

The Common Representative Intermediates Mechanism version 2 in the United Kingdom Chemistry and Aerosols Model

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Key Points:

- The CRI-Strat mechanism has been integrated into the UKCA model, greatly increasing the complexity of VOC chemistry compared to StratTrop.
- CRI-Strat simulates higher surface ozone compared to StratTrop due to greater production, but tropospheric ozone burden is similar.
- The ozone and oxidised nitrogen budgets when running with the CRI-Strat mechanism show high sensitivity to the input NMVOC emissions.

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Abstract

We document the implementation of the Common Representative Intermediates Mechanism version 2, reduction 5 (CRIv2-R5) into the United Kingdom Chemistry and Aerosol model (UKCA) version 10.9. The mechanism is merged with the stratospheric chemistry already used by the StratTrop mechanism, as used in UKCA and the UK Earth System Model (UKESM1), to create a new CRI-Strat mechanism. CRI-Strat simulates a more comprehensive treatment of non-methane volatile organic compounds (NMVOCs) and provides traceability with the Master Chemical Mechanism (MCM). In total, CRI-Strat simulates the chemistry of 233 species competing in 613 reactions (compared to 87 species and 305 reactions in the existing StratTrop mechanism). However, while more than twice as complex than StratTrop, the new mechanism is only 75% more computationally expensive. CRI-Strat is evaluated against an array of *in situ* and remote sensing observations and simulations using the StratTrop mechanism in the UKCA model. It is found to increase production of ozone near the surface, leading to higher ozone concentrations compared to surface observations. However, ozone loss is also greater in CRI-Strat, leading to less ozone away from emission sources and a similar tropospheric ozone burden compared to StratTrop. CRI-Strat also produces more carbon monoxide than StratTrop, particularly downwind of biogenic VOC emission sources, but has lower burdens of nitrogen oxides as more is converted into reservoir species. The changes to tropospheric ozone and nitrogen budgets are sensitive to the treatment of NMVOC emissions, highlighting the need to reduce uncertainty in these emissions to improve representation of tropospheric chemical composition.

Plain Language Summary

To understand the climate and predict how it will change in the future, we need to understand its chemical composition - the trace gases and small particles that exist in tiny quantities in the atmosphere. A key tool we use to do this are computer models which simulate the atmosphere and processes within it. Key processes include the formation of ozone, a harmful pollutant and greenhouse gas in the lower atmosphere. However, the chemistry involved in forming ozone is very complicated, so computer simulations of the atmosphere must greatly simplify the chemistry. These simple schemes may introduce errors in the model. We also have much more complex chemical mechanisms which simulate our best understanding of all chemical reactions, but these complex schemes require too much computational power to be used when simulating the whole atmosphere. In this paper, we describe the implementation of a chemical mechanism that sits between these levels of complexity, realistically simulating the formation and destruction of ozone without being too slow to run. We compare this new mechanism against measurements taken of the atmosphere and the preexisting, simpler chemical mechanism and show that the new mechanism greatly enhances the amount of ozone that is produced.

1 Introduction

Understanding chemical processes in the lower atmosphere is of vital importance for tackling the problems of air pollution and making accurate projections of how the Earth system will change due to human activity (Sillman, 1999; Akimoto, 2003; Von Schneidemesser et al., 2015; Monks et al., 2015; Boucher et al., 2013). However, the chemistry of the troposphere is extremely complicated because of the wide variety of non methane volatile organic compounds (NMVOCs) whose structures are diverse and whose lifetimes and abundances cover many orders of magnitude (Atkinson, 1990; M. Jenkin et al., 1997; Goldstein & Galbally, 2007). Two particularly challenging aspects of tropospheric chemistry regard the understanding of the formation and destruction of tropospheric ozone and the impacts of aerosols, with much of the difficulty in both of these research topics stemming from the importance of NMVOCs. The rate of production of ozone is non-

linearly dependent on the combination of nitrogen oxides ($\text{NO}_x = \text{NONO}_2$) and NMVOCs levels, with high rates of ozone production occurring when levels of both are high and net ozone destruction occurring when there is a large excess of NMVOCs (NO_x -limited regime) or an excess of NO_x (VOC-limited regime) (Sillman, 1999; Monks et al., 2015). Larger NMVOC molecules, such as monoterpenes, aromatic compounds and long chain n-alkanes ($\text{C}_{>10}$ – where C indicates the number of carbon atoms in the NMVOC) play an important role in the generation of secondary organic aerosol (SOA). SOA can make up over 50% of submicron aerosol mass (Jimenez et al., 2009), but we still have significant uncertainty in the exact chemical makeup, and models still fail to accurately simulate it (Tsigaridis et al., 2014; Hodzic et al., 2020). Unfortunately, many thousands of different NMVOC species have been identified in the atmosphere, and many more are yet to be discovered, making a complete representation of all NMVOC species and their chemistry in a model an impossible task (Goldstein & Galbally, 2007; Heald & Kroll, 2020).

The key mapping between the input NMVOCs and their effects on ozone and SOA is their oxidation mechanism. Such mechanisms are well known from laboratory experiments for the simplest NMVOCs ($\text{C}_{<5}$). Based on the wealth of experimental data (McGillen et al., 2020), structure activity relationships (SARs) have been derived to fill in the gaps (M. E. Jenkin et al., 2018b, 2018a; M. E. Jenkin, Valorso, et al., 2019) and to extrapolate our understanding of NMVOC oxidation to cover a wide range of structures and configurations. Aumont et al. (2005) have shown that the oxidation process in the atmosphere can be treated as a geometric problem, with the number of species produced during the oxidation of an alkane NMVOC with n carbon atoms being given by:

$$\Omega \approx \sum_{i=2}^n \frac{1}{2} (11)^2 (7)^{i-2}. \quad (1)$$

For a C_5 compound, this equation leads to $\approx 10^5$ species forming, presenting a huge amount of complexity.

To approach the problems of complexity relating to NMVOC chemistry in the atmosphere, researchers have typically followed one of two routes for developing chemistry mechanisms depending on the research questions being tackled and relevant spatial and temporal scales. Detailed, explicit mechanisms (Aumont et al., 2005) and near-explicit mechanisms, such as the Master Chemical Mechanism (MCM; Saunders et al. (2003); M. E. Jenkin et al. (2003)), comprise of the amalgamation of known relevant chemistry as measured from laboratory chemical kinetic studies (Atkinson et al., 2006; S. P. Sander et al., 2011) and SARs (M. E. Jenkin et al., 2018b, 2018a; M. E. Jenkin, Valorso, et al., 2019) and evaluated against field and chamber experiments (M. E. Jenkin et al., 2012; Novelli et al., 2018). These mechanisms represent our best understanding of chemical processes in the atmosphere, often comprising of thousands of species and many times more reactions, and are continuously updated and expanded as new data and understanding comes to light. However, this makes them computationally very expensive to run and hence these explicit mechanisms are mostly used in box model studies (M. E. Jenkin et al., 2015; Derwent, 2017).

An alternative design approach is to construct mechanisms that are as simple as possible but as complicated as necessary, the aim being to represent the key chemical processes and their interactions with as few chemical species and reactions as possible. Such atmospheric chemistry schemes typically have 10s of species and dozens to hundreds of reactions (e.g., the StratTrop scheme which simulates 81 species, 291 reactions (Archibald et al., 2020)). Chemical processes are still informed by the best available data, but similar species are lumped together or represented by surrogate species and more complex but important processes are parameterised to reduce the mechanism complexity. Mechanisms may also become quite specialist, with those designed to simulate urban air pollution (e.g., (Stockwell et al., 1990)) differing greatly from those intended to simulate the whole atmosphere at a coarse resolution (e.g., (Archibald et al., 2020)), as

125 they focus on different aspects of the chemistry system occurring in the real atmosphere.
126 These approximations are necessary to run interactive chemistry in 3D models that must
127 also simulate other key processes such as transport, deposition, clouds and radiation. Even
128 so, the chemistry component is often the most computationally expensive part of a 3D
129 model (Esenturk et al., 2018).

130 These different approaches leave open a gap between our most comprehensive mech-
131 anisms and simpler ones. Simpler schemes can perform well for the photochemical con-
132 ditions they were designed for, but may perform poorly when simulating other regions,
133 or fail to properly represent how chemical conditions should change in response to changes
134 in emissions or climate. Due to their complexity, it is impractical to use near-explicit chem-
135 istry schemes in the same 3D model setup as simpler mechanisms to see how they would
136 perform instead. While they can be evaluated against comprehensive schemes in box model
137 simulations, when results differ it is difficult to pinpoint which aspect of the chemistry
138 is causing the differences, due to the lack of traceability, and it can be unclear whether
139 the simpler mechanisms respond realistically in photochemical conditions outside of those
140 evaluated in box model experiments. There is therefore a clear need for intermediate com-
141 plexity mechanisms - ones which are fully traceable to more comprehensive schemes and
142 are known to respond similarly to changes in forcings, but are still simple enough to be
143 used in 3D interactive models.

144 The Common Representative Intermediates (CRI) mechanism (M. E. Jenkin et al.,
145 2002; M. Jenkin et al., 2008) is just such an intermediate complexity mechanism. The
146 number of species and reactions are reduced by over an order of magnitude compared
147 with the MCM. However, through a rigorous development process, in which it was fully
148 and systematically evaluated against the more complex MCM mechanism at each stage
149 of complexity reduction and no change that significantly degraded representation of ozone
150 production was allowed through (Watson et al., 2008), the scheme is fully traceable to
151 the MCM. The end result is a mechanism that is simple enough to be run in a 3D model,
152 but which we can be confident responds to changes in emissions and conditions accord-
153 ing to our best understanding as represented in the MCM. Intermediate complexity mech-
154 anisms offer enormous benefits when used in 3D models as a research tool to study the
155 importance of chemical processes which are ignored in simpler schemes and as a bench-
156 mark against which to test, evaluate and inform development of these simpler schemes.
157 By having more confidence in the representation of gas-phase chemistry, it is possible
158 to attribute remaining model biases to other structural components of the model.

159 The CRI mechanism has been used in several models now, including the STOCHEM
160 Lagrangian global chemical transport model (S. Utembe et al., 2010; M. A. H. Khan et
161 al., 2015), the Weather Research and Forecasting model with chemistry (WRF-Chem),
162 an online regional coupled model (Archer-Nicholls et al., 2014; Lowe et al., 2015; M. A. Khan
163 et al., 2019) and in a regional nested configuration of the European Monitoring and Eval-
164 uation Programme chemical transport model for the UK (EMEP4UK) (Hood et al., 2018).
165 However, to the best of our knowledge, it has never been used in a global chemistry-climate
166 model. In this paper we document the implementation of the CRIv2-R5 mechanism in
167 the United Kingdom Chemistry and Aerosol (UKCA) model (Morgenstern et al., 2009;
168 O'Connor et al., 2014), as used in the Met Office Unified Model (UM) and the UK's Earth
169 System Model (UKESM1) (Sellar et al., 2019), evaluate it against a suite of observations
170 and rigorously compare the new mechanism to the existing chemical mechanism (Strat-
171 Trop; Archibald et al. (2020)). We note that there are differences in the reaction rate
172 coefficients in the two mechanisms that reflect their independent development and re-
173liance on different assessments of kinetic parameters, which has a bearing on model sim-
174 ulations (Newsome & Evans, 2017). We also explore how differences in the allocation of
175 NMVOC emissions in the two mechanisms contributes to the differences between them.

176 The UKESM1 model is used for quantifying and understanding climate forcing, in-
177 cluding as part of the Coupled Model Intercomparison Project Phase 6 (CMIP6) (Eyring

178 et al., 2016), making projections of future air quality (including crop yields and human
179 health impacts), and increasingly being used to quantify impacts of mitigation. The im-
180 plementation of the CRI mechanism into the UKCA, UM and UKESM1 models repre-
181 sents a step change the potential for simulating the complex chemistry-climate interac-
182 tions between ozone and NMVOCs in the coupled Earth System across many chemical
183 environments and multi-century timescales.

184 2 Model Description

185 2.1 The UKCA model

186 The United Kingdom Chemistry and Aerosols (UKCA) model is a sub-model of
187 the Met Office Unified Model (UM) and is designed to simulate atmospheric composi-
188 tion for weather and climate modelling. UKCA is a part of the UKESM1 (Sellar et al.,
189 2019) Earth system model, and uses the Chemistry of Stratosphere and Troposphere (Strat-
190 Trop) chemical mechanism (Archibald et al., 2020), which merges the Stratospheric and
191 Tropospheric chemical mechanisms described by (Morgenstern et al., 2009) and (O’Connor
192 et al., 2014), respectively. The UKCA model provides ozone, methane, and nitrous ox-
193 ide fields to the UM radiation scheme, as well as calculating oxidant fields that are used
194 to drive the GLOMAP-mode aerosol scheme (Mann et al., 2010; Mulcahy et al., 2018,
195 2020).

196 The StratTrop chemical mechanism and its implementation in UKESM1 is described
197 in detail by Archibald et al. (2020). It uses the Fast-JX photolysis scheme (Neu et al.,
198 2007), which was implemented in UKCA as described by Telford et al. (2013). The ASAD
199 chemical mechanism framework (Carver et al., 1997) is used to provide a flexible and ex-
200 tendable approach to mechanism development and to enable a choice of numerical inte-
201 gration schemes. A sparse-matrix Newton-Raphson chemical solver (Wild & Prather,
202 2000) is used here, applying the quasi-Newton approximations recommended by Esenturk
203 et al. (2018) to reduce run time. Wet deposition of soluble chemical compounds is pa-
204 rameterised following Giannakopoulos et al. (1999) and dry deposition is based on the
205 resistance type model of Wesley (1989), as described by O’Connor et al. (2014).

206 2.2 The Common Representative Intermediates Mechanism

207 The Common Representative Intermediates (CRI) Mechanism has been described
208 in detail in previous papers (M. E. Jenkin et al., 2002; M. Jenkin et al., 2008; Watson
209 et al., 2008; S. R. Utembe et al., 2009), so this section only covers a brief overview of the
210 design philosophy. CRI is a fully traceable, reduced complexity representation of the Mas-
211 ter Chemical Mechanism (MCM) (M. E. Jenkin et al., 2002; Saunders et al., 2003), with
212 CRI version 2 optimised against the MCM version 3.1. The chemistry of inorganic com-
213 pounds and smaller initial organic molecules (such as methane, ethane and ethene) is
214 functionally identical in CRI to the full MCM. However, the CRI scheme substantially
215 reduces the total number of species and reactions by lumping the intermediate oxida-
216 tion products of larger NMVOC species based on an index defined as the total number
217 of carbon-carbon (C – C) and carbon-hydrogen (C – H) bonds, counting double bonds
218 (C = C) as two. This “CRI index” can be understood as the maximum potential num-
219 ber of O_x molecules ($O_x = O_3 + NO_2$) generated by the VOC in question, assuming com-
220 plete oxidation to CO_2 and H_2O , with every HO_2 and RO_2 molecule created convert-
221 ing one NO molecule to NO_2 , thereby generating one ozone molecule when the NO_2 is
222 photolysed (M. E. Jenkin et al., 2002). This means that every primary NMVOC species
223 in CRI will produce the same number of ozone molecules as its equivalent in the MCM,
224 even though the mechanism is greatly simplified by lumping together similar interme-
225 diate species. The intermediate species are named according to their structure, CRI in-
226 dex and functional group as explained in the supplementary Table S1. Using this lump-
227 ing method, the number of species and reactions in the CRIv2 mechanism is reduced to

228 434 species and 1183 reactions from 4500 species and ≈ 12600 reactions in MCMv3.1, cov-
 229 ering the degradation of the same 115 emitted NMVOC species without compromising
 230 the mechanism's ability to simulate ozone production.

231 The CRIV2 scheme also underwent several further stages of complexity reduction
 232 by lumping together emitted species, with the reduced mechanisms evaluated against the
 233 MCM at each stage to preserve ozone forming potential, as described by Watson et al.
 234 (2008). The version implemented in the UKCA model is the simplest of these reductions,
 235 reduction number five (CRIV2-R5) with 196 species, the same version as implemented
 236 into the WRF-Chem model (Archer-Nicholls et al., 2014) and STOCHEM-CRI model
 237 (S. Utembe et al., 2010; M. A. H. Khan et al., 2015). This version includes the full CRIV2
 238 degradation of isoprene, α -pinene, β -pinene and several aromatic species known to con-
 239 tribute to secondary organic aerosol (SOA) production, as described by S. R. Utembe
 240 et al. (2009).

241 3 Implementation of CRIV2-R5 in UKCA

242 The CRIV2-R5 chemical mechanism was designed for use in boundary layer/urban
 243 air quality scenarios. Compared to the StratTrop mechanism used in UKCA, it repre-
 244 sents the chemistry of the following primary NMVOCs and their products which either
 245 do not exist or are heavily parameterised in StratTrop:

- 246 1. C4 alkane chemistry via n-butane (C4H10)
- 247 2. C2-C4 alkene chemistry via ethene, propene and trans-2-butene (C2H4, C3H6 and
 248 TBUT2ENE).
- 249 3. Alkyne chemistry via C2H2.
- 250 4. Expanded biogenic volatile organic compound (BVOC) chemistry with explicit iso-
 251 prene (C5H8) degradation and additional monoterpene chemistry via α -pinene and
 252 β -pinene (APINENE and BPINENE).
- 253 5. Aromatic chemistry via benzene, toluene and o-xylene (BENZENE, TOLUENE
 254 and OXYLENE).
- 255 6. Higher oxidised NMVOC chemistry with the addition of Ethanol (EtOH), propanal
 256 (EtCHO) and methyl ethyl ketone (MEK).

257 In addition, it expands on several key chemical processes, for example all peroxy rad-
 258 icals (RO_2) can react with all other peroxy radicals ($\text{RO}_2 + \text{R}'\text{O}_2$) and most can form
 259 organonitrates from $\text{RO}_2 + \text{NO}$ reactions. However, CRIV2-R5 lacks key photolysis re-
 260 actions and chemical species which are important for the chemistry of the upper atmo-
 261 sphere. The UKCA chemistry-aerosol model is designed to be used in the UK Met Of-
 262 fice Unified Model (UM) and UK Earth System Model (UKESM1) configurations (Sellar
 263 et al., 2019), which simulate the whole atmosphere up to a model top of 85 km in the
 264 lower mesosphere. Hence, some changes were made to make the CRIV2-R5 scheme suit-
 265 able for simulating stratospheric chemical conditions as well as the troposphere. In all,
 266 the following development tasks were made to import the CRIV2-R5 mechanism into the
 267 UKCA model such that it was suitable for use in a 3D Earth System model configura-
 268 tion:

- 269 1. Capability for peroxy radical self- and cross-reactions with summed RO_2 pool.
- 270 2. Adding stratospheric chemistry, using the same species and reactions as used in
 271 the StratTrop mechanism.
- 272 3. Coupling with the GLOMAP aerosol mechanism.
- 273 4. Linking with online Fast-JX photolysis.
- 274 5. Wet and dry deposition of species added.

