblin\(^4\), D. K. Sing\(^1\), D. S. Amundsen\(^5,6\), and D. Acreman\(^1\)

\(^1\) Astrophysics Group, University of Exeter, Exeter EX4 2QL, UK; bdrummond@exeter.ac.uk

\(^2\) Met Office, Exeter EX1 3PB, UK

\(^3\) Maison de la simulation, CEA, CNRS, Ens de Lyon, Univ Lyon1, CNRS, CRAL, UMR5574, F-69007 Lyon, France

\(^4\) Department of Applied Physics and Applied Mathematics, Columbia University, New York, NY 10025, USA

\(^5\) NASA Goddard Institute for Space Studies, New York, NY 10025, USA

\(^6\) NASA Ames Research Center, Mail Stop 248-6, Moffett Field, CA 94035, USA

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Abstract

We present a study of the effect of wind-driven advection on the chemical composition of hot-Jupiter atmospheres using a fully consistent 3D hydrodynamics, chemistry, and radiative transfer code, the Met Office Unified Model (UM). Chemical modeling of exoplanet atmospheres has primarily been restricted to 1D models that cannot account for 3D dynamical processes. In this work, we couple a chemical relaxation scheme to the UM to account for the chemical interconversion of methane and carbon monoxide. This is done consistently with the radiative transfer meaning that departures from chemical equilibrium are included in the heating rates (and emission) and hence complete the feedback between the dynamics, thermal structure, and chemical composition. In this Letter, we simulate the well studied atmosphere of HD 209458b. We find that the combined effect of horizontal and vertical advection leads to an increase in the methane abundance by several orders of magnitude, which is directly opposite to the trend found in previous works. Our results demonstrate the need to include 3D effects when considering the chemistry of hot-Jupiter atmospheres. We calculate transmission and emission spectra, as well as the emission phase curve, from our simulations. We conclude that gas-phase nonequilibrium chemistry is unlikely to explain the model–observation discrepancy in the 4.5 μm Spitzer/IRAC channel. However, we highlight other spectral regions, observable with the James Webb Space Telescope, where signatures of wind-driven chemistry are more prominent.

Key words: planets and satellites: atmospheres – planets and satellites: composition – planets and satellites: gaseous planets

1. Introduction

Gas-phase chemical composition is crucial in determining the circulation, temperature, and observable properties of planetary atmospheres. The opacity, determined by the composition, controls the heating, which, in turn, drives the dynamics.

Chemical modeling of exoplanet atmospheres has largely been restricted to 1D chemical kinetics codes (e.g., Moses et al. 2011; Venot et al. 2012; Zahnle & Marley 2014; Drummond et al. 2016; Tsai et al. 2017), showing that vertical transport-induced quenching can lead to departures from chemical equilibrium.

Hot-Jupiter atmospheres have large horizontal temperature gradients, due to intense irradiation of the dayside, that drive fast horizontal wind velocities. Therefore, nonequilibrium chemistry due to horizontal transport has been speculated to be at least as important as vertical mixing in hot-Jupiter atmospheres (Showman et al. 2009; Moses et al. 2011) and has been suggested to explain the model–observation discrepancy found in thermal phase curves for several hot Jupiters (Knutson et al. 2012; Zellem et al. 2014; Wong et al. 2016).

Using a 3D atmosphere model with temperature and chemical relaxation schemes, Cooper & Showman (2006) investigated the effect of wind-driven advection on the chemistry of methane, carbon monoxide, and water. They found vertical quenching from a deep level of the atmosphere leads to a horizontally uniform composition at lower pressures. Cooper & Showman (2006) conclude that vertical (not horizontal) mixing is dominant in determining the chemical composition.

Agúndez et al. (2014) considered a 1D chemical kinetics model with a time-varying temperature profile (a “pseudo-2D model”), representing a column of atmosphere rotating around the equator. Agúndez et al. (2014) conclude that horizontal quenching is more important than vertical quenching, with the nightside atmosphere being “contaminated” by the composition of the dayside.

We investigate the effect of wind-driven advection on the atmospheric composition using a 3D general circulation model consistently coupled with a flexible radiative transfer scheme and a chemical relaxation scheme. We simulate the atmosphere of HD 209458b and find significant quantitative and qualitative differences with both Cooper & Showman (2006) and Agúndez et al. (2014). We conclude that a combination of horizontal and vertical transport determines the 3D chemical composition of hot-Jupiter atmospheres. We also identify particular wavelength regions where wind-driven chemistry can cause observable differences in the transmission and emission spectra, compared with what is predicted assuming chemical equilibrium.

