Abstract. We present the first incorporation of the Common Representative Intermediates version 2.2 tropospheric chemistry mechanism, CRI v2.2, combined with stratospheric chemistry, into the global chemistry–climate United Kingdom Chemistry and Aerosols (UKCA) model to give the CRI-Strat 2 mechanism. A rigorous comparison of CRI-Strat 2 with the earlier version, CRI-Strat, is performed in UKCA in addition to an evaluation of three mechanisms, CRI-Strat 2, CRI-Strat and the standard UKCA chemical mechanism, StratTrop v1.0, against a wide array of surface and airborne chemical data.

CRI-Strat 2 comprises a state-of-the-art isoprene scheme, optimized against the Master Chemical Mechanism v3.3.1, which includes isoprene peroxy radical isomerization, HO₃, recycling through the addition of photolabile hydroperoxy aldehydes (HPALD), and isoprene epoxy diol (IEPOX) formation. CRI-Strat 2 also features updates to several rate constants for the inorganic chemistry, including the reactions of inorganic nitrogen and O(¹D).

The update to the isoprene chemistry in CRI-Strat 2 increases OH over the lowest 500 m in tropical forested regions by 30%–50% relative to CRI-Strat, leading to an improvement in model–observation comparisons for surface OH and isoprene relative to CRI-Strat and StratTrop. Enhanced oxidants also cause a 25% reduction in isoprene burden and an increase in oxidation fluxes of isoprene and other biogenic volatile organic compounds (BVOCs) at low altitudes with likely impacts on subsequent aerosol formation, atmospheric lifetime, and climate.

By contrast, updates to the rate constants of O(¹D) with its main reactants relative to CRI-Strat reduces OH in much of the free troposphere, producing a 2% increase in the methane lifetime, and increases the tropospheric ozone burden by 8%, primarily from reduced loss via O(¹D) + H₂O.
The changes to inorganic nitrogen reaction rate constants increase the NO$_x$ burden by 4% and shift the distribution of nitrated species closer to that simulated by StratTrop.

CRI-Strat 2 is suitable for multi-decadal model integrations and the improved representation of isoprene chemistry provides an opportunity to explore the consequences of HO$_x$ recycling in the United Kingdom Earth System Model (UKESM1). This new mechanism will enable a re-evaluation of the impact of BVOCs on the chemical composition of the atmosphere and further probe the feedback between the biosphere and the climate.

1 Introduction

Isoprene (2-methyl-1,3-butadiene) makes up 70% of all non-methane biogenic volatile organic compound (BVOC) emissions, with annual average emissions of $594 \pm 34$ Tg C yr$^{-1}$ over the period 1980–2010 (Sindelarova et al., 2014). Isoprene’s rapid chemical oxidation in the atmosphere by OH, O$_3$, and NO$_3$ directly affects the tropospheric oxidizing capacity, ozone burden, and the processing of other trace gases like methane (e.g. Archibald et al., 2011; Khan et al., 2020), while also serving as an important source of secondary organic aerosol (SOA) (e.g. Scott et al., 2014; Kelly et al., 2018; Claeyts and Maenhaut, 2021). Thus, isoprene has substantial effects on the radiative balance of the atmosphere, both directly via production of SOA and ozone and indirectly via its changes to the oxidizing capacity of the atmosphere, influencing methane lifetime and production of other aerosol species, such as from oxidation of monoterpenes and SO$_2$ (Unger et al., 2014; Makonnen et al., 2012; Sporre et al., 2020). An accurate representation of isoprene chemistry in climate models is essential to understanding the feedbacks between the biosphere and the rest of the Earth system and thus capturing isoprene’s climatic impact.

However, the treatment of isoprene in the chemistry schemes of many climate models is outdated or oversimplified (e.g. Squire et al., 2015). The last decade has seen significant advances in our understanding of the isoprene oxidation pathway, principally the concept of rapid, intramolecular hydrogen shifts (H-shifts), also termed isomerization reactions, in the isoprene hydroxy peroxy radicals (frequently termed ISOPOO). Predictions from theoretical work (Peeters et al., 2009; Peeters et al., 2014) and observations (Crain et al., 2011; Teng et al., 2017; Wennberg et al., 2018) have established this pathway to be competitive with the traditional bimolecular reactions of the peroxy radical with NO, NO$_3$, HO$_2$, and RO$_2$ in certain conditions such as low NO$_x$ (= NO+NO$_2$) environments. These H-shift reactions lead to the production of HO$_x$ (= OH+HO$_2$) either directly or indirectly following the degradation of the isomerization products (e.g. Archibald et al., 2011; Jenkin et al., 2015; Wennberg et al., 2018).

This process, termed HO$_x$ recycling, has been shown to be important for low-NO$_x$, high-isoprene regions of the atmosphere (Butler et al., 2008; Lelieveld et al., 2008). By adding a simple, fixed-yield OH production pathway from ISOPOO to represent OH production from hydroperoxy aldehydes (HPALDs), Archibald et al. (2011) simulated an 8%–18% increase in tropospheric O$_3$ burden, while the tropospheric OH burden increased by 17% in the present day (PD) and by 50% in a pre-industrial (PI) atmosphere featuring 1860 emissions of key chemical species such as NO$_x$, CO, and isoprene. Consequently, the lifetime of methane was predicted to decrease by between 11% (in a future climate scenario) and 35% (in the PI). This illustrated the significant impact that such a process could have on our understanding of the PI atmosphere (and the radiatively active components therein) and thus the PD–PI change and climate sensitivity. While the greatest change to the chemistry was simulated in the boundary layer (BL), convection of isoprene and its oxidation products into the free troposphere resulted in this added chemistry having global impacts.

The effect on oxidants from HO$_x$ recycling influences the lifetimes of isoprene and other BVOCs such as monoterpenes and thus the extent of their dispersion and the location of the subsequent SOA formation. Karset et al. (2018) found that when lower oxidant fields were applied to the PI atmosphere, isoprene, monoterpenes, SO$_2$, and other key aerosol precursors were more dispersed from their sources, reaching higher altitudes and enhancing particle number concentration in the remote free troposphere. The radiative impact of the resulting aerosols was greater due to their enhanced lifetime (from slower deposition) and the highly non-linear relationship between aerosol number and cloud forcing, where the addition of a given concentration of aerosol has a much greater impact in remote regions where the background concentration of aerosol is smaller (Chen et al., 2016). The importance of oxidants to BVOCs and aerosol was also shown in Sporre et al. (2020) where models with an interactive oxidant scheme simulated a BVOC-driven depletion of oxidants and attendant greater dispersion of BVOCs and their oxidation products (including SOA precursors). In contrast, a prescribed oxidant approach saw BVOC oxidation confined far more to source regions, reducing dispersion.

Changes in oxidant fields also perturb the oxidation pathways of SO$_2$. In the United Kingdom Chemistry and Aerosols (UKCA) model, SO$_2$ can be oxidized in the gas phase by OH (to yield H$_2$SO$_4$) or in the aqueous phase by O$_3$ or H$_2$O$_2$ (Mulcahy et al., 2020). This has consequences for the aerosol mass and number distributions because only H$_2$SO$_4$ can nucleate new particles in UKCA; therefore, amplifying the gas-phase pathway over the aqueous pathways leads to a greater number of smaller aerosols. Thus, uneven changes to these pathways can alter the size and number distribution of the aerosol population, affecting the radiative properties of aerosols and clouds. Decreases in OH in other UKCA studies (Weber et al., 2020a; O’Connor et al.,
have resulted in simulated reductions in particle number concentration and cloud droplet number concentration. The resulting negative cloud radiative forcing is smaller in magnitude, as the lower cloud droplet number concentration (CDNC) makes the clouds less “bright” (Twomey, 1974). The impact of different oxidant schemes on the burden and lifetime of dimethylsulfide (DMS), an important SO2 precursor, and the impact on sulfate aerosol transport are both highlighted by Mulcahy et al. (2020).

While Archibald et al. (2011) used a relatively simple approach to simulate HOx recycling, further advances in the chemical understanding have led to a near-explicit representation of HOx recycling being incorporated into comprehensive mechanisms, including the Master Chemical Mechanism (MCM v3.3.1) (Jenkin et al., 2015) and the CalTech isoprene scheme (Wennberg et al., 2018). However, such mechanisms are far too large for use in global chemistry–climate models.

There exist a few reduced mechanisms featuring this state-of-the-art isoprene chemistry suitable for use in chemistry–climate models, including the CalTech reduced isoprene scheme (Bates et al., 2019), the MAGRITTE v1.1 model (Müller et al., 2019), the Mainz Organic Mechanism (Sander et al., 2019), the updated ECHAM-MESSy (Novelli et al., 2020), and the Common Representative Intermediates mechanism v2.2 (CRI v2.2) (Jenkin et al., 2019a), the latter of which is the focus of this work. The CRI v2.2 is an update to the Common Representative Intermediate v2.1 mechanism (Jenkin et al., 2008; Utembe et al., 2010; Watson et al., 2008) and was developed from the fully explicit Master Chemical Mechanism (MCM) version 3.3.1 (Jenkin et al., 2015), which describes the degradation of organic compounds in the troposphere. In the CRI framework, species are lumped together into surrogate molecules whose reactivity is optimized against the fully explicit MCM. A description of CRI v2.2 is given in Jenkin et al. (2019a). The CRI v2.1, along with the corresponding stratospheric chemistry, has already been incorporated into UKCA as CRI-Start (CS) (Archer-Nicholls et al., 2021) as an alternative to the simpler but more widely used STRAT-TROP (ST) chemistry scheme (Archibald et al., 2020a), the scheme used for UKESM’s contributions to CMIP6 (e.g. Sellar et al., 2020; Thornhill et al., 2021).

Using the reduced Caltech Isoprene Mechanism, which includes H-shifts of ISOPOO in GEOS-CHEM, Bates et al. (2019) simulated significant increases in OH (> 100 %) and HO2 (up to 50 %) over the Amazon and other forested tropical regions as a result of the HOx recycling. After implementing updated rate constants for isoprene H-shifts in GEOS-CHEM, Møller et al. (2019) also found that globally around 30 % of all isoprene peroxy radicals undergo at least one H-shift reaction, resulting in an OH yield of 47 % per isoprene molecule, and that adding all isoprene H-shift reactions increased boundary layer OH by up to a factor of 3 in the Amazon. Using CESM and CAM-CHEM and the MOZART-TS2 mechanism, Schwantes et al. (2020) showed reasonable agreement for some isoprene oxidation products over the southeastern USA.

Jenkin et al. (2019a), using CRI v2.2 in the STOCHEM Lagrangian chemistry transport model, showed the significant influence of HOx recycling in CRI v2.2 simulating a 6.4 % increase in the tropospheric OH burden relative to the CRI v2.1 and increases in surface OH of 20 %–50 % over much of the forested tropical regions. Khan et al. (2020), using the same setup, also simulated enhanced surface OH and attendant decreases in methane lifetime (0.5 years) and isoprene burden (17 %).

However, while the reduced mechanisms featuring HOx-recycling chemistry have been tested in chemistry–climate models, less work has been done in terms of multi-species comparison to observations and detailed analysis of the effect to global atmospheric composition. This study introduces the CS2, based on CRIv2.2 and expanded with stratospheric chemistry, as a mechanism in UKCA, evaluates its performance against observational data, and compares its output and key processes to the related CS mechanism and the well-established ST mechanism. By providing a wide-ranging comparison to observations and a detailed description of the changes CS2 causes in global and regional atmospheric chemistry, this current work builds on the existing literature to further develop our understanding of the consequences of HOx recycling.

