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Analysis of UK and European NOx and VOC emission scenarios in the Defra model intercomparision exercise

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1 ANALYSIS OF UK AND EUROPEAN NO_X AND VOC EMISSION SCENARIOS IN THE DEFRA MODEL

2 INTERCOMPARISON EXERCISE

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Highlights

- Emission scenarios were implemented in eight ozone air quality models
- NOx- and VOC sensitivities for peak ozone levels were highly variable between days
- Filtering by model performance minimised apparent conflicts between models

Abstract

Simple emission scenarios have been implemented in eight United Kingdom air quality models with the aim of assessing how these models compared when addressing whether photochemical ozone formation in southern England was NOx- or VOC-sensitive and whether ozone precursor sources in the UK or in the Rest of Europe (RoE) were the most important during July 2006. The suite of models included three Eulerian-grid models (three implementations of one of these models), a Lagrangian atmospheric dispersion model and two moving box air parcel models. The assignments as to NOx- or VOC-sensitive and to UK- versus RoE-dominant, turned out to be highly variable and often contradictory between the individual models. However, when the assignments were filtered by model performance on each day, many of the contradictions could be eliminated. Nevertheless, no one model was found to be the 'best' model on all days, indicating that no single air quality model could currently be relied upon to inform policymakers robustly in terms of NOx- versus VOC-sensitivity and UK- versus RoE-dominance on each day. It is important to maintain a diversity in model approaches.

19 1. Introduction

20 Air quality models play an important role in air quality policy development by simulating and 21 visualising the conversion of ozone precursor emissions into ground-level ozone levels. Policy makers 22 formulate abatement strategies which aim to reduce ozone levels by reducing ozone precursor 23 emissions. Strategies can be evaluated to determine whether any emission reductions have been 24 stringent enough to achieve acceptable air quality in terms of internationally-accepted air quality 25 standards, guidelines and targets. Strategies may not necessarily be judged as pass or fail but may be 26 evaluated side-by-side with other strategies or against a do-nothing scenario. Increasingly policy 27 makers are using cost-benefit analysis in which the costs of an abatement strategy may be set 28 against the benefits of any environmental improvement as predicted by air quality models.

30 A huge range of air quality models address ground-level ozone and almost all of them have been 31 used in Europe in a policy context (see Kukkonen et al., 2012). Here the ability of a number of the 32 ground-level ozone models used by the Department for Environment, Food and Rural Affairs (Defra) 33 for its policy support and development to respond to policy-relevant questions, is addressed. The 34 model predictions for a given emission scenario differed widely and we try to explain why. For 35 simplicity, we focus on an episode of peak ozone in southern England and two policy-relevant 36 questions in the context of this one episode: is it better to reduce nitrogen oxide (NO_x) emissions or 37 volatile organic compounds and is better for any reductions to be undertaken concertedly across 38 Europe or unilaterally within the UK to reduce peak ozone levels? This study addresses the potential 39 conflicts that may arise when several models are employed to provide support and advice to policy 40 makers regarding emission control strategies to reduce episodic peak ozone in the UK. Potential 41 conflicts are illustrated with reference to NO_x and VOC emission sensitivities and to controlling emissions from different geographical areas. This study does not try to formulate such policy advice 42 43 and support but rather focusses on the difficulties inherent when conflicting results are available 44 from eight air quality models.

45

46 2. Methodology

The models employed in this study have all been employed to describe photochemical ozone
formation across north-west Europe and across the UK. Full details of the eight distinct models are
given in the Supplementary Information. They include 3-dimensional Eulerian grid models, a
Lagrangian atmospheric dispersion model and moving box trajectory-based models and employ a
range of chemical mechanisms to describe photochemical ozone formation from VOC and NO_x
emissions. A brief summary of the models is as follows:

29

- Community Multi-scale Air Quality (CMAQ) model (with 3 distinct implementations)
- Air Quality Unified Model (AQUM),
- European Monitoring and Evaluation Programme for the UK (EMEP4UK) model,
- Numerical Atmospheric dispersion Model Environment (NAME) model,
- Ozone Source Receptor Model (OSRM),
- Photochemical Trajectory Model (PTM).

To reduce the scope and complexity of the study to a level which was tractable, detailed attention was given to the behaviour of ground level ozone during July 2006 at the long-established EMEP rural air quality monitoring station at Harwell, Oxfordshire, UK. This station is located about 80 km due west of London and is surrounded by agricultural fields and a large campus of research establishments. The location of this site is considered typical of much of rural south-east England.

64

The weather across the UK generally during July 2006 was notable because of its high pressure and 65 66 high frequency of southerly winds. It was very warm and increasingly humid during the first six days of July 2006, with temperatures of 30 - 32 °C recorded daily in southern England. From the 14^{th} 67 onwards, the weather was sunny and increasingly hot, with daily maximum temperatures above 68 32°C from 16th – 27th (Eden, 2006). Ozone observations for Harwell were taken from the UK National 69 70 Air Quality Archive (http://www.airguality.co.uk/archive/data and statistics.php) and converted from μ g m⁻³ to ppb units using the factor 0.50. These data demonstrated the occurrence of 71 photochemical ozone episodes producing hourly ozone levels in excess of 50 ppb on $1^{st} - 4^{th}$, 6^{th} , 15^{th} 72 - 22nd, 24th - 27th July. The peak hourly ozone level of 106 ppb was recorded on 18th July see Figure 73 1. Also shown on Figure 1 are the daily advection regimes (as compass bearings N through NW) 74 75 based on Lamb Weather type (LWT) (Jenkinson and Collinson, 1977) where A refers to anticyclonic 76 and C cyclonic, on NILU FLEXTRA trajectories (Stohl et al., 1995) for Harwell and on the NAME model

(see Supplementary Information) air history maps (Manning et al., 2011) where EU refers to
advection from a large area of north-west Europe.

79

80 Intentionally, no attempt was made to harmonise the input data to the models. Necessarily, the models have used comparable sources for the emission inventory data, for example, based on 81 82 European Monitoring and Evaluation Programme (EMEP) emissions and the UK National 83 Atmospheric Emission Inventory (NAEI) (for further details, see the Supplementary Information), 84 with VOC speciation data from the NAEI. However, no attempt was made to harmonise the hourly, 85 weekly and seasonal time profiles, gridding or speciation profiles assumed. The models have used 86 different meteorological archives and descriptions of meteorological processes and meteorological 87 models to drive the different parameterisations of boundary layer processes, deposition, 88 atmospheric transport and dispersion. Again, no attempt was made to harmonise the chemical 89 mechanisms employed despite the known sensitivity of ozone predictions for North America to 90 chemical mechanism choice (Luecken et al., 2008) nor the biogenic VOC emission inventories and 91 their speciation.