Table 1. Comparison of gas-phase chemical mechanisms: the StratTrop chemical mechanism as described by (Archibald et al., 2020); the CRIv2-R5 mechanism used as the basis of development as described by M. Jenkin et al. (2008); Watson et al. (2008); S. R. Utembe et al. (2009); CRI-Strat which is CRIv2-R5 combined with species and reactions needed for simulating the stratosphere taken from the StratTrop mechanism; and CRI-Strat+GLOMAP-mode which further includes species and reactions needed for aerosol production.

	StratTrop + GLOMAP-mode	CRI-v2-R5	CRI-Strat	CRI-Strat + GLOMAP-mode
No. Species	87	198	219	233
No. Tracers	83	146	167	181
No. Non transported prognostics	4	52	52	52
No. Peroxy radicals*	9	47	47	47
No. Emitted species	23	27	27	38
No. Photolysis reactions	60	100	124	128
No. Bimolecular reactions	212	451	536	554
No. Termolecular reactions	25	29	36	39
No. Heterogeneous reactions	8	0	5	8
No. Wet deposited species	34	74	80	83
No. Dry deposited species	41	124	128	131

* Peroxy radicals are transported tracers in the StratTrop mechanism,

275 Table 1 gives a summary of the chemical mechanisms as integrated into the UKCA
 276 model, with the CRI-Strat mechanism with coupling to GLOMAP-mode aerosol being
 277 the full mechanism described and analysed in this paper. The CRI-Strat mechanism was
 278 implemented into UKCA using the ASAD framework (Carver et al., 1997), meaning it
 279 can now easily be ported to other models which share the same framework such as TOM-
 280 CAT/SLIMCAT chemical transport model (CTM) (Chipperfield, 2006), the offline GLObal
 281 Model of Aerosol Processes (GLOMAP) (Spracklen et al., 2006) or the Frontier Research
 282 System for Global Change (FRSGC) version of the University of California, Irvine (UCI)
 283 global CTM(CTM) (Wild et al., 2000). The full CRI-Strat+GLOMAP-mode mechanism
 284 has over twice the number of species (83 to 181) and reactions (305 to 729) as the Strat-
 285 Trop mechanism. However, the run time is only about 75% longer, hence achieving a greater
 286 than doubling of complexity for less than double the cost (details in Section S1 of the
 287 supplement). While still computationally expensive and not designed to replace Strat-
 288 Trop for all model studies, multi-century Earth System model simulations with the mech-
 289 anism are plausible.

290 The CRIv2-R5 mechanism used to build CRI-Strat was originally optimised against
 291 the MCMv3.1 (M. E. Jenkin et al., 2003; Saunders et al., 2003), which drew heavily on
 292 kinetic parameters evaluated by the International Union of Pure and Applied Chemistry
 293 (IUPAC) Task Group on Atmospheric Chemical Kinetic Data Evaluation (e.g., Atkinson
 294 et al. (1997, 2004)). In contrast, the StratTrop scheme (Archibald et al., 2020) draws on
 295 a mixture of data from the MCMv3.2 website, the IUPAC Task Group web pages and
 296 the NASA JPL Evaluation No. 17 (S. P. Sander et al., 2011). The reaction rate coef-
 297 ficients for common reactions therefore do not always agree, including for some reactions
 298 which are extremely important for tropospheric composition. In some cases CRI-Strat
 299 is out of date, in others the mechanisms are simply different as they have different sources.
 300 Key differences in rate coefficients are documented in Section S4 of the supplement. The
 301 CRI version 2.2 mechanism was recently released (M. E. Jenkin, Khan, et al., 2019) and,
 302 among other developments, had some of its reaction rates coefficients revised to match

303 recent assessments to be consistent with MCMv3.3.1 (M. E. Jenkin et al., 2015). Un-
304 fortunately, this release was too late to be integrated into the development cycle for the
305 mechanism presented in this paper. However, updating the mechanism to CRI version
306 2.2 is part of ongoing work.

307 The remainder of this section describes the developments and implementation of
308 the CRI-Strat mechanism in detail.

309 **3.1 Peroxy Radical Chemistry**

310 Peroxy radicals (RO_2), a class of short-lived compounds formed during oxidation
311 of VOCs, play a crucial role in the formation of tropospheric ozone (Lightfoot et al., 1992)
312 and secondary organic aerosols (Bianchi et al., 2019; Mcfiggans et al., 2019). While in
313 high- NO_x environments they tend to react with NO , forming NO_2 and ozone (M. Jenkin
314 & Clemitshaw, 2000; Monks, 2005), in low- NO_x environments they usually react either
315 with HO_2 to form hydroperoxides (ROOH), themselves (self-reactions) or other RO_2 species
316 (cross-reactions) (Tyndall et al., 2001; Orlando & Tyndall, 2012). However, because in-
317 individually simulating the reaction of each RO_2 species with every other RO_2 species for
318 the 47 RO_2 species in CRIv2-R5 would be prohibitively expensive, the approximation
319 described by M. Jenkin et al. (1997) is used, whereby each peroxy radical undergoes a
320 reaction with the summed total of all peroxy radical species (the “ RO_2 pool”). The sec-
321 ond order reaction rate coefficient is calculated as the geometric mean of the self-reaction
322 rate of the peroxy radical species in question and the $\text{CH}_3\text{O}_2 + \text{CH}_3\text{O}_2$ reaction rate. The
323 RO_2 pool is not consumed by these reactions, hence these RO_2 -permutation reactions
324 can be thought of as pseudo-unimolecular reactions with a first order rate coefficient pro-
325 portional to the total RO_2 concentration.

326 To enable this chemistry within the framework of the UKCA model, peroxy rad-
327 icals were defined as a unique type - that of a non transported prognostic field. Within
328 the ASAD framework and chemical solver, peroxy radicals are treated like other chem-
329 ical tracers, but in the rest of the model they are not transported as this is unnecessary
330 for such short lived species. Avoiding the transportation of the 47 peroxy radical species
331 offers substantial savings as some of the largest computational costs of the UKCA model
332 come from tracer transport (Esenturk et al., 2018). The definition of peroxy radicals as
333 a unique type also provides a simple method of tagging the species for their concentra-
334 tions to be summed when calculating the total RO_2 pool. The RO_2 pool is defined as
335 a field that can affect the rates of reactions but does not have its concentration directly
336 changed from chemical reactions. Instead, its concentration is calculated for each grid-
337 cell and is updated as the concentrations of constituent RO_2 species change. Because
338 ASAD uses an implicit backward euler solver with Newton Raphson iteration (Carver
339 et al., 1997; Wild et al., 2000), the RO_2 pool is recalculated with each iteration of the
340 solver, not just at each timestep.

341 Overall, this method enables efficient representation of peroxy radical chemistry
342 without adding an excessive number of reactions or transported tracers. This framework
343 can be built on further, for example to parameterise the formation of RO_2 accretion prod-
344 ucts which are important for the formation of highly oxidised organic material (HOM)
345 (Weber et al., 2020).

346 **3.2 Merging with Stratospheric Chemistry**

347 As both the MCM and CRI mechanisms are designed to simulate polluted bound-
348 ary layer chemistry, they lack many reactions which are not relevant in the boundary layer
349 but are important in the upper troposphere and stratosphere due to the differing pho-
350 tochemical conditions at different altitudes. The UKCA model is used as part of UKESM1
351 (Sellar et al., 2019) with a model top at 85 km in the standard configuration, therefore

352 it is essential that the chemical mechanism can adequately represent both tropospheric
353 and stratospheric chemistry.

354 The CRIv2-R5 scheme was merged with the Stratospheric chemistry in UKCA, de-
355 scribed by Morgenstern et al. (2009) and updated into the StratTrop mechanism by Archibald
356 et al. (2020), to produce the new CRI-Strat mechanism (see Table 1). In total, 20 species
357 and 121 reactions that are important for stratospheric chemistry were added. Some of
358 the added reactions involve chemical species already present in the CRIv2-R5 mecha-
359 nism. In all cases, imported reactions used the same rate coefficients or cross sections
360 as the equivalent reactions in StratTrop. Full details of the CRI-Strat mechanism are
361 given in Tables S2-S5 of the supplement.

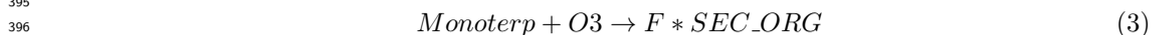
362 Photolysis reactions were added for methyl peroxy nitrate ($\text{CH}_3\text{O}_2\text{NO}_2$, MeO_2NO_2
363 in CRI-Strat) using the same cross sections as for HO_2NO_2 (Browne et al., 2011). The
364 species $\text{CH}_3\text{O}_2\text{NO}_2$ is not part of the StratTrop mechanism but is an important reser-
365 voir for NO_x in cold temperatures and so photolysis of this species is needed to prevent
366 accumulation in the upper troposphere and stratosphere. Reactions forming water vapour
367 with a sizeable flux in the upper atmosphere were adjusted to ensure that H_2O was spec-
368 ified as a product; while chemical production of water vapour is typically neglected in
369 the troposphere it forms an important part of the budget in the stratosphere.

370 3.3 Coupling with GLOMAP Aerosols

371 The UKCA model incorporates a modal representation of aerosol size distribution
372 using the two-moment aerosol microphysics scheme from the Global Model of Aerosol
373 Processes (GLOMAP-mode) (Mann et al., 2010) for all aerosol components (sulphate,
374 sea-salt, black carbon and organic carbon, but not currently ammonium nitrate), with
375 the exception of mineral dust which employs a bin scheme as described by Woodward
376 (2001). GLOMAP-mode is fully coupled with the StratTrop mechanism in UKCA (Archibald
377 et al., 2020) as described by (Mulcahy et al., 2020).

378 To couple the CRI-Strat mechanism to GLOMAP-mode aerosol, the standard Strat-
379 Trop couplings have been mimicked wherever possible, except for secondary organic aerosol
380 (SOA) and dimethyl sulphide (DMS) chemistry as discussed below. Because the oxidant
381 fields are different in CRI-Strat, the production of aerosols may differ greatly between
382 CRI-Strat and StratTrop. The species and reactions in CRI-Strat needed for coupling
383 with GLOMAP-mode are marked with an ^A in supplementary Tables S2-S6. As the main
384 focus of this paper is to evaluate changes in the gas phase due to implementation of the
385 CRI-Strat mechanism, here we only document how the aerosol is coupled. Evaluating
386 the changes to the aerosol fields and their impacts on atmospheric composition and cli-
387 mate will be the subject of a follow up paper.

388 Formation of SOA is parameterised in GLOMAP-mode through formation of the
389 SEC_ORG tracer, which represents the extremely low volatility products of biogenic volatile
390 organic compound (BVOC) oxidation. SEC_ORG does not undergo any further chem-
391 istry, and is either permanently condensed to the aerosol phase as organic aerosol or is
392 lost via deposition processes. When the StratTrop mechanism is coupled with GLOMAP-
393 mode aerosol, SEC_ORG is formed via the reactions:



399 with rate coefficients equal to the equivalent oxidation reactions of α -pinene with OH,
400 O_3 and NO_3 respectively. The factor F represents the yield of SEC_ORG from Monoterp
401 and is set at runtime to be equal to 26% for all of the above reactions, based on a 13%
402 yield from α -pinene (Tunved et al., 2004; Spracklen et al., 2006) and doubled to account

403 for a lack of SOA production from isoprene and anthropogenic species (Mulcahy et al.,
404 2020).

405 To simulate formation of SEC_ORG in the CRI-Strat mechanism, all of the AP-
406 INENE and BPINENE initial oxidation reactions produce SEC_ORG at the same yield
407 as the StratTrop reactions with Monoterp, in addition to all of the gas-phase products
408 which are important for ozone chemistry (see supplementary Table S5). This method
409 is not ideal as it does not conserve carbon. However, it is designed to produce a simi-
410 lar amount of SEC_ORG as the StratTrop mechanism in order to enable fair compar-
411 ison of the gas-phase chemistry between the mechanisms and it serves as a placeholder
412 until a more complete coupling between gas-phase chemistry and aerosol can be devel-
413 oped. Working on more explicit couplings between the organic gas-phase chemistry and
414 aerosol routines is ongoing and will build on CRI version 2.2 (M. E. Jenkin, Khan, et
415 al., 2019) and CRIV2.2 with Highly Oxygenated Organic Material (CRI_HOM)(Weber
416 et al., 2020) versions of the mechanism. In the long term, this potential for more real-
417 istic chemical coupling between gas-phase organic chemistry and SOA formation is one
418 of the key advantages of using a semi-explicit mechanism such as CRI-Strat over a sim-
419 pler mechanism such as StratTrop.

420 Dimethyl sulphide (DMS) is a sulphur containing compound emitted in large quan-
421 tities by natural sources and is a critically important source of SO₂, particularly in the
422 preindustrial atmosphere, which can be oxidised to form sulphate aerosols (Andreae, 1990).
423 The DMS chemistry used here is the same as in the CRIV2-R5 implementation in the
424 WRF-Chem model (Archer-Nicholls et al., 2014) and uses the von Glasow and Crutzen
425 (2004) DMS chemistry scheme. This description of multi-generational DMS oxidation
426 is more comprehensive than that used in StratTrop (Archibald et al., 2020) and fits bet-
427 ter with the complexity of the rest of the CRI-Strat mechanism. Evaluating and updat-
428 ing the DMS chemistry in both the CRI-Strat and StratTrop mechanisms is part of on-
429 going work.

430 3.4 Coupling with FastJ-X photolysis

431 The Fast-JX photolysis scheme implemented in UKCA calculates photolysis rates
432 (“j” rates) for each reaction based on experimentally determined cross sections and quan-
433 tum yields at a range of wavelength bins (Telford et al., 2013). The CRI-Strat mecha-
434 nism includes many more photolysis reactions than StratTrop (128 compared to 60). How-
435 ever, most of the added species do not have experimentally determined photolysis cross
436 sections and quantum yields. In the original CRI scheme, used in a box model, the pho-
437 tolysis rates for these species were calculated using a two stream isotropic photolysis scheme
438 described by Hayman (1997). When the mechanism was ported to WRF-Chem (Archer-
439 Nicholls et al., 2014), photolysis cross section and quantum yield data were adopted from
440 those of species that were already available in WRF-Chem. However, for new species,
441 a box model was used to generate photolysis rate profiles as a function of solar zenith
442 angle. Profiles with similar shapes generated in WRF-Chem were then scaled to match
443 the profiles from the box model. These scaling factors were then applied to the corre-
444 sponding cross section and quantum yield data to calculate surrogate photolysis rates
445 for the new species.

446 For this implementation, cross sections already available in UKCA are used for added
447 photolysis reactions wherever possible, otherwise the surrogate cross sections and scal-
448 ing factors used in the WRF-Chem implementation are applied. References for the pho-
449 tolysis cross-section data are given in Telford et al. (2013). Details of all photolysis re-
450 actions in the CRI-Strat mechanism are provided in supplementary Table S3.

3.5 Wet and dry deposition

There are many new species in the CRIV2-R5 mechanism which are efficiently wet and dry deposited but would otherwise have long chemical lifetimes in the atmosphere. The deposition of new species not included in the StratTrop mechanism is based on the functional group(s) of the species in question. The dry deposition rates and Henry's law coefficients used are described in more detail by O'Connor et al. (2014) and Archibald et al. (2020). Appropriate dry deposition velocities and Henry's Law coefficients were taken from those of existing species in the StratTrop mechanism with a similar functional group or structure; Table 2 shows which surrogate species are used for deposition rate coefficients for each additional species in CRI-Strat. In the case of a species fitting in more than one category, the class with the fastest deposition rates was used. This ensures that all species undergoing deposition are removed at an appropriate rate, but data for specific species can be updated in the future when experimental data becomes available.

4 Methods

4.1 Description of Model Setup

The experiments conducted in this study use the UM version 10.9. The model configuration is based on UKESM1, with 85 vertical levels using terrain-following hybrid height coordinates up to 85 km and a horizontal resolution of $1.25^\circ \times 1.875^\circ$ (N96) (Walters et al., 2019). All of the simulations are run using nudging of wind and temperature fields and prescribed sea surface temperatures (Telford et al., 2008) to the ERA-interim re-analysis product by ECMWF (Dee et al., 2011) to constrain all simulations to observed meteorological evolution so the evaluation can focus on how changes to the chemical mechanism affect atmospheric composition (Zhang et al., 2014). The model simulations run from 1 September 2008 to 1 January 2019, with the analysis period from 1 January 2010 to the end of the runs. Well mixed greenhouse gases are not emitted, rather carbon dioxide levels are set as a constant field while methane, nitrous oxide and CFCs are prescribed with constant lower boundary conditions, all at 2014 levels (Archibald et al., 2020).

4.2 Emissions

The emissions used in this study 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), and can be downloaded from <http://www.globalchange.umd.edu/ceds/ceds-cmip6-data/>. All of the experiments use repeated 2014 emissions as the closest to present day available in the inventories. Anthropogenic emissions are based on the Emission Database for Global Atmospheric Research (EDGARv4.3.1) (<http://edgar.jrc.ec.europa.eu/overview.php?v=431>) across the globe, incorporating more detailed regional datasets where available (Hoesly et al., 2018), while biomass burning emissions for the modern period are largely based on Global Fire Emissions Database version 4 with small fires (GFED4s) (Van Der Werf et al., 2017) and merged with other datasets as described by Van Marle et al. (2017). Combination of these datasets onto a unified grid is described by Feng et al. (2020). Offline biogenic emissions are derived from the Model of Emissions of Gases and Aerosols from Nature (MEGAN) version 2.1 (Guenther et al., 2012). Oceanic BVOC emissions are included from the POET inventory (Olivier et al., 2003).

Table 3 shows the mappings used to link the CEDS emissions to CRI-Strat. Lumping of raw NMVOC species to CRIV2-R5 speciation is based on methods described in (Watson et al., 2008). We also include equivalent mappings to the StratTrop scheme, as well as total global emissions for 2014 in TgC yr⁻¹.

Table 2. CRI-Strat species which undergo wet or dry deposition but do not exist in StratTrop do so using coefficients based on surrogate StratTrop species.