2 Development of CS2 – incorporation of CRI v2.2 into UKCA

It is important to note that the CRI v2.2 mechanism, like the CRI v2.1 mechanism, is strictly a tropospheric chemistry scheme. In developing the whole atmosphere mechanism CS, Archer-Nicholls et al. (2021) merged the CRI v2.1 mechanism with the stratospheric chemistry scheme (Morgenstern et al., 2009) in UKCA (Table 1) to allow this scheme to be used within UKESM1 (Sellar et al., 2019). The same approach was taken in this work with the stratospheric scheme unchanged and tropospheric scheme switched from CRI v2.1 to CRI v2.2. Therefore, to differentiate the “CRI v2.2” mechanism used in UKCA in this work from the solely tropospheric CRI v2.2 mechanism described on the CRI v2.2 website (http://cri.york.ac.uk/, last access: 10 July 2021), the UKCA mechanism will henceforth be referred to as CRI-Strat 2 (CS2) (Table 1). A full description of the changes made to CS to update it to CS2 is given Sect. 1.1 in the Supplement, and a summary of the changes will now be discussed.

CS2 features a significant update to isoprene oxidation chemistry relative to CS with the incorporation of 1.6 and 1.4 H-shift reactions of isoprene peroxy radicals and an update to the organonitrate scheme (as detailed in Jenkin et al., 2019a). To the best of our understanding, CS2 also features updates to multiple reaction rate constants (which were out
Table 1. Comparison of the CRI-STRAT and CRI-STRAT 2 chemical mechanisms.

<table>
<thead>
<tr>
<th></th>
<th>CRI-STRAT (CS)</th>
<th>CRI-STRAT 2 (CS2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tropospheric chemistry scheme</td>
<td>CRI v2.1 (Jenkin et al., 2008; Watson et al., 2008; Utembe et al., 2010)</td>
<td>CRI v2.2 (Jenkin et al., 2019a)</td>
</tr>
<tr>
<td>Stratospheric chemistry scheme</td>
<td>Stratospheric chemistry (Morgenstern et al., 2009; Archibald et al., 2020)</td>
<td>Stratospheric chemistry (Morgenstern et al., 2009; Archibald et al., 2020)</td>
</tr>
<tr>
<td>No. of species</td>
<td>219</td>
<td>228</td>
</tr>
<tr>
<td>No. of bimolecular reactions</td>
<td>536</td>
<td>582</td>
</tr>
<tr>
<td>No. of termolecular reactions</td>
<td>36</td>
<td>44</td>
</tr>
<tr>
<td>No. of photolysis reactions</td>
<td>128</td>
<td>140</td>
</tr>
</tbody>
</table>

Table 2. Species added and removed from the CS mechanism in the development of the CRI-Strat 2 mechanism.

<table>
<thead>
<tr>
<th>Added species</th>
<th>Species functionality</th>
<th>MCM v3.3.1 equivalent</th>
</tr>
</thead>
<tbody>
<tr>
<td>HPUCARB12</td>
<td>Hydroperoxy aldehyde (HPALD)</td>
<td>C5HPALD1, C5HPALD2</td>
</tr>
<tr>
<td>HUCARB9</td>
<td>Unsaturated hydroxy carbonyl</td>
<td>HMVK, HMAC</td>
</tr>
<tr>
<td>IEPOX</td>
<td>Isoprene epoxide diol</td>
<td>IEPOXA, IEPOXB, IEPOX</td>
</tr>
<tr>
<td>HMML</td>
<td>Hydroxymethyl-methyl-a-lactone</td>
<td>HMML</td>
</tr>
<tr>
<td>DHCPCARB9</td>
<td>Dihydroperoxy carbonyl</td>
<td>DHPMEK, DHPMAL</td>
</tr>
<tr>
<td>DHP12OOH</td>
<td>Trihydroperoxy carbonyl</td>
<td>C536OOH</td>
</tr>
<tr>
<td>DHCARB9</td>
<td>Dihydroxy carbonyl</td>
<td>HO12CO3C4</td>
</tr>
<tr>
<td>RU12NO3</td>
<td>Hydroxy carbonyl nitrate</td>
<td>C57NO3, C58NO3, C58ANO3</td>
</tr>
<tr>
<td>RU10NO3</td>
<td>Hydroxy carbonyl nitrate</td>
<td>MVKNO3, MACRNO3</td>
</tr>
<tr>
<td>DHPR12O2</td>
<td>Dihydroperoxy carbonyl peroxy radical</td>
<td>C536O2, C537O2</td>
</tr>
<tr>
<td>MACO3</td>
<td>Unsaturated acyl peroxy radical</td>
<td>MACO3</td>
</tr>
<tr>
<td>RU10AO2</td>
<td>Hydroxy carbonyl peroxy radical</td>
<td>MACRO2</td>
</tr>
<tr>
<td>Removed species</td>
<td></td>
<td>MCM v3.2 equivalent</td>
</tr>
<tr>
<td>RU12PAN</td>
<td>PAN-type species with at least one hydroxy group</td>
<td>CSPAN19</td>
</tr>
<tr>
<td>TNCARB11</td>
<td>Alkyl carbonyl</td>
<td>N/A</td>
</tr>
<tr>
<td>TNCARB12</td>
<td>Alkyl carbonyl</td>
<td>N/A</td>
</tr>
</tbody>
</table>

of date in CS, Archer-Nicholls et al., 2021) as documented in the IUPAC Task Group on Atmospheric Chemical Kinetic Data Evaluation (http://iupac.pole-ether.fr/, last access: 9 April 2021). Changes were implemented to the rate constants of the reactions of O(1D) with H2O, O2, and N2; rate constants of multiple inorganic nitrogen reactions such as those forming species with the peroxy acyl nitrate moiety, termed PAN-type, and HONO; and the HO2 + NO reaction and the rate constants of organic peroxy radicals (RO2) with NO and NO3. These updates ensure consistency between the CS2 mechanism incorporated in UKCA and that described on the CRI v2.2 website (http://cri.york.ac.uk/, last access: 10 July 2021). The photolysis of glyoxal, formaldehyde, and propionaldehyde was also updated (see Sect. S6 in the Supplement).

CS2 has 9 more species than CS (Tables 1, 2) as well as 46 additional bimolecular reactions, 12 additional photolysis reactions, and 8 additional unimolecular and termolecular reactions (Table 1). This leads to a modest increase in runtime (6%) compared with CS, whose runtime was already ~75% greater than ST. Incorporation of CS2 into UKCA involved extensive use of the UM-UKCA virtual machine environment (Abraham et al., 2018).

The main update to the isoprene chemistry is the inclusion of 1,6 and 1,4 H-shift reactions of the isoprene peroxy radical (termed RU14O2 in CRI nomenclature). The 1,6 H-shift process is well studied (Peeters et al., 2009; Crounse et al., 2011; Teng et al., 2017; Wennberg et al., 2018) and follows the $k_{bulk,1,6H}$ rate coefficient described in Jenkin et al. (2019a), capturing the dependence of isomerization on both temperature and the rates of reaction of RU14O2 with the standard bimolecular partners (NO, NO3, HO2, and RO2). This pathway yields hydroperoxy aldehydes (HPALDs, termed HPU-CARB12 in CS2) and dihydroperoxy carbonyls peroxy radicals (DPHR12O2). The photolysis of the highly photolabile HPALD (HPUCARB12) and its product HUCARB9 (unsaturated hydroxy carbonyl) are key routes for HOx regeneration.

The production of the isoprene epoxide diol (IEPOX) from the isoprene hydroperoxide (RU14OOH) and the hydroxymethyl-methyl-a-lactone (HMML) also represent important updates (Jenkin et al., 2019a). IEPOX and HMML are known SOA precursors (Nguyen et al., 2014, 2017; Allan et al., 2014), and thus their addition may enable a more ex-
explicit representation of SOA formation within the CRI framework, as opposed to the current framework whereby SOA formation is represented by the condensation on existing aerosol of a single inert tracer, Sec_Org, which is made from monoterpene oxidation (Mann et al., 2010; Mulcahy et al., 2020). This is beyond the scope of this paper but will be a focus of future work.

The introduction of HPUCARB12 and HUCARB9 necessitates a careful update to the FASTJX photolysis scheme used by UKCA (Telford et al., 2013). The cross-sectional dependence of wavelength for HPALDs is assumed to be the same as methacrolein (Peeters et al., 2009; Wennberg et al., 2018; Schwantes et al., 2020) but with a significantly larger quantum yield (QY). Prather (2015) recommends a QY of 0.003 for methacrolein, and Liu et al. (2017) recommends a QY of 0.55 for HPALDs (both used by Wennberg et al., 2018). To implement the photolysis of these new species, the photolysis frequency of HPUCARB12 was taken to be the photolysis frequency for methacrolein scaled by the ratio of the QY of HPALDs to the QY of methacrolein, the same approach used by Schwantes et al. (2020) for 6 HPALDs. A scaling of 0.5 was applied to the photolysis frequency of HUCARB9 in agreement with the MCM v3.3.1.

In addition to the updates to isoprene chemistry, CRIv2.2 has had the rate coefficients for many organic and inorganic reactions updated to bring the mechanism into agreement with the MCM v3.3.1 and IUPAC. These affect the overall chemistry in three major ways. The first involves the major reactions of the excited oxygen radical, O(1D). The rate constants of O(1D) with H2O, O2, and N2 set to values in CS1, decreased by 4%–15% respectively, to bring them into agreement with the current IUPAC values (http://iupac.pole-ether.fr, last access: 9 April 2021). This also means the rate constant of O(1D) with N2 became much closer (within ±1.5%) to that used in ST (Archibald et al., 2020a) and that rate constants for the reactions with O, O, ONO, and MeONO + OH set to values in CS1 also move closer to those used in ST. The result of this is a reduction in the fraction of O(1D) reacting with H2O by 10%–15%, thus lowering OH production while also reducing O3 loss via this pathway.

The second involves multiple inorganic reactions of nitrate species. The formation rate constants for PAN-type species (species with peroxyacyl nitrate functionality), HONO2, H2O2NO2, and N2O5 changed by around −45%, −15%, −45%, and +50%–75% in the troposphere, respectively. The change for PAN brought its formation rate constant much closer to that used in ST (within ±7%), and this was also the case for HONO2 and H2O2NO2 formation. The rate constant of HO2 + NO, the single biggest production source of O3, decreased by 4%.
Table 5. Location, reference, time period, and species measured in observational data sets and the corresponding modelling approach. For the ZF2 Brazil, ATTO, Borneo, GABRIEL, FAAM and SE4CRS data sets, model data was filtered to select only the same days as observational data.