2. Model Description

We use the Met Office Unified Model (UM), based on a dynamical core that solves the deep-atmosphere, non-hydrostatic...
Navier–Stokes equations (Mayne et al. 2014a, 2017; Wood et al. 2014). The model uses the open-source SOCRATES\textsuperscript{7} radiative transfer scheme (Edwards 1996; Edwards & Slingo 1996) to calculate the radiative flux and heating rates in each column (Amundsen et al. 2014, 2016, 2017). The UM has previously been applied to the atmospheres of hot Jupiters (Mayne et al. 2014b, 2017; Amundsen et al. 2016), sub-Neptunes (Drummond et al. 2018), and terrestrial exoplanets (Boutle et al. 2017).

2.1. Chemical Relaxation Scheme

Previous UM simulations of hydrogen-dominated atmospheres (e.g., Amundsen et al. 2016) have assumed local chemical equilibrium. Here we relax this assumption by consistently coupling a chemical relaxation scheme to the UM, following the method of Cooper & Showman (2006).

The chemical relaxation scheme parameterizes the kinetic interconversion of chemical species. In this study, we focus on the major gas-phase absorbers methane, carbon monoxide, and water. The material derivative of the carbon monoxide mole fraction $\dot{f}_{CO}$ is

$$ \frac{Df_{CO}}{Dt} = \frac{\partial f_{CO}}{\partial t} + \frac{u}{r \cos \phi} \frac{\partial f_{CO}}{\partial \lambda} + \frac{v}{r} \frac{\partial f_{CO}}{\partial \phi} + \frac{w}{\partial \tau}, $$

where $t$ is time, $r$ is radius, $\phi$ is latitude, $\lambda$ is longitude, and $u$, $v$, and $w$ are the zonal, meridional, and vertical wind velocities, respectively. The first term on the right describes changes of $f_{CO}$ with time due to local processes (such as chemical conversion), while the remaining terms describe the advection processes in each direction.

Using the chemical relaxation method, we write

$$ \frac{\partial f_{CO}}{\partial t} = - \frac{f_{CO} - f_{CO,eq}}{\tau_{chem}}, $$

where $f_{CO,eq}$ is the chemical equilibrium mole fraction of carbon monoxide and $\tau_{chem}$ is a chemical timescale. The advection of the tracer is performed by the extensively tested, semi-implicit, semi-Lagrangian advection scheme of the UM (see Wood et al. 2014 for details).

Equation (2) describes the “relaxation” of $f_{CO}$ toward a prescribed equilibrium profile $f_{CO,eq}$ on a given timescale $\tau_{chem}$. We use the same calculation of $\tau_{chem}$ as Cooper & Showman (2006).

Assuming all carbon is contained in carbon monoxide and methane and all oxygen is contained in carbon monoxide and water the mole fractions of methane $f_{CH_4}$ and water $f_{H_2O}$ can be calculated from $f_{CO}$ via mass balance,

$$ f_{CH_4} + f_{CO} = A_C = 4.57 \times 10^{-4}, \quad (3) $$

$$ f_{H_2O} + f_{CO} = A_O = 8.31 \times 10^{-4}, \quad (4) $$

where $A_C$ and $A_O$ are constants based on the solar elemental abundances (Asplund et al. 2009).\textsuperscript{8}

The above assumption is a good one for the high temperature, solar composition atmospheres of hot Jupiters, where other carbon and oxygen containing species are generally several orders of magnitude less abundant (e.g., Burrows & Sharp 1999; Moses et al. 2011). However, the assumption will break down for metallicities greater than solar, where species such as HCN and CO$_2$ become more abundant (e.g., Moses et al. 2013). We have compared the equilibrium methane mole fraction calculated via the mass balance approach with that from a full Gibbs energy minimization method and found negligible differences.

We use the same method as Cooper & Showman (2006) to calculate the equilibrium mole fractions of methane, carbon monoxide, and water, assuming fixed mole fractions of hydrogen $f_{H_2} = 0.853$ and helium $f_{He} = 0.145$.