Compound class	Functional group(s) / molecule	Dry/wet/ both	CRIv2-R5 species	StratTrop surrogate
Alcohols	-OH	Both	EtOH, i-PrOH, n-PrOH, AROH14, AROH17, ARNOH14, ARNOH17	MeOH
Aldehydes	-CHO	Dry	HOCH2CHO, CARB14, CARB17, CARB11A, UCARB10, UCARB12, NUCARB12, UDCARB8, UDCARB11, UDCARB14, TNCARB26, TNCARB10, TNCARB12, TNCARB11, CCARB12, TNCARB15, TXCARB24, ANHY, TXCARB22, UDCARB17	MeCHO
Carboxylic Acids	-CO ₂ H	Both	RCOOH25	MeCO ₂ H
Glyoxals	2x -C(O)-	Both	CARB3, CARB6, CARB9, CARB12, CARB15	MGLY
Hydroxy-ketones	-OH and R(O)R'	Both	CARB7, CARB10, CARB13, CARB16	HACET
Hydroxy-nitrates	-OH and -ONO ₂	Dry	HOC2H4NO ₃ , RN9NO ₃ , RN12NO ₃ , RN15NO ₃ , RN18NO ₃ , RU14NO ₃ , RTN28NO ₃ , RTX28NO ₃	ISON
Monoterpenes	C ₁₀ H ₁₆	Dry	APINENE, BPINENE	Monoterp
Nitrate-carbonyls	-ONO ₂ and -C(O)-	Both	NOA, NUCARB12, RTX24NO ₃ , RTN25NO ₃ , RTX22NO ₃ , RTN23NO ₃	NALD
Peroxides	-OOH	Both	HOC2H4OOH, RN10OOH, RN13OOH, RN16OOH, RN19OOH, RN8OOH, RN11OOH, RN14OOH, RN17OOH, RU14OOH, RU12OOH, RU10OOH, NRU14OOH, NRU12OOH, RN9OOH, RN12OOH, RN15OOH, RN18OOH, NRN6OOH, NRN9OOH, NRN12OOH, RA13OOH, RA16OOH, RA19OOH, RTN28OOH, NRTN28OOH, RTN26OOH, RTN25OOH, RTN24OOH, RTN23OOH, RTN14OOH, RTN10OOH, RTX28OOH, RTX24OOH, RTX22OOH, NRTX28OOH	EtOOH
Peroxy Acids	-CO ₃ H	Both	EtCO ₃ H, HOCH ₂ CO ₃ H	MeCO ₃ H
Peroxyacyl Nitrates	-PAN	Dry	PHAN, RU12PAN, RTN26PAN	PAN

Table 3. Mapping of raw CMIP6 NMVOC emissions to CRI-Strat and StratTrop mechanisms, with total emitted carbon mass for 2014 from Anthropogenic, biomass burning and biogenic sources.

Raw emission classes	Anthropogenic TgC yr ⁻¹	Biomass Burning TgC yr ⁻¹	Biogenic TgC yr ⁻¹	CRIv2-R5 Species	StratTrop Species
VOC1: Alcohols	0.4 3.	3.5 0.1	48.5 9.5	MeOH EtOH	MeOH to MeOH
VOC2: Ethane	5.3	2.8	1.0	C2H6	C2H6
VOC7: Ethene	4.9	4.1	25.8	C2H4	to C2H6
VOC9: Ethyne	3.1	1.1	–	C2H2	to C2H6
VOC3: Propane	5.5	0.6	1.0	C3H8	C3H8
VOC8: Propene	3.0	3.5	14.	C3H6	to C3H8
VOC4-6: Butanes and higher alkanes	4.8	0.4	0.1	C4H10	N/A
VOC10: Isoprene	–	0.6	588	C5H8	C5H8
VOC11: Monoterpenes	–	1.2	94.7	67% to APINENE* 33% to BPINENE*	Monoterp [†]
VOC12: Other Alkenes and Alkynes	6.5	0.8	2.6	TBUTZENE	N/A
VOC13: Benzene	6.1	2.0	–	BENZENE	N/A
VOC14: Toluene	7.0	3.9	86.5	TOLUENE	N/A
VOC15-17: Xylenes and higher aromatics	3.1	1.1	–	oXYLENE	N/A
VOC21: Formaldehyde	1.0	2.1	1.8	HCHO	HCHO
VOC22: Other Aldehydes	0.5 0.6	3.4 0.8	10.0 2.0	MeCHO EtCHO	MeCHO to MeCHO
VOC23: Ketones	1.5 1.0	1.1 0.9	22.9 0.5	Me2CO MEK	Me2CO to Me2CO
VOC24: Acids	– 4.4	0.5 7.1	1.4 1.9	HCOOH MeCO2H	N/A [‡]
Total CRI:	70.5	40.6	900.6		
Total StratTrop:	27.9	23.9	710.6		

* Two to one split between APINENE and BPINENE emissions according to Guenther et al. (2012) and Sindelarova et al. (2014).

[†] In the StratTrop mechanism Monoterp only forms Sec.Org and does not contribute to ozone formation. Therefore, emissions mapped to Monoterp are not included in the total NMVOC emissions for the StratTrop mechanism.

[‡] Acids have historically not been mapped to StratTrop mechanism in UKESM1 (Archibald et al., 2020), even though HCOOH and MeCO2H are existing species.

500 The CRI-Strat mechanism utilises a wider breadth of input NMVOC emissions, using
501 almost all of the available data from the CEDS emission database, therefore has a
502 greater amount of total carbon mass emitted compared to an equivalent StratTrop run.
503 For anthropogenic emissions, the most significant changes are due to the addition of aromatic
504 species and C₄ alkane/alkene emissions. For biogenic emissions, the largest change
505 is that monoterpenes are mapped to the chemically reactive APINENE and BPINENE
506 tracers in CRI-Strat, which undergo oxidation and contribute to ozone formation, whereas
507 in StratTrop they are mapped to the MONOTERP tracer which is only considered as
508 a precursor of SOA and does not contribute to ozone production). The total NMVOC
509 emissions for the year 2014 using the CEDS emissions inventory are given in Table 3.

510 4.3 Model Simulations

511 Two base simulations are conducted, labelled StratTrop and CRI-Strat (Table 4).
512 These use the emissions associated with their respective mechanism, as described in Table
513 3. The StratTrop simulation uses a slightly modified version of the mechanism, where
514 the reactions of NO₃ + DMS and NO₃ + Monoterp were adjusted so that they conserved
515 nitrogen, when previously they did not. These changes were made to enable a fair comparison
516 between the mechanisms for Section 5.6. These changes have a minimal impact
517 on the overall chemical composition in StratTrop and are explained in detail in Section S3
518 of the Supplement.

519 The treatment of emissions can be seen as an intrinsic part of a chemical mechanism.
520 However, the emissions of additional NMVOC species in the CRI-Strat simulation which
521 are not represented in StratTrop pose a dilemma when comparing the two mechanisms:
522 are the differences between the simulations due to the different approaches in representing
523 chemistry, there being more reactive carbon available in CRI-Strat, or a combination of
524 these factors? Two additional simulations were therefore designed to better understand the
525 effects of additional NMVOC emissions. A CRI-Strat run was conducted with identical
526 emissions to the StratTrop base run (CRI_Emiss_ST) to isolate the changes just due to
527 the chemical mechanism without any changes in emissions. The final CRI-Strat simulation
528 only uses emissions from NMVOC emission classes that are also used by StratTrop, but
529 these are mapped to the appropriate CRI-Strat species, hence it can be used to identify
530 the effects of changing the speciation of NMVOC emissions without increasing the total
531 carbon mass. The key difference is that C₂ and C₃ species are mapped into different
532 species, hence this scenario is called speciated C2-C3 emissions (CRI_Emiss_C2C3).
533 For example, emissions of ethane, ethene and ethyne are mapped to C2H6, C2H4 and
534 C2H2 respectively in CRI_Emiss_C2C3, rather than being lumped to C2H6 as in the
535 StratTrop and CRI_Emiss_ST scenarios. The MONOTERP tracer is unreactive in StratTrop
536 (it can only be oxidised to form SEC_ORG and cannot contribute to ozone production),
537 therefore all monoterpene emissions are mapped to MONOTERP in CRI_Emiss_ST and
538 CRI_Emiss_C2C3, with the respective reactions copied over from StratTrop. The
539 simulations are summarised in Table 4.

540 5 Results

541 In this paper, we focus on understanding how changes in chemistry affect gas-phase
542 species which are important for describing the global tropospheric composition as drivers
543 to climate: ozone, carbon monoxide, hydroxy radicals, methane lifetime and nitrogen oxides.
544 We present a short evaluation against surface and remote sensing products, but the main
545 focus is on comparing the performance of the CRI-Strat with the StratTrop mechanism,
546 which is already well evaluated (Archibald et al., 2020), and on understanding how
547 these changes are sensitive to the treatment of NMVOC emissions. While the changes
548 in chemistry and oxidant fields are also expected to affect formation of aerosols, these
549 influences will be explored in more detail in follow up studies.

Table 4. Summary of simulations.

Scenario Name	Chemical Mechanism	Emissions	Purpose
CRI-Strat	CRI-Strat	Standard CRI	Base CRI scenario
StratTrop	StratTrop	Standard ST	Base StratTrop scenario
CRI_Emiss_ST	CRI-Strat	Standard ST	Isolating effect of chemical representation without changes to emissions
CRI_Emiss_C2C3	CRI-Strat	Speciated C2-C3 emissions	Isolating effect of NMVOC speciation without changing total emitted mass

550 5.1 Model evaluation

551 5.1.1 Surface ozone

552 Lowest model level ozone concentrations from the base CRI-Strat and StratTrop
 553 model simulations are evaluated against the global dataset of rural surface ozone sen-
 554 sors compiled for the Tropospheric Ozone Assessment report (TOAR) (Schultz et al., 2017).
 555 The TOAR dataset compiles ozone measurements from surface station-based observa-
 556 tions around the globe including data from regional and national air quality networks,
 557 as well as larger global collaborations.

558 The data used in this work are the gridded surface ozone datasets, which include
 559 mean surface ozone concentrations on a $2^\circ \times 2^\circ$ global grid. Schultz et al. (2017) also
 560 provide gridded station means separated by urban or rural classification, based on the
 561 population density, nightlight intensity, and OMI satellite NO_2 column. The model low-
 562 est level ozone concentration was evaluated against the rural mean surface ozone grid-
 563 ded dataset, because the relatively large grid size in the UKCA model ($1.25^\circ \times 1.875^\circ$,
 564 approximately 135 km horizontal grid spacing at the equator) means that urban-scale
 565 chemistry is not well resolved. In all comparisons with TOAR gridded data, the model
 566 output was regridded to the same $2^\circ \times 2^\circ$ grid as the TOAR data before analysis.

567 Figure 1 shows comparison of CRI-Strat and StratTrop model simulations with TOAR
 568 observations across the globe, grouped by region. Further plots comparing model out-
 569 put with TOAR data are included in the supplement Figures S6-S9. The CRI-Strat sim-
 570 ulation has consistently higher surface ozone values across almost the entire world, with
 571 this increase most pronounced in populated regions such as Europe and East Asia. In
 572 more remote regions, the difference between the CRI-Strat and StratTrop simulations
 573 is much smaller. The CRI-Strat and StratTrop simulations follow similar seasonal trends,
 574 showing this variation is driven more by the parent model and shared traits, such as sea-
 575 sonal variation in emissions. In many regions, both simulations are low compared to ob-
 576 servations in winter months and high in summer months. Due to CRI-Strat having higher
 577 ozone in general, the global summer high bias is greater in CRI-Strat (+12.6 ppbv) com-
 578 pared to StratTrop (+7.4 ppbv), but has a smaller negative bias in winter (-4.3 ppbv com-
 579 pared to -9.2ppbv). These results indicate that the CRI-Strat mechanism has a higher
 580 ozone production efficiency than StratTrop.

581 The high bias in surface summer ozone and low bias in winter are likely due to struc-
 582 tural issues in using a coarse, global model as has been documented elsewhere (Young
 583 et al., 2013, 2018; Archibald et al., 2020). For example, the coarse horizontal resolution
 584 results in the emissions being smoothed and less heterogeneous, particularly around ur-
 585 ban areas and large point sources, leading to greater mixing of NO_x and NMVOCs and
 586 ozone production, particularly in summer (Wild & Prather, 2006; Stock et al., 2014; Fenech
 587 et al., 2018). The low bias in winter, when local production is lower, may be due to in-

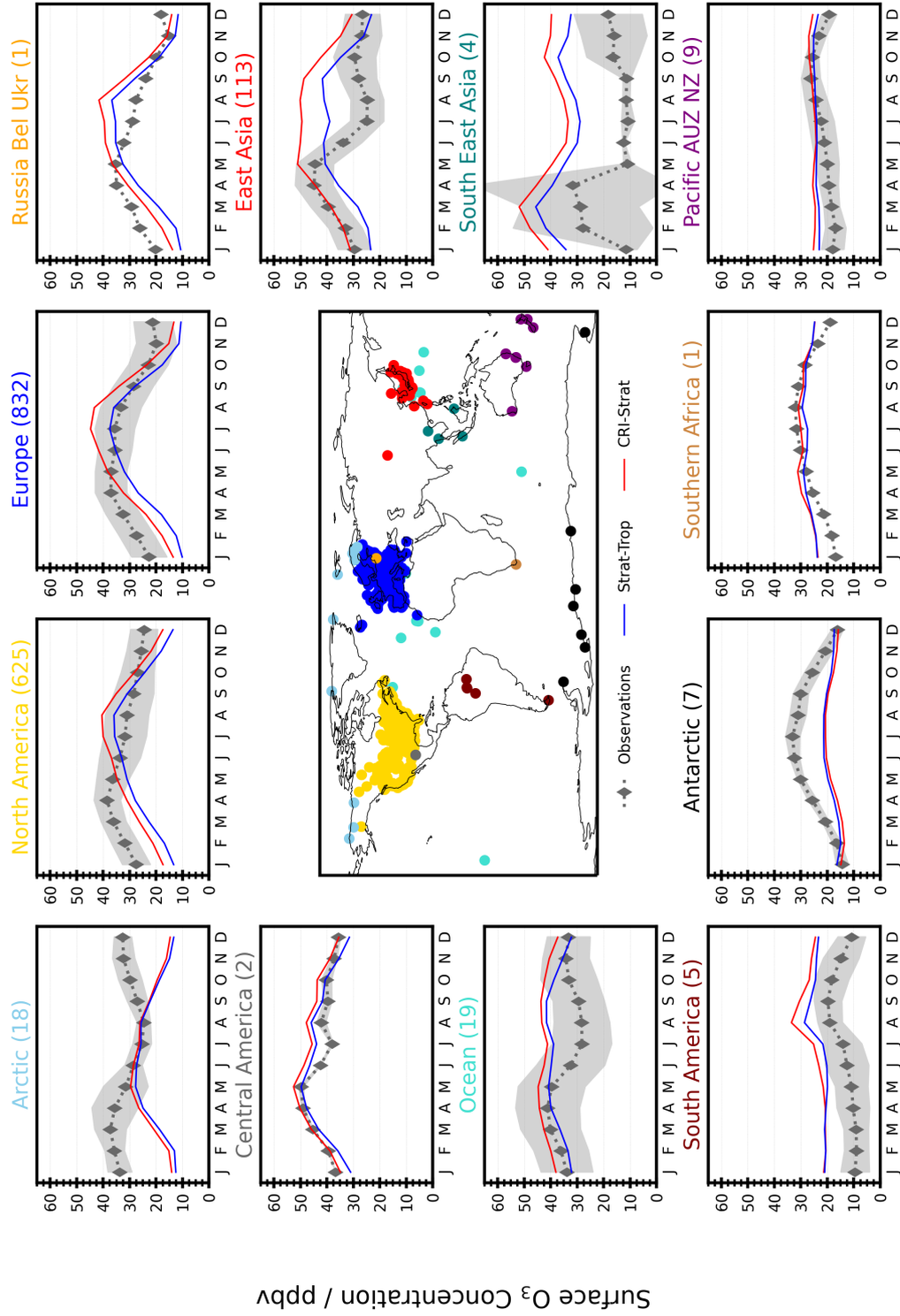


Figure 1. Monthly average surface ozone concentrations from low elevation rural sites on the TOAR network using data from 2010 to 2018. Data is grouped by region, with number of stations in brackets, and compared with corresponding locations from StratTrop (blue) and CRI-Strat (red) model simulations.

588 sufficient long range transport of ozone or loss due to deposition being too great. In some
 589 regions, such as East Asia, the seasonal cycle in the model simulations is out of phase,
 590 with the model runs showing peak ozone in the summer when the observations are at
 591 their lowest. This is likely indicative of missing model processes, such as heterogeneous
 592 chemistry or poor representations of local weather processes, such as monsoon cycles.
 593 Any biases caused by such structural weaknesses in the parent model are common to both
 594 mechanisms, but may be accentuated in CRI-Strat due to greater ozone production when
 595 photochemical conditions allow. Given these issues, the higher bias in CRI-Strat com-
 596 pared to the TOAR surface ozone observations likely more clearly reveal other biases in
 597 the model which compensate for errors associated with the less accurate description of
 598 the underlying chemistry in StratTrop.

599 **5.1.2 Surface carbon monoxide**

600 Modeled carbon monoxide (CO) surface fields are evaluated against the National
 601 Oceanic and Atmospheric Administration (NOAA) climate Monitoring and Diagnostics
 602 Laboratory (CMDL) dataset (Pétron et al., 2002), as shown in Figure 2. The NOAA data
 603 are derived from regular *in situ* flask samples and screened for local pollution events. The
 604 observational data evaluated here are calculated as a monthly average climatology de-
 605 rived from observations made between 2010-15. The StratTrop simulation shows a low
 606 bias at most sites in the Northern hemisphere, which are more influenced by anthropogenic
 607 pollution, but is close to observed values in the tropics and over the southern hemisphere
 608 which are more remote and dominated by biogenic and biomass burning emissions. These
 609 effects have been documented elsewhere (Archibald et al., 2020). The CRI-Strat simu-
 610 lation has greater CO at all sites and over all seasons compared to StratTrop. This re-
 611 duces the size of the negative bias at northern hemisphere sites but creates a positive
 612 bias over the remote southern hemisphere sites. The seasonal trends in both simulations
 613 are similar, showing that these are more sensitive to model dynamics and seasonal vari-
 614 ation in emissions than chemical mechanism. The higher CO in CRI-Strat is due to sec-
 615 ondary production of CO from oxidation of the additional NMVOCs but this has mixed
 616 effects on the comparison with observations, improving biases in the northern hemisphere
 617 whilst making them worse in the southern hemisphere.