<table>
<thead>
<tr>
<th>Dataset (location/coordinates)</th>
<th>Reference</th>
<th>Dates of measurement</th>
<th>Measurement details</th>
<th>Species considered</th>
<th>Corresponding model run (Table 3 unless stated)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ZF2 Brazil field campaign, Amazon (−2.60°, −60.21°, 60 km NNW of Manaus)</td>
<td>See Sect. S4 in the Supplement</td>
<td>22 June 2016–5 July 2016</td>
<td>1 min interval measurements at 30 m above ground (above tree canopy)</td>
<td>O₃, CO, SO₂, NO₂, isoprene, monoterpenes, benzene</td>
<td>ZF2 Brazil</td>
</tr>
<tr>
<td>Instant ATTO tower, Amazon (−2.14°, −59.00°, 150 km NE of Manaus)</td>
<td>Yáñez-Serrano et al. (2015)</td>
<td>February 2013, September 2013, and February 2014</td>
<td>16 min interval measurements at multiple heights above ground (0.05, 0.5, 4, 12, 25, 38, 53 and 79 m)</td>
<td>Isoprene, monoterpenes, methyl vinyl ketone (MVK), methacrolein (MACR), isoprene hydroperoxide (ISOPOOH), acetone (All PTRMS)</td>
<td>ATTO</td>
</tr>
<tr>
<td>GAW station, Borneo (5.0°, 117.5°)</td>
<td>Hewitt et al. (2010), Whalley et al. (2011), Edwards et al. (2013)</td>
<td>April–July 2008</td>
<td>10 min intervals</td>
<td>OH, HO₂ (both FAGE), O₃ (Thermo Environmental Instruments 49C UV absorption instrument) isoprene, monoterpane (both PTRMS), HCHO (Aerolaser Hantzsch), CO (Aerolaser AL5002), MeCHO, acetone MACR, MVK (both GC-FID), PAN (GC-MS), NO₂ (Thermo environmental instruments 42C)</td>
<td>Borneo</td>
</tr>
<tr>
<td>GABRIEL aircraft campaign (Suriname, Guyana, and French Guiana)</td>
<td>Butler et al. (2008)</td>
<td>October 2005</td>
<td>Daytime aircraft measurements sampling ∼0.3–8 km at 30 s intervals</td>
<td>O₃, NO (both ECOEX), HCHO, CO (both MPIC TRISTAR), acetone, isoprene, MACR, MVK (all PTRMS)</td>
<td>GABRIEL</td>
</tr>
<tr>
<td>FAAM aircraft campaign, Borneo</td>
<td>Hewitt et al. (2010)</td>
<td>July 2008</td>
<td>Daytime aircraft measurements sampling ∼0.3–7 km at 5 min intervals</td>
<td>O₃ (TECO 49), isoprene (PTRMS), CO (AERO AL5002)</td>
<td>FAAM</td>
</tr>
<tr>
<td>SE4CRS flight campaign (Southeast US)</td>
<td>Toon et al. (2016)</td>
<td>August–September 2013</td>
<td>Daytime aircraft measurements sampling up to 12 km at 1 min intervals</td>
<td>O₃ (ERSL), CO (DACOM), Isoprene (WAS), ISOPOOH, HPALDs, IEPOX, isoprene nitrate (all CIT)</td>
<td>SE4CRS</td>
</tr>
<tr>
<td>Global isoprene columns</td>
<td>Wells et al. (2020)</td>
<td>January, April, July, and October 2013</td>
<td>Global monthly mean isoprene column values</td>
<td>Isoprene</td>
<td>Isoprene column</td>
</tr>
</tbody>
</table>

Finally, the rate constants for most RO₂ + NO and RO₂ + NO₃ reactions have been changed by +12.5 % and −8 %, respectively, while maintaining the same temperature dependence. This is likely to have a smaller impact than the other chemistry changes but at the margins will make reactions with NO more competitive with the isomerization reactions of the ISOPOO.

The implementation of CRI v2.2 by Khan et al. (2020) in the STOCHEM model, while including the updates to isoprene chemistry and the RO₂ + NO and RO₂ + NO₃ reactions, did not feature updates to the rate constants for O(¹D) with H₂O, O₂, and N₂ or the inorganic nitrogen reactions. Therefore, even in low-altitude terrestrial conditions where isoprene HOₓ recycling tends to dominate the change in OH, comparison between Khan et al. (2020) and the results of this work must be caveated with the changes to the inorganic chemistry.

In addition to the chemistry changes, updates are made to the photolysis of several species. Two additional photolysis reactions of glyoxal (CARB3 in the CRI mechanisms) were added in addition to updates to the photolysis parameters for HCHO and EtCHO (propionaldehyde). The wavelength bins of the product of the cross section and quantum yield used by FAST-JX (Telford et al., 2013) were updated to the v7.3 values from Prather (2015) for HCHO and EtCHO. The photolysis of CARB3, which had previously been estimated in...
CS by a scaling of HCHO photolysis (Archer-Nicholls et al., 2021), is replaced with the glyoxal photolysis for 999 hPa from v7.3 of Prather (2015). This reaction does exhibit a modest pressure dependence, but this has not been incorporated into FAST-JX at the current time.

In addition to the changes to the chemistry and photolysis, updates to the wet deposition scheme were implemented to both CS and CS2 schemes. The previous approach of applying parameters for a standard surrogate for other species with the same functional groups (e.g., EtOOH was used for most hydroperoxides), as described in Archer-Nicholls et al. (2021), was updated to use either data for the precise species (taken from Schwantes et al., 2020) or a more closely related surrogate. The changes to the wet deposition parameters are detailed in Table S1 in the Supplement. As they were applied to both CS and CS2 mechanisms, they are unlikely to have a significant influence on the inter-mechanism difference. No changes were made to the dry deposition scheme in this work.

3 Model runs

All model runs were performed using the United Kingdom Chemistry and Aerosols Model (UKCA), run at a horizontal resolution of $1.25^\circ \times 1.875^\circ$ with 85 vertical levels up to 85 km (Walters et al., 2019), and the GLOMAP-mode aerosol scheme, which simulates sulfate, sea salt, BC, organic matter, and dust but does not simulate currently nitrate aerosol (Mulcahy et al., 2020). In this setup, the inert chemical tracer Sec_Org, which condenses irreversibly onto existing aerosol, was produced at a 26% yield solely from reactions of $\alpha$-pinene and $\beta$-pinene with O$_3$, OH, and NO$_3$ with the enhanced yield applied to account for a lack of SOA formation from isoprene or anthropogenic species (Mulcahy et al., 2020).

The runs in this work fell into two distinct categories. Firstly, short runs (generally 1–2 months, Table 3) with higher-frequency (hourly) output using the ST, CS, and CS2 chemical mechanisms were performed to evaluate each mechanism’s performance against the observational data. Secondly, longer runs (2–5 years, Table 4) with monthly output using the CS and CS2 chemical mechanisms (or variants of CS2 for sensitivity tests) were conducted to facilitate a rigorous comparison of the global chemical composition (Table 4).

Temperature and horizontal wind fields were nudged (Telford et al., 2013) in all model runs to atmospheric re-analyses from ECMWF (Dee et al., 2011) to constrain the simulations to consistent meteorology, thus preventing diverging meteorology from adding to the differences resulting from the chemical mechanisms and replicating the atmospheric conditions experienced when the observations were recorded as closely as possible. Nudging only occurred above $\sim$1200 m in altitude, and thus the majority of the planetary boundary layer was not nudged. The model runs were atmosphere-only runs with prescribed sea surface temperatures (SSTs). CO$_2$ is not emitted but set to a constant field, while methane, CFCs, and N$_2$O are prescribed with constant lower boundary conditions, all at 2014 levels (Archibald et al., 2020a).

The emissions used in this study are the same as those from Archer-Nicholls et al. (2021) and are those developed for the Coupled-Model Intercomparison Project 6 (CMIP6) (Collins et al., 2017). Anthropogenic and biomass burning emissions data for CMIP6 are from the Community Emissions Data System (CEDS), as described by Hoesly et al. (2018). For the short runs, time series anthropogenic and biomass burning emissions were used for all ST runs and all CRI runs up to 2015. For the runs done for the purpose of comparison to the observational data recorded at the ZF2 site near Manaus in 2016 (see Tables 3, 5), time slice 2014 emissions were used due to a lack of post-2015 CRI emissions, but the impact of the difference is expected to be minimal.

All longer runs used time slice 2014 emissions for anthropogenic and biomass burning emissions. Oceanic emissions were from the POET 1990 data set (Olivier et al., 2003), and all biogenic emissions except isoprene and monoterpenes (see Sect. 3.3) were based on 2001–2010 climatologies from Model of Emissions of Gases and Aerosols from Nature under the Monitoring Atmospheric Composition and Climate project (MEGAN-MACC) (MEGAN) version 2.1 (Guenther et al., 2012) and are discussed further in Sect. 3.3. A full description of the emission sources for each emitted species is given in Table S2 in the Supplement.

All mechanisms used the same raw emissions data. However, the additional emitted species required by CS and CS2 means the total mass of emitted organic compounds is greater in CS and CS2, and the lumping of species for emissions is also different. The approach and consequences are discussed in Archer-Nicholls et al. (2021).

3.1 Short runs for model–observation comparisons

The runs performed for comparison to observations are detailed in Table 3 and correspond to an observational data set described in Sect. 4 and Table 5. All runs were spun up for a minimum of 3 months. For most of the runs, hourly model output was used so as to allow for detailed comparison with observations. The only exceptions were the runs performed for the comparison to the isoprene column data (“isoprene column” Table 3) for which monthly means were used.

3.2 Longer runs for mechanistic intercomparison

The longer runs (Table 4) were designed with the primary aim of examining the consequences of the mechanism changes between CS and CS2 and followed an approach similar to that used by Archer-Nicholls et al. (2021). These runs also served a secondary purpose as they enabled longer-term...
comparisons to observations for several species. We ran two 5-year nudged runs (1-year spin up, 4-year analysis) with the CS and CS2 mechanisms. In addition, five 2-year sensitivity runs (1-year spin up, 1-year analysis) were performed to analyse the impact of the individual changes to the isoprene scheme, O(1D) reactions, inorganic nitrogen reactions, RO2 + NO/NO2 reactions, and photolysis reactions, as discussed in Sect. 2. These sensitivity tests featured mechanisms based on the CS2 mechanism, but each had a different feature that was reverted to that found in CS.

CS2_O1D used the old rate constants from CS for the reaction of O(1D) with N2, O2, and H2O. CS2_inorgN used the rate constants from CS for the formation of HONO2, HO2NO2, PANs, HONO, and N2O4, as well as for the reactions of HO2 + NO, OH + MeONO2, OH + PAN, and OH + MPAN.

CS2_isoprene followed the isoprene reactions from CS as closely as possible, with the major change being the omission of the isomerization reactions of RU14O2 and subsequent production of HPALDs and other species that are key for HOx recycling.

In CS2_RO2_N, the rate constants for the RO2 + NO and RO2 + NO3 reactions were reverted to those used in CS, which led to a 12.5% decrease and 8% increase, respectively, for the vast majority of these reactions. Where branching ratios changed between CS and CS2, the CS2 branching ratios were maintained and the rate constants scaled accordingly.

Finally, CS2_photo used the parameters and reactions from CS for the photolysis of CARB3 (glyoxal), HCHO, and EtCHO and was performed to evaluate the impact of update to photolysis (see Sect. S6 in the Supplement).

Each sensitivity test, when compared to the CS2 run, provides information as to the impact of the change of the respective section of the mechanism (when taken in isolation); for example, the impact of the changes to the rate coefficients of O(1D)’s reactions is examined by comparing the CS2 and CS2_O1D runs.

A full description of the changes to reactions and rate constants for each sensitivity test is given Sect. S2 in the Supplement. The changes to the photolysis were found to have a minimal effect on atmospheric composition compared with the other sensitivity tests and are described in their entirety in the Supplement. The analysis of the longer runs is discussed in Sect. 5.

3.3 Biogenic emissions

This work used the interactive biogenic volatile organic compound (iBVOC) emissions system (Pacifico et al. 2012) for isoprene and monoterpenes, which is the standard approach for UKESM’s contributions to CMIP6 (Sellar et al., 2019). Emissions of isoprene and monoterpenes are calculated interactively based on temperature, photosynthetically active radiation (PAR), and plant functional type for each grid cell. While a diel cycle for isoprene is standard in UKESM, iBVOC has the advantage of also simulating a diel cycle of emissions for monoterpenes, leading to improved model performance relative to observation (see Sect. 4). The dependence on temperature and PAR means that emissions of BVOCs differ slightly between runs and thus between mechanisms. However, nudging considerably inhibits divergence of surface temperature between comparative runs, and thus the differences between emissions were < 5% and typically 1%–2%, significantly smaller than the differences caused by the mechanisms.

Monoterpane emissions were speciated in a 2 : 1 α-pinene : β-pinene ratio as used in Archer-Nicholls et al. (2021).