We have tested our model by reproducing the results of Cooper & Showman (2006), using a similar temperature relaxation scheme. Qualitatively, we find very similar results: wind-driven advection leads to a vertically and horizontally uniform composition, with almost all carbon in carbon monoxide, for pressures less than 1 bar. In chemical equilibrium, we also reproduce the large methane abundance on the nightside. Quantitative differences are small (~1%) but expected to exist due to many differences between the two models, e.g., model discretization, resolution, numerical methods, and diffusion schemes.

2.2. Model Setup and Simulation Parameters

We use the same simulation parameters for HD 209458b as Amundsen et al. (2016) with 144 points in longitude and 90 in latitude, and 66 vertical levels and dynamical and radiative timesteps of 30 s and 150 s, respectively. We refer the reader to Amundsen et al. (2014, 2016, 2017) for a complete description of the radiative transfer method, calculation of the opacities and sources of the line-lists.

Simulated observables are calculated directly from the UM, using a higher spectral resolution than when computing the heating rates (see Boutle et al. 2017). The transmission spectrum is computed from the 3D model using all the columns on the nightside of the limb. Columns are treated independently using spherical shell geometry and the resultant spectra are averaged to give a global mean. A full description is given in Lines et al. (2018b).

We use the advected $f_{CO}$, and corresponding $f_{CH_4}$ and $f_{H_2O}$ from mass balance, to calculate the opacity due to these species. Therefore, the radiative transfer is computed consistently with the advected chemistry of these molecules, allowing feedback between the local composition, the thermal structure, and the circulation.

For other chemical species that are included as opacity sources (Amundsen et al. 2016), we assume chemical equilibrium abundances. The equilibrium mole fraction of molecular nitrogen is calculated in an analogous way to carbon monoxide (Cooper & Showman 2006), using the net reaction

$$ N_2 + 3H_2 \leftrightarrow 2NH_3, $$

where ammonia is then found via mass balance with $N_2$,

$$ 2f_{N_2} + f_{NH_3} = A_N = 1.16 \times 10^{-4}. $$

The mole fractions of the alkali species are estimated using the same parameterizations as those in Amundsen et al. (2016).

We present results from two simulations that are identical, except that the first assumes local chemical equilibrium, while

\textsuperscript{7} \url{https://code.metoffice.gov.uk/trac/socrates}

\textsuperscript{8} Several previous publications (Tremblin et al. 2015; Drummond et al. 2016, 2018; Goyal et al. 2018) erroneously referred to Caffau et al. (2011) for the elemental abundances. The correct reference for the elemental abundances is Asplund et al. (2009).
the second includes advection and chemical relaxation of $f_{\text{CO}}$. We refer to these simulations throughout the rest of this Letter simply as the “equilibrium” and “relaxation” simulations.

The simulations are initialized with zero wind velocities and a horizontally uniform pressure–temperature ($P-T$) profile, as in Amundsen et al. (2016). The carbon monoxide tracer is initialized to chemical equilibrium corresponding to the initial $P-T$ profile, as in Cooper & Showman (2006).

We integrate the model for 1000 Earth days, to compare with Cooper & Showman (2006), by which point the atmosphere has reached a steady state for pressures less than $10^6$ Pa. The deep atmosphere, on the other hand, continues to evolve in Amundsen et al. 2016b, 2017; Amundsen et al. 2016 after this time and requires integration times that are infeasible with current models.

3. Dynamics, Thermal Structure, and Composition

Figure 1 compares the mole fraction of methane between the equilibrium and relaxation simulations. Figure 2 shows the dynamical and thermal structure from the relaxation simulation, which are very similar to the results of Amundsen et al. (2016).

The equilibrium abundance of methane clearly traces the temperature structure (compare Figures 1 and 2), as the chemistry is only dependent on the local temperature and pressure, for a given elemental composition. The equilibrium methane mole fraction varies with latitude and longitude by several orders of magnitude with larger abundances, where the atmosphere is cooler.

Advection homogenizes the methane abundance, both horizontally and vertically, across a large pressure range. Generally, the methane abundance in the relaxation simulation is larger than that in the equilibrium simulation. The winds that drive this homogenization (Figure 2) are characterized by a strong equatorial zonal jet, upwelling on the dayside and downwelling on the nightside and alternating regions of poleward and equatorward flow in the meridional direction.