618 **5.1.3 Tropospheric ozone column**

619 Although surface ozone is a pressing concern for air quality, the influence of ozone
 620 on climate is more dependent on ozone in the upper troposphere and the tropospheric
 621 ozone burden (Bowman et al., 2013). We use two different monthly mean gridded satel-
 622 lite data products to evaluate tropospheric column ozone (TCO) in the CRI-Strat and
 623 StratTrop simulations. The OMI-MLS TCO monthly gridded data, determined by sub-
 624 tracting the Microwave Limb Sounder (MLS) stratospheric column ozone (Waters et al.,
 625 2006) from the Ozone Monitoring instrument (OMI) total column ozone (Dobber et al.,
 626 2006), is available between 60°S-60°N with a horizontal resolution of $1^\circ \times 1.25^\circ$, as de-
 627 scribed by Ziemke et al. (2006, 2019). For comparison with OMI-MLS data (2010-2018),
 628 the modelled TCO is calculated by vertically integrating the model ozone between the
 629 surface and the tropopause (defined as $380K + 2$ PV; Hoerling et al. (1993)). The OMI
 630 data (Boersma et al., 2007) were produced by the Remote-Sensing Group at Rutherford
 631 Appleton Laboratory using a profile retrieval scheme developed first for GOME-2 (Miles
 632 et al. 2015). Individual profiles were gridded on a monthly basis with a horizontal res-
 633 olution of $1.5^\circ \times 1.5^\circ$ and a correction applied in each layer for bias with respect to ozoneson-
 634 des which had been derived as a function of month of year and latitude (R. Siddans pri-
 635 vate communication). For comparison with OMI data (2010-2017) in the surface-450 hPa
 636 and 450-170 hPa layers, we use monthly, gridded averaging kernels and *a priori* infor-
 637 mation to minimise vertical sampling differences between OMI and UKCA. Although
 638 potential issues with using monthly mean rather than individual averaging kernels can

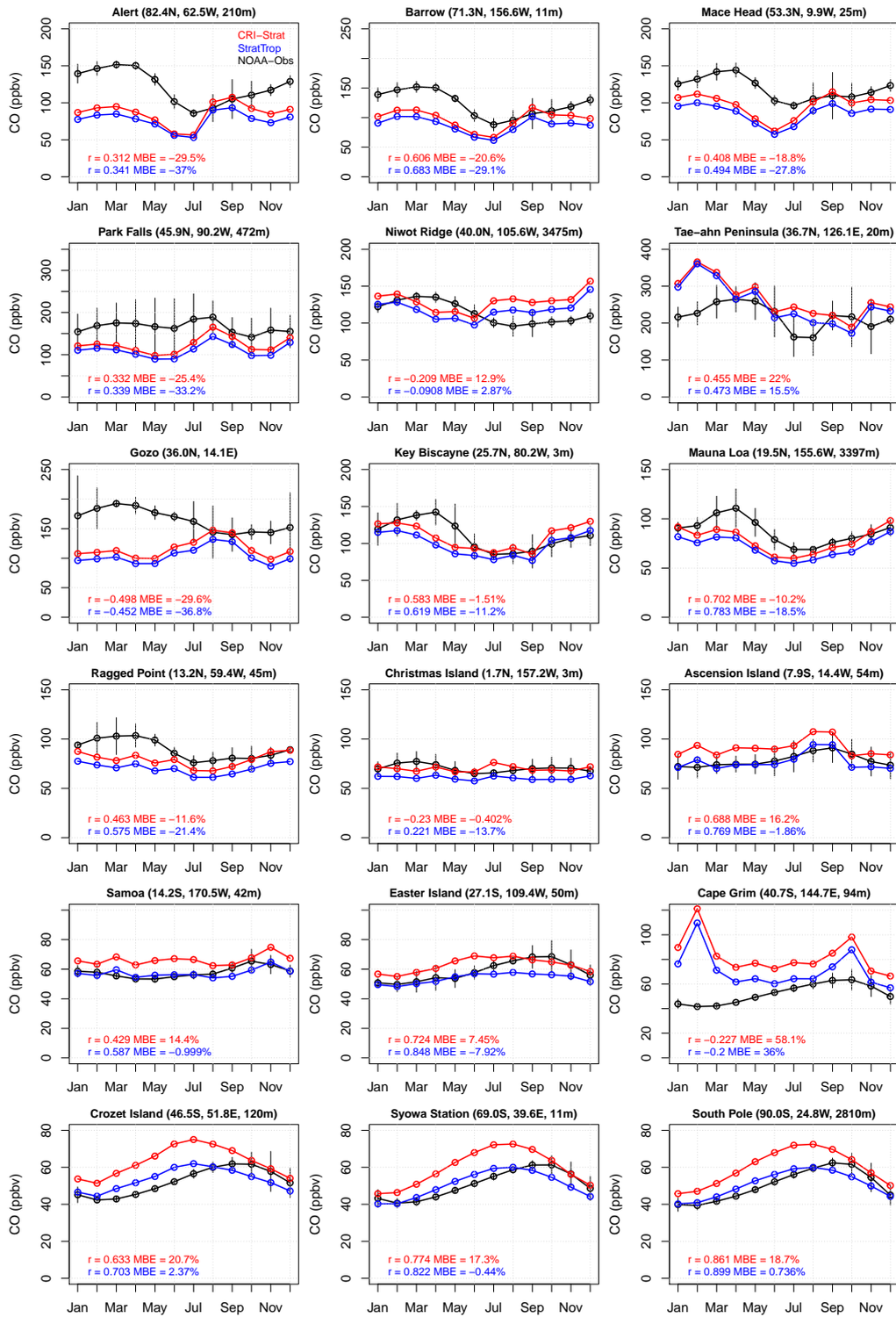


Figure 2. Average surface CO concentrations from CMDL network (black) compared with CRI-Strat (red) and StratTrop (blue) model simulations, showing average seasonal variation, correlation coefficient and mean bias error over 2010-2018 evaluation period.

639 arise for certain species and instruments (von Clarmann & Glatthor, 2019), agreement
 640 between model and observation are found to be improved substantially through appli-
 641 cation of monthly mean averaging kernels in this analysis.

642 In Figure 3, average tropospheric ozone column is compared between the OMI-MLS
 643 satellite product and the CRI-Strat and StratTrop simulations. Differences in tropospheric
 644 ozone column between each of the model simulations are presented in supplementary Fig-
 645 ure S10. The tropospheric ozone column is remarkably similar in the CRI-Strat and Strat-
 646 Trop simulations, both showing a similar high bias over the tropics and a low bias at high
 647 latitudes (Figure 3 (c, e)). The CRI-Strat and StratTrop simulations also have similar
 648 tropospheric ozone burden between 60°S and 60°N (303 Tg and 308 Tg respectively, com-
 649 pared to 301 Tg from OMI-MLS).

650 The seasonal zonal mean ozone in Figure 3 (b, d, f) is also more similar in the two
 651 model simulations compared to OMI-MLS. Both mechanisms show a clear high bias over
 652 the tropics throughout the year, with a strong peak between 0 and 30°N between March
 653 and June. In contrast, the satellite product has higher ozone column values around 40°N
 654 in June-July, with a slightly smaller peak around 30°S in October-November, a pattern
 655 not represented in the model simulations. Considering the large differences in surface
 656 ozone between the two mechanisms (Figure 1) it is perhaps surprising how similar the
 657 total tropospheric ozone is between the two mechanisms, and this is probably a sign that
 658 common model deficiencies (such as errors in emissions or transport) cause similar bi-
 659 ases in both simulations. However, these tropospheric column comparisons may hide sig-
 660 nificant differences in vertical distribution of ozone.

661 The high bias in tropospheric ozone column seen over equatorial regions in both
 662 model simulations (Fig. 3 c-f) is absent in the lower troposphere (Fig. 4 c, e), but ap-
 663 pears in the upper troposphere (Fig. 4 d, f). As the mechanisms show a similar bias in
 664 this region of the atmosphere, similar structural weaknesses must be causing the bias in
 665 both simulations, likely contenders being errors in lightning NO_x emissions (Banerjee
 666 et al., 2014) or convective transport (Hoyle et al., 2011). CRI-Strat has higher ozone columns
 667 than StratTrop above polluted regions such as India and downwind of Europe, mostly
 668 in the lower troposphere, but has reduced ozone compared to StratTrop in the less pol-
 669 luted southern hemisphere (4 g, h).

670 **5.2 Comparison of the ozone budget and processes in CRI-Strat and Strat-** 671 **Trop**

672 In this section, we investigate why tropospheric chemical composition differs be-
 673 tween the CRI-Strat and StratTrop mechanisms. This is assessed using the base CRI-
 674 Strat and StratTrop simulations, and the two CRI-Strat simulations with modified NMVOC
 675 emissions (CRI_Emiss_ST and CRI_Emiss_C2C3). Collectively, these simulations allow
 676 us to explore the impact of changing chemistry without changing emissions, the differ-
 677 ence attributable to explicit speciation of emissions, and the impact of the additional chem-
 678 istry of C₄ alkanes and alkenes, aromatics and monoterpene. Given that the StratTrop
 679 mechanism does not use a large fraction of the reactive carbon mass included in the in-
 680 put inventory, this analysis may also illustrate the causes of certain biases in the Strat-
 681 Trop mechanism and how these might be improved if it were to be modified to use a wider
 682 selection of NMVOC species, informing future development of this mechanism.

683 **5.2.1 Ozone comparison**

684 Boundary layer ozone is much higher over land in CRI-Strat compared to Strat-
 685 Trop, but is lower over remote oceans (Figure 5). Ozone levels are mostly lower in CRI_Emiss_ST
 686 and CRI_Emiss_C2C3 than StratTrop, but are higher over polluted regions. The vari-
 687 ability in ozone is greater in CRI_Emiss_C2C3 than in CRI_Emiss_ST, and there are higher

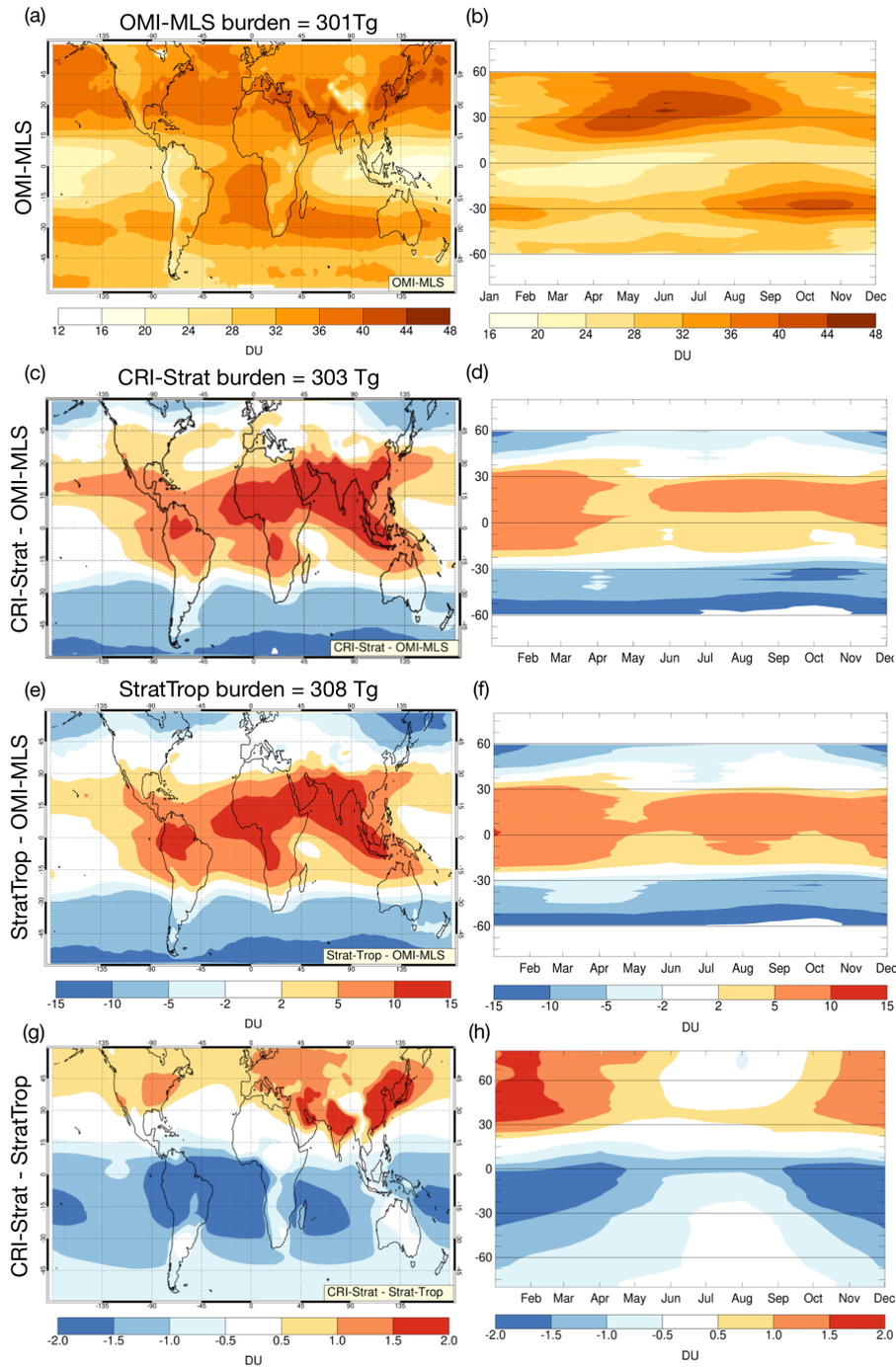


Figure 3. Map of tropospheric ozone column (DU) from OMI-MLS (a) and annual variation in zonal means (b), averaged between 2010-2018. Difference in tropospheric ozone column between CRI-Strat and OMI-MLS (c, d), between StratTrop and OMI-MLS (e, f), and between CRI-Strat and StratTrop noting different colour scale (g, h). Burdens given above panels a, c and e are calculated by summing ozone over the troposphere between 60°S to 60°N as this is the range covered by the OMI-MLS product.

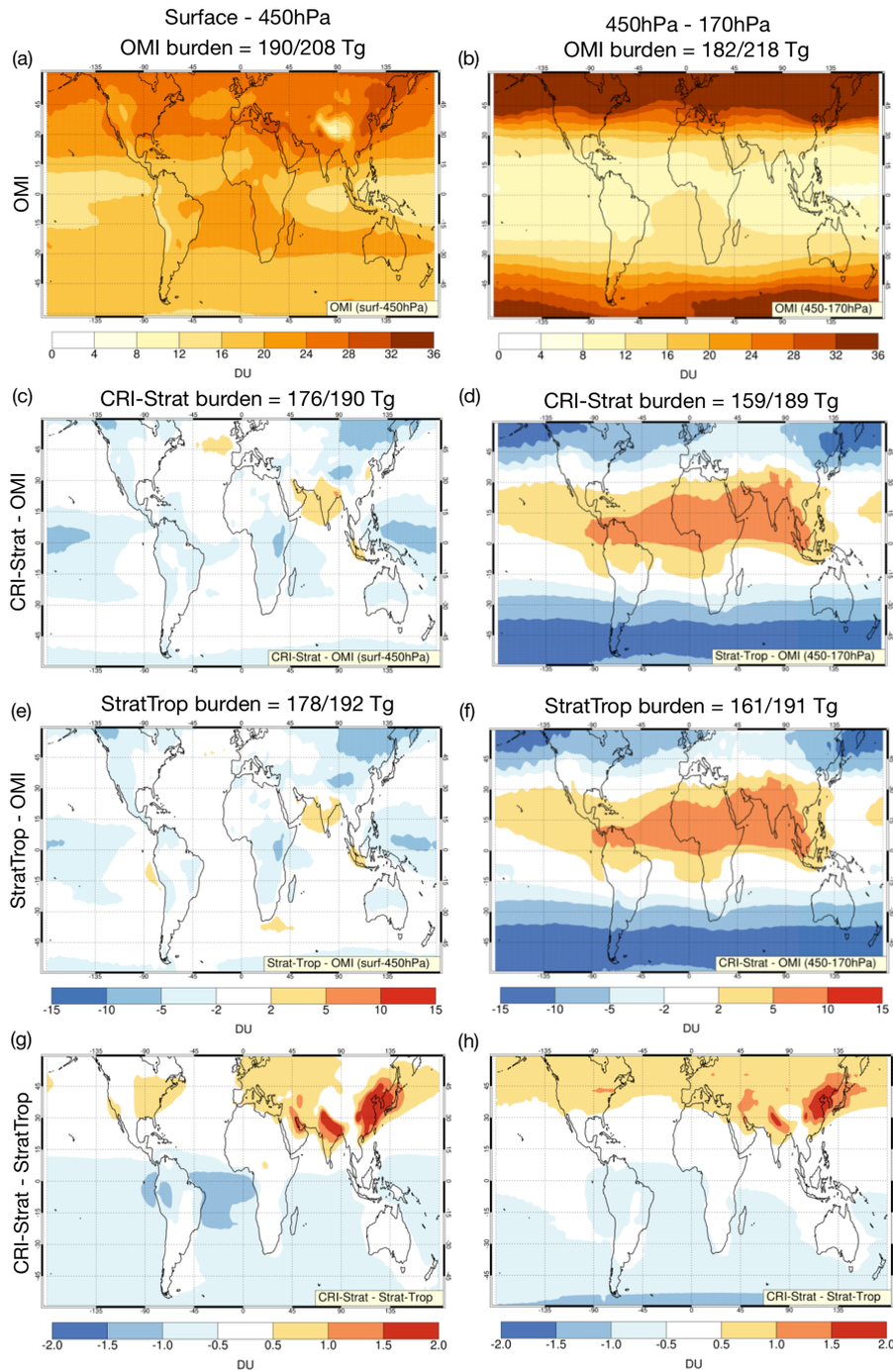


Figure 4. Ozone column between surface and 450 hPa from OMI satellite product as map (a) and from 450 to 170 hPa (b), averaged between 2010-2017. Difference in ozone column between CRI-Strat and OMI over respective pressure ranges (c, d), between StratTrop and OMI (e, f), and between CRI-Strat and StratTrop, noting different colour scale (g, h). Burdens are given over the respective pressure ranges for between 60°S to 60°N then over 90°S to 90°N.

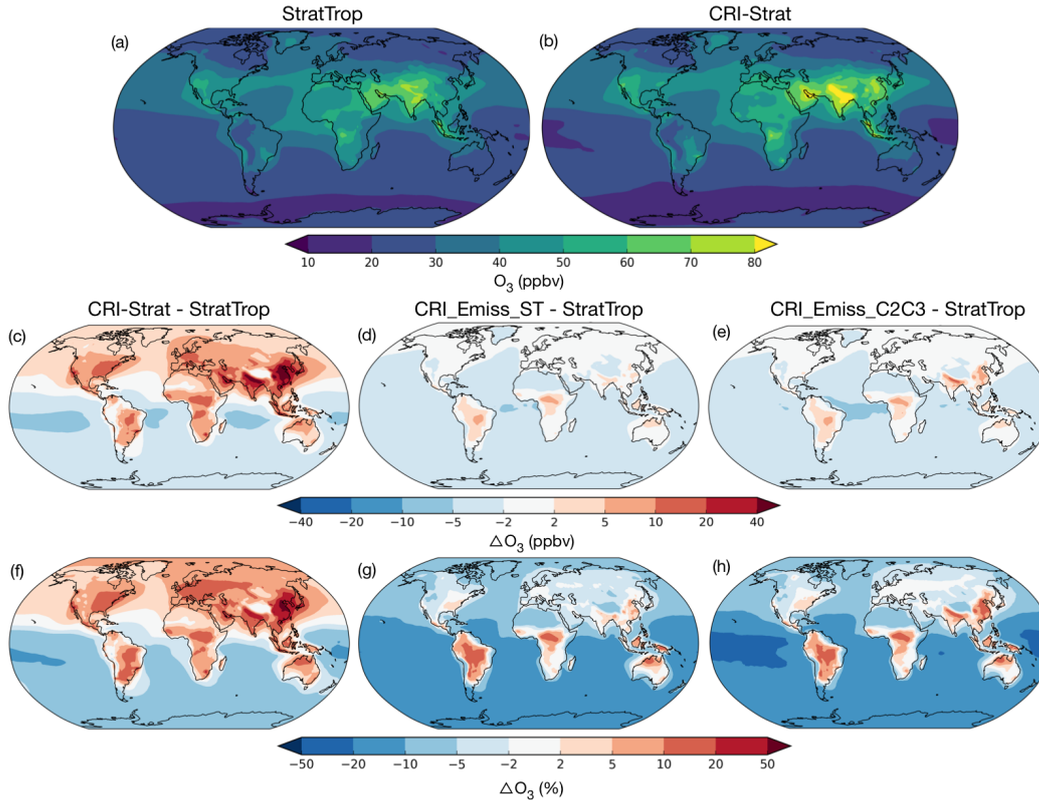


Figure 5. Mean ozone averaged over lower 1 km of the atmosphere in StratTrop (a) and CRI-Strat (b). Absolute difference (c-e) and relative difference (f-h) in ozone over lower 1 km of atmosphere between CRI-Strat and StratTrop (c, f), CRI_Emiss_ST - StratTrop (d, g), and CRI_Emiss_C2C3 - StratTrop (e, h).