There are temporal and spatial disparities between using iBVOC emissions and offline emissions, such as the MEGAN-MACC data set (Sindelarova et al., 2014, as used by Archer-Nicholls et al., 2021), which could affect conclusions about mechanism observational biases. These differences are discussed in more detail in Sect. S3 in the Supplement. In short, for the ZF2 Brazil, ATTO, and Borneo sites for the periods considered, the isoprene and monoterpene (MT) emissions were higher when using the iBVOC approach than for MEGAN-MACC (Figs. S1 and S2 in the Supplement).

4 Comparison with observations

The shorter UKCA models runs listed in Table 3 were used to evaluate mechanism performance against six high frequency observational data sets (three surface or near-surface data sets and three aircraft campaigns) from the Amazon, Borneo, and the southeastern USA, all important regions for BVOC production. In addition, satellite-derived isoprene columns (Wells et al., 2020) were compared to model output (Isoprene Column, Table 3). Monthly mean data from the longer CS and CS2 runs (Table 4) for O3, CO, and HONO2 were also compared to a range of observational data. A summary of the observation data sets is given in Table 5, and locations of the surface and airborne campaigns are shown in Fig. S3 in the Supplement.

Diel profiles for multiple species were calculated from the three surface and near-surface sites, and the vertical profiles were calculated from the ATTO site.

The three flight campaigns considered were the October 2005 Amazon GABRIEL campaign (Butler et al., 2008), the July 2008 Borneo Facility for Airborne Atmospheric Measurements (FAAM) (Hewitt et al., 2010), and the Studies of Emissions and Atmospheric Composition, Clouds and Climate Coupling by Regional Surveys (SEAC4RS) flight campaign over the southeastern USA in August–September 2013 (Toon et al., 2016). Hourly model output corresponding to the days and times of the flights was used for the mechanism–observation comparison for each campaign. Model and ob-
observational data were binned into 250 m/500 m altitude bins and median values for the variables of interest across the whole region for a given altitude bin were considered. For the SEAC4RS comparison, observational data were also filtered to exclude urban plumes (NO$_2$ > 4 ppb), fire plumes (acetonitrile > 0.2 ppb), and stratospheric air (O$_3$/CO > 1.25), while missing data were not used, and data flagged as a lower limit of detection were set to zero as previously done in Schwantes et al. (2020). Estimated limits of detection are shown for relevant species for the GABRIEL and FAAM campaigns. The performance of each mechanism is now described for the key species, e.g. O$_3$, HO$_x$, isoprene, certain isoprene oxidation products, and monoterpene. A brief commentary about other species including HONO$_2$, CO, PAN, HCHO, MeCHO, EtCHO, and acetone is given in the Supplement.

4.1 Ozone

CS2 exhibits a modest increase in O$_3$ (∼1–4 ppb) over CS at all surface sites (Fig. 1), exacerbating the existing high surface bias of CS, whose drivers were discussed in Archer-Nicholls et al. (2021), and the smaller high bias of ST. On a diel basis, the mechanisms are able to replicate the shape of the diel cycle at the ZF2 site (with similar diel profiles at the ATTO site) but perform less well in Borneo, simulating pronounced diel cycles with a high bias compared to much more muted cycles from observation.

An increase of ∼1–4 ppb relative to CS is also exhibited by CS2 for monthly mean O$_3$ when both mechanisms are compared to observational data at 10 locations from pole to pole at four pressure levels (250, 500, 750, and 900 hPa) (Fig. S4 in the Supplement). CS2 reduces the low bias in polar regions but increases the CS’s high bias in the tropics and eastern US.

Model high biases are also observed from flight data comparisons (Figs. 2b,f, and S6a in the Supplement). In the Amazon, where the observed and modelled NO vertical profiles agree well (Fig. S6e in the Supplement), there is little difference between the three mechanisms. Each exhibits the highest high bias at low and a smaller high bias in the free troposphere. CS2 exhibits a high bias of 15–20 ppb for the SEAC4RS campaign (Fig. S6d in the Supplement), with perhaps some influence from the low-altitude NO$_2$ model high bias. In Borneo, all mechanisms exhibit a roughly consistent high bias of ∼20 ppb for ST, increasing to 30 ppb for CS2. Interestingly, all the mechanisms simulate a significant low bias for NO$_2$ (Fig. S6f in the Supplement), which may indicate biomass burning events which are not simulated, something which might be expected to promote higher ozone concentrations.

4.2 HO$_x$

Modelled surface OH increases in all locations from ST through CS to CS2, with a significant increase in midday OH from CS to CS2 (Fig. 1). In Borneo, OH is consistently biased low in the three mechanisms, but the best comparison is exhibited by CS2 where the mean diel bias compared to ST and CS decreases by 43 %–50 % and 24 %–40 %, respectively, over the period considered. The drivers of the HO$_x$ change are explored further in Sect. 5.

Surface HO$_2$ was also simulated to increase in all locations from ST to CS to CS2. Significant high bias was simulated in Borneo (the only observational data set) (Fig. S7 in the Supplement) for the CRI mechanisms, including at night. The simulated ratio of HO$_2$ to OH is highly biased in all mechanisms. However, it is best simulated in CS2, indicating that the increase in OH is much larger than that for HO$_2$. It should be noted that none of the mechanisms at present include the heterogenous reactions of HO$_2$ and their inclusion, which will be addressed in future work, should reduce the HO$_2$ high bias.

The comparison of modelled HO$_x$ to observation is complicated by large discrepancies in key reaction partners. Relative to observed values of 100–130 ppb, CO in ST in Borneo is highly biased by 13 and 27 ppb, while CO in the CRI mechanisms exhibits larger biases of ∼35–50 and ∼50–60 ppb during April–May and June–July, respectively (Fig. S7). These high biases would enhance modelled HO$_2$ at the expense of OH, potentially explaining the modelled low biases in OH. Indeed, the OH model low bias is greater in the June–July period. This highlights the complexity of model–observation comparisons: the CRI mechanisms may well simulate secondary CO production from isoprene more accurately than ST but other model biases, for example in emissions of CO, NO, and isoprene, can lead to the CRI mechanisms appearing worse. Nevertheless, if the CO high bias is reduced in future, we might reasonably assume that the modelled OH will improve still further.

4.3 Isoprene

Modelled isoprene from all three mechanisms was compared to surface observations, flight campaign data, and isoprene columns measured by satellite.

4.3.1 Isoprene surface measurements

CS2 yields the best model–observation comparison for surface isoprene on a daily basis in all locations (Fig. 1k–o). CS2 reduces the high bias in the diel profiles by 50 %–60 % relative to ST and 20 %–40 % to CS at the ZF2, ATTO, and Borneo sites, driven by the elevated OH concentrations.

In most locations the model simulates, to a greater or less extent, a “twin peak” isoprene profile with a sharp rise around 07:00 LT and a second, smaller peak at 19:00 LT. This was
most pronounced in the Amazon dry season (ATTO Sept 2013). The morning peak is likely to be due to a combination of the sharp rise in simulated isoprene emissions which starts at 06:00–07:00 LT, outweighing the concurrent rise in OH, and an underestimation in the model of the rate of BL height growth, which can trap isoprene close to the surface, causing a buildup. By contrast, observed isoprene concentrations exhibit a much slower morning growth, reaching a peak in early afternoon. While the “out-of-phase” nature of the profiles is unlikely to be the sole driver of model–observation difference, it will play a role since isoprene chemistry occurs on the timescale of ~1–2 h and atmospheric oxidizing capacity varies throughout the day.

Over the lowest 80 m at the ATTO site, all mechanisms are biased high in the daytime (09:00–15:00 LT) and nighttime (21:00–03:00) and has larger diel variation than day (Fig. S8a–d in the Supplement). This is consistent with the smallest bias, but produce similar isoprene vertical gradients to observations. The effect of boundary layer height was further considered by looking separately at the periods 06:00–08:00 and 17:00–19:00 LT (Fig. S8e–h in the Supplement).
Supplement). In contrast to the daytime and nighttime periods, during the 06:00–08:00 LT period the simulated isoprene gradient is significantly more negative than the observation, indicating less vertical mixing, and similar results are seen with the MT profile (Fig. S8m–p in the Supplement). This is most noticeable in September where the largest morning peak is seen in the diel profile for both species and lends support to the theory that the simulated BL height is not increasing as quickly as in reality, leading to more isoprene and MT being trapped at low altitude. Smaller differences between observed and simulated isoprene and MT vertical gradients are seen during 17:00–19:00 LT, coinciding with smaller evening peaks in the diel profiles. This suggests the reduction in BL height is more accurately simulated than the morning increase.

The major drivers of the remaining model–observation difference are likely to be the concentrations of oxidants (despite the increases seen in CS2, OH remains low biased in Borneo) and the emissions of isoprene (including the modelled vs. actual diel cycle). The concentrations of isoprene and other species also vary significantly through and above the tree canopy, as shown by the ATTO measurements (Fig. S8), and the global model resolution is not high enough to resolve the vertical gradient of species in the canopy. When testing the CRI v2.2 in STOCHEM-CRI with isoprene emissions from the MEGAN-MACC inventory, Khan et al. (2020) noted that halving the isoprene emissions reduced the model–observation disagreement significantly and attributed the model high bias in their work to high biases in the emissions of isoprene.

4.3.2 Isoprene flight measurements

Model–observation comparisons of isoprene vertical profiles extending into the boundary layer and into the free tropo-
spherically reveal quite a different story from the surface analysis (Fig. 2a, e, and h).

Despite being biased high at the surface and at low altitude, simulated isoprene vertical profiles over the Amazon and Borneo rapidly show a low bias as altitude increases. There are likely two reasons for this. The first is the vertical mixing, already discussed in relation to the isoprene and MT surface diel cycles. Secondly, for the Amazon and Borneo campaigns only estimated detection limits (0.1 ppb in both cases) could be used. This has the effect of biasing the median of the observational data to higher values as very low values are ignored. In the SEAC4RS campaign, all data points flagged as below the detection limit were set to zero, mitigating this issue. The enhanced oxidative capacity of CS2 at low altitude results in the lowest simulated vertical concentrations among the three mechanisms, but the general low bias above the surface is an issue faced by all mechanisms, suggesting it is not just down to modelling of the chemistry.

4.3.3 Isoprene columns

To consider isoprene on a global scale, monthly modelled isoprene columns for all mechanisms are compared to satellite observations from January, April, July, and October 2013 (Wells et al., 2020) (Fig. 3).

Significant variation in model bias is exhibited between the mechanisms with ST exhibiting the highest isoprene columns and CS2 the lowest. In South America, CS2 exhibits the smallest bias, while the ST columns are over double the observed values for April and July. CS and CS2 exhibit the smallest biases in Africa and Southeast Asia, respectively. The low biases in North America (∼0.7–1.5×10^15 molec.cm^{-2}), Europe (∼0.5–2.7×10^15 molec.cm^{-2}), and Central Asia (∼0.1–1.1×10^15 molec.cm^{-2}) are quite consistent across the mechanisms and are in some cases almost equal in magnitude to the observed columns, which suggests the bias is driven more by insufficient emissions rather than the chemistry scheme in these locations.

CS and CS2 yield lower isoprene columns and generally smaller model biases than ST. This comparison highlights the significant influence of the different chemical schemes on the simulated isoprene column and thus the considerable challenges of determining isoprene emissions via top-down approaches using back-calculation from observed concentrations or column values: different chemistry schemes will lead to different emission estimates.

4.4 Isoprene oxidation products

During the GABRIEL flight campaign, the major isoprene oxidation products MACR and MVK were measured via PTRMS. At the ATTO tower, isoprene oxidation products were also measured via PTRMS but in this case were defined as the sum of MACR, MVK, and ISOPOOH (Yáñez-Serrano et al., 2015) and to avoid confusion we refer explicitly to the isoprene oxidation products as either MVK + MACR (for Gabriel) or MVK + MACR + ISOPOOH (ATTO). In each case, the observational data are compared with model data.