Figure 3 shows vertical profiles of the mole fractions of methane, carbon monoxide, and water for a series of longitude points around the equator. In the relaxation simulation, methane is homogenized vertically and/or horizontally, depending on the pressure. The mole fractions of both carbon monoxide and water, which are approximately constant with pressure (for $P > 10^6$ Pa) and longitude in equilibrium, are not significantly affected.

Comparing the relaxation and equilibrium simulations, moving from high to low pressures, the abundance of methane deviates from equilibrium at $\sim 3 \times 10^5$ Pa. Over a relatively narrow pressure range, the methane abundance becomes uniform with longitude, with a value corresponding to the profile with the smallest equilibrium methane mole fraction. Here, zonal transport is driving the chemistry away from equilibrium.

Between $10^5$ and $10^4$ Pa, the methane abundance increases with decreasing pressure, to values significantly above those shown in the equilibrium simulation. This is due to meridional transport of gas from higher latitudes, where methane is more abundant in equilibrium. The predominant meridional component of the wind velocity throughout the atmosphere is poleward; however, small regions of equatorward flow, at low latitudes, act to transport gas toward the equator (see Figure 2).

For pressures less than $10^4$ Pa, the methane abundance becomes approximately uniform with pressure due to vertical quenching, with a quenched mole fraction of $\sim 3 \times 10^{-6}$, larger than the values predicted by chemical equilibrium.

In summary, at the equator, we find that the mole fraction of methane is sequentially affected by zonal, meridional, and vertical transport, moving from high to low pressures. To first order, we find that vertical quenching is the most important process and determines the methane abundance for all pressures less than $P \sim 10^4$ Pa. However, importantly, the abundance at the vertical quench point is determined by horizontal transport. This shows the importance of including 3D transport when considering transport-induced quenching of chemical species in exoplanet atmospheres.

Since we include interaction between the dynamics, radiative transfer, and chemistry, we can quantify the effect of wind-driven chemistry on the circulation and temperature fields. We find modest differences ($\sim 1\%$). This is not surprising, in this particular case, as the important absorbers, water and carbon monoxide, show negligible departure from chemical equilibrium. In addition, though methane varies from the equilibrium mole fraction by several orders of magnitude, its abundance remains small ($f_{\text{CH}_4} \lesssim 3 \times 10^{-6}$).

Figure 3 shows the estimated dynamical and chemical timescales from our simulations. The dynamical timescale is split into the zonal $\tau_{\text{dyn}}^u$, meridional $\tau_{\text{dyn}}^v$, and vertical $\tau_{\text{dyn}}^w$ components, which we estimate as

$$\tau_{\text{dyn}}^u = \frac{L}{u} = \frac{2\pi R_p}{u}$$

$$\tau_{\text{dyn}}^v = \frac{L}{v} = \frac{\pi R_p}{2v}$$

$$\tau_{\text{dyn}}^w = \frac{H}{w}$$

where $L$ is the relevant horizontal length scale, $R_p$ is the planetary radius, and $H$ is the vertical scale height. The relevant chemical timescale here is for the interconversion of carbon monoxide and methane, discussed in Section 2.1. We have assumed dayside averages of $u$, $v$, and $T$, for $\tau_{\text{dyn}}^u$, $\tau_{\text{dyn}}^v$, and $\tau_{\text{chem}}$, respectively. For $\tau_{\text{dyn}}^w$, we assumed the dayside average of $v$ over the northern hemisphere ($0^\circ < \phi < 90^\circ$) only.

Where $\tau_{\text{chem}} < \tau_{\text{dyn}}^w$, we expect chemical equilibrium to hold, while for $\tau_{\text{chem}} > \tau_{\text{dyn}}^w$ we instead expect advection to drive the local composition. Each of the directional components of the dynamical timescale cross $\tau_{\text{chem}}$ at approximately $10^5$ Pa, which agrees well with the pressure level at which methane diverges from equilibrium. We note that the timescales presented here serve as order of magnitude estimates, which depend on the assumed horizontal or vertical length scale and on the sampled wind velocity and temperature. In reality, it is the ratio of the local chemical and dynamical timescale that is of importance. The timescales shown in Figure 3 are therefore not expected to predict the precise pressure level of the zonal, meridional, and vertical quench points in our simulation.