92

93 Each of the 8 models was set up with their respective base case conditions for July 2006 and run in 94 their standard configurations as described in the Supplementary Information. The highest hourly 95 ozone levels predicted each day by each model are plotted together with the corresponding 96 observations in Figure 1. All of the models were able to account satisfactorily for the observed day-97 to-day variations in daily peak ozone levels in that they exhibited elevated levels during the periods $1^{st} - 4^{th}$, $15^{th} - 20^{th}$ and $24^{th} - 27^{th}$ July with relatively lower, background levels between $7^{th} - 14^{th}$ 98 99 and 28th – 31st July. Some of the observed episode days, however, were missed by some of the 100 models. Individual normalised mean biases (NMBs) for daily ozone maxima for July 2006 spanned

the range from -0.18 to -0.04. In the context of the simple evaluation criterion of NMB being in the
range -0.2 < NMB <0.2, proposed by Derwent et al., (2010), model performance was considered
entirely satisfactory for all eight models for July 2006 at Harwell.

104

105 NMBs were negative for all models for July daily maximum ozone levels at Harwell, largely because of poor model performance for July 18th and 19th, see Figure 1. Only one model simulated over 100 106 107 ppb for the daily maximum ozone level on these days and seven models gave less than 90 ppb. 108 Model performance was therefore generally poor for these days with highest ozone levels. It is 109 conceivable that the observations were strongly influenced by ozone precursor emissions associated 110 with the 2006 heat-wave which are not adequately represented in the emission inventories 111 employed in the standard model configurations. Air quality during much of the spring and summer 112 of 2006 was influenced by wild-fires in the Russian Federation (Saarikoski et al., 2007; Witham and Manning, 2007; Anttila et al., 2008; Niemi et al., 2009) and it is possible that this influence 113 specifically impacted upon the observed ozone levels at the Harwell station during July 18th and 19th. 114

115

Model performance against observations is the subject of further study (Carslaw, 2013) and is not considered further here. It is enough to note that the performance of all eight models during July 2006 as a whole was considered satisfactory and all of the models were able to account satisfactorily for the observed day-to-day variations in the daily peak ozone levels. Because the performance of each model was considered satisfactory, there was no reason to distinguish one set of model results from another and accordingly we have anonymised the models. Each set of model predictions was considered an equally plausible set of possible answers to the policy-relevant questions:

123

Do the models agree on the sensitivities to peak O₃ levels to NO_x and VOC emissions?

- Do the models agree on the relative importance of UK precursor emissions to those in the
 rest of Europe?
- Do the levels of agreement improve if those models and days that had poorer matches
 between models and observations were excluded?
- 128

129 3. NO_x- versus VOC-sensitivity

An important issue in developing strategies for amelioration of ground-level O₃ is whether to reduce
 NO_x emissions or VOC emissions or both. To address this issue, attention has been focussed in the
 modelling on the impact of four simple NO_x and VOC emission scenarios, keeping all other emissions
 constant:

- S1: 30% reductions in man-made NO_x emissions across Europe,
- S2: 30% reductions in man-made VOC emissions across Europe,
- S3: 30% reductions in man-made NO_x and VOC emissions across Europe,
- S4: 30% reductions in man-made NO_x and VOC emissions across the UK.

138 The choice of 30% is arbitrary. It is nevertheless comparable to the scale of emission reductions that 139 policy-makers commonly consider. It has been chosen because it is neither too small nor too large 140 and to be consistent with a large literature on photochemical ozone model sensitivity to VOC and 141 NO_x emissions, see for example, Sillman (1999) and Sillman and He (2002). To assess the impact of 142 30% across-the-board reductions in man-made NO_x and VOC emissions relative to the 2006 base 143 case, each model ran the S1 and S2 emission scenario cases. The maximum hourly ozone levels 144 simulated for the base case and the two scenario cases for each day of July 2006 were determined for each model. 145

146 Overall impacts on July-mean O₃ levels

The impact of the 30% reductions in NO_x emissions carried out across the UK and the Rest of Europe (RoE) (Scenario S1) on the July mean daily maximum ozone levels varied considerably between the eight models. O₃ responses (base case minus scenario case) covered the range from -2.0 ppb to +2.0 ppb, with three models producing an increase (-ve response) and five models producing a decrease (+ve response). Figure 2 presents a 'box and whisker' plot of the eight model responses. The interquartile range, shown as a shaded box, confirms that the median model response of +0.4 ppb was not statistically different from zero.

154

In contrast, Figure 2 shows that the impact on the July mean daily O₃ maximum of 30% reductions in man-made VOC emissions (Scenario S2) was a decrease (+ve response) for all eight models, with responses spanning the range from +0.4 to +3.2 ppb. The median response of +1.2 ppb was statistically significantly different from zero. These model simulations showed that VOC reductions always produced an improvement is air quality, in contrast to the mixed results for NO_x reductions using the July mean daily maximum O₃ as an index.

161 <u>Daily assignments of NO_x- versus VOC sensitivity</u>

162 The responses to the 30% NO_x emission reduction and the 30% VOC emission reduction carried out 163 across the UK and the RoE were analysed by considering the model responses on individual days 164 rather than for the month as a whole. If the O_3 response to a 30% NO_x reduction was greater than 165 that to a 30% VOC reductions, then that day was assigned as NO_x-sensitive. Conversely, if the O₃ 166 response to a 30% VOC reduction was greater than that to a 30% NO_x reduction, then that day was 167 assigned as VOC-sensitive. Table 1 shows the VOC- versus NO_x-sensitive assignments for each day of 168 July for each of the eight models. There was complete agreement on the assignments on only six 169 days, with differing levels of disagreement on the remaining 25 days. However, all models showed 170 how the NO_x- versus VOC-sensitivity could switch on a daily basis from NO_x-sensitive to VOC-

sensitive and back again during the month. The question is which model is giving the correctassignment when there are apparent contradictions.

173

174	Figure 3 presents a scatter plot of the O_3 responses to 30% NO_x reductions against the O_3 responses
175	to 30% VOC reductions for the eight models and for the 15 50-ppb episode days. Also shown is the
176	1:1 correspondence line representing the locus of equal responses. Points above the line have
177	responses to 30% VOC reductions that are greater than to 30% $\rm NO_x$ reduction and so have been
178	assigned as VOC-sensitive. Points below the line have been assigned as NO _x -sensitive. The vast
179	majority of points are located above the x-axis showing that almost all of the points show positive
180	responses to 30% VOC reductions and hence that air quality improves. In contrast, there are a small
181	but significant number of points to the left of the y-axis, showing that some models show negative
182	responses to 30% NO _x reductions, implying that air quality deteriorates.