At the ATTO site, all mechanisms are largely biased high for MVK + MACR + ISOPOOH, but CS2 produces the best comparison to observations for both diel and vertical profiles (Fig. 1 and Figs. S9 and S11 in the Supplement). CS2 also yields the smallest high bias for the ratio of MVK + MACR + ISOPOOH to isoprene (a metric less sensitive to discrepancies between actual and modelled isoprene emissions) in the Amazon (Figs. 1, S9, 11). Despite the greater oxidizing capacity of the planetary boundary layer (PBL) in the CS2 simulations, the MVK + MACR + ISOPOOH concentrations are lower. This is attributed to the fact that in the relatively low-NOx environment around the ATTO tower, the isomerization reactions of the isoprene peroxy radical are particularly important and favour the production of HPALDs and other species over MACR, MVK, and ISOPOOH.

Relative to the GABRIEL flight data (Fig. 2d), the ratio of MVK + MACR to isoprene is biased high in all mechanisms, albeit with the CRI mechanisms exhibiting a smaller bias than ST.

4.5 Isoprene nitrate, IEPOX, and HPALDs

The isoprene oxidation products HPALDs and IEPOX, unique to the CS2 mechanism in this study, are compared, along with isoprene, ISOPOOH, and isoprene nitrate (Fig. S6), to observational data from the SEAC4RS campaign over the southeastern USA. Modelled isoprene (Fig. 2h) exhibits a significant low bias, in line with the isoprene column analysis (Fig. 3), and is attributed to insufficient emissions. Unsurprisingly, ISOPOOH (Fig. 2i), isoprene nitrate (Fig. S4c), and HPALDs (Fig. 2j) are also biased low. However, IEPOX (Fig. 2j) compares favourably to observations.

The apparent good performance of IEPOX, despite the significant low biases of isoprene and its direct precursor ISOPOOH, is likely to be due to a missing sink to the aerosol phase. IEPOX is readily lost to aerosol by reactive uptake (Nguyen et al., 2014, 2015; Allan et al., 2014); a process featured in Schwantes et al. (2020) (who simulated lower IEPOX concentrations) but not in UKCA. The rate constant for IEPOX’s production from ISOPOOH is ∼30 % lower than that used by a mechanism of similar complexity, MOZART TS2 (Schwantes et al., 2020), while IEPOX’s loss via OH has a similar rate constant to MOZART TS2. Including reactive uptake of IEPOX in future updates may reduce this high bias. The processing of IEPOX is unlikely to affect HOx recycling as much as HPALDs; however, its importance to SOA formation means it will be a focus of future work.

The low bias of HPALDs, also simulated to a lesser extent in Schwantes et al. (2020), who used isoprene emissions from the MEGAN v2.1, is important given its role in HOx re-
Figure 3. Monthly mean isoprene column values from the global isoprene column observational data set (Wells et al., 2020) for (a) January, (b) April, (c) July, and (d) October 2013. Model bias (model–observation) using (d–h) ST, (i–l) CS, and (m–p) CS2. Numbers in (a–d) show the area-weighted mean model column values and in (e–p) show the model bias for individual terrestrial regions (the number in the North Atlantic refers to Europe and that in the South Atlantic refers to Africa).

There remains uncertainty in HPALD photolysis frequencies. In this work simulated HPALD destruction is dominated by reaction with OH and photolysis, which are roughly equal ascending to 2.5 km, whereupon OH’s importance grows rapidly at the expense of photolysis. To test the impact of photolysis uncertainty on the bias, two further runs were performed with the photolysis frequency of HPALDs scaled by 0.5 and 3, respectively. These tests change HPALD concentrations in the lowest 2 km by $+30\%$ and $-50\%$ (Fig. 2k), respectively, suggesting concentration of HPALDs is dependent on the photolysis frequency of HPALDs, which is not currently well constrained.

Interestingly, these scaling tests only change low-altitude OH by $\sim 2\%–3\%$ in the southeastern USA, suggesting the uncertainty in HPALD photolysis from the current approach may not have a huge impact on oxidants in this region, although this may in part be due to the modelled isoprene and HPALD low biases (Fig. 2h, k). Furthermore, the fact that the modelled photolysis frequency of methacrolein here is low biased by a factor of 2.5–3 (not shown) suggests that if further changes to the HPALD frequency are made in future, any potential reductions in methacrolein frequency should be scrutinized carefully. Nevertheless, constraining HPALD photolysis further will be a key focus of future work. A lack of OH measurements prevents attempts to constrain the OH loss pathway.

Evaluating HPALD production is also challenging since observations of ISOPO2 were not measured. Over the relevant temperatures, the rate constant for HPALD production in CS2 is 6–14 times greater than the equivalent used by Schwantes et al. (2020), which would, if anything, make a low bias less likely. The sensitivity of HPALD production to the concentrations of the bimolecular reaction partners of ISOPO2 (e.g. NO) can also lead to resolution issues with the model: regions with high and low NO concentrations are treated as a single region within the model (model grids can be up to $\sim 125$ km wide at the Equator) with moderate [NO], suppressing HPALD formation (see Schwantes et al., 2020). A commentary on the global distribution of HPALDs and IEPOX is given in Sect. 5.

4.6 Monoterpenes (MT)

Simulated surface diel monoterpene profiles (Fig. 1) are characterized by early morning and evening peaks that are not
present in observations. As discussed in relation to the isoprene diel cycle, the morning peak is probably caused by a combination of the simulated emissions increasing too early and a delayed evolution of the simulated BL height, trapping large quantities of monoterpenes close to the surface (Fig. S8). The evening peak coincides with a reduction of simulated OH to near zero and therefore is probably driven by oxidant reduction and a reduction in the BL height. Around midday the mechanisms do a better job in most locations, with the lower values in the CRI mechanisms driven by the greater oxidant concentrations. In four of the five locations, CS2 yields the smallest model bias, although it is acknowledged that other issues, such as the BL dynamics, need attention.

5 Comparison to CRI-STRAT

The performance of the CS mechanism compared to the simpler ST mechanism was discussed in detail in Archer-Nicholls et al. (2021). Here we describe chemical composition of the atmosphere simulated by CS2 relative to that from CS using the longer model runs summarized in Table 2. Particular attention is paid to O$_3$ and its production and loss fluxes, HO$_x$, isoprene and monoterpenes, the isoprene oxidation productions IEPOX and HPALDs, nitrated species (NO$_y$), and the potential impacts to aerosols. Changes to CO and HCHO are discussed in Sect. S5 in the Supplement.

5.1 O$_x$

As in Archer-Nicholls et al. (2021), the change to O$_3$ was analysed by considering the sum of odd oxygen, NO$_2$ and its reservoir species, termed O$_x$, defined in Eq. (1).

\[
O_x = O + O_3 + NO_2 + 2N_2O_5 + 3NO_3 + HONO_2 + HO_2NO_2 + PANs
\]  

Tropospheric O$_3$ burden increases by 8% from 328 Tg in CS to 354 Tg in CS2. Much of the free troposphere exhibits increases of 2–6 ppb (≈ 6%–14%) in O$_3$, with large parts of the tropical troposphere increasing by more than 4 ppb (Fig. 1). This increase is driven chiefly by a 1.3% decrease in O$_3$ chemical destruction, resulting in a 12% increase in net chemical O$_3$ production. The sensitivity tests (Table S4) reveal the update to the isoprene mechanism only has a minor effect on O$_3$ burden (≈ 2 Tg decrease), while the changes to O(1D) and inorganic nitrogen reactions each yield increases of 17 Tg (when considered in isolation) with greater impacts in the lower and upper troposphere, respectively (Fig. S16 in the Supplement). The changes to the O$_3$ burden in the sensitivity tests do not sum to the total 26 Tg increase from CS to CS2, which indicates a degree of interplay between the diff-
different updates, an unsurprising result given O₃’s central role in tropospheric chemistry.

O₃ lifetime, defined as the ratio of O₃ burden (BO₃) to the sum of chemical (L_O₃) and physical (D_O₃) O₃ loss fluxes (Eq. 2) (Young et al., 2018; Archibald et al., 2020b), increase by 8% to 18.84 in CS2, while ozone production efficiency (OPE), defined as moles of O₃ produced per mole of NOₓ emitted (E_NO) (Eq. 3) (Archer-Nicholls et al., 2021), increases negligibly from 33.74 to 33.78.

\[
\tau_3 = \frac{BO_3}{L_O_3 + D_O_3}
\]  
\[
OPE = \frac{P_{O_3}}{E_{NO}}
\]

O₃ below 500 m increases across almost the entire globe with increases of 2–4 ppb (~5%–7.5%) over much of Europe, Africa, and the Americas and 4–5 ppb over India and China (Fig. 4), exacerbating the existing high bias in CS (Archer-Nicholls et al., 2021). The sensitivity tests allow this change to be partially decomposed into the different drivers (Fig. S13 in the Supplement). The update to isoprene chemistry produces localized increases in O₃ over the tropical forested regions of South America, Africa, and East Asia of 2–4 ppb: the increase in O₃ production via HO₂+NO and MeO₂+NO outweighs the reduction in the non-methyl peroxy radical (RO₂+NO) pathway (discussed later). While comparison to Khan et al. (2020) is difficult given the multiple mechanistic differences, O₃ production from RO₂+NO also decreased in their study. The changes to O(1D) also yield an increase in >1 ppb across the entire globe (due to reduced O₃ loss via O(1D)+H₂O) with a larger increase (2–3 ppb) encompassing ~20°S–40°N. The change to inorganic nitrogen also leads to terrestrial increases of 2–4 ppb from increased O₃ production via HO₂+NO and RO₂+NO.

5.1.1 O₃ Budget

O₃ production and loss fluxes for CS and CS2 are given in Table 6 and the breakdown for the sensitivity tests is given in Table S4. O₃ production decreases in CS2 in much of the tropical and Southern Hemisphere (SH) BL and lower free troposphere but increases in the Northern Hemisphere (NH) mid-latitude BL and tropical high troposphere, while O₃ loss decreases strongly in the tropical BL and lower free troposphere (Fig. 5). Despite the modest changes to total O₃ production and loss fluxes, the story is more complicated than it first appears due to offsetting changes to the key chemical production and loss fluxes.

5.1.2 O₃ production

The HO₂+NO pathway represents the largest absolute increase of O₃ production (3.2%, Table 6) with particular increases in the NH tropics and mid-latitude boundary layer and tropical upper troposphere (Fig. S12 in the Supplement). The drivers of this change are complex: the low altitude increases are driven by the significant increases in HO₂ (Fig. 6), which exceed 5% in places, while at higher altitudes the increase is attributed to a localized 15%–20% rise in NO. The sensitivity tests suggest the change to the isoprene scheme (CS2_isoprene) is a key driver in the rise of low-altitude HO₂ (and thus the flux), while the change to the inorganic nitrogen reactions (CS2_inorgN) also contributes to the increased flux at low altitudes and is chiefly responsible for the increase at higher altitudes.

However, the increase in HO₂+NO is offset by a decrease in the NO+RO₂ flux (15.4%, Table 6) where RO₂ comprises all peroxy radicals except the methyl peroxy radical, MeO₂. This reduction is strongest in the tropical BL and low free troposphere and driven by a significant decrease in the RO₂ burden (32%). This burden reduction arises from the isomerization pathways that inhibit the conversion of the isoprene-derived peroxy radical, RU14O₂, to the other peroxy radicals RU12O2 and RU10O2 (via reactions with standard partners such as NO and NO₃) by providing competing routes that yield other species whose degradation pathways do not produce further RO₂ (Khan et al., 2020). For example, the HPALDs produced are photolyzed to hydroxy acetone and unsaturated hydroxy carbonyls that further degrade producing mostly closed-shell products and HO₂. This rapid reaction pathway for RU14O2 sees its burden decrease by 35% in CS2 compared to CS, and tropical low-altitude mixing ratios decline by over 30%. Similar declines in the RO₂+NO flux (15%) and RO₂ burden (33%) are seen for CS2 relative to the CS2_isoprene sensitivity test, providing strong evidence that the change to isoprene is driving the change in RO₂. Khan et al. (2020) also simulated a reduction in RO₂ burden (and a corresponding drop in O₃ production via this pathway), although their decrease of 6.5% is less than half the equivalent value (including MeO₂) of 15% in this work, likely due to the other differences between the mechanisms used in their work and this study (see Sect. 1).