688 ozone levels over highly populated regions such as the Indo-Gangetic plain but lower lev-
 689 els in remote regions. This is due to CRI_Emiss_C2C3 having emissions of alkenes such
 690 as C_3H_6 which are more reactive and have a higher ozone production efficiency than equiv-
 691 alent alkanes such as C_3H_8 , but have shorter lifetimes and contribute less to ozone pro-
 692 duction downwind of sources (M. E. Jenkin et al., 2002).

693 Looking at the vertical distribution of ozone, high ozone levels in CRI-Strat are lo-
 694 calised to the lower atmosphere in the northern hemisphere (Figure 6 (a, d)). Ozone con-
 695 centrations are lower in the upper tropical troposphere and across the southern hemi-
 696 sphere. CRI-Strat has higher ozone in the lower stratosphere, but these differences are
 697 relatively small ($< 2\%$) and the largest fractional differences are in the lower atmosphere.
 698 In the CRI_Emiss_ST and CRI_Emiss_C2C3 scenarios, ozone levels are lower through-
 699 out most of the atmosphere, and the only region that shows similar zonal average ozone
 700 is the boundary layer between 30N and 60N (Figure 6 (b,c, e, f)). This demonstrates that
 701 the CRI-Strat mechanism is more efficient at producing ozone near emission sources, but
 702 that losses are greater in remote regions of the atmosphere.

703 The relationship between surface ozone and emissions is shown in the ozone iso-
 704 leths in Figure 7. Ozone concentrations in all of the simulations are lower in NO_x -limited
 705 and VOC-limited regimes (in the bottom-right and top-left of each panel respectively),
 706 and drop slightly faster in CRI-Strat than StratTrop as NO_x emissions increase. Peak
 707 ozone concentrations in StratTrop occur in regions with lower VOC and NO_x emissions,

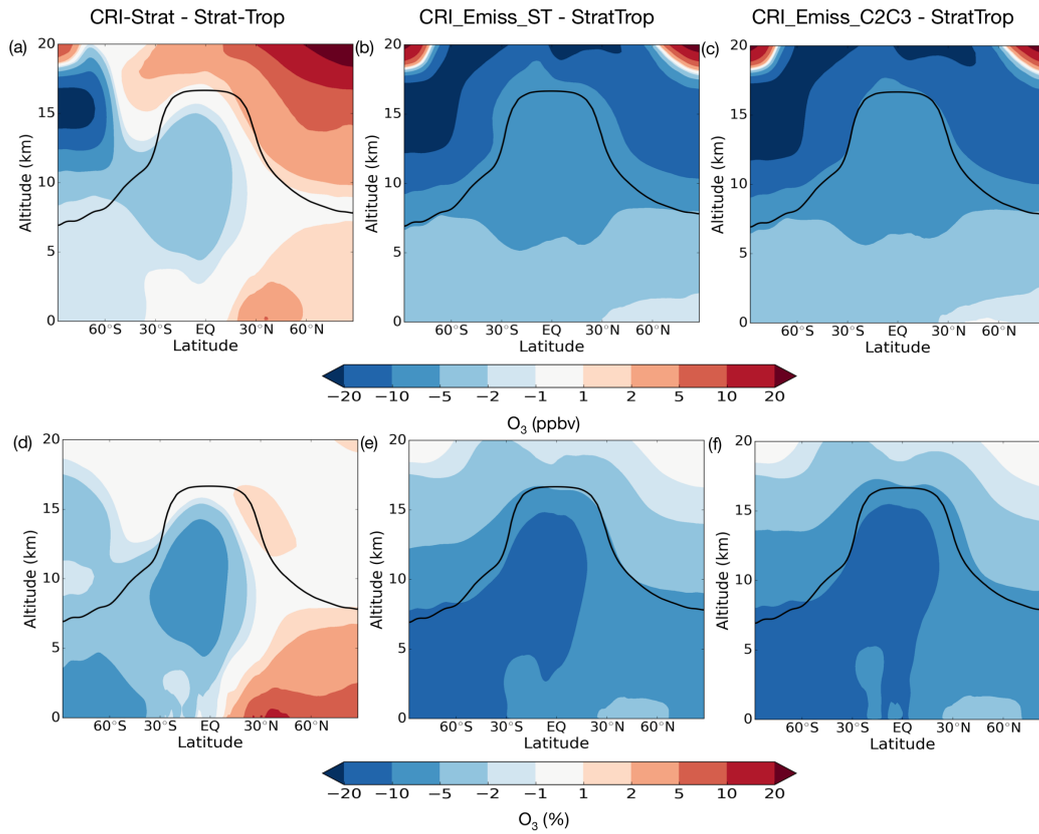


Figure 6. Zonal mean absolute difference (a-c) and percentage difference (d-f) in ozone between CRI-Strat and StratTrop (c, f), CRI_Emiss_ST - StratTrop (d, g), and CRI_Emiss_C2C3 - StratTrop (e, h). Black lines show average tropopause height over simulation period.

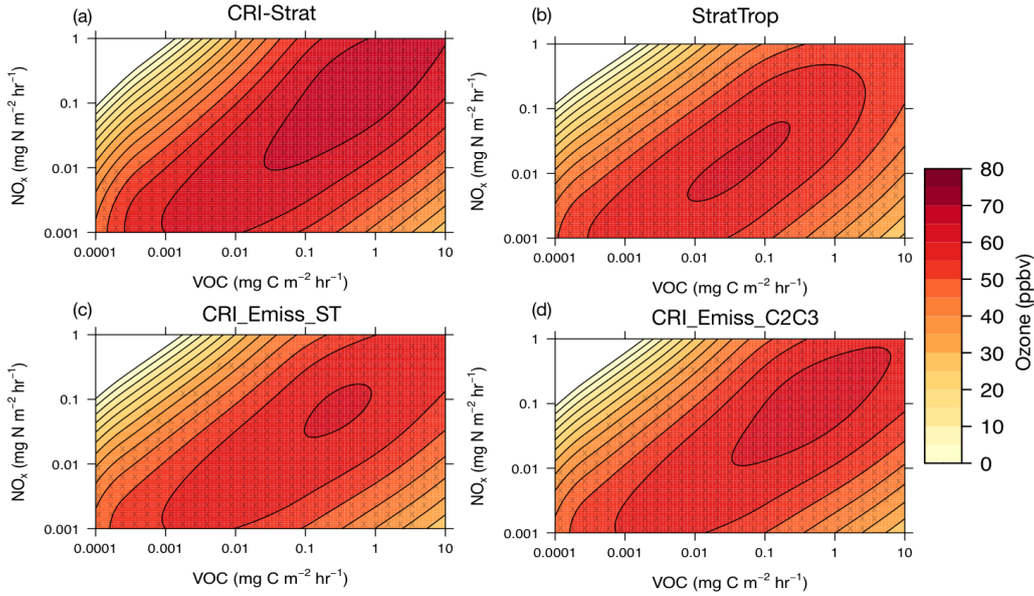


Figure 7. Ozone isopleths showing average surface ozone levels with respect to surface emissions of VOCs and NO_x in StratTrop (a), CRI-Strat (b), CRI_Emiss_ST (c) and CRI_Emiss_C2C3 (d) model simulations. Isopleths have been drawn by mapping average surface ozone concentrations from each gridcell of each of the model simulations with NO and NMVOC emissions at each gridcell, then interpolating results to give a smooth field. Stippling shows regions of the phase space which have sufficient data from the model output.

708 between 0.01-0.1 mg C m⁻²hr⁻¹ and 0.01-0.1 mg N m⁻²hr⁻¹ respectively), and then drop
 709 off at high emission regions (Fig. 7 (a)). The CRI-Strat simulation has the highest ozone
 710 levels, with the increased NMVOC emissions spurring ozone production in more regions
 711 with high NO_x emissions (Fig. 7 (b)). The CRI_Emiss_ST scenario shows a similar dis-
 712 tribution to StratTrop but peak ozone levels occur in regions with higher emissions (Fig.
 713 7 (c)). CRI_Emiss_C2C3 has peak ozone at even higher emission levels and the peak in
 714 ozone covers a broader region of the phase space, demonstrating that increased specia-
 715 tion of NMVOC emissions allows for greater ozone production in polluted regions as
 716 more is emitted as short lived, reactive compounds (Fig. 7 (d)).

717 5.2.2 Production and loss of ozone

718 To investigate these differences in ozone between the simulations, we present a full
 719 budget analysis of tropospheric ozone. Tropospheric production and loss of ozone is cal-
 720 culated using diagnostics which track the odd oxygen family, including NO₂ and its reser-
 721 voir species, collectively known as O_x (Wang et al., 1998):

$$722 \quad O_x = O_3 + O + O(^1D) + NO_2 + 2NO_3 + 3N_2O_5 + HONO_2 + HO_2NO_2 + PANs. \quad (5)$$

723 This definition of O_x is based on the principle that the rate limiting step for ozone
 724 production following NO₂ photolysis is the conversion from NO to NO₂. Hence produc-
 725 tion (P_{O_x}) is primarily via the pathways HO₂ + NO, CH₃O₂ + NO and R'O₂ + NO (where
 726 R'O₂ is the sum of all peroxy radicals apart from CH₃O₂). Chemical loss of ozone (L_{O_x})
 727 is defined as the sum of all chemical pathways which result in net loss of ozone, primar-

ily via reaction of $O(^1D)$ with water vapour and catalytic loss of O_3 with HO_x but also via minor reaction pathways involving NO_3 . Net ozone production is therefore defined as the difference between chemical production and loss:

$$Net_{O_x} = P_{O_x} - L_{O_x} \quad (6)$$

O_x is also lost via deposition (D_{O_x}), both directly through dry deposition of O_3 and indirectly via deposition of NO_2 and its reservoir species. The final part of the budget is from transfer of ozone from the stratosphere to the troposphere (S_{O_x}), which can be inferred from the excess loss of ozone assuming no long-term changes in tropospheric burden.

$$S_{O_x} = L_{O_x} + D_{O_x} - P_{O_x} \quad (7)$$

Finally, the lifetime of O_x in the troposphere is calculated by dividing the total tropospheric burden by the total loss via chemical sinks and deposition (Young et al., 2018).

$$\tau_{O_x} = B_{O_x} / (L_{O_x} + D_{O_x}) \quad (8)$$

In the CRI-Strat simulation, ozone production is much higher than StratTrop in the lower atmosphere, but also in most of the troposphere between $60^\circ S$ and $90^\circ N$ (Fig. 8 (a) and supplementary Figure S11). The only region where StratTrop has faster production is in the southern upper tropical troposphere. In CRI_Emiss_ST and CRI_Emiss_C2C3, ozone production is even more concentrated in the boundary layer, with much less production than StratTrop in the upper troposphere (Fig. 8 (b, c)). However, all simulations using the CRI-Strat mechanism have greater ozone loss over most of the troposphere (Fig. 8 (d-f)), so that the main region of greater net ozone production is in the boundary layer and there is a layer of net loss immediately above (Fig. 8 (g-i)). As shown in Supplementary Figure S12, the difference in distribution of L_{O_x} is almost identical to the flux through the $O(^1D) + H_2O$ reaction, one of the reactions that has different reaction rate coefficients between the mechanisms. This reaction drives a large component of the ozone changes between the mechanisms.

Table 5 gives an overview of the tropospheric ozone budget, production and loss terms. While CRI-Strat and StratTrop have similar total tropospheric ozone burdens of 331.8 Tg and 336.8 Tg respectively, the burdens of CRI_Emiss_ST and CRI_Emiss_C2C3 are both around 30 Tg lower. Compared to the TOAR model intercomparison (Young et al., 2018), the tropospheric ozone burdens in the StratTrop and CRI-Strat simulations are within the model interquartile range (320-370 Tg), whereas CRI_Emiss_ST and CRI_Emiss_C2C3 are below that range. Gaudel et al. (2018) calculate total ozone burdens between 333 and 345 Tg for the 2010-2014 period from satellite products which can observe all latitude bands, overlapping with StratTrop and slightly above CRI-Strat but much higher than CRI_Emiss_ST and CRI_Emiss_C2C3. Ozone production efficiency (OPE, defined as moles of O_x produced per mole of NO_x emitted) is also higher in CRI-Strat (31.4) than StratTrop (27.2).

The lifetime of ozone in all of the simulations using CRI-Strat chemistry is much shorter (17.3-17.5 days) than with StratTrop (19.8 days), hence the lower ozone in CRI_Emiss_ST and CRI_Emiss_C2C3 likely reflects the shorter lifetime of O_x . However, the CRI-Strat simulation has a similar ozone burden to StratTrop because increased production compensates for the increased losses. In fact, the lifetime of ozone in all of these simulations is short compared to previous analysis (e.g. 22.3 ± 2.0 days from Young et al. (2013), although these were for the year 2000 rather than 2014). Note also that the lifetime calculation is dependent on the definition of O_x , for example Bates and Jacob (2020) find O_3 lifetimes of approximately 75 days as they do not consider the $O(^1D) + H_2O$ reaction to cause net loss of O_x . The short lifetimes could partly explain why both CRI-Strat and Strat-Trop have lower ozone column values at high latitude than OMI-MLS (Figure 3), as insufficient ozone is transported from the tropics, where production is highest.

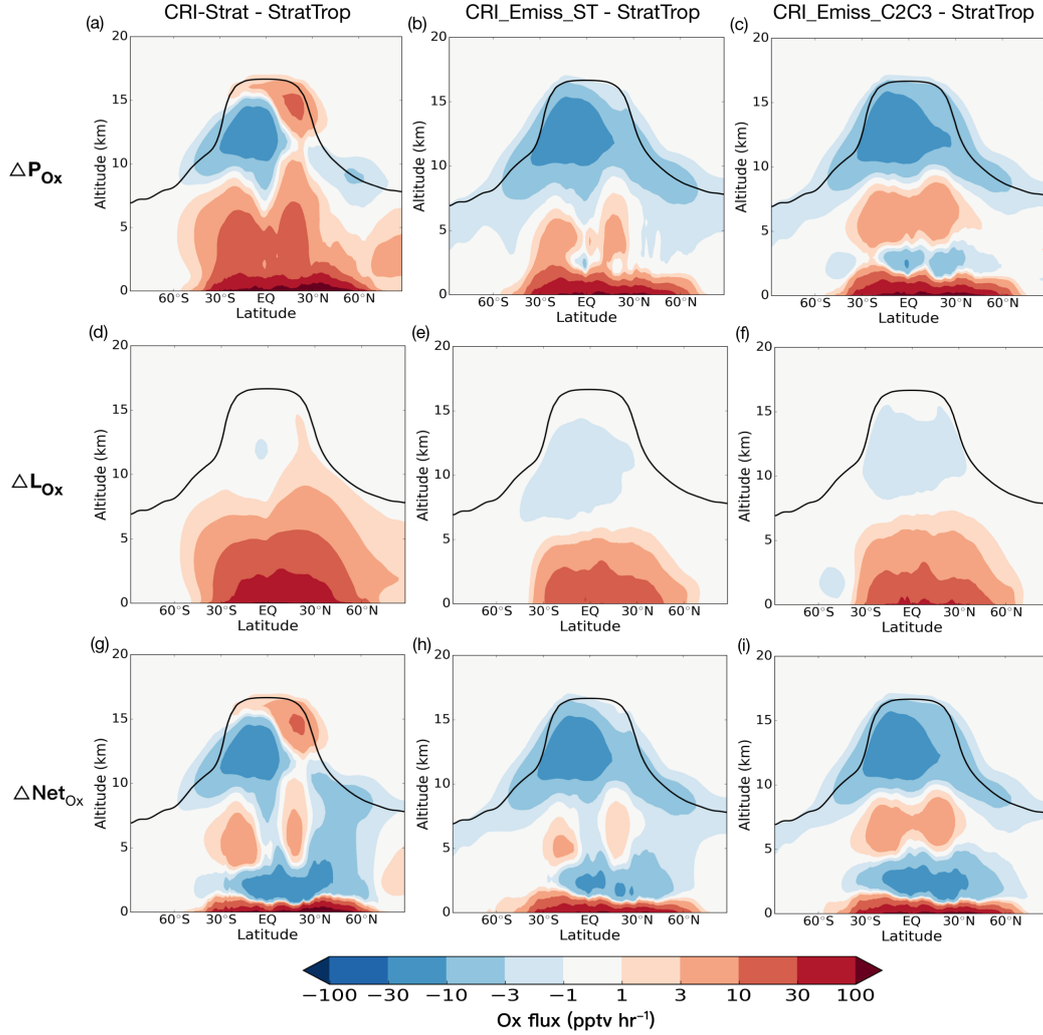


Figure 8. Zonal mean difference in chemical production of Ox between CRI-Strat and StratTrop (a), CRI_Emiss_ST and StratTrop (b), and CRI_Emiss_C2C3 and StratTrop (c) over 2010-2018 simulation period. Zonal mean difference in chemical loss of Ox between CRI-Strat and StratTrop (d), CRI_Emiss_ST and StratTrop (e), and CRI_Emiss_C2C3 and StratTrop (f). Zonal mean difference in net chemical production of Ox between CRI-Strat and StratTrop (g), CRI_Emiss_ST and StratTrop (h), and CRI_Emiss_C2C3 and StratTrop (i). Black lines show average tropopause height over simulation period.

Table 5. Overview of O_x burden, lifetime, ozone production efficiency (OPE), chemical production, chemical loss, deposition and inferred stratosphere to troposphere transfer. Values in brackets give fraction of total chemical production for the production terms and fraction of total losses (L_{Ox} + D_{Ox}) for the chemical loss and deposition terms.