183

184 The majority of the points in Figure 3 form a 'wedge-shaped' pattern. The apex of the wedge is at 185 the right-hand side of the plot, at the high NOx-response – low-VOC response and widens towards 186 the left-hand side of the plot. There is a tendency for VOC-responses to be smallest when NOx-187 responses are greatest and VOC-responses to be greatest when NOx-responses are negative. This 188 characteristic tendency has its origins in the theory underpinning NO_x- and VOC-sensitivity as 189 demonstrated by Sillman (1999) and Sillman and He (2002). Superimposed on this characteristic 190 tendency is the impact of model uncertainty which is manifest in terms of the relative scatter 191 between the sets of model points. The axis of the wedge-shaped pattern is almost perpendicular to 192 the 1:1 correspondence line. As a consequence, the characteristic tendency and the model 193 uncertainty strongly impact on the location of the points relative to the 1:1 correspondence line and 194 hence on the NO_x- versus VOC-sensitivity assignments. There are 62 points out of the 120 that are

VOC-sensitive and 58 points that are NO_x-sensitive, indicating a slight preponderance in favour of
 VOC-sensitivity for the episode days.

197

198 The above analysis has shown that there can be a considerable level of disagreement between model assignments of policy-relevant characteristics for O₃ during July 2006. Policy-makers expect 199 200 that all models used in their support are able to reproduce real-world behaviour. So now we check 201 to see if, by setting a benchmark for such comparisons, we are able to disregard some model results 202 and to focus only on those that deliver good model performance against observations (for this 203 particular test case). Accordingly we set a benchmark of ± 0.1 for the NMB for each day and 204 disregard model results outside this range. This benchmark is set at an arbitrary level and has been 205 tightened to \pm 0.05 specifically for the PTM model because some information about observed O₃ 206 levels has been used in the selection of the results from multiple replicates using different back-track 207 trajectories, (see the Supplementary Information for further details). The setting of the benchmark 208 level is a compromise: set too low and all model results would be filtered out and set too high and 209 the situation would not substantially change from that in Table 1.

210

Table 2 presents the NO_x- versus VOC-sensitivity assignments for only those models that achieved the benchmark NMB of ± 0.1 (± 0.05 for the PTM) on a given day during July 2006. Comparing Tables 1 and 2 shows how setting a benchmark for model performance on each day could drastically reduce the number of table entries. However, there was also a marked reduction in the number of contradictory assignments. Those models that performed better against observations on particular days appeared to give more robust assignments in terms of VOC- versus NO_x-sensitivity. The refinement process in moving from Table 1 to Table 2 has led to a decrease in the proportion of

assigned days from 25 out of 31 to 7 out of 21, thereby increasing the level of consensus betweenthe models.

220

221	Nevertheless, Table 2 shows that selecting for better model performance did not remove all
222	conflicts. Of the 31 days in July 2006, no conflicts were recorded for 20 days, conflicts were recorded
223	on 7 days and no assignments were possible for 4 days. Of the days with conflicts, 4 days were non-
224	episode days with observed maximum hourly O_3 levels below 50 ppb, leaving only 3 days where the
225	conflict in assignment may have some policy significance. Of the days when a clear-cut assignment
226	could be made, twice as many days were assigned to the NO_{x} -sensitive category than to the VOC-
227	sensitive category. Generally speaking then, the 'best' models indicated that actions to control $\ensuremath{NO_x}$
228	emissions rather than VOC emissions would be the more effective approach to reducing episodic
229	peak O₃ levels at Harwell during July 2006.

230

231 All of the four days at Harwell during July 2006 when no assignments were made were episode days, including 18^{th} July on which O_3 levels exceeded 100 ppb. All models had difficulty in simulating O_3 232 233 mixing ratios approaching these levels. It is possible that the observed O_3 levels on this and on the 234 other three days were strongly influenced by O₃ precursor sources that were omitted from or were 235 inadequately included in the emission inventories. Possible candidate sources include agricultural 236 burning and forest fires as explained in Section 2. Equally well, there may be difficulties in describing 237 meteorological conditions during these episode days. In any case, filtering by model performance 238 removed the NO_x- versus VOC-sensitivity assignments that may have been based on possibly 239 inadequate evidence.

The conclusion from Table 2 is that there are fewer contradictory NO_x- versus VOC-sensitivity assignments when model performance is used to select the 'better' or 'best' models on each day. The 'best' model changed from day to day and no single model was 'best' on all days. The choice of benchmark based on a daily NMB in the range \pm 0.1 (and \pm 0.05 for the PTM) was arbitrary and the selection of a different benchmark would change the character of Table 2. However, two conclusions would still stand, namely: selecting 'best' models reduces apparently contradictory assignments and no one model would always be the 'best' model on all days.

248

249 4. UK- versus Rest of Europe dominance

A further important issue for UK policymakers has been whether the balance of effort in terms of O_3 precursor emission reductions should be focussed on UK emissions or on emissions from the Rest of Europe (RoE). To assess this issue, attention has been directed to the simple emission scenarios S3 and S4, which focus on the influence of O_3 precursor sources in the UK versus those across Europe as a whole. Figure 2 presents a 'box and whisker' plot of the eight model responses to precursor emissions reductions carried out across Europe as a whole (S3) and across the UK (S4).

256

Since the UK emissions were included in the European emissions, an estimate of the impact of the
RoE emissions could be obtained by subtraction of the UK impacts from the European (UK+RoE)
impacts. Therefore if, for a given day and given model, the O₃ response to the 30% reduction in UKonly VOC and NO_x emissions was greater than the difference in response between European
emissions reductions and UK emission reductions, then that day was assigned as UK-dominant.
Conversely, if the response to the reductions in UK emissions was less than the difference in
responses between the European and UK emissions reductions, then that day was assigned as RoE-

264 dominant. This subtraction assumes that O_3 responses are linear and additive, a reasonable working 265 assumption for these relatively small percentage reduction in precursor emissions.

266

Table 3 shows the UK- versus RoE-dominance assignments for each model and for each day in July
2006. There was complete agreement on UK- and RoE-dominance on only five days and some
disagreement on the remaining 26 days. Again, it was apparent that assignment of the major source
regions, whether UK or RoE, varied from day to day and so again the question is which of the model
assignments is correct for each day.