To illustrate the transport of material to the equatorial region, we perform simulations, initialized from the equilibrium simulation at 1000 days, that include a tracer that is fixed at unity in a “source” region ($20^\circ > |\phi| > 90^\circ$ and $P \gtrsim 10^6$) and initialized to zero elsewhere. As the tracer departs the “source” region, its value decays exponentially to
zero with time (with a half-life of 1000 days). Figure 4 shows snapshots from this simulation, revealing transport from \(|\phi| > 20^\circ\) into the equatorial region, over a timescale of several hundred days, due to equatorward meridional winds (Figure 2), and subsequent upwelling. This experiment highlights the importance of the combined effect of horizontal and vertical transport that can only be captured using 3D models.
4. Simulated Observables

Figure 5 shows the synthetic observations from our simulations, derived directly from the 3D model: the emission phase curve in several spectral regions, and secondary eclipse and transmission spectra. The observed 4.5 μm phase curve (Zellem et al. 2014) and simulated JWST observations (using PandExo Batalha et al. 2017) are both overlaid. The PandExo simulations were performed for the NIRSpec G395H and MIRI...
Figure 3. Left: vertical abundance profiles of methane, carbon monoxide, and water from the relaxation (solid) and equilibrium (dashed) simulations at a series of longitude points around the equator ($\phi = 0^\circ$). Right: estimated dynamical and chemical timescales.

Figure 4. Snapshots showing the zonal-mean of the tracer value. The regions enclosed by the red dashed lines indicate (approximately) where the tracer is fixed to a value of unity: the "source" regions. Elsewhere, the value of the tracer exponentially decays with a half-life $t_{1/2} = 1000$ days. Green arrows illustrate the direction of flow.
LRS modes, using a single eclipse with equal in to out of transit observation time, noise floor of 50 ppm, detector saturation of 80% full well and stellar and planetary parameters from the TEPCAT database. All instrument-related parameters were kept at the PandExo defaults.

We find a negligible difference between the emission of the equilibrium and relaxation simulations within the 4.5 $\mu$m Spitzer/IRAC channel, suggesting that gas-phase nonequilibrium chemistry is unlikely to explain the model–observation discrepancy. More prominent signatures of wind-driven chemistry are apparent within the 3–4 $\mu$m and 7–10 $\mu$m spectral regions (including the 3.6 $\mu$m Spitzer/IRAC channel), corresponding to methane absorption features. Clear differences between the equilibrium and relaxation simulations can be seen in both transmission and emission, particularly between 3–4 $\mu$m for the NIRSpec G395H mode on the James Webb Space Telescope. The decreased flux and increased transit depth both result from enhanced methane abundances that increase the opacity in these spectral regions.

Figure 5. Top: emission phase curves in several spectral bands. The observed 4.5 $\mu$m Spitzer/IRAC channel curve (Zellem et al. 2014) is included (black) with 1σ uncertainty. Middle and bottom: secondary eclipse emission and transmission spectra with PandExo simulated observations for the NIRSpec G395H (circles) and MIRI LRS (squares) modes, binned to a resolution of $R \sim 60$ and $R \sim 30$, respectively.

5. Discussion and Conclusions

We have developed the first fully consistent 3D dynamics, radiative transfer, and chemistry model applied to exoplanets. We have focussed on the effect of wind-driven advection on methane, carbon monoxide and water, the major gas-phase absorbers. Using a chemical relaxation scheme, based on Cooper & Showman (2006), we have shown that methane is generally enhanced above what is expected from chemical equilibrium, for the specific case of HD 209458b. This is directly opposite to the trend found by previous studies (Cooper & Showman 2006; Agúndez et al. 2014). We find that methane is homogenized, horizontally and vertically, for pressures less than 10^4 Pa, through a combination of horizontal and vertical transport.

Cooper & Showman (2006) conclude that quenching in the vertical direction is more important than in the horizontal. They argue that vertical quenching transports gas from high pressures, where carbon monoxide is favored over methane, ultimately leading to a horizontally uniform composition for lower pressures. In contrast, our simulations show that a combination of horizontal and vertical quenching leads to an increase in the methane abundance, compared with chemical equilibrium.