	CRI-Strat	StratTrop	CRI-Emiss_ST	CRI-Emiss_C2C3
O ₃ burden (Tg)	331.9	336.8	308.0	306.8
O _x lifetime (days)	17.3	19.8	17.5	17.4
NMVOC emissions (Tg C year ⁻¹)	1012	762	762	762 [†]
OPE (mole _{O₃} mole _{NO_x} ⁻¹)	31.4	27.2	28.7	28.9
Chemical production (P_{Ox})	6624	5725	6057	6092
Total	4152 (62.7%)	3853 (67.3%)	3886 (64.2%)	4001 (65.7%)
HO ₂ + NO	1540 (23.3%)	1285 (22.5%)	1452 (24.0%)	1419 (23.3%)
CH ₃ O ₂ + NO	882 (13.3%)	545 (9.5%)	676 (11.2%)	629 (10.3%)
R'O ₂ + NO	64.5 (0.7%)	41.3 (0.7%)	41.5 (0.7%)	43.2 (0.7%)
Other*				
Chemical loss (L_{Ox})	5853	5128	5355	5380
Total	3196 (45.5%)	2660 (42.9%)	3022 (47.1%)	3005 (46.6%)
O(¹ D) + H ₂ O	1713 (24.4%)	1596 (25.7%)	1498 (23.4%)	1513 (23.5%)
HO ₂ + O ₃	708 (10.1%)	714 (11.5%)	667 (10.4%)	680 (10.6%)
OH + O ₃	160 (2.3%)	96.5 (1.6%)	115 (1.79%)	129 (2%)
O ₃ + Alkene	76.3 (1.1%)	61.5 (1.0%)	53.1 (0.8%)	52.6 (0.8%)
Other [‡]				
Deposition (D_{Ox})	1164	1081	1061	1068
Total	974 (13.9%)	896 (14.4%)	872 (13.6%)	878 (13.6%)
O ₃ dry dep	190.3 (2.7%)	185 (3.0%)	189 (2.95%)	190 (2.94%)
NO _y dep				
Inferred STT (Tg O ₃ year ⁻¹)	394	483	361	355

[†] Same carbon emission mass as StratTrop but different speciation.

*“Other” O_x production channels are the sum of inorganic acid oxidation, RONO₂ oxidation and RONO₂ photolysis.

[‡]“Other” O_x loss channels are the sum of O(³P) + O₃, O³P + NO₂, N₂O₅ + H₂O and NO₃ chemical losses.

779 The ozone production and loss terms for CRI-Strat (6624 and 5853 Tg yr⁻¹ respec-
 780 tively) are both much higher than for StratTrop (5725 and 5128 Tg yr⁻¹), and are higher
 781 than most models from the TOAR assessment (Young et al., 2018). It has been noted
 782 before that models with representation of higher aromatics and monoterpenes have high
 783 P_{O_x} and L_{O_x} terms (Porter et al., 2017), but the calculation of the net chemical tendency
 784 (P_{O_x} - L_{O_x}) shows that the CRI-Strat results in a greater overall propensity to form
 785 ozone in spite of the emissions of NMVOCs (c.f. StratTrop and CRI_Emiss_ST exper-
 786 iments Table 4). The P_{O_x} and L_{O_x} terms in CRI_Emiss_ST and CRI_Emiss_C2C3
 787 are very similar to each other, in both cases higher than StratTrop and lower than the base
 788 CRI-Strat run. There is a general trend for “hotter” chemistry - more ozone is produced
 789 and lost - in all of the simulations using the CRI-Strat chemical mechanism. We can also
 790 say that the addition of higher NMVOC emissions has a larger impact on the ozone bud-
 791 get (CRI-Strat vs CRI_Emiss_ST and CRI_Emiss_C2C3) than the improved speciation
 792 of NMVOC emissions (CRI_Emiss_ST vs CRI_Emiss_C2C3).

793 The increase in ozone production in CRI-Strat occurs over all channels, but the largest
 794 relative increase is via R'O₂ + NO (882 Tg yr⁻¹), which is over 50% larger in CRI-Strat
 795 than in StratTrop (545 Tg yr⁻¹), partly because there is more R'O₂ in CRI-Strat. The
 796 CRI_Emiss_ST simulation shows its largest increase in production compared to Strat-
 797 Trop via the R'O₂ + NO channel, as well as the CH₃O₂ + NO channel, whereas in CRI_Emiss_C2C3
 798 it is via the HO₂ + NO channel. Note though that the increased production via HO₂ + NO
 799 and CH₃O₂ + NO is also linked to oxidation of larger NMVOCs, as many reactions pro-
 800 duce secondary HO₂ and CH₃O₂ as larger molecules are oxidised.

801 Chemical loss of O_x via the O(¹D) + H₂O channel is greater in CRI-Strat (3196 Tg yr⁻¹)
 802 than in StratTrop (2660 Tg yr⁻¹). In addition, the CRI_Emiss_ST and CRI_Emiss_C2C3
 803 scenarios have similar losses via this channel (3022 Tg yr⁻¹ and 3005 Tg yr⁻¹ respec-
 804 tively) to the base CRI-strat simulation, scaling linearly with ozone burden. The O(¹D) + H₂O
 805 pathway is the primary driver behind why O_x lifetime is much shorter in the CRI-Strat
 806 chemical mechanism compared to StratTrop. This is down to both the reaction rate co-
 807 efficients for the hydroxyl radical producing O(¹D) + H₂O → 2OH reaction being greater
 808 in CRI-Strat than in StratTrop, and the stabilising reaction O(¹D) + M → O(³P) + M
 809 is smaller, which collectively result in excited O(¹D) being about 25% more likely to re-
 810 act with H₂O than stabilise with M for a given concentration of water vapour (see Fig-
 811 ure S2 and Section S4 in the supplement for details).

812 5.3 Hydroxyl and Hydroperoxyl Radicals

813 The hydroxyl radical (OH) is the major oxidant in the troposphere for almost all
 814 NMVOC species. Together with the hydroperoxyl radical (HO₂) it forms the HO_x fam-
 815 ily (HO_x = OH + HO₂). Zonal means of OH and HO₂ from the StratTrop simulation and
 816 differences to other simulations are shown in Figure 9. All of the simulations with CRI-
 817 Strat chemistry show a bimodal pattern in OH of high levels in the lower troposphere
 818 and low levels in the upper troposphere compared to StratTrop (Fig. 9 (c-e)), and a pat-
 819 tern of more HO₂ in both the lower and upper troposphere (Fig. 9 (f-h)). Total HO_x,
 820 which is roughly equal to HO₂, is much higher in all of the simulations using CRI-Strat
 821 chemistry than StratTrop (Fig. 9 (f-h)), which is unsurprising given the main source is
 822 the O(¹D) + H₂O reaction which we already know has a much higher flux in CRI-Strat
 823 than StratTrop (Table 5).

824 In the lower atmosphere, which has higher O₃ in the CRI-Strat simulation and is
 825 abundant with water vapour, we have greater HO_x production and more OH and HO₂.
 826 There are also more O₃ + Alkene reactions in CRI-Strat, which produce OH, and HO₂
 827 is produced as a byproduct of many VOC oxidation and photolysis reactions. Together,
 828 these differences lead to the “hotter” chemistry in CRI-Strat.

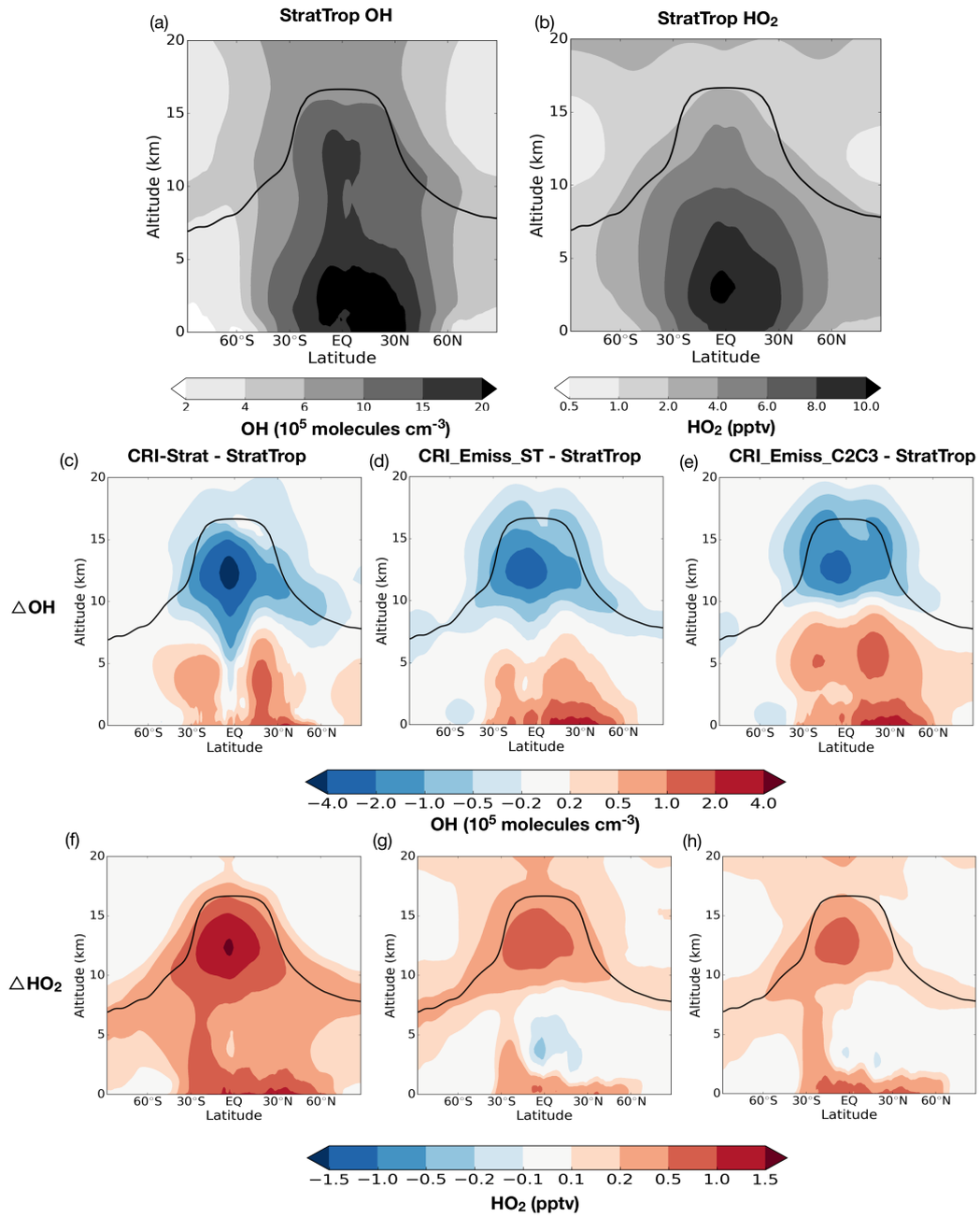


Figure 9. Zonal mean OH (a) and HO₂ (b) in StratTrop simulation. Differences in zonal mean OH and HO₂ between CRI-Strat and StratTrop (c, f), CRI_Emiss_ST and StratTrop (d, g), and CRI_Emiss_C2C3 and StratTrop (e, h). Black lines show average tropopause height over simulation period.

Table 6. Overview of mass weighted annual average tropospheric HO_x concentrations, OH northern hemisphere to southern hemisphere ratio, OH : HO₂ ratio, methane lifetime with respect to OH and HO₂ + HO₂ flux.

	CRI-Strat	StratTrop	CRI_Emiss_ST	CRI_Emiss_C2C3
[OH] (10 ⁶ molecules cm ⁻³)	1.335	1.339	1.348	1.375
OH NH:SH ratio	1.38	1.35	1.4	1.4
[HO ₂] (pptv)	6.27	5.90	6.02	6.06
OH : HO ₂ ratio (%)	1.49	1.67	1.61	1.63
CH ₄ lifetime W.R.T. OH (years)	7.77	8.13	7.71	7.60
HO ₂ + HO ₂ flux (P mole year ⁻¹)	60.5	32.2	38.8	39.9

829 In the upper troposphere, OH concentrations are much lower in CRI-Strat com-
830 pared to StratTrop for a number of reasons. It is less moist than the lower atmosphere
831 and away from emission sources of short-lived alkenes (which can make OH), hence pri-
832 mary OH production is lower. However, CRI-Strat has more formaldehyde than Strat-
833 Trop and other carbonyls which can be photolysed in the upper troposphere to produce
834 HO₂. CRI-Strat has more long-lived NMVOCs and CO in the upper troposphere which
835 primarily react with OH converting it into HO₂. Ozone levels in the upper troposphere
836 are also lower in CRI-Strat. Hence CRI-Strat has more HO_x in the upper troposphere,
837 but a greater proportion is as HO₂ and there is less OH than in StratTrop.

838 Overall, CRI-Strat has slightly lower average OH concentration than StratTrop,
839 whereas the CRI_Emiss_ST and CRI_Emiss_C2C3 simulations have more OH overall than
840 the base CRI-Strat and StratTrop simulations (Table 6). Hence, given the same emis-
841 sions the CRI-Strat mechanism produces more HO_x than StratTrop, but this is coun-
842 tered by the additional NMVOCs in the full CRI-Strat mechanism which primarily re-
843 act with OH, converting it into HO₂, leading to a lower OH : HO₂ ratio in CRI-Strat.
844 The total HO₂ + HO₂ flux, a major sink of HO_x in the atmosphere, is almost twice as
845 high in CRI-Strat than in StratTrop due to both the increase in HO₂ and a faster re-
846 action rate (see supplementary Figures S2 and S4).

847 The methane (CH₄) lifetime is slightly shorter in CRI-Strat than in StratTrop, even
848 though OH is higher in StratTrop. The reason is because the OH + CH₄ reaction rate
849 coefficient is slightly faster in CRI-Strat (see Figure S5 in the supplement). While this
850 difference has little impact on the concentration of CH₄, because this is set by surface
851 boundary conditions, it influences tropospheric chemistry because CH₄ is a major source
852 of the methyl peroxy radical (CH₃O₂), the most abundant peroxy radical and an impor-
853 tant component of the tropospheric ozone forming process (Table 5).

854 5.4 Carbon Monoxide

855 Carbon monoxide (CO) is important as the main chemical sink for OH, convert-
856 ing it into HO₂, and as a toxic air pollutant in its own right. It is produced from both
857 primary emission sources, generally from incomplete combustion of biomass and fossil
858 fuels, and secondary production from the oxidation of VOCs (Grant et al., 2010).

859 The total tropospheric burden of CO is over 50 Tg higher in CRI-Strat compared
860 to StratTrop, and the CRI_Emiss_ST and CRI_Emiss_C2C3 simulations also have higher
861 CO burdens (Table 7). The CO lifetime varies linearly with OH concentration between
862 the simulations (Table 6), as the OH + CO reaction occurs at the same rate in both mech-
863 anisms, and so is slightly longer in CRI-Strat as it has slightly less OH overall compared

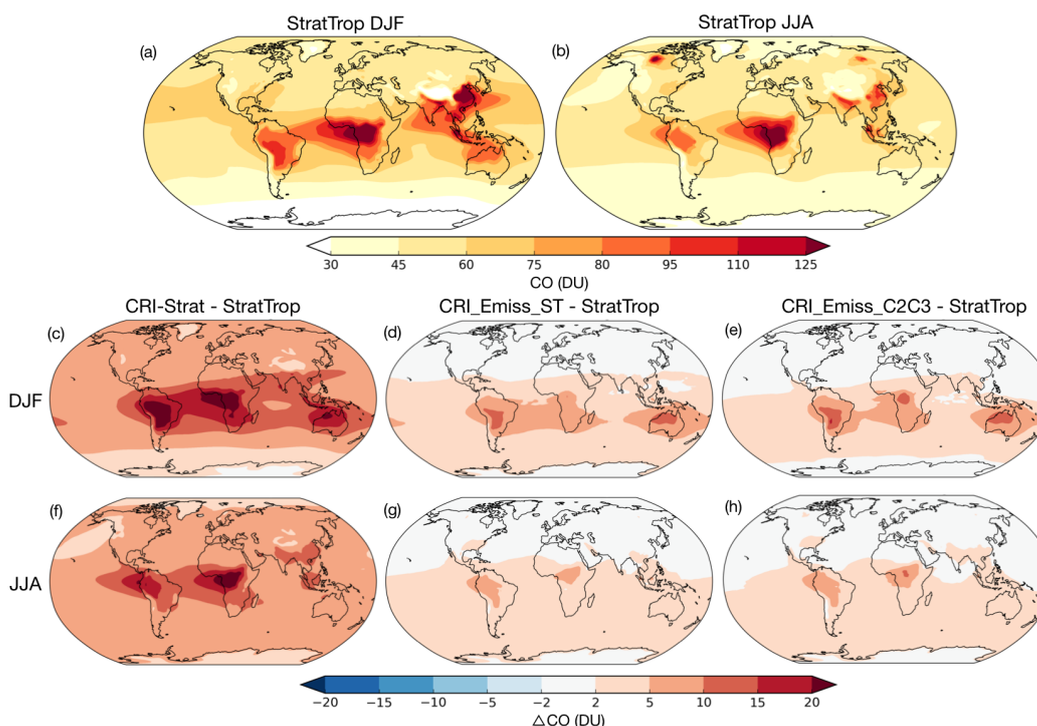


Figure 10. Average tropospheric column CO in StratTrop simulations for DJF (a) and JJA (b). Differences in tropospheric CO column in DJF and JJA between CRI-Strat and StratTrop (c, f), CRI_Emiss_ST and StratTrop (d, g), and CRI_Emiss_C2C3 and StratTrop (e, h).

864 to StratTrop. As all of the simulations use the same primary CO emissions, the differ-
 865 ences in CO burden are due to secondary production and oxidant profiles. Although most
 866 of this secondary production occurs via formaldehyde (HCHO) oxidation and photolysis
 867 channels, there is a large increase in contribution from other channels in CRI-Strat
 868 compared to StratTrop (these being oxidation and photolysis of larger carbonyls, O_3 +alkene
 869 reactions and PAN oxidation). The increase in CO production from HCHO and other
 870 channels is also apparent to a lesser extent in the CRI_Emiss_ST and CRI_Emiss_C2C3
 871 scenarios. The more explicit NMVOC degradation chemistry in the CRI-Strat mech-
 872 anism produces more HCHO and other oxidised NMVOCs, which go on to form CO as
 873 they are further oxidised and undergo photolysis.