272

A detailed analysis of UK- versus RoE-dominance is hampered by a lack of simple rules such as those 273 274 that exist for NO_{x} -versus VOC-sensitivity. However, a simple scatter plot provides a suitable 275 introduction to the UK- versus RoE-dominant assignments. Accordingly, Figure 4 presents a scatter 276 plot of the O₃ responses to 30% reductions in both NO_x and VOC emissions carried out across the UK 277 and RoE versus the responses to 30% reductions carried out across the UK only, for all models and all 278 15 episode days. Also shown is the 1:1 correspondence line which represents the locus of points 279 where the responses across the UK and the RoE are equal to those across the UK only. Figure 4 280 shows that a small fraction of points lie above the line and that the vast majority of points lie below 281 the line. That is to say, most models indicate that the O_3 levels on most episode days at this location 282 are dominated by ozone precursor sources in the RoE and that the levels on only a few days are 283 dominated by precursor sources in the UK. Subtracting the O₃ responses to the emission reductions 284 in the UK only from the responses to the reductions carried out in the UK + RoE, yields an estimate 285 for the response to the emission reductions carried out in the RoE only. The greater the response to 286 emission reductions carried out across the RoE, the further the points move below the 1:1 line in 287 Figure 4. Responses to RoE-only emission reductions are thus seen to be relatively large compared

with responses to UK-only emission reductions on all episode days and with all models.

289 Nevertheless, the considerable amount of scatter in this figure mean that it is not straightforward to

draw robust conclusions about UK- versus RoE-dominance on specific days using specific models.

291

Over all the episode days and all the models, the average O_3 response to 30% emission reductions in both NO_x and VOC in the UK was 0.0 ± 1.5 ppb. Whereas, that to reductions carried out across the RoE was considerably greater at 2.7 ± 0.7 ppb. Episode days were highly likely to be RoE-dominant and this conclusion was robust to choice of model. It was associated with the preponderance of transport from north-west Europe during July 2006 as noted in Figure 1.

297

To reduce the conflicts between UK- versus RoE-dominance assignments, filtering by model performance against observations was undertaken as shown in Table 4 using the same benchmarks as for Table 2. Again, the number of contradictory assignments has been drastically reduced. Of the 31 days in July 2006, cross-model agreement as to UK- versus RoE-dominance has been reached on 18 days, contradictory assignments on 9 days and no assignments on four days. The possible reasons for the lack of assignments on the four episode days have been highlighted above.

304

Contradictory assignments were found on nine days compared with seven days for NO_x- versus VOCsensitivity. This suggests that UK- versus RoE-dominance is somewhat less robust compared with NO_x- versus VOC-sensitivity. Nevertheless, on the basis of Table 4, it is concluded that the 'best' models gave less contradictory assignments, that the 'best' model changed from day to day and that no model was designated as 'best' model on all days. Generally speaking, the 'best' models indicated that daily maximum O₃ levels at Harwell during July 2006 were impacted more by precursor emission sources in the RoE than by sources within the UK.

313 5. Implementing an ENSEMBLE approach

314 In the field of atmospheric dispersion modelling, conflicting realisations of air quality forecasts are 315 increasingly being resolved through the use of ensembles (Potempski and Galmarini, 2009). In the field of air quality modelling, Van Loon et al. (2007) and Vautard et al. (2009) employed ensembles 316 317 extensively in their study of O₃ and nitrogen dioxide (NO₂) levels across Europe using seven regional 318 air quality models. Following their lead, the arithmetic mean of all eight sets of model results and 319 their sensitivity cases were calculated to develop a synthetic set of model results, ENSEMBLE, which 320 were processed in an analogous manner as the set of eight model results. The benchmark of NMB of 321 \pm 0.1 for each day was applied and the results for the ENSEMBLE were added to Tables 2 and 4.

322

Looking at the ENSEMBLE results in Table 2 for NO_x- versus VOC-sensitivity, there appeared to be no clear advantage from the ENSEMBLE results over the individual models A – H in terms of the number of days with NMBs passing the benchmark. The models A – H showed between 4 and 14 entries, whereas the ENSEMBLE showed 12 entries. The ENSEMBLE confirmed the assignment to NO_xsensitive on five days and added to the conflicting assignments on the remaining seven days. On this basis, it was concluded that the ENSEMBLE approach did not add significantly to the assignment of NO_x- versus VOC-sensitivity for episodic peak O₃ at Harwell, Oxfordshire during July 2006.

330

The ENSEMBLE results for UK- versus RoE-dominance following the implementation of the NMB benchmark, confirmed the assignments based on the individual models A – H on five days and added to the conflicting assignments on the remaining seven days. The ENSEMBLE approach did not add significantly to the assignment of UK- versus RoE-dominance.

336 6. Discussion and conclusions

337 One of the main purposes of air quality modelling in Europe is to assist and support policymakers in 338 the formulation of robust and cost-effective strategies for the control of the transboundary 339 formation and transport of O_3 . Because a number of O_3 precursor emission sources have already 340 been effectively controlled, the remaining policy options tend to be expensive or complex. Options 341 for the further control of VOC emissions involve tackling solvent emissions, industrial emissions or evaporation from the gasoline distribution chain. Those for NO_x emissions involve tackling diesel 342 343 vehicle exhausts and large and small combustion sources. Policymakers in the UK can reasonably ask 344 the modelling community whether the balance of future effort should be focussed on VOC or NO_x 345 emissions, or both, and, in view of the evidence for transboundary O_3 formation and transport, 346 whether future efforts should be focussed on domestic precursor sources or on foreign sources. 347 These considerations have driven the formulation of this present study and its focus on the categoric 348 assignments as to whether the episodic peak O_3 levels in south east England in July 2006 are NO_x- or 349 VOC-sensitive and whether they are dominated by precursor sources within the UK or in the RoE.

350

351 In this study, attention has been focussed on the EMEP monitoring station located at Harwell, 352 Oxfordshire in the rural south east UK. This location was chosen because of its relative remoteness 353 from large population centres. Other candidate stations were ruled out because of their coastal 354 locations (Lullington Heath, Rochester, St Osyth and Sibton) which would have unduly biased the 355 results in favour of transboundary sources rather than local formation and transport. Some stations 356 are too close to London (Teddington, Hillingdon and London Eltham) and roadside stations were 357 ruled out because they would be subject to the influence of local O₃ destruction rather than local 358 formation.