The fluxes of NO with HO₂, MeO₂, and RO₂ account for over 99.5% of total O₃ production in both mechanisms, and the changes in other pathways are an order of magnitude smaller in absolute terms. The reduction in the rate constant for OH+MeONO₂ (Sect. 4.1) reduces O₃ production from organic nitrate oxidation significantly while also driving the increase in O₃ production from organic nitrate photolysis. The addition of the photolysis of isoprene hydroxy nitrate and the other nitrates RU12NO3 and RU10NO3 make smaller contributions.

5.1.3 O₃ loss

The change in O₃ chemical destruction is dominated by the reduction in O(1D)+H₂O reaction (7.2%), which accounts for 54% of O₃ loss in CS but only 49% in CS2. In the sensitivity run CS2_O1D, which uses the same O(1D) rate constants as CS, the O(1D)+H₂O flux accounts for 54%
Table 6. Annual mean O₃ diagnostics for CRI-STRAT and CRI-STRAT 2 and the difference between the mechanisms (percentage changes in parentheses). UKESM1 CMIP6 1995–2004 using ST: chemical production = 5315 Tg yr⁻¹; chemical loss = 4476 Tg yr⁻¹; dry deposition = 867 Tg yr⁻¹ (Griffiths et al., 2021). Entries in bold show total production, loss and deposition, and entries in plain text show individual components of this production, loss and deposition.

<table>
<thead>
<tr>
<th></th>
<th>CS</th>
<th>CS2</th>
<th>CS2 – CS</th>
</tr>
</thead>
<tbody>
<tr>
<td>O₃ burden (Tg)</td>
<td>328</td>
<td>354</td>
<td>26 (7.9 %)</td>
</tr>
<tr>
<td>O₃ lifetime (d)</td>
<td>17.4</td>
<td>18.8</td>
<td>1.4 (8.0 %)</td>
</tr>
<tr>
<td>OPE</td>
<td>33.74</td>
<td>33.78</td>
<td>0.05 (0.1 %)</td>
</tr>
<tr>
<td><strong>Chemical production (Tg yr⁻¹)</strong></td>
<td><strong>6572</strong></td>
<td><strong>6582</strong></td>
<td><strong>10 (0.1 %)</strong></td>
</tr>
<tr>
<td>HO₂ + NO</td>
<td>4099</td>
<td>4322</td>
<td>132 (3.2 %)</td>
</tr>
<tr>
<td>MeOO + NO</td>
<td>1573</td>
<td>1583</td>
<td>10 (0.6 %)</td>
</tr>
<tr>
<td>NO + RO₂</td>
<td>849</td>
<td>717</td>
<td>−131 (−15.4 %)</td>
</tr>
<tr>
<td>Other</td>
<td>51</td>
<td>49</td>
<td>−1 (−2.8 %)</td>
</tr>
<tr>
<td><strong>Chemical loss (Tg yr⁻¹)</strong></td>
<td><strong>5834</strong></td>
<td><strong>5757</strong></td>
<td><strong>−77 (1.3 %)</strong></td>
</tr>
<tr>
<td>O(¹D) + H₂O</td>
<td>3157</td>
<td>2928</td>
<td>−229 (−7.2 %)</td>
</tr>
<tr>
<td>HO₂ + O₃</td>
<td>1666</td>
<td>1819</td>
<td>152 (9.1 %)</td>
</tr>
<tr>
<td>OH + O₃</td>
<td>740</td>
<td>796</td>
<td>57 (7.6 %)</td>
</tr>
<tr>
<td>O₃ + Alkene</td>
<td>166</td>
<td>101</td>
<td>−65 (−39.2 %)</td>
</tr>
<tr>
<td>Other</td>
<td>105</td>
<td>113</td>
<td>8 (10.1 %)</td>
</tr>
<tr>
<td><strong>Deposition (Tg yr⁻¹)</strong></td>
<td><strong>1133</strong></td>
<td><strong>1207</strong></td>
<td><strong>76 (6.5 %)</strong></td>
</tr>
<tr>
<td>O₃ dry deposition</td>
<td>942</td>
<td>1018</td>
<td>77 (8.0 %)</td>
</tr>
<tr>
<td>NO₃ deposition</td>
<td>191</td>
<td>189</td>
<td>−3 (1.3 %)</td>
</tr>
<tr>
<td>Inferred STT (Tg yr⁻¹)</td>
<td>395</td>
<td>384</td>
<td>−13 (−3.3 %)</td>
</tr>
</tbody>
</table>

Figure 5. Annual zonal mean change in (a) total O₃ production flux, (b) total O₃ chemical loss flux, and (c) net O₃ chemical production flux. The purple line indicates mean tropopause height.

of O₃ chemical loss. As this reaction involves water, the change is strongest in the tropical BL and low free troposphere (Fig. S14 in the Supplement).

The increase in O₃ loss via HO₂ + O₃ (9.1 %, Table 6) is driven predominantly by changes to the inorganic nitrogen and O(¹D) reactions, while the isoprene scheme is simulated to have little impact. O₃ loss via OH + O₃ also increases (7.6 %) despite the decrease in free troposphere HO₂ with the new isoprene chemistry and revised inorganic nitrogen reactions simulated to play important roles. O₃ destruction from O₃ + alkene reactions decline significantly (39 %) yet increase at very low altitudes (< 500 m) before decreasing at higher altitudes. This altitude dependence may arise from the enhanced O₃ low altitude driving a greater O₃ + alkene flux, but at higher altitudes the depletion of the volatile organic compounds (VOCs) by O₃ and the elevated OH mean O₃ destruction is lower.

5.2 HOₓ

The change to OH shows significant spatial and altitudinal variation, increasing at low altitude over land but decreasing...
Table 7. Tropospheric average HO$_x$ parameters for CS and CS2.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>CS</th>
<th>CS2</th>
<th>CS2 − CS</th>
</tr>
</thead>
<tbody>
<tr>
<td>[OH]/10$^6$ cm$^{-3}$</td>
<td>1.355</td>
<td>1.334</td>
<td>−0.021 (1.5 %)</td>
</tr>
<tr>
<td>[HO$_2$]/10$^8$ cm$^{-3}$</td>
<td>0.990</td>
<td>0.988</td>
<td>−0.002 (0.2 %)</td>
</tr>
<tr>
<td>[OH]/[HO$_2$] (%)</td>
<td>1.369</td>
<td>1.349</td>
<td>−0.02 (1.5 %)</td>
</tr>
<tr>
<td>CH$_4$ lifetime w.r.t. OH/years</td>
<td>7.43</td>
<td>7.60</td>
<td>−0.17 (2.3 %)</td>
</tr>
</tbody>
</table>

Figure 6. Annual zonal mean changes in (a) OH and (b) HO$_2$ between CS2 and CS, (c) absolute and (d) percentage in change in OH in lowest ∼500 m of atmosphere, (e) the change in OH in lowest 500 m between the CS2 and CS2_isoprene sensitivity test, and (f) HO$_x$ production flux from HPUCARB12 and HUCARB9 photolysis as a percentage of HO$_x$ from O$_3$(1D) + H$_2$O. Dark purple lines in (a) and (b) indicate the average height of tropopause.
over the oceans and in much of the free troposphere. This stems from the different drivers of OH concentrations and their relative importance in different regions.

At low altitude, the terrestrial increases in OH (Fig. 6c, d) are revealed by the sensitivity tests to be driven predominantly by the isoprene scheme: a clear illustration of impact of the HOX-recycling chemistry (Fig. S15 in the Supplement). The inorganic nitrogen changes make a smaller contribution to the low-latitude OH increase, while the O(1D) changes reduce low-altitude OH, but this effect is only noticeable over the oceans.

This significant increase in low terrestrial altitude OH is of particular interest in the context of BVOCs and their impact on the chemical composition of the atmosphere. Concentrations in the lowest 500 m increase by 2–3 \times 10^5 \text{ cm}^{-3} (30\%–50\%) in much of the Amazon, with similar changes seen in other tropical regions and the southeastern USA, the regions with the greatest emissions of isoprene and BVOCs. The boreal forest regions in North America and Eurasia exhibit modest increases of up to 10\% in places since isoprene emissions are lower (Fig. S15). The influence of the updated isoprene chemistry is further apparent when the HOX production flux from two of the key new HOX-recycling pathways, i.e. photolysis of the HPALD and hydroxy unsaturated carbonyl (HUCARB9) species, is compared to that from O(1D)+H_2O (Fig. 6f). Over the Amazon and other tropical regions, HOX,
Figure 8. Annual mean mixing ratios for (a) HPALDs and (b) IEPOX over lowest ~100 m. DJF (December–January–February) and JJA (June–July–August) zonal mean mixing ratios for HPALDs (c, d) and IEPOX (e, f); note the differing scales for HPALD and IEPOX plots and log scales for (c–f).

flux from this pathway amounts to more than 40 % of that from O(^1D) + H_2O. The difference in BVOC-driven depletion of oxidant concentrations at low altitudes will be even more pronounced when CS2 is compared with ST, which exhibited even lower tropical low-altitude HOx (e.g. Fig. 9, Archer-Nicholls et al., 2021).

However, in much of the free troposphere, OH decreases by 2 %–10 % relative to CS due to the changes made to the O(^1D) rate constants (Fig. S20 in the Supplement), which reduces the fraction of O(^1D) reacting with H_2O (Fig. 6a). This general decline is reversed in the upper tropical troposphere (10–15 km), where OH increases by up to 15 %, driven by an increase in NO stemming from the update to inorganic nitrogen reactions and a smaller contribution from the updated isoprene chemistry (Fig. S16). These free troposphere changes partially reverse the changes simulated between CS and ST (Fig. 6, Archer-Nicholls et al., 2021). In that comparison, tropical free troposphere OH (~2–6 km) increased in CS relative to ST by 0.5–2 × 10^5 cm^{-3}, while here CS2 yields a decrease in the same location of 0.25–1 × 10^5 cm^{-3} compared to CS. In the upper tropical troposphere, CS decreased OH by 1–4 × 10^5 cm^{-3} relative to ST, while CS2 exhibits an increase of 0.25–1.5 × 10^5 cm^{-3} in the same region compared to CS. Thus, the distribution of free troposphere OH in CS2 is more similar to that in ST than the CS distribution is.
Overall, the reduction in the free troposphere OH outweighs the increases elsewhere with the tropospheric air mass-weighted concentration and burden of OH decreasing in CS2 by 1.5% and 0.49%, respectively. This is in sharp contrast to the 6.4% increase in burden simulated by Khan et al. (2020). However, in the CS2_O1D sensitivity test the OH burden increases by 6.6% relative to CS, allowing us to be confident that the discrepancy between Khan et al. (2020) and this work is predominantly down to the differing O(1D) rate constants. Despite the increase in surface OH, the net reduction in tropospheric OH yields a 2.3% increase in methane lifetime from 7.43 to 7.60 years (Table 7), also in contrast to the 0.5 years decrease in methane lifetime simulated by Khan et al. (2020). However, the isolated change to isoprene chemistry, given by the comparison of CS2 and CS2_isoprene, causes a methane lifetime decreases of 2.2% due to the enhanced low-altitude OH.