874 The bulk of the additional CO burden is in the southern hemisphere downwind of
 875 regions with high BVOC emissions, such as the Amazon, Central Africa and Australia
 876 (Fig. 10). In CRI_Emiss_ST and CRI_Emiss_C2C3, increases in CO column are localised
 877 to these regions and downwind of them (Fig. 10 (d, e, g, h)), implying that the primary
 878 cause of increased CO is the more explicit, multigenerational representation of isoprene
 879 degradation in the CRI-Strat mechanism. The base CRI-Strat simulation has much more
 880 CO over these BVOC dominated regions and a background increase in column CO of
 881 around 5-10 DU across almost the entire world, including the northern hemisphere (Fig.
 882 10 (c, f)). This is due to additional CO production from degradation of higher NMVOCs
 883 emitted by anthropogenic sources, as well as from the explicit APINENE and BPINENE
 884 chemistry in CRI-Strat.

885 Tropospheric CO in the StratTrop mechanism was evaluated against multiple datasets
 886 by Archibald et al. (2020). They found significant negative biases in column CO of 10-
 887 20 DU over much of the Northern hemisphere, but similar values across the Southern

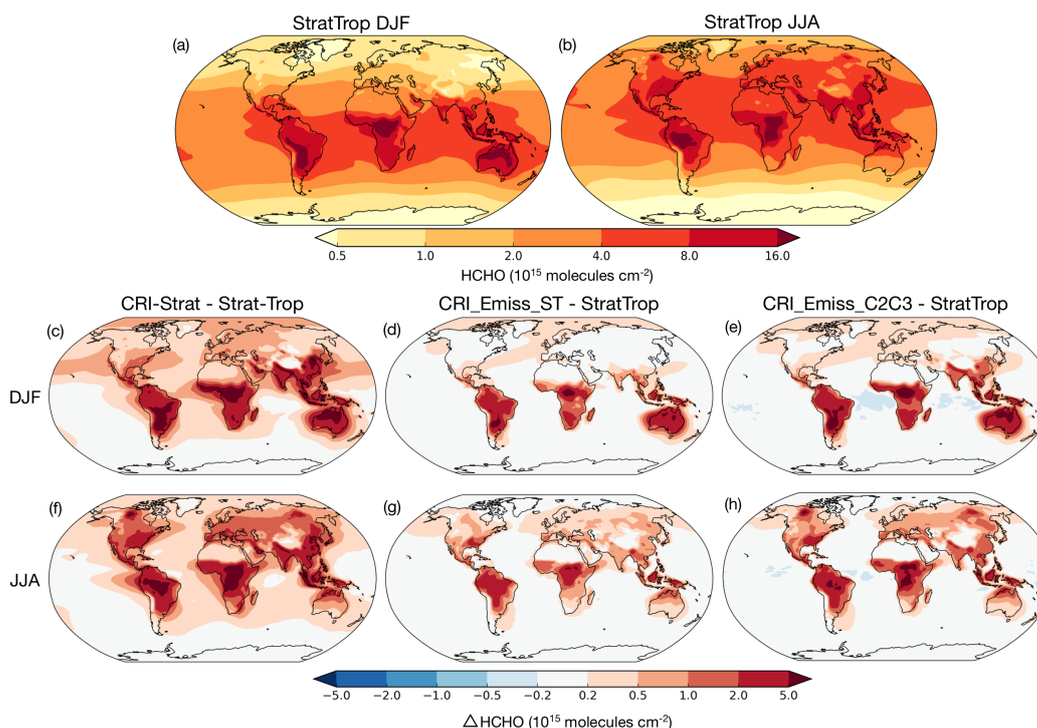


Figure 11. Average tropospheric column HCHO in StratTrop simulations for DJF (a) and JJA (b). Differences in tropospheric HCHO column density in DJF and JJA between CRI-Strat and StratTrop (c, f), CRI.Emiss.ST and StratTrop (d, g), and CRI.Emiss.C2C3 and StratTrop (e, h).

888 hemisphere and a high bias over BVOC rich Central Africa and Amazonia compared to
 889 the MOPITT satellite product. The CRI-Strat simulation has a lot more CO in the North-
 890 ern hemisphere, reducing the negative bias there, but far too much CO in the southern
 891 hemisphere particularly downwind of BVOC sources. Although the CRI-Strat mecha-
 892 nism captures secondary production of CO better than StratTrop, it highlights that too
 893 much CO production occurs in the southern hemisphere, likely due to errors in BVOC
 894 emissions. In addition, the model setup either lacks some key CO or CO precursor emis-
 895 sions in the Northern hemisphere, CO lifetime is too short in the model and/or not enough
 896 CO is being transported from low to high latitudes.

897 This importance of HCHO in CO production is emphasised in Figure 11, which shows
 898 much higher HCHO column densities above high BVOC emission regions in all of the
 899 simulations that use CRI-Strat chemistry. As HCHO is the dominant source of secondary
 900 CO, this will lead to higher CO burdens downwind. The more explicit representation
 901 of isoprene and monoterpene degradation chemistry clearly leads to greater secondary
 902 production of HCHO and as a result CO.

903 5.5 Peroxy Radicals

904 Peroxy radicals (RO_2) are formed from the oxidation of VOCs and are important
 905 but short-lived intermediates in tropospheric ozone formation. The largest differences
 906 between the two chemical mechanisms centre around treatment of RO_2 species, as dis-
 907 cussed in section 3.1. The CRI-Strat has a total of 47 RO_2 species (as opposed to 9 in
 908 StratTrop), all of which undergo reactions with NO, NO_3 , HO_2 and the summed RO_2

Table 7. Overview of average tropospheric CO burden and key fluxes for CO production/loss and for peroxy radical reactions

Flux	CRI-Strat	StratTrop	CRI-Emiss_ST	CRI-Emiss_C2C3
CO burden (Tg)	354.9	300.2	317.2	315.2
CO lifetime (days)	39.1	38.6	38.1	37.2
CO production (Tg CO year ⁻¹)	3402	2915	3121	3171
Emissions	1111 (32.7%)	1111 (38.1%)	1111 (35.6%)	1111 (35%)
HCHO + OH	580.3 (17.1%)	492 (16.9%)	551 (17.7%)	566 (17.8%)
HCHO + $h\nu$	1293 (38%)	1076 (36.9%)	1163 (37.3%)	1177 (37.1%)
Other Chem	125 (3.7%)	71.5 (2.5%)	101 (3.25%)	107 (3.4%)
Other Phot	293 (8.6%)	164 (5.6%)	194 (6.23%)	211 (6.7%)
CO Loss (Tg CO year ⁻¹)	3311	2841	3041	3090
CO + OH	3157 (95.4%)	2704 (95.2%)	2900 (95.4%)	2948 (95.4%)
CO Dry Dep	153 (4.6%)	137 (4.8%)	141 (4.7%)	142 (4.6%)
RO ₂ fluxes (P mole year ⁻¹)	111.5	75.8	91.5	87.9
RO ₂ + NO	18.4 (16.5%)	11.4 (15.0%)	14.1 (15.4%)	13.1 (14.9%)
RC(O)O ₂ + NO ₂ (PAN Prod)	20.6 (18.5%)	19.6 (25.9%)	14.7 (16.1%)	14.0 (16.0%)
RO ₂ + NO ₃	0.3 (0.3%)	0.2 (0.3%)	0.3 (0.3%)	0.2 (0.3%)
RO ₂ + HO ₂	60.4 (54.2%)	41.2 (54.4%)	52.4 (57.3%)	51.0 (58.0%)
RO ₂ + RO ₂	11.8 (10.6%)	3.4 (4.5%)	10.1 (11.04%)	9.5 (10.8%)

radical pool in ways that conserve the number of potential ozone forming steps. Acetyl peroxy radicals ($\text{RC}(\text{O})\text{O}_2$) are also crucial as they form thermally stable peroxyacetyl nitrate (PAN) compounds, important NO_y reservoir species. CRI-Strat has multiple addition PAN species from degradation of higher NMVOC species. Table 7 gives a summary of the key RO_2 reaction fluxes in the troposphere. The CRI-Strat simulation has a 50% higher RO_2 flux compared to StratTrop. The CRI_Emiss_ST and CRI_Emiss_C2C3 simulations have total fluxes 20% and 16% higher than StratTrop respectively, showing that a large fraction of the additional flux is a result of the more explicit multigeneration chemistry, and not just from the additional NMVOC emissions in CRI-Strat.

There are big differences in the respective fates of RO_2 species. In the CRI-Strat mechanism, RO_2 species are more than twice as likely to react with another RO_2 species than in StratTrop, because the full range of these cross-reactions are parameterised for all RO_2 species. The CRI-Strat mechanisms also have a greater flux through the $\text{RO}_2 + \text{HO}_2$ branches, which form ROOH species which are then often lost via wet and dry deposition, due to higher HO_2 and RO_2 concentrations in CRI-Strat (see Table 6). There is a much larger flux through PAN forming reactions in StratTrop. The MPAN forming reaction in StratTrop is much faster than the equivalent reaction in CRI-Strat (see supplementary Figure S5), and hence the production of MPAN and PAN-type species is greater in StratTrop even though all other PAN-type species (including PAN) are formed in greater abundance in CRI-Strat. However, as PAN-type species rapidly dissociate back to the original reactants in high temperature conditions, the higher flux through the $\text{RC}(\text{O})\text{O}_2 + \text{NO}_2$ pathway does not necessarily lead to permanent oxidation, unlike via the other branches.

5.6 Nitrogen Oxides and their reservoirs

There are stark differences in nitrogen oxides between the different model simulations, as shown in Figure 12. In this analysis, we use the StratTrop simulation as a baseline to evaluate CRI-Strat, as StratTrop in UKESM1 has been thoroughly evaluated by Archibald et al. (2020). In the StratTrop simulation, the tropospheric column density of NO_2 is highest over polluted regions, such as China and Europe, and is higher in local winter than summer, due to higher emissions, and lower oxidising capacity and photolysis rates in winter (Fig. 12 (a, b)). Archibald et al. (2020) found a high bias downwind of these polluted regions in winter of up to 5×10^{15} molecules cm^{-2} (approximately 50%) compared to the OMI satellite product, but a small low bias over most of the rest of the world (see Archibald et al. (2020) Fig. 18). The CRI-Strat simulation has much lower column density of NO_2 across the Northern Hemisphere continental regions in winter such as over China where CRI-Strat has column densities more than 5×10^{15} molecules cm^{-2} lower than StratTrop (Fig. 12 (c)). CRI_Emiss_ST has lower NO_2 column than StratTrop directly over high emission regions but is similar downwind (Fig. 12 (d, g)). CRI_Emiss_C2C3 is somewhere between the base CRI-Strat and CRI_Emiss_ST simulations, with a lower column density of NO_2 over polluted regions and downwind of them over continents, but not to the same degree as CRI-Strat (Fig. 12 (e, h)).

Nitrogen is conserved through chemical reactions (noting that some minor changes were made to the StratTrop mechanism in order for it to conserve nitrogen, see Section S3 in the supplement for details) and all simulations have exactly the same NO_x emissions, therefore if NO_2 is lower it must have been converted into some other form and/or lost via deposition. It is conventional to use the shorthand NO_x as the sum of nitrogen oxides NO and NO_2 , NO_z is the sum of their reservoir species, and NO_y is the sum of all oxidised nitrogen compounds:

$$\text{NO}_x = \text{NO} + \text{NO}_2 \quad (9)$$

$$\text{NO}_z = \text{HONO}_2 + \text{NO}_3 + 2 * \text{N}_2\text{O}_5 + \text{HO}_2\text{NO}_2 + \text{XONO}_2 + \text{PANs} + \text{RONO}_2 + \text{CH}_3\text{O}_2\text{NO}_2 + \text{Nitrophenols} \quad (10)$$

$$\text{NO}_y = \text{NO}_x + \text{NO}_z \quad (11)$$

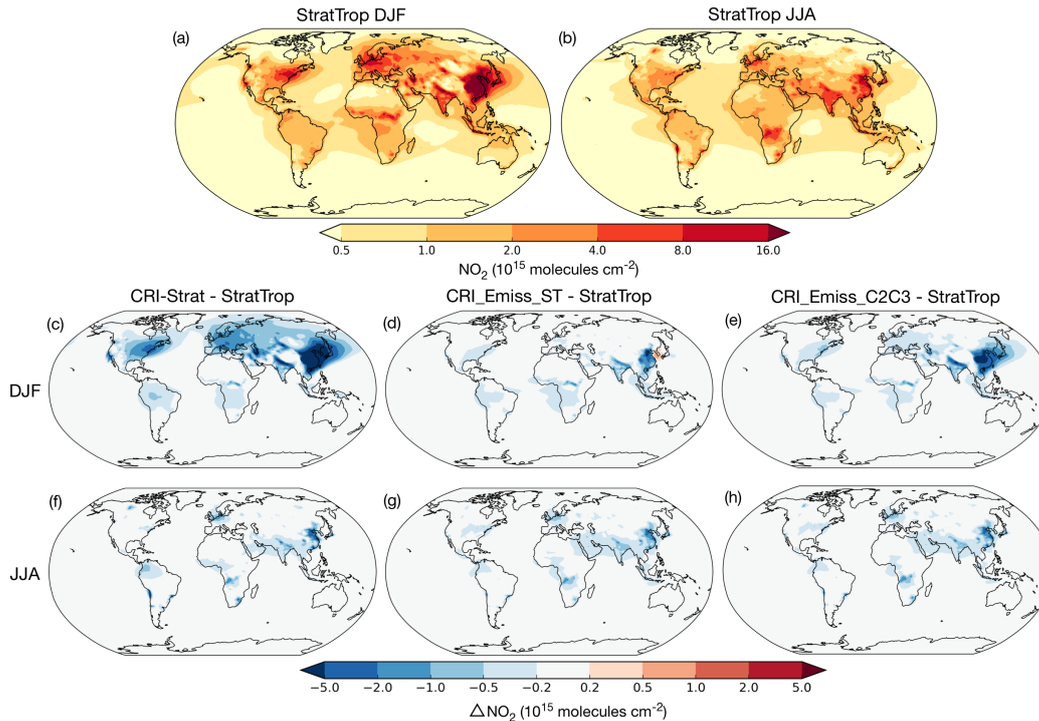


Figure 12. Average tropospheric column NO₂ in StratTrop simulations for DJF (a) and JJA (b). Differences in tropospheric NO₂ column in DJF and JJA between CRI-Strat and StratTrop (c, f), CRI_Emiss_ST and StratTrop (d, g), and CRI_Emiss_C2C3 and StratTrop (e, h).

961 The conversion of NO_x to NO_z can inhibit the ozone forming process. The most com-
 962 ponent of NO_z is nitric acid (HONO₂), which is efficiently dry and wet deposited and
 963 so acts as a sink for reactive nitrogen in the troposphere. However, some reservoir species
 964 (such as PAN) can also aid overall ozone production if they are transported, releasing
 965 NO_x into other regions of the troposphere where the ozone forming potential per NO_x
 966 molecule is greater.

967 When analysing how oxidised nitrogen differs between these model simulations, its
 968 important to understand some key differences between the two chemical mechanisms.
 969 The CRI-Strat mechanism has the same inorganic nitrogen species as StratTrop, but it
 970 has many more organic nitrogen containing species which contribute to NO_z. Firstly, in
 971 CRI-Strat almost every RO₂ + NO reaction has a minor branch which forms an organon-
 972 itrate (RONO₂), whereas in StratTrop only CH₃ONO₂ and ISON (which represents organon-
 973 itrates from isoprene oxidation) exist. Peroxy radicals with nitrate groups are also formed
 974 from NO₃ initiated reactions with alkenes, which can also go on to form stable organon-
 975 itrates. These organonitrates can act as NO_x reservoirs, transported long distances be-
 976 fore undergoing further oxidation or photolysis to release NO_x or getting deposited out
 977 of the atmosphere. CRI-Strat also has several more PAN-type species, formed from ther-
 978 mal equilibrium between peroxyacyl radicals and NO₂. Finally, there are some species
 979 in CRI-Strat for which there is no equivalent in StratTrop: CH₃O₂NO₂ is formed
 980 from the thermal equilibrium between CH₃O₂ and NO₂ and can be an important reservoir species
 981 in the upper troposphere (Browne et al., 2011), while nitrated phenols are formed dur-
 982 ing the oxidation of benzene and toluene in the presence of NO_x (Harrison et al., 2005).

983 Zonal mean differences in these summed species are shown in Figure 13. There is
 984 reduced NO_x in the CRI-Strat mechanism compared to StratTrop (Fig. 13 (a)), partic-

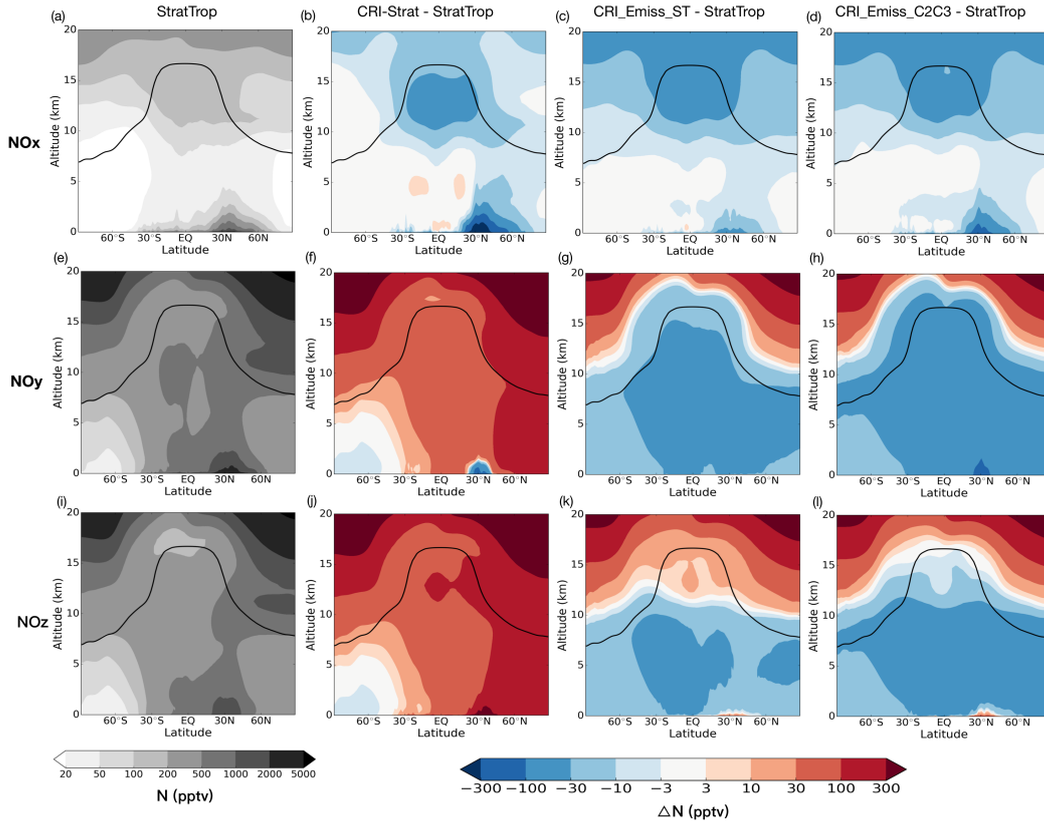


Figure 13. Average zonal mean NO_x (a), NO_y (e) and NO_z (f) in StratTrop run from 2010-2018. Zonal mean differences in NO_x , NO_y and NO_z between CRI-Strat and StratTrop (b, f and j), CRI_Emiss_ST and StratTrop (c, g and k), and CRI_Emiss_C2C3 and StratTrop (d, h, l) respectively. Black lines show average tropopause height over simulation period.