360 However, the focus on a specific station for the analysis may not necessarily play to the strengths of 361 the Eulerian models compared with the Lagrangian or moving parcel models. A strength of grid-362 based models is that they can yield maps showing how O₃ levels and O₃ responses vary spatially over 363 entire regions, such as the south east UK. However, because of potential uncertainties in defining 364 horizontal transport within a spatial resolution of a few km, spatial mismatch may occur between 365 gridded model output and the actual grid square containing an individual monitoring station, i.e. the 366 model may fail to reproduce high O₃ at one particular site on a given day (the criterion used in this 367 study to define good and poor model performance) for a reason unrelated to its skill in general at 368 capturing VOC-NOx-ozone photochemistry over larger spatial and temporal domains. This will 369 potentially be an issue where a large spatial gradient in O_3 occurs in the vicinity of the monitoring 370 site chosen for observation-model comparison. Figure 5 shows the simulated daily maximum hourly O_3 level for the 6th July across the whole of southern England from one of the grid models in this 371 372 intercomparison which illustrates the strong spatial gradient in maximum ozone across the location 373 of the Harwell monitoring station (marked by the black circle in the figure). We therefore note that 374 our approach of utilising data from a single monitoring station to evaluate model performance may 375 somewhat have favoured Lagrangian over Eulerian model approaches if our sole aim had been to 376 evaluate model performance. However, the aim of this study has been to illustrate the issues 377 involved in using models in support of air quality policy formulation rather than the selection of the 378 'best' model.

379

By setting a benchmark in terms of model performance against observations, we have been able to filter the policy-relevant assignments made with eight air quality models of NO_x- versus VOCsensitivity and UK- versus RoE-dominance to obtain a more robust understanding of the origins of the O₃ episodes observed in the south east of England during July 2006. There were fewer

contradictory assignments when model performance against observations was used to select the
'best' model out of the eight models on each day. The 'best' model changed from day to day and no
one model was always designated the 'best' model on all days. The choice of benchmark for the
daily NMB was arbitrary and selection of a different benchmark could change the character of the
analysis.

389

390 In this study, the use of an ensemble approach has been assessed following the suggestions of Van 391 Loon et al. (2007) and Vautard et al. (2009). Both studies reported advantages of using ensembles 392 for the assessment of long-term O_3 levels using annual mean and SUMO35 metrics. There appeared 393 to be little advantage in using ensembles for the assessment of NO₂ levels because the ensemble 394 failed to represent the highest peak values. Our conclusion is that the ensemble approach did not 395 add significantly to the analysis of emission sensitivities at Harwell during July 2006. Our focus was 396 on episodic peak O₃, a metric that is generally underpredicted in models. This may go a long way 397 towards explaining why the ensemble approach offered little advantage in this study.

398

399 These conclusions will need to be extended by further work in the future to cover different regions 400 of north-west Europe and to different months and years with their different advection regimes and 401 hence source-receptor relationships. We urgently need to understand the differences in model 402 formulation that have led to the observed conflicts in model responses, whether these lie in 403 meteorological datasets, biogenic VOC emissions or different temporal profiles in emissions. This 404 work shows that we currently do not have access to a single air quality model that is guaranteed to 405 deliver the most likely outcomes to policy makers in terms of emission sensitivities on each day. It is 406 important to maintain a diversity in model approaches to further the development of our 407 understanding of O₃ transboundary formation and transport in north west Europe. We need a wide

- 408 diversity of models, not because it would guarantee a more accurate ensemble, but because it
- 409 would give more chances for model results to be acceptable and robust for policy purposes.

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Model	А	В	С	D	Е	F	G	Н	
1 st	VOC								
2 nd	NO _x								
3 rd	NOx	VOC	VOC	NOx	VOC	VOC	NOx	NOx	
4 th	NOx	VOC	VOC	VOC	VOC	NOx	NOx	VOC	
5 th	NO_x	VOC	VOC	VOC	NOx	NO_x	NOx	VOC	
6 th	VOC	VOC	NOx	VOC	NO_x	NO_x	NO_x	NO_x	
7 th	NO_x	VOC	VOC	VOC	VOC	NO_x	NO_x	VOC	
8 th	NO_x	VOC	NOx	NO _x	NO_x	NO_x	NO_x	NO_x	
9 th	NO_x	NOx	NOx	NO _x	NO_x	NO_x	NO_x	VOC	
10^{th}	NO_x	VOC	VOC	NOx	NOx	VOC	VOC	VOC	
11 th	NO_x	NOx	NOx	VOC	NO_x	NO_x	NO_x	NO_x	
12 th	NO _x	VOC							
13 th	VOC								
14 th	NO _x	VOC	VOC	NO _x	NO _x	VOC	NO _x	NO _x	
15 th	NO _x	VOC	VOC	VOC	NOx	VOC	VOC	NO _x	
16 th	VOC	NO _x							
17 th	VOC	VOC	VOC	VOC	VOC	VOC	NO _x	VOC	
18 th	VOC	VOC	VOC	VOC	VOC	NOx	NOx	VOC	
19 th	NO _x	VOC	VOC	NO _x	VOC	NOx	VOC	NO _x	
20 th	NOx	VOC	VOC	NO _x	NOx	NOx	NOx	NOx	
21 st	NO _x								
22 nd	VOC	VOC	VOC	NO _x	NO _x	VOC	NO _x	NO _x	
23 rd	VOC	VOC	VOC	NOx	VOC	NOx	NOx	VOC	
24 th	VOC	VOC	VOC	VOC	VOC	VOC	NOx	VOC	
25 th	NOx	VOC	VOC	VOC	NOx	NOx	NOx	NOx	
26 th	VOC	VOC	VOC	VOC	NOx	NOx	NOx	NOx	
27 th	VOC	VOC	VOC	NOx	NOx	VOC	NOx	NOx	
28 th	NOx	VOC	VOC	NO _x	VOC	VOC	NOx	NOx	
29 th	NO _x								
30 th	NO _x	NOx							
31 st	NO _x	VOC	VOC	NO _x	NO_x	NO _x	NO _x	NO _x	

Table 1. Assignments of NO_x - or VOC-sensitivity for each model A-H for each day of July 2006.

Notes: highlighting denotes days when all assignments agree.

	Obs,									
Model	ppb	А	В	С	D	Е	F	G	Н	ENS ^a
1 st	82							VOC		
2 ^{na}	80			NO _x						
3 ^{ru}	81							NO _x		
4 ¹¹	79									
5"'	38	NO _x								
6 th	60	VOC		NO _x		NO _x	NO _x		NOx	VOC
/ th	29			VOC						
8 th	34	NO _x		NO		NO _x				NO _x
9 th	32	NOx		NO _x		NOx				
10 11 th	21			VUC				NO		NO
11 13 th	39	NO		VOC						
12 12 th	55 22	NOx		VUC	VOC	VOC		VOC		VUC
13 1 4 th	33	VUC			VUC	VUC		VUC		
14 1 r th	4Z						VOC			
15 16 th	51 75						VUC	VOC		
17 th	75							VUC		
18 th	106									
19 th	103	NO.								
20 th	58	NO.						NO.		NO.
21 st	61				NO			NO		
22 nd	56				NO _v			NO _v	NO,	voc
23 rd	43	VOC			- *		NO _x	NO	- X	VOC
24 th	72						- ^	- ^		
25 th	69		VOC			NO _x				NO _x
26 th	65			VOC	VOC	NOx				VOC
27 th	63		VOC							
28 th	43		VOC		NO _x				NO _x	VOC
29 th	36					NO _x	NO _x	NO _x	NOx	NO _x
30 th	36	NOx	NO _x		NOx	NOx				NOx
31 st	43	~	VOC		NOx	~	NO _x	NO _x		

Table 2. Assignments of NO_x - or VOC-sensitivity for each model A-H for each day of July 2006 having filtered the results on the basis of model performance for each day using a NMB threshold of ± 0.1 (± 0.05 for the PTM model), together with the observed maximum hourly mean ozone concentration.