HO2 also increases at low altitude (up to 6%–8% at the surface, Fig. 6b), driven primarily by the new isoprene chemistry, but this increase extends much further into the free troposphere than OH, reaching nearly 5 km above the Equator. HO2 decreases in the rest of the free troposphere, partially from O(1D) changes, and does not exhibit the high increase shown by OH, instead declining by 6%–8% in the tropical high troposphere, resulting in a burden decrease of 0.7%. The greater increase in low-altitude HO2 (than for OH) is likely to be due in part to co-located increases in CO of 3–6 ppb (see the Supplement and Fig. S21a).

5.3 BVOCs

The interactive nature of iBVOC emissions led to average isoprene emissions being 0.36 Tg yr\(^{-1}\) (0.06%) lower in CS2, while monoterpene emissions were 0.05 Tg yr\(^{-1}\) (0.05%) lower. However, these differences are dwarfed by the reductions in the burdens of isoprene, α-pinene, and β-pinene of 26%, 18%, and 24%, respectively.

Isoprene mixing ratios averaged over the lowest \(\sim 100\) m decrease by 1–3 ppb (\(\sim 10\%–30\%\)) in large parts of South America, Africa, and Southeast Asia (Fig. 7). The greater terrestrial low-altitude HO\(_2\) increases the OH-initiated oxidative flux of isoprene by 3.5%, attributable almost entirely to the updated isoprene scheme. However, this is actually outweighed by a 23% decrease in isoprene destruction by O\(_3\), while oxidation via NO\(_3\) increased by 3.7%. Despite the modest global increase, isoprene oxidation is confined even more to low-altitude regions (Fig. 7), a feature also simulated by Karset et al. (2018) (Fig. 8). This also results in lower mixing ratios throughout the whole troposphere (Fig. 7).

α-pinene’s chemical destruction by OH, O\(_3\), and NO\(_3\) changed by 7.5%, −6.3%, and −0.8%, respectively, leading to a total flux increase of 0.05 Tg yr\(^{-1}\) (+0.05%). The corresponding changes for β-pinene with OH, O\(_3\), and NO\(_3\) were 3.2%, −4.3%, and −0.2%, with a total increase of 0.70 Tg yr\(^{-1}\) (1.5%).

The reductions to these BVOC burdens are greater than those simulated by Khan et al. (2020) of 17%, 4%, and 9% for isoprene, α-pinene and β-pinene, respectively. However,
Khan et al. (2020) simulated a reduction in O$_3$ over tropical regions and a much smaller increase in NO$_3$ burden (1 %), which would have resulted in significantly lower BVOC destruction fluxes, particularly for the monoterpenes. As discussed in Sect. 4, CS2 simulates a reduction in the model high bias of surface isoprene and (to a lesser extent) monoterpenes when compared to CS and ST.

5.4 HPALDs and IEPOX

While a comparison cannot be made between CS and CS2 for HPALDs and IEPOX, their importance for HO$_x$ recycling and SOA formation, respectively, means that examining their global distribution is still useful. Both species follow the surface distribution of isoprene closely (Fig. 8) with IEPOX concentrations typically an order of magnitude greater than HPALDs, something also reflected in their burdens (0.39 and 0.02 Tg, respectively). As discussed in Sect. 4, loss of IEPOX to aerosol via reactive uptake is not currently modelled, and simulated concentrations will decrease once this process is included. Indeed, accurate modelling of IEPOX and its contribution to SOA has been suggested to be important in future climate scenarios (Jo et al., 2021), which highlights the benefits of including IEPOX in CS2 but also the need for careful consideration of how aerosol uptake is modelled. Simulated advection up to the upper tropical troposphere is clearly seen in the DIF zonal means, with potentially important consequences for the IEPOX-derived SOA that has been observed in the lower troposphere in flight campaigns (e.g. Allan et al., 2014).

5.5 NO$_y$

The distribution of nitrated products (NO$_y$) between reactive (NO$_x$) and reservoir species (NO$_z$) changes between CS and CS2 and is detailed in Table 8. Here we use the standard definitions of NO$_x$, NO$_z$, and NO$_y$ (Archer-Nicholls et al., 2021) (Eqs. 4–6):

\[
\begin{align*}
\text{NO}_x &= \text{NO} + \text{NO}_2, \\
\text{NO}_z &= \text{NO}_3 + 2\text{N}_2\text{O}_5 + \text{HONO}_2 + \text{HO}_2\text{NO}_2 \\
&\quad + \text{ClONO}_2 + \text{BrONO}_2 + \text{PANs} + \text{RONO}_2 \\
&\quad + \text{CH}_3\text{O}_2\text{NO}_2 + \text{Nitrophenols}, \\
\text{NO}_y &= \text{NO}_x + \text{NO}_z.
\end{align*}
\]

RONO$_2$ comprises alkyl nitrates, hydroxy nitrates, and hydroperoxy nitrates, while PANs comprise all species with peroxyacyl nitrate functionality.
The NO\textsubscript{y} burden decreases by 4.8 % (in terms of mass of N), driven primarily by a 20 % decline in PANs. However, the NO\textsubscript{x} burden increases by 4 % with the widespread increase in the tropical high troposphere of 10–20 ppt (up to 25 %), outweighing the reduction in the NH mid-latitude PBL (10–50 ppt, 1 %–2.5 %) (Fig. 9a). The increase in NO\textsubscript{x} and the reduction in NO\textsubscript{y} leads to the fraction of NO\textsubscript{y} as reactive nitrogen increasing by 9 %, and the associated increases to the O\textsubscript{3} production, particularly in the free troposphere, are identified in Sect. 5.1. The sensitivity tests revealed the high-altitude NO\textsubscript{x} rise to be driven predominantly by the change to the inorganic nitrogen, with a smaller contribution from the updated isoprene scheme (Fig. S17 in the Supplement).

The 6 % reduction in NO\textsubscript{y} burden is dominated by the decrease in PANs, which exceeds 40 ppt in most of the 40° N–40° S troposphere (Fig. 10e). The decrease in the PAN formation rate constant discussed in Sect. 2 is not the principal driver of this reduction despite reducing by 40 % in much of the troposphere. For the single year used for the sensitivity tests, the PANs burden in CS2_inorgN (featuring the larger formation rate constant) (0.292 Tg N) is much closer to that in CS2 (0.290 Tg N) than in CS (0.364 Tg N). A more important factor is the reduction in the PAN-precursor acyl peroxy radical (MeCO\textsubscript{3}), driven by the updated isoprene chemistry, whose burden decreases by over 20 % in both CS2 and the sensitivity test CS2_inorgN. This dependency is clearly illustrated by the fact that the isolated change to the inorganic nitrogen reactions (CS2_inorgN) only produces a small decrease to low-altitude PANs, while the change to isoprene scheme (CS2_isoprene) yields a much larger decrease in PANs in spatial agreement with the CS2–CS difference (Fig. S18 in the Supplement). The PANs burden of 0.317 Tg N in the CS2_isoprene test is also closer to that in CS. However, the change in PANs between CS2 and CS is still larger than that simulated from the isolated isoprene chemistry change alone, which suggests there are some synergistic effects occurring.

The 0.4 % increase in HONO\textsubscript{2}, including increases of up to 30 ppt in the tropical mid-troposphere (Fig. 9d), is driven more by the update to the isoprene scheme than the change to inorganic nitrogen reactions (Fig. S19 in the Supplement).

The 59 % increase in RONO\textsubscript{2} burden in CS2 is predominantly due to the significant reduction in the rate constant for the OH-initiated destruction of MeONO\textsubscript{2}, the principal organonitrate, which brings CS2 into agreement with STRAT-TROP and the most recent IUPAC value. At 290 K, the rate constant is 18 times lower in CS2 and at 250 K it is 50 times lower, yielding a 3-fold MeONO\textsubscript{2} burden increase.
The update to the isoprene scheme, when isolated, actually reduces RONO2, despite the introduction of the two new organic nitrates (RU12NO3 and RU10NO3). As discussed in the context of the RO2 burden, this is driven by the added competition from the RU14O2 isomerization reactions: the flux of the RU14O2 + NO reaction is 15 % lower in CS2 than CS. The increase in RONO2 is simulated to be 10–20 ppt in the tropical lower altitude and 2–10 ppt for the rest of the troposphere (Fig. 9f, S20).

With the significant drop in PANs as a fraction of NOx (34 % to 28 %) and the increase in HONO2 and NOx, the breakdown of NOy in CS2 is closer to that in ST (Archer-Nicholls et al., 2021). The increase in RONO2 is the only major exception to this since ST, which only has two organonitrate species (isoprene nitrate and MeONO2), has a lower RONO2 burden than CS.

5.6 Impacts on aerosols

A key area of future research with the CRI mechanisms will be on their influence on aerosols. The spatial changes to oxidants are likely to influence secondary organic aerosol (SOA) formation, as discussed in Sect. 1. In UKCA, SOA is produced from the tracer Sec_Org, a surrogate for the oxidized products of α-pinene and β-pinene that adds to existing organic aerosol with an optional boundary layer nucleation scheme involving Sec_Org and H2SO4 based on Metzger et al. (2010) also available. The Sec_Org burden decreases by 7 % in CS2 with noticeable annual variation (DJF −10 %, JJA −4 %). Despite the burden decrease, within the lowest 500 m in the tropics Sec_Org mass concentration increases by 2 %–10 %, driven by an increase its production from α-pinene and β-pinene (Fig. 10a, b). Above this region, Sec_Org production and mass concentration decrease, and thus it appears the greater low-altitude oxidative capacity in CS2 leads to greater production of Sec_Org within the boundary layer but lower concentrations above it. This is likely to have an impact on SOA distribution (and lifetime) since deposition and loss to the aerosol phase is greater in the boundary layer due to the steep decline in aerosol surface aerosol density with altitude. Further detailed analysis involving the fluxes of Sec_Org to aerosol and the resulting changes to size and number distributions are beyond the scope of this work, but examining wider consequences for SOA, in the context of the BVOC-mediated feedback between the biosphere and climate, will form a key area of future research. It is also worth noting an even more pronounced perturbation to SOA may be seen if isoprene is allowed to produce Sec_Org, which is a more realistic approach to simulating SOA (e.g. Scott et al., 2014) and will be explored in future work.

The global perturbation to the oxidation pathways of SO2, another important aerosol precursor, are more modest. From CS to CS2, the oxidative fluxes of SO2 with OH, H2O2, and O3 change by +0.9 %, +0.02 %, and 1.7 %, respectively, while the tropospheric sulfate aerosol burden decreases by just 2.3 %. However, as with isoprene oxidation and Sec_Org production, the burden change belies the more complex perturbations occurring. The increased oxidants at lower altitudes and reduction at greater altitudes result in gas-phase SO2 oxidation increasing by 2.5 %–10 % in the tropical and mid-latitude PBL but decreasing at higher altitudes (Fig. 10c, d). This effect is expected to be even more pronounced when CS2 is compared to ST, which simulates even lower low-altitude OH than CS (Archer-Nicholls et al., 2021) and has been the standard mechanism for investigations into aerosol–oxidant coupling in UKCA (Thornhill et al., 2021; Weber et al., 2020; O’Connor et al., 2021). Therefore, the mechanism-driven changes to oxidants are likely to have consequences for both SOA and sulfate aerosol. While a full investigation into oxidant aerosol coupling is beyond the scope of this paper, it will form a central part of future work with the CRI mechanisms.

6 Summary and synthesis

The key changes between CS and CS2, driven by the multiple chemistry changes, can be summarized as follows.

1. O3 production increases marginally in CS2, but a larger decrease in O3 destruction, driven by a significant reduction in the O(1D) + H2O flux, leads to a greater O3 tropospheric burden and mixing ratios.