985 ularly in the lower atmosphere in the northern hemisphere and in the upper troposphere,
 986 that is visible in the base CRI-Strat simulation (Fig. 13 (b)) and in the runs with same
 987 total NMVOC mass emissions to StratTrop (Fig. 13 (c and d)). In contrast, total ox-
 988 idised nitrogen NO_y is generally higher in CRI-Strat compared to StratTrop (Fig. 13 (e
 989 and f)), except in the lower atmosphere around 30°N (specifically over East Asia and
 990 India, see supplementary Figure S13), but lower almost everywhere in CRI_Emiss_ST
 991 and CRI_Emiss_C2C3 (Fig. 13 (g and h)). The increase in NO_y is shown to be due to
 992 there being more NO_z almost everywhere in the CRI-Strat model simulation compared
 993 to StratTrop (Fig. 13 (j)). However, in the CRI-STEMiss and CRI_Emiss_C2C3 simu-
 994 lations higher NO_z is only found in the lower atmosphere northern hemisphere down-
 995 wind of high emitting regions (Fig. 13 (k and l)). Hence the increased NO_z in the CRI-
 996 Strat simulation is clearly tied in with the extra NMVOC emissions. Without these added
 997 organic species, the CRI-Strat chemical mechanism has greater propensity to produce
 998 more NO_z in polluted regions but loose oxidised nitrogen mass overall.

999 The base CRI-Strat simulation has more tropospheric NO_y than StratTrop (1.112 Tg N
 1000 vs. 1.018 Tg N), but has less of the total nitrogen as NO_x (10.3% vs 14.9%) and more
 1001 as NO_z reservoirs (89.7% vs 85.1%), as shown in Table 8. However, CRI-Strat has a smaller
 1002 fraction of NO_y as HONO_2 (45.8% vs 50.4%), the single largest component of NO_z . In-
 1003 stead, a greater fraction is stored as PANs and RONO_2 , as well as $\text{CH}_3\text{O}_2\text{NO}_2$ and ni-
 1004 trophenol species which are not in the StratTrop chemical mechanism.

Table 8. Overview of tropospheric oxidised nitrogen burdens (fraction of total NO_y in brackets), tropospheric oxidised nitrogen emission and deposition fluxes, stratosphere-troposphere transfer (STT) and NO_y lifetime in the troposphere (fraction of total NO_y deposition in brackets).

	CRI-Strat	StratTrop	CRI_Emiss_ST	CRI_Emiss_C2C3
NO_y Burden (Tg N)	1.112	1.018	0.950	0.910
NO_x Burden (Tg N)	0.115 (10.3%)	0.152 (14.9%)	0.136 (14.3%)	0.131 (14.4%)
NO_z Burden (Tg N)	0.998 (89.7%)	0.866 (85.1%)	0.814 (85.7%)	0.779 (85.6%)
HONO_2 Burden (Tg N)	0.509 (45.8%)	0.513 (50.4%)	0.506 (53.2%)	0.512 (56.2%)
Other inorganic NO_z (Tg N)	0.021 (1.9%)	0.018 (1.7%)	0.018 (1.9%)	0.019 (2.1%)
PANs (Tg N)	0.367 (33.0%)	0.296 (29.1%)	0.245 (25.8%)	0.206 (22.7%)
RONO_2 (Tg N)	0.061 (5.5%)	0.039 (3.9%)	0.038 (4.0%)	0.035 (3.9%)
$\text{CH}_3\text{O}_2\text{NO}_2$ (Tg N)	0.008 (0.7%)	N/A	0.007 (0.7%)	0.007 (0.7%)
Nitrophenols (Tg N)	0.031 (2.8%)	N/A	0.0	0.0
Total NO_x Emissions (Tg N year ⁻¹)	61.5	61.5	61.5	61.5
Total NO_y Deposition (Tg N year ⁻¹)	63.0	62.9	63.0	63.0
Inferred STT (Tg N year ⁻¹)	1.46	1.40	1.43	1.44
NO_x Dry deposition (Tg N year ⁻¹)	6.83 (10.8%)	7.70 (12.2%)	7.36 (11.7%)	7.25 (11.5%)
HONO_2 Wet deposition (Tg N year ⁻¹)	29.0 (46.0%)	30.1 (47.8%)	30.1 (47.8%)	30.0 (47.7%)
HONO_2 Dry deposition (Tg N year ⁻¹)	22.0 (34.9%)	21.6 (34.3%)	22.3 (35.5%)	22.4 (35.6%)
Other inorganic NO_z deposition (Tg N year ⁻¹)	1.21 (1.9%)	0.97 (1.6%)	1.04 (1.6%)	1.08 (1.7%)
PANs dry deposition (Tg N year ⁻¹)	1.70 (2.7%)	1.28 (2.0%)	0.894 (1.4%)	0.918 (1.5%)
RONO_2 deposition (Tg N year ⁻¹)	2.09 (3.3%)	1.30 (2.1%)	1.22 (2.0%)	1.28 (2.0%)
Nitrophenol deposition (Tg N year ⁻¹)	0.22 (0.4%)	N/A	0.0	0.0
NO_y deposition lifetime (days)	6.44	5.91	5.51	5.28
HONO_2 deposition lifetime (days)	3.65	3.62	3.53	3.57
PANs deposition lifetime (days)	78.9	84.5	100.1	82.0
RONO_2 deposition lifetime (days)	10.7	11.0	11.4	9.99

1005 Many of these differences can be attributed to the added NMVOC emissions in the
 1006 CRI-Strat simulations. The greater total quantity of NMVOC emissions, added larger
 1007 NMVOCs and the explicit depiction of their multigenerational degradation provides more
 1008 opportunities for NO_z reservoir species to form. In the CRI_Emiss_ST simulation, the
 1009 total tropospheric NO_y is 0.95 Tg N, showing that the CRI-Strat mechanism has lower
 1010 NO_y when if run with the same NMVOC emissions as StratTrop (Table 8). The frac-
 1011 tion of NO_x (14.3%) is slightly lower than StratTrop, but still higher than CRI-Strat.
 1012 In the CRI_Emiss_C2C3 simulation, NO_y is lower still, at 0.91 Tg N. The driving fac-
 1013 tor behind these differences appears to be a greater fraction of total NO_y as HONO_2 in
 1014 CRI_Emiss_ST (53.2%) and CRI_Emiss_C2C3 (56.2%) compared to StratTrop (50.4%)
 1015 or the base CRI-Strat run (45.8%). However, CRI_Emiss_ST and CRI_Emiss_C2C3 both
 1016 have slightly less HONO_2 than StratTrop in absolute terms.

1017 The importance of these differences in oxidised nitrogen species burdens can be bet-
 1018 ter understood by analysing the sources and sinks of oxidised nitrogen in the troposphere.
 1019 In all of the simulations, total NO_x emissions are identical, and total NO_y deposition al-
 1020 most identical, as shown in Table 8. Total NO_y deposition is greater than total emis-
 1021 sions because of net transfer of NO_y species (mostly HONO_2) from the stratosphere to
 1022 the troposphere, with this additional NO_y originating from reaction of N_2O and $\text{O}(^1\text{D})$
 1023 in the stratosphere. From the tropospheric NO_y burden and sum of all deposition sinks,
 1024 we can calculate the mean tropospheric NO_y lifetime, which ranges from around 6.4 days
 1025 in CRI-Strat to 5.9 days in StratTrop, 5.5 days in CRI_Emiss_ST and only 5.3 days in
 1026 CRI_Emiss_C2C3. By decomposing the channels by which NO_y is deposited, we can see
 1027 that less is deposited as NO_x or HONO_2 in CRI-Strat compared to StratTrop, and a larger
 1028 fraction is deposited as organic nitrogen containing species. In general these organic nitrogen-
 1029 containing species deposit less efficiently than HONO_2 (R. Sander, 2015) and have longer
 1030 lifetimes with respect to deposition, effectively extending the NO_y lifetime and allow-
 1031 ing total NO_z to accumulate (Table 8). They are also more effective at transporting NO_x
 1032 from polluted regions and releasing it in cleaner regions which are more NO_x sensitive,
 1033 going a long way to explaining why the CRI-Strat simulation has the highest overall rate
 1034 of ozone production (Table 5).

1035 Compared to the StratTrop model run, the CRI_Emiss_ST and CRI_Emiss_C2C3
 1036 simulations have similar wet deposition loss of HONO_2 but greater loss via dry depo-
 1037 sition. This is because there is more HONO_2 production in the boundary layer (see sup-
 1038plementary Figure S14), where it can be rapidly lost via dry deposition before it has time
 1039 to be transported away from emission sources. Hence, total tropospheric NO_z and NO_y
 1040 levels, and NO_y lifetime, are overall lower in CRI_Emiss_ST and CRI_Emiss_C2C3 com-
 1041pared to StratTrop.

1042 5.7 Summary and synthesis

1043 Collating all of the information covered in this analysis, we can understand that
 1044 the variations in tropospheric ozone are largely driven by the variations in the NO_y fields
 1045 between the simulations:

- 1046 1. In polluted regions, ozone production is more efficient in the CRI-Strat mecha-
 1047 nism due to the more explicit representation of multigenerational NMVOC- NO_x
 1048 chemistry, particularly when the amount of NMVOC available is increased. How-
 1049 ever, loss of ozone is also more rapid, particularly due to the increased proportion
 1050 of O_x lost via the $\text{O}(^1\text{D}) + \text{H}_2\text{O}$ reaction.
- 1051 2. The enhanced HONO_2 production in CRI-Strat leads to greater NO_y removal in
 1052 the boundary layer via dry deposition, reducing the amount entering the free tro-
 1053 sphere.

- 1054 3. In the base CRI-Strat run, faster HONO₂ production is compensated by greater
 1055 production of organic nitrogen-containing species which extend the lifetime of NO_y
 1056 and enhance ozone production in remote regions.
- 1057 4. These competing effects roughly balance over the whole troposphere, leading to
 1058 similar tropospheric ozone burdens in the CRI-Strat and StratTrop simulations,
 1059 albeit with more ozone near the surface in CRI-Strat.
- 1060 5. The CRI_Emiss_ST and CRI_Emiss_C2C3 simulations are less able to form organon-
 1061 itrates than the CRI-Strat simulation so lose more nitrogen to HONO₂ deposition
 1062 and therefore have lower NO_y burdens. Combined with the shorter ozone lifetime,
 1063 these two simulations have the lowest tropospheric ozone burdens despite having
 1064 higher ozone production rates than StratTrop in the polluted boundary layer.

1065 Overall, we find a strong sensitivity between emissions of NMVOCs and the oxi-
 1066 dised nitrogen budget, which leads to considerable knock on effects to the tropospheric
 1067 ozone burden and oxidising capacity of the atmosphere. This has important implications
 1068 for the need to improve emissions of NMVOCs (which are typically poorly constrained
 1069 (Huang et al., 2017)) and how these emissions are treated in simpler mechanisms such
 1070 as StratTrop. The dependence of NO_x to emissions of NMVOCs is also relevant for how
 1071 we interpret comparisons of NO_x between models and observations.

1072 The CRI mechanism has now been implemented in a number of models, including
 1073 STOCHEM (S. Utembe et al., 2010) and WRF-Chem (Archer-Nicholls et al., 2014). The
 1074 STOCHEM implementation also saw an increase in ozone production and loss compared
 1075 to its existing mechanism, but this led to much higher ozone burdens across the tropo-
 1076 sphere, particularly over the oceans. In contrast, CRI-Strat has similar tropospheric ozone
 1077 burdens to StratTrop and lower ozone concentrations across much of the world's oceans.
 1078 We believe the higher tropospheric ozone columns did not occur in UKCA because of
 1079 the shorter O_x lifetime in CRI-Strat compared to StratTrop due to the faster flux through
 1080 the O(¹D) + H₂O reaction in CRI-Strat. The WRF-Chem implementation also saw an
 1081 increase in ozone production but little overall increase in ozone over the model domain
 1082 compared to the CBM-Z mechanism. However, in a regional model a species such as ozone,
 1083 with a typical lifetime of a few weeks, is very sensitive to the effects of boundary con-
 1084 ditions. In Archer-Nicholls et al. (2014), boundary conditions were prescribed by the MOZART
 1085 model for both CBM-Z and CRIv2-R5. Combining WRF-Chem with boundary condi-
 1086 tions driven by UKCA simulations, with both models using the same CRI mechanism,
 1087 could offer a solution to the problem of how to account for the impact of boundary con-
 1088 ditions.

1089 6 Conclusions

1090 The CRIv2-R5 chemical mechanism has been integrated into the UKCA model, merged
 1091 with the existing stratospheric chemistry and coupled to GLOMAP-mode aerosol to cre-
 1092 ate the CRI-Strat mechanism. The mechanism is constrained to reliably reproduce the
 1093 ozone forming potential from MCMv3.1, enabling traceability to our best understand-
 1094 ing and providing a benchmark to evaluate simpler mechanisms and test other aspects
 1095 of model setup. This new mechanism marks a step change in chemical complexity and
 1096 comprehensiveness, improving representation of important tropospheric processes, such
 1097 as BVOC chemistry, peroxy radicals and organonitrates, while remaining sufficiently af-
 1098 fordable for use in an Earth System Model (approximately 75% longer runtime compared
 1099 to an equivalent run with StratTrop). CRI-Strat can now be used in UKESM1, a flag-
 1100 ship model used for CMIP6 (Eyring et al., 2016), opening up new potential scientific en-
 1101 quires. As the implementation has been done using the ASAD framework (Carver et
 1102 al., 1997), it can also be ported to other models which share the same framework.

1103 In this paper we critically evaluate and compare CRI-Strat to the well-established
 1104 StratTrop mechanism, highlighting some key differences:

- 1105 1. CRI-Strat has higher surface ozone and CO compared to StratTrop, improving
 1106 some biases compared to observations but worsening others.
- 1107 2. CRI-Strat chemistry is generally “hotter” than StratTrop: it has much higher pro-
 1108 duction and loss of ozone, more HO_x and more secondary production of CO.
- 1109 3. Total tropospheric ozone burden and ozone column density are surprisingly sim-
 1110 ilar between the simulations, given the large differences in production rate.
- 1111 4. CRI-Strat partitions a greater fraction of nitrogen into reservoir forms (NO_z) with
 1112 a lower fraction in the reactive form (NO_x).
- 1113 5. Many differences are related to the speciation of NMVOCs. Significantly more emit-
 1114 ted species are included in CRI-Strat than in StratTrop. When using exactly the
 1115 same emissions, CRI-Strat has 8.6% lower tropospheric ozone burden than Strat-
 1116 Trop.

1117 Some of these differences reflect differences in reaction rate coefficients for key reactions,
 1118 a number of which are out of date in CRIv2-R5 and therefore CRI-Strat. The faster pro-
 1119 duction of ozone that occurs in CRI-Strat when photochemical conditions allow means
 1120 that it is more sensitive to model structural uncertainties than StratTrop, particularly
 1121 relating to emissions, model resolution and parameterisations such as for lightning-NO_x
 1122 emissions. The more complex CRI-Strat mechanism is not designed to be a replacement
 1123 for StratTrop, but provides a new tool that expands the possible scientific questions that
 1124 can be tackled with the model and a benchmark to evaluate against.

1125 The tests and evaluation described in this paper set out to fully characterise the
 1126 CRI-Strat mechanism against the StratTrop mechanism (the reference mechanism for
 1127 UEKSM1) for the gas-phase composition of species relevant for the climate. This pro-
 1128 vides information that is essential to understand and make use of the new mechanism.
 1129 However, the experiments performed do not use CRI-Strat to its full potential. We ex-
 1130 pect it to exceed the capabilities of StratTrop when run at higher spatial resolution and
 1131 in evaluation against field campaigns with a focus on oxidants. Future work will also fo-
 1132 cus on highly polluted environments or those dominated by BVOCs and production of
 1133 SOA, for which CRI has been shown to provide a robust framework for simulating (S. Utembe
 1134 et al., 2010). We also plan to run experiments for different climate and emission regimes
 1135 such as the pre-industrial atmosphere; these experiments (combined with multi model
 1136 analyses) will enable us to understand if we can be confident that UKESM1 represents
 1137 the changes in composition and chemistry-climate feedbacks from pre-industrial to the
 1138 present day realistically. This evaluation has also neglected analysis of aerosols, whose
 1139 formation rates will differ with CRI-Strat due to changes in oxidant fields, and will be
 1140 properly evaluated in the future. There is also scope to improve the coupling between
 1141 CRI-Strat and the GLOMAP-mode scheme, for example updating reaction rates and ex-
 1142 tending the isoprene chemistry with inclusion of CRIv2.2 (M. E. Jenkin, Khan, et al.,
 1143 2019), improving representation of BVOC environments, OH recycling and further ex-
 1144 pansion with HOM chemistry (Weber et al., 2020).

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1163 Due to intellectual property right restrictions, we cannot provide either the source
 1164 code or documentation papers for the UM. The Met Office Unified Model is available
 1165 for use under licence. The functionality described here is fully available in the UM trunk
 1166 from version 11.8. A number of research organisations and national meteorological ser-
 1167 vices use the UM in collaboration with the UK Met Office to undertake basic atmospheric
 1168 process research, produce forecasts, develop the UM code and build and evaluate Earth
 1169 system models. For further information on how to apply for a licence; see [https://www](https://www.metoffice.gov.uk/research/approach/modelling-systems/unified-model)
 1170 [.metoffice.gov.uk/research/approach/modelling-systems/unified-model](https://www.metoffice.gov.uk/research/approach/modelling-systems/unified-model) (last
 1171 access: 24 November 2020). Data tables of the full CRI-Strat mechanism as described
 1172 in this paper are included in the supplement.

1173 UM simulations are compiled and run in suites developed using the Rose suite en-
 1174 gine (<http://metomi.github.io/rose/doc/html/index.html>, last access: 24 Novem-
 1175 ber 2020) and scheduled using the Cylc workflow engine (<https://cylc.github.io/>,
 1176 last access: 24 November 2020). Both Rose and Cylc are available under version 3 of the
 1177 GNU General Public License (GPL). In this framework, the suite contains the informa-
 1178 tion required to extract and build the code as well as configure and run the simulations.
 1179 Each suite is labelled with a unique identifier and is held in the same revision-controlled
 1180 repository service in which we hold and develop the model’s code. This means that these
 1181 suites are available to any licensed user of the UM. The input emissions data are avail-
 1182 able from Input4MIPs (<https://esgf-node.llnl.gov/projects/input4mips/>, last ac-
 1183 cess: 24 November 2020).

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