^a ENS refers to the ENSEMBLE

Table 3. Assignments of UK- or Rest of Europe-dominance for each model A-H for each day of July 2006.

Model	А	В	С	D	Е	F	G	Н
1 st	RoE	UK	RoE	UK	RoE	RoE	RoE	RoE
2 nd	RoE							
3rd	RoE							
4 th	RoE	UK	RoE	RoE	RoE	RoE	RoE	RoE
5 th	RoE	RoE	RoE	RoE	UK	RoE	RoE	RoE
6 th	UK	RoE	RoE	RoE	UK	RoE	RoE	RoE
7 th	UK	RoE	RoE	RoE	RoE	UK	UK	RoE
8 th	RoE	UK						
9 th	UK	RoE	RoE	RoE	RoE	UK	RoE	RoE
10^{th}	UK	UK	RoE	RoE	RoE	UK	RoE	RoE
11 th	UK	UK	UK	RoE	UK	UK	UK	UK
12 th	UK	RoE						
13 th	RoE							
14 th	UK	RoE	RoE	RoE	UK	RoE	UK	UK
15 th	UK	UK	RoE	RoE	UK	RoE	RoE	UK
16 th	RoE							
17 th	RoE	RoE	RoE	RoE	UK	RoE	RoE	RoE
18 th	RoE							
19 th	RoE	UK						
20 th	RoE	RoE	UK	RoE	RoE	RoE	RoE	RoE
21 st	RoE	RoE	UK	RoE	RoE	RoE	UK	UK
22 nd	UK	UK	RoE	RoE	RoE	RoE	RoE	RoE
23 rd	UK	RoE	RoE	UK	RoE	UK	UK	RoE
24 th	UK	UK	RoE	RoE	RoE	UK	UK	RoE
25 th	RoE	UK	RoE	RoE	RoE	RoE	RoE	RoE
26 th	RoE	RoE	RoE	UK	RoE	RoE	UK	RoE
27 th	UK	RoE	RoE	RoE	RoE	RoE	UK	UK
28 th	UK	RoE	RoE	RoE	RoE	RoE	UK	UK
29 th	RoE	UK	RoE	RoE	RoE	RoE	RoE	RoE
30 th	UK	RoE	UK	UK	RoE	UK	UK	UK
31 st	RoE	RoE	RoE	RoE	RoE	RoE	UK	RoE

Notes: highlighting denotes days when all assignments agree.

Table 4. Assignments of UK- or RoE-dominance for each model A-H for each day of July 2006 having filtered the results on the basis of model performance for each day using a NMB threshold of \pm 0.1 (\pm 0.05 for the PTM model).

Model 1 st	A	В	С	D	Е	F	G RoF	Н	ENS ^a
2 nd			RoE						
3 rd							RoF		
4 th									
5 th	RoE								
6 th	UK		RoE		UK	RoE		RoE	RoE
7 th			RoE						
8 th	RoE				RoE				RoE
9 th	UK		RoE		RoE				
10^{th}			RoE						
11 th							UK		UK
12 th	UK		RoE				RoE		RoE
13 th	RoE			RoE	RoE		RoE		
14 th							UK		
15 th						RoE	RoE		
16 th							RoE		
17 th									
18 th									
19 th	RoE								
20 th	RoE						RoE		RoE
21 st				RoE			UK		
22 rd				RoE			RoE	RoE	RoE
23 ^{ru}	UK					UK	UK		RoE
24 th									
25 th		UK			RoE				RoE
26 th			ROE	UK	ROE				ROE
27 th		ROE							
28 th		ROE		ROE	D . F	D . F		UK	ROE
29" 20 th		Def			KOE	KOE	KOE	KOE	KOE
30°	UK	KOE		UK	KOE	DeF			KOE
51		KOE		KOE		KOE	UK		

^a ENS refers to the ENSEMBLE



Figure 1. Daily maximum hourly ozone concentrations for all eight models A-H and the observations for each day of July 2006 at the rural Harwell, Oxfordshire site. Also shown at the bottom of the figure are the daily advection regimes as shown by Lamb Weather types (LWT), NILU FLEXTRA trajectories (NILU) and NAME air history maps (NAME, see text.



Figure 2. Box-whisker plots of the changes in July mean daily maximum ozone concentration across the eight models, for the S1 - S4 scenarios. Shaded box: interquartile range, black square: median.



Figure 3. Scatter plot of the eight model O_3 responses to 30% NO_x reductions versus 30% VOC reductions for the episode days of July 2006. Also shown is the 1:1 correspondence line above which points indicate VOC-sensitive model simulations and below which they indicate NO_x-sensitive simulations.



Figure 4. Scatter plot of the O_3 responses on episode days for the eight models to 30% reductions in NO_x and VOC emissions carried out across the UK and the RoE versus the O_3 responses to 30% reductions in NO_x and VOC emissions carried out across the UK only.



Figure 5. Simulated maximum hourly ozone across southern England on the 6th July 2006 from one of the Eulerian grid models in the model intercomparison. The black circled cross symbol marks the location of the Harwell monitoring site.

SUPPLEMENTARY INFORMATION

ANALYSIS OF UK AND EUROPEAN NO_x AND VOC EMISSION SCENARIOS IN THE DEFRA MODEL INTERCOMPARISON EXERCISE

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1. Details of the Models

1.1 CMAQ – AEA

The CMAQ – AEA model is an application of the United States Environmental Protection Agency (EPA) Community Multiscale Air Quality (CMAQ) modelling system which is a third-generation air quality model available online at www.cmaq-model.org . CMAQ is designed for applications ranging from regulatory and policy analysis to understanding the complex interactions of atmospheric chemistry and physics. It is a three-dimensional Eulerian (i.e., gridded) atmospheric chemistry and transport modelling system that simulates ozone, particulate matter (PM), toxic airborne pollutants, visibility, and acidic and nutrient pollutant species throughout the troposphere. Designed as a "oneatmosphere" model, CMAQ can address the complex couplings among several air quality issues simultaneously across spatial scales ranging from local to hemispheric. The CMAQ source code is highly transparent and modular to facilitate the model's extensibility through community development by members of the air quality modelling community. CMAQ was first developed in the late 1990's, the latest version 4.7.1 released in 2010. In the CMAQ – AEA implementation, the model has been run at horizontal resolutions of 48km (Europe) and 12km (UK) for this study. A new version at 50km and 10km is currently used for the forecast. The 48+12km simulation uses a 26 layer vertical structure with 12 layers below 800m and a lowest layer of 9 m. The 50+10km forecast uses 19 layers, the lowest at 18m this increases the stability of the weather forecast. For limited studies the resolution was reduced to 4km.