2. The update to the isoprene chemistry increases low-altitude tropical HOx, but the reduction in OH production from O(1D) + H2O results in lower HOx concentrations in much of the free troposphere, increasing methane lifetime.

3. The update to the inorganic nitrogen reactions increases NOy as a fraction of NOy, with a significant increase in the upper tropical free troposphere and a co-located increase in OH. The PAN burden decreases by 20 %.

4. The increase in boundary layer oxidative capacity reduces the burden of BVOCs and confines their oxidation even more to low altitudes, with likely consequences for aerosol production and lifetime.

7 Conclusions

The radiative impact of isoprene, via its influence on atmospheric chemical composition and organic aerosol, means an accurate description of its chemistry is crucial for advancing our understanding of pre-industrial, present-day and future atmospheres. In this study we describe the incorporation of the Common Representative Intermediates chemistry scheme version 2.2 (CRI v2.2), along with accompanying stratospheric chemistry, into the global chemistry–climate model.
UKCA to create the mechanism CRI-Strat 2 (CS2). The introduction of CS2 into UKCA facilitates a semi-explicit description of HOx-recycling chemistry during isoprene oxidation via the isomerization of isoprene peroxy radicals to produce HPALDs that yield HOx upon photolysis. This is a key process for reconciling the model low bias of HOx in low-NOx, BVOC-rich regions. In addition, CS2 also features updates to the rate constants of the reactions of O(1D), inorganic nitrogen, and organic peroxy radicals with NO and NO3, bringing the mechanism into agreement with the most recent IUPAC values. CS2 is one of the first mechanisms with this functionality suitable for long-term climate integrations.

A rigorous comparison using UKCA with CS2 and two other chemical mechanisms, STRAT-TROP (ST) (the standard chemistry mechanism used in UKESM1’s contributions to CMIP6 experiments) and CRI-STRAT (CS) (which has tropospheric chemistry from an earlier version of the CRI, CRI v2.1), is performed against high-frequency surface and airborne observational data from BVOC-rich regions for multiple chemical species, including O3, HOx, HO2, and isoprene, and monoterpenes and isoprene oxidation production. The HOx recycling in CS2 results in significantly enhanced surface diel OH (up to 50 % higher than CS at midday) in the Amazon and Borneo (improving model low bias), leading to improved modelling of diel and vertical isoprene profiles and reducing the mean 24 h bias by 50 %–60 % and 20 %–40 % relative to ST and CS, respectively, across the locations considered. However, CRI-Strat 2 exacerbates the existing isoprene model low bias away from the surface, suggesting potential issues with model vertical convection. CS and CS2 yield smaller isoprene column biases compared to observations than ST, in line with the surface and free troposphere observational comparisons, while also illustrating the significant influence the chemical mechanism has on modelled column. This comparison also highlights the significant influence that the different chemistry schemes have on the simulated isoprene column and thus the considerable challenges of determining isoprene emissions via back-calculation.

The low-altitude high biases for O3 in CS increase modestly (1–2 ppb) in CS2. Simulated monoterpenes concentrations are biased high at the surface at most of the locations considered, with CS2 returning the smallest bias. As with isoprene, simulated monoterpenes display sharp morning and evening peaks that are believed to be due to boundary layer height issues. Modelled high bias of IEPOX and the low bias of HPALDs suggest further investigation of the key processes of loss to aerosol for IEPOX and HPALD photolysis frequency are needed.

In addition to observational comparisons, a detailed comparison of UKCA model output using CS2 is performed, complementing the earlier comparison of ST and CS (Archer-Nicholls et al., 2021). Sensitivity tests are also performed to help isolate the drivers of the differences between CS and CS2. CS2 simulates an 8 % increase in tropospheric O3 burden this is primarily driven by reduced O3 loss as the changes to rate constants of O(1D) with H2O, O2, and N2 mean that a smaller fraction of O(1D) reacts with H2O to produce OH. Low-altitude O3 increases by 2–4 ppb over much of the globe, driven predominantly by changes to the O(1D) and inorganic nitrogen reactions. More broadly, the widespread influence of the changes to the rate constants of O(1D) and multiple inorganic nitrogen species highlights the importance of having accurate information for these parameters.

Relative to CS, low-altitude OH increases over terrestrial regions, exceeding 50 % in some tropical forested regions, primarily due to the influence of HOx recycling from isoprene. However, OH decreases over the oceans and in much of the free troposphere, driven by updates to the rate constants of the reaction of O(1D) with H2O, O2, and N2. As a result, methane lifetime increases by 1.9 %, which is in stark contrast to previous studies using CRI v2.2 in the STOCHEM model that did not make changes to O(1D) and inorganic nitrogen reactions. When the changes to isoprene chemistry are isolated, methane lifetime decreases by 2.2 %, which is qualitatively in line with previous studies. The addition of isomerization pathways in the updated isoprene scheme reduces the methyl (7 %) and non-methyl peroxy (36 %) radical burdens.

The distribution of nitrated species (NOx) in CS2 is closer to that simulated in ST than CS is with a significant reduction (20 %) in the burden of PANs that is driven by a reduction in the precursor RO2. The NOx burden increases by 4 %.

The increase in low-altitude OH reduced the burdens of isoprene (25 %) and monoterpenes (11 %–18 %) and the extent of their dispersion: more oxidation takes place in the boundary layer, where loss of oxidation products such as the lumped SOA precursor Sec_Org to existing aerosol is likely to be greater. Enhanced SO2 oxidation in the boundary layer is also simulated. These changes are likely to have implications for SOA and sulfate aerosol, particularly as CS has already been shown to have a more highly oxidizing boundary layer than ST. Therefore, the difference between CS2 and ST (the mechanism used to explore chemical–aerosol coupling in UKESM1 in CMIP6 experiments) is likely to be significant and will be the subject of future work.

The addition of CS2 also lays the groundwork for the incorporation of a novel chemistry scheme that describes the formation of the highly oxidized organic molecules (HOMs) derived from biogenic species such as α-pinene (e.g. CRI-HOM, Weber et al., 2020b). HOMs are crucial for new particle formation without sulfuric acid (Kirkby et al., 2016; Simon et al., 2020), a process which is an important source of new particles in the Amazonian free troposphere (Zhao et al., 2020) and has been simulated to have consequences for our understanding of pre-industrial aerosol burden (Gordon et al., 2016). The influence of isoprene in HOM production (Klindner-Schaar et al., 2009; McFiggans et al., 2019; Heinritzi et al., 2020) can also be captured by the addition of CRI-HOM, making UKCA one of the very first global
chemistry–climate models to feature a semi-explicit representation of HOMs and enabling further investigation into the climatic impact of the interaction between BVOCs. The addition of long-chain terpenes to CS2 is also planned, including sesquiterpenes, which may reduce the surface ozone high bias, form HOMs, and make improvements to the uptake of oxidized species to plant surfaces.

While certain elements of the CRI-STRAT 2 mechanism in UKCA such as the ozone high bias remain problematic, its incorporation represents a major step forward in our ability to simulate isoprene chemistry in low-NOx environments. The simulated changes to oxidants in CRI-Strat 2 will affect the atmosphere’s radiative balance by perturbing certain greenhouse gases and aerosols, and investigating the impact of this will be a major topic of future work. In particular, the feedback between the biosphere and climate, mediated by BVOCs, will be evaluated using multiple mechanisms to assess their influence. CRI-Strat 2 can be taken up for use, alongside other mechanisms, to further our understanding of the wide-ranging impact BVOCs have on climate.

Code availability. Due to intellectual property right restrictions, we cannot provide either the source code or documentation papers for the UM. The Met Office United Model is available for use under licence. A number of research organizations and national meteorological services use the UM in collaboration with the UK Met Office to undertake basic atmospheric process research, produce forecasts, develop the UM code, and build and evaluate Earth system models. For further information on how to apply for a licence, see https://www.metoffice.gov.uk/research/approach/model-modelling-systems/unified-model (last access: 24 November 2020).

Data availability. The description of the ZF2 field campaign is given in Sect. S4, and the observational data are available at https://doi.org/10.17863/CAM.65133 (Weber and Bannan, 2021).

The observational data from the SEAC4RS flight campaign are available at https://www-air.larc.nasa.gov/cgi-bin/ArcView/seac4rs?MERGE=1#60_SECOND.DC8_MRG (Chen, 2016).

The observational data from the ATTO tower are available to download at https://www.attodata.org (last access: 9 April 2021). Specific data sets used were https://www.attodata.org/ddm/data/Showdata/72, https://www.attodata.org/ddm/data/Showdata/73, https://www.attodata.org/ddm/data/Showdata/74, and https://www.attodata.org/ddm/data/Showdata/77 (Yáñez-Serrano, 2020a, b, c, d).

The observational data from the FAAM aircraft are available at http://data.ceda.ac.uk/badc/op3/data/op3-aircraft (last access: 27 March 2021), and the Borneo data can be found at http://data.ceda.ac.uk/badc/op3/data (last access: 27 March 2021; Natural Environment Research Council et al., 2010).

Data tables of the full CRI-Strat 2 mechanism and the mechanisms used in the sensitivity test described in this paper are included in the Supplement. The CRI v2.2 mechanism can be viewed and downloaded from http://cri.york.ac.uk (Jenkin et al., 2019b).

Model data and analysis code are available from James Weber on request.

Supplement. The supplement related to this article is available online at: https://doi.org/10.5194/gmd-14-5239-2021-supplement.

Author contributions. Mechanism incorporation was carried out by JaW with support from SAN and NLA and advice from ATA, YMS, MJ, MAHK, and DES. Observational comparison experiments were designed and carried out by JaW with advice from SAN, NLA, and ATA. Mechanism intercomparison experiments were designed by JaW with advice from ATA, NLA, and SAN and executed by JaW, TJB, CJP, AB, and PA compiled and supplied the ZF2 Brazil observational data, and TJB wrote the field campaign description in the Supplement. RHS advised on the SEAC4RS data and analysis. JaW interpreted the ZF2 Brazil, Borneo, ATTO, FAAM, GABRIEL, and SEAC4RS observational data with advice from SAN, ATA, and JoW. JaW wrote the paper. All co-authors discussed the results and commented on the paper.

Competing interests. The authors declare that they have no conflict of interest.

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Acknowledgements. This work used Monsoon2, a collaborative high-performance computing facility funded by the Met Office and the Natural Environment Research Council. This work used JASMINE, the UK collaborative data analysis facility.

We are grateful to Horst Fischer, Hartwig Harder, and Pete Edwards for their assistance and advice and to Jason Surratt for providing the calibration standards for the ZF2 Brazil study. We are grateful for the field support from the LBA central office at INPA in Manaus.

Financial support. James Weber has been funded by a Vice-Chancellor’s Award from the Cambridge Trust. Scott Archer-Nicholls and Alex T. Archibald have been funded by NERC PROMOTE (grant no. NE/P016383/1). Nathan Luke Abraham and Alex T. Archibald are supported by NERC and NCAS through the ACSIS project. Youngsub M. Shin has been funded by NERC through the University of Cambridge ESS-DTP. M. Anwar H. Khan and Dudley E. Shallcross are funded by NERC (grant no. NE/K004905/1), Bristol ChemLabS, and the Primary Science Teaching Trust. Thomas J. Bannan, Carl J. Percival, Asan Baca, and Paulo Artaxo acknowledge funding from FAPESP – Fundação de Amparo à Pesquisa do Estado de São Paulo (grant no. 2017/17047-0).
Review statement. This paper was edited by Tim Butler and reviewed by two anonymous referees.

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Geosci. Model Dev., 14, 5239–5268, 2021

https://doi.org/10.5194/gmd-14-5239-2021


https://doi.org/10.5194/gmd-14-5239-2021

Geosci. Model Dev., 14, 5239–5268, 2021


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