When using a similar temperature relaxation scheme, rather than deriving the heating rates via radiative transfer calculations, our model gives very similar results to Cooper & Showman (2006). The temperature relaxation scheme used by Cooper & Showman (2006) results in a significantly warmer dayside and cooler nightside compared with the radiative transfer method (Showman et al. 2009; Amundsen et al. 2016); the larger day–night contrast then drives a faster equatorial jet. Different locations of the quench points, and different equilibrium abundances at the quench points, due to differing temperature and wind velocity fields, explain the contrasts between our results and those of Cooper & Showman (2006).

We also find significant differences with Agúndez et al. (2014) who concluded that a combination of vertical and zonal transport leads generally to a decrease in the methane abundance, compared with chemical equilibrium. While our results suggest, alternatively, that 3D transport acts to increase
the methane abundance above equilibrium, the approach of our model and theirs is significantly different.

Agúndez et al. (2014) take a highly simplified approach to horizontal transport (solid-body rotation around the equator) but include a full chemical kinetics scheme. On the other hand, our present model considers 3D transport due to the resolved wind but with a highly simplified scheme. We do not expect the uncertainties inherent to the chemical relaxation method to be the reason for the opposite trend compared with Agúndez et al. (2014). We suspect that their approximate treatment of horizontal and vertical advection are the main reasons for our opposing results. To confirm this, and to better understand the opposite trends predicted by the two methods, requires coupling a chemical kinetics scheme to a 3D dynamical model, a work that is under progress.

Our results clearly demonstrate the need to include 3D dynamical processes when considering the effect of transport on the composition of exoplanet atmospheres. While vertical quenching dominates over a large pressure range, horizontal transport is important for higher pressures and sets the vertically quenched mole fraction of methane, in the equatorial region. Such processes simply cannot be included consistently in 1D, or even 2D, atmosphere models.

We found small (~1%) changes to the wind velocity and temperature structure due to the interaction between the composition, the radiative transfer and, subsequently, the dynamics. Interaction between the wind-driven chemistry and the thermal and dynamical structure may be more important for cooler atmospheres, which are closer to the CO–CH₄ equilibrium profile, or planets with metallicities greater than solar, where important absorbers such as carbon dioxide and hydrogen cyanide are more abundant.

We find that wind-driven advection of methane, carbon monoxide, and water is unlikely to explain the model–observation discrepancy in the 4.5 μm Spitzer/IRAC channel emission phase curve. Alternative explanations may involve clouds or hazes (Lines et al. 2018a) or non-solar abundances (Kataria et al. 2014; Drummond et al. 2018), which can alter the dynamics, thermal structure, and opacity.

We find signatures of wind-driven chemistry in the spectral regions 3–4 μm and 7–10 μm due to enhanced methane absorption. These wavelength regions will be accessible with the upcoming James Webb Space Telescope.

Our simulations include advection due to the resolved, large-scale wind. However, we do not account for unresolved motions that may result in additional transport. An important future work would be to include and quantify this process. We also neglect photochemical processes. However, the main effects of photochemistry are usually restricted to pressures less than 1 Pa (e.g., Moses et al. 2011; Drummond et al. 2016), beyond the domain of our simulations.

We use a chemical relaxation scheme (Cooper & Showman 2006) that parameterizes the kinetic interconversion of methane and carbon monoxide and relies on an approximated timescale. The accuracy of the method therefore depends on the estimation of this timescale. To improve on this, we are coupling a chemical kinetics scheme to the UM, which does not rely on such approximations. Coupling a flexible chemical kinetics scheme to a GCM is crucial to enable the study of a larger number of chemical species and a wider range of chemical scenarios, particularly where the chemical relaxation method may not be accurate or relevant.

In summary, our results indicate that wind-driven chemistry, due to 3D dynamics, leads to a significant increase in the abundance of methane, compared with chemical equilibrium, for HD 209458b-like atmospheres. The increased methane absorption is predicted to significantly effect the emission and transmission spectra, detectable with the James Webb Space Telescope.

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**ORCID iDs**

B. Drummond @ https://orcid.org/0000-0001-7589-5484
N. J. Mayne @ https://orcid.org/0000-0001-6707-4563
É. Hebrard @ https://orcid.org/0000-0003-0770-7271
P. Tremblin @ https://orcid.org/0000-0001-6172-3403
D. K. Sing @ https://orcid.org/0000-0001-6050-7645
D. S. Amundsen @ https://orcid.org/0000-0002-5612-7321
D. Acreman @ https://orcid.org/0000-0002-4881-7584

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