European emissions are based on the 2006 EMEP emissions. UK emissions are based on the 2006 NAEI. Temporal profiles were used for the main emission SNAP sectors. Natural emissions are based on the Biogenic Potential Inventory. Numerical weather data are produced using WRFv3 on the same scale as CMAQ. Boundary and forcing conditions are provided by ECMWF for 2006 and GFS forecast is used for the daily AQ forecast. The chemical mechanism used for the AQ forecasting is Carbon Bond 05 with extensions for Cl, aqueous and aerosol chemistry. The alternative chemical mechanisms available in CMAQ v4.7 is SAPRC-99. CB-IV and RADM2 are available in earlier versions. Dry deposition currently runs within the MCIP (Meteorology Chemistry Interface Processor) and uses a surface exchange aerodynamic method using surface resistance, canopy resistance, and stomatal resistance to compute dry deposition velocities.

1.2 CMAQ – King's College London

The CMAQ – King's College London is an application of a 3-D Eulerian grid air quality model. CMAQ was released to the public in June 1998 by the United States EPA. The primary goals of the model are to improve 1) the environmental management community's ability to evaluate the impact of air quality management practices for multiple pollutants at multiple scales and 2) the scientist's ability to better probe, understand, and simulate chemical and physical interactions in the atmosphere. The CMAQ modelling system is set up at the ERG for both current and future policy assessment. Currently, the model is used as part of health impact assessment research at the ERG (MRC centre).

Domain setting: Domains with 4 nested level (23 vertical levels) Dom1: 81km grid spacing, 47 x 44 cells Dom2: 27km grid spacing, 39x39 cells Dom3: 9km grid spacing, 66x108 cells Dom4: 3km grid spacing, 72x72 cells Dom5: 1km grid spacing, 62x51 cells In the present study, European emissions were based on EMEP and UK NAEI emissions. Meteorological data were based on WRF3.1. The chemical mechanism used was Carbon Bond-05 with aerosol and aqueous chemistry. The dry deposition scheme was based on a surface exchange aerodynamic method which uses surface resistance, canopy resistance and stomatal resistance to compute dry deposition velocities.

1.3 CMAQ – University of Hertfordshire

The CMAQ modelling system configuration is as used by Appel et al. (2012) for AQMEII for the European domain using a horizontal grid spacing of 18 km. A detailed description of the anthropogenic emissions used is available in Pouliot et al. (2012). Biogenic emissions of isoprene and terpene, calculated using the Model of Emissions of Gases and Aerosols from Nature (MEGAN; Guenther and Wiedinmyer, 2007; Sakulyanontvittaya et al., 2008), are included on the same resolution as the anthropogenic emissions. The fire emissions were bases on 2006 daily fire estimates from the Moderate Resolution Imaging Spectroradiometer (MODIS) fire radiative power product (Sofiev et al., 2009).The calculations used 34 vertical layers. Model options employed include the CB05 chemical mechanism with chlorine chemistry extensions, the AERO5 aerosol module, the Asymmetric Cloud Model 2 (ACM2) PBL scheme. The simulations utilised boundary concentrations from the GEOS-Chem global model (see Schere et al., 2012). The meteorological fields were obtained from the Weather Research and Forecasting (WRF) model (see Vautard et al., 2012). For the WRF model run, the initial conditions and lateral boundary conditions were derived from the European Centre for Medium-range Weather Forecasts (ECMWF) gridded analyses.

1.4 EMEP4UK – Centre for Ecology and Hydrology

The EMEP4UK model (Vieno et al., 2010) is a Eulerian grid model based on the EMEP Unified model (Simpson et al., 2012). The development of the EMEP4UK model first started in 2006 by Massimo Vieno (University of Edinburgh, CEH Edinburgh), and Peter Wind and David Simpson (Norwegian Meteorological institute).

EMEP4UK is a nested model run at a spatial resolution of 50 km x 50 km (170 x 133 grid) over the full EMEP extended European regional domain and at a finer resolution of 5 km x 5 km (222 x 260 grid) over a British Isles domain for the main model results.

NAEI emissions data have been used for the UK and EMEP emissions data have been used

everywhere else. Meteorological data have been obtained from the WRF model versions 2.2, 3.1.1, and 3.2. The EMEP Unified model chemistry scheme has been used although more chemical schemes are going to be available with the new version of the EMEP Unified model. The EMEP Unified model deposition scheme has been used to treat dry deposition.

1.5 Ozone Source Receptor Model (OSRM)

The OSRM is a Lagrangian trajectory model whose development has been led by AEA working through an enduring consortium of leading UK experts under contract to Defra (and previous Departments) since 1999. Following the initial design of the model in a research and development stage, various features of the model were enhanced to improve model performance, to take account of further developments in the underlying science and to make the model more suitable for direct application to Defra air quality policy. Since around 2005, the emphasis has shifted from development to maintenance and application of the model as a policy tool for examining the response of the UK ground-level ozone climate to changes in precursor emissions in the UK and Europe.

OSRM uses NAEI 1x1km emissions data for NO_x, VOCs, CO and SO₂ grouped into 8 source sectors for the UK. Over the Rest of Europe in the EMEP domain: EMEP 50x50km emissions data are used in combination with country totals for scaling to years up to 2020. Temporal profiles for man-made emission sources are employed for the different sectors. The NAEI VOC speciation profile is used and the assignment of the 664 individual VOCs in the NAEI speciated inventory to the 13 VOCs in the OSRM is based on reactivity and structural considerations. Gridded emissions for shipping are based on the Entec studies. An emission term is added to the emission rate of isoprene to represent the natural biogenic emissions from European forests and agricultural crops. The emission estimates can either be the same as those used in the UK PTM or from the biogenic inventory produced using the PELCOM land cover dataset and the TNO tree species inventory.

The UK Met Office provides meteorological datasets derived from the NAME model. 30 boundary layer meteorological parameters are provided at 6-hourly resolution over a year, covering a domain from 30°W to 40°E and 20° to 80°N at 1° spatial resolution. These data are used to derive 96-hour back trajectories to specified receptors. The OSRM now has meteorological data in this form for each calendar year from 1999 to 2009.

The current version of the OSRM uses an updated version of the mechanism in STOCHEM: 70 chemical species involved in 195 thermal and photochemical reactions. An experimental version of the OSRM (Version 25) has been using the most reduced form of the latest CRIv2 (CRIv2-R5) mechanism (196 chemical species, 555 reactions) which is linked to the Master Chemical Mechanism.

Dry deposition processes are represented using a conventional resistance approach, in which the rate of dry deposition is characterised by a deposition velocity. Different deposition velocities are used over land and sea. The ozone deposition velocity over land has an imposed diurnal and seasonal cycle. The OSRM works in conjunction with a surface ozone flux model which has been updated recently with the latest parameterisations from the SEI DO₃SE model and treatment of dry deposition is currently being modified to give reduced deposition during dry periods.

1.6 NAME – Met Office

The NAME model was originally developed by the Met Office's Atmospheric Dispersion Group following the Chernobyl incident to simulate medium and long range transport and wet and dry deposition of radionuclides. NAME is three-dimensional a Lagrangian dispersion model that simulates the dispersion, chemistry and deposition processes occurring in the atmosphere. The model runs employs three-dimensional meteorological fields from the Met Office Unified Model. The model is well documented and has numerous applications, for example modelling volcanic eruptions, accidental releases of radionuclides, the spread of foot and mouth disease and air quality. A detailed description of the NAME model physics can be found in Ryall and Maryon (1998) and a description of the atmospheric chemistry model applications can be found in Redington et al., (2009).

Pollutant emissions are represented by releasing millions of air parcels, each able to represent the released mass of many different species. The air parcels are carried by the three-dimensional wind field obtained from the Unified Model (UM). Local turbulent motion is simulated using a random walk technique which requires a diffusion coefficient calculated from the local turbulent velocity variance and the local turbulent timescale. Above the boundary layer these two quantities are fixed, but within the boundary layer they are defined in terms of the local atmospheric stability and local surface quantities. The UM provides direct output of boundary layer height for use in NAME.

NAME's chemistry scheme is based on that of the Met Office's global STOCHEM model. NAME's dry deposition scheme is based on the concept of a deposition velocity and has various degrees of

sophistication. In its simplest form, a fixed deposition velocity for a given species is specified. More generally, a resistance analogy is used to calculate a species dependent deposition velocity. The surface resistance term, denoting the resistance to capture by the surface itself, for a given species can either be a simple fixed value or a more explicit parameterisation dependent on land surface properties. The laminar sub-layer resistance term, representing the resistance to transport through the thin quasi-laminar layer adjacent to the surface, is parameterised according to gaseous or aerosol species, and for aerosol species is dependent on the particle size. The deposition scheme is applied to all air parcels within the boundary layer.

The model domain was 14°W-19.9°E, 42°N-62°N with chemistry and output grid set to ~10km x 10km (0.15° longitude, 0.09° latitude). The model was run using emissions data for 2006 from the National Atmospheric Emissions Inventory (NAEI) over the UK (http://www.naei.org.uk) and from the European Monitoring and Evaluation Programme (EMEP) over the rest of Europe (http://www.emep.int). All emissions were assumed to be constant throughout the year at the annual rate. A daily cycle, varying according to the day of the week, was applied to pollutants emitted by road traffic. Over the UK the NAEI emissions were split into large point sources (containing specific release height information) and small area sources (4 km x 4 km) and large area sources (20 km x 20 km) with release heights of 0–20 m for traffic sources and 0–50 m for other sources. The EMEP emissions data was released from 0-100m.

NAME was run using meteorological data provided by the Met Office Unified Model in the form of three dimensional three hourly met fields, with a horizontal resolution of 0.375° latitude by 0.5625° longitude (~40 x 40 km over the UK), and thirty three vertical levels.

1.7 Air Quality in the Unified Model AQUM – Met Office

AQUM is a limited area configuration of the Met Office Unified Model (MetUM) which uses the UKCA chemistry scheme. The MetUM is a sophisticated system capable of modelling regions from limited areas to globally and with timescales from less than hourly to climate scales. UKCA development first began in 2003 as part of a joint project initially comprising the Met Office and the universities of Cambridge and Leeds, with the aim of building a chemistry and aerosols sub-model within the Met Office's Unified Model for use in climate modelling. Since 2005, AQUM (Air Quality in the Unified Model) has been developed by the Met Office as a configuration of UKCA for modelling regional air quality. AQUM is run online, as part of the Met Office Unified Model, which is an Eulerian meteorological model.

For modelling air quality in the United Kingdom, the following emissions data sets are typically used: NAEI emissions at 1km x 1 km resolution over the UK, ENTEC - 5km x 5km emissions (2007) for shipping surrounding the UK and EMEP emissions at 0.5° x 0.5° over the remainder of Europe. AQUM uses the RAQ (Regional Air Quality) scheme, which is an updated version of the STOCHEM chemical mechanism. Dry deposition is based on a Wesely scheme.

1.8 Photochemical Trajectory Model (PTM) – rdscientific

The PTM model is a moving air parcel trajectory model that is used to describe photochemical ozone and fine particle formation in north west Europe. The PTM model is used to quantify the contribution made by each VOC species and each VOC source category to the long-range transboundary formation and transport of ozone and PM across North West Europe. These contributions are defined in terms of Photochemical Ozone Creation Potentials POCPs and SOAPs. This is the only European model able to evaluate the role of a wide range of VOCs and their sources in ozone policy formation. POCPs are widely used in a wide range of policy analyses and in life cycle analyses.

The PTM uses SO₂, NO_x, NH₃, VOCs, CO and CH₄ emissions taken from 2010 version of NAEI for the UK and SO₂, NO_x, NH₃, VOCs, CO and CH₄ emissions for the rest of Europe were taken from the EMEP webdab (2010). Isoprene emissions were taken from EMEP. Terpene emissions were taken from Hope Stewart and Nick Hewitt for UK and GEIA for Europe.

4-day 3-D back-track trajectories from Met Office Unified model providing latitude, longitude, altitude, boundary layer depth, temperature were used to describe the meteorological processes. Between 30 and 1,000 equal probability trajectories arriving at each arrival point between 15:00 and 15:15 z each day from Met Office NAME model were used in the present study. A Wesely dry deposition velocity scheme was used but no treatment was given for wet deposition. All model results were obtained using the CRIv2 chemical mechanism. Details of the model description are given in Derwent et al., (2009).

The PTM was run with each of the 30 equal probability trajectories for each day. The trajectory that gave the closest results to the observations for ozone for each day was selected and these results were used in the Defra model intercomparison.