Predictions of UK Regulated Power Station Contributions to Regional Air Pollution and
 Deposition: A Model Comparison Exercise

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16 ABSTRACT

17 Contributions of the emissions from a UK regulated fossil-fuel power station to regional air 18 pollution and deposition are estimated using four air quality modeling systems for the year 2003. 19 The modeling systems vary in complexity and emphasis in the way they treat atmospheric and 20 chemical processes, and include the Community Multiscale Air Quality (CMAQ) modeling 21 system in its versions 4.6 and 4.7, a nested modeling system that combines long- and short-range 22 impacts (referred to as TRACK-ADMS), and the Fine Resolution Atmospheric Multi-pollutant 23 Exchange (FRAME) model. An evaluation of the baseline calculations against UK monitoring 24 network data is performed. The CMAO modeling system version 4.6 dataset is selected as the 25 reference dataset for the model footprint comparison. The annual mean air concentration and 26 total deposition footprints are summarized for each modeling system. The footprints of the power 27 station emissions can account for a significant fraction of the local impacts for some species (e.g. more than 50% for SO₂ air concentration and non-sea-salt sulfur deposition close to the source) 28 29 for 2003. We calculate the spatial correlation and the coefficient of variation of the root mean 30 square error (CVRMSE) between each model footprint and that calculated by the CMAQ 31 modeling system version 4.6. The correlation coefficient quantifies model agreement in terms of

32 spatial patterns, and the CVRMSE measures the magnitude of the difference between model

33 footprints. Possible reasons for the differences between model results are discussed. Finally,

34 implications and recommendations for the regulatory assessment of the impact of major

35 industrial sources using regional air quality modeling systems are discussed in the light of results

36 from this case study.

37

38 IMPLICATIONS

39 Modeling tools are required to assess the contribution of industrial sources to ambient levels of 40 air pollution, acid deposition, and eutrophication. This study evaluates the performance 41 characteristics of regional air quality modeling systems in predicting contributions of the 42 emissions from a UK regulated fossil-fuel power station to regional air pollution and deposition. 43 It contrasts acid deposition modeling approaches used in the UK and demonstrates the sensitivity 44 of the modelling systems to large emission changes. This work suggests considering an ensemble 45 average of model calculations to provide an estimate of the uncertainty associated with an 46 industrial source footprint.

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48 INTRODUCTION

49 Despite large reductions in terms of absolute emission levels since the 1990s, the power generation sector remains a significant contributor to pollutant emissions in the UK.¹ The 50 51 pollutants emitted by power stations include sulfur dioxide (SO_2) , oxides of nitrogen (NO_y) , 52 and particulate matter smaller than 10 μ m in aerodynamic diameter (PM₁₀). The power generation sector contributed 48%, 24%, and 7% to the UK emissions of SO_2 , NO_x , and PM_{10} 53 respectively, in 2007.¹ These three air pollutants are associated with negative effects on human 54 55 health (e.g. respiratory problems) and damage to the environment. Deposition of sulfur and 56 nitrogen can lead to critical loads for acidity levels being exceeded in sensitive terrestrial and 57 aquatic ecosystems. Additionally, nitrogen deposition can cause eutrophication of ecosystems. Emissions of SO_2 , NO_x , and PM_{10} from power stations are regulated by the 'EC Directive 58 59 2001/80/EC on the limitation of emissions of certain pollutants into the air from large combustion plants.² The European Union (EU) has set air quality limit values for a range of air 60 61 pollutants, which are specified by the 'EC Directive 2008/50/EC on ambient air quality and cleaner air for Europe.³ In the UK, national air quality standards and objectives have been set to 62

meet these legal limit values.⁴ Recognizing that air pollutants cross national borders, national 63 ceilings for emissions of key pollutants (including SO₂ and NO_x) have been put in place at the 64 EU level as part of the Convention on Long-Range Transboundary Air Pollution, and set for 65 66 2010 in the 1999 Gothenburg Protocol. In this context, the regulatory assessment of power 67 stations (and more generally large industrial sources) is an important factor to include in the 68 design of a cost effective strategy to meet emission-ceiling targets and to reduce air pollution, 69 acidification and eutrophication of ecosystems, and climate change impacts. Such an assessment 70 requires appropriate modeling tools.

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72 A number of air quality modeling systems have already been applied for regulatory purposes in 73 the UK. These modeling systems include a nested modeling system (referred to as TRACK-ADMS,⁵ hereafter), used for national annual audits,⁵ and the Fine Resolution Atmospheric Multi-74 pollutant Exchange (FRAME) model,⁶ used for national assessment of acid deposition.^{6,7,8} 75 76 Recently, the UK Environment Agency has been considering using more advanced (in the way 77 they treat atmospheric and chemical processes) air quality modeling systems, such as the Community Multiscale Air Quality (CMAQ) modeling system,^{9,10} as one of its primary 78 79 regulatory assessment tools. Hence, a model comparison exercise has been setup to examine the 80 performance characteristics of regional air quality modeling systems in relation to regulatory use, 81 and more specifically the response of those modeling systems to large emission changes. For the 82 purpose of this exercise, contributions of the emissions from a UK regulated fossil-fuel power 83 station to regional air pollution and deposition are quantified using the CMAQ modeling system 84 in its versions 4.6 and 4.7, TRACK-ADMS, and FRAME, for the year 2003. 85

86 **BASELINE CALCULATIONS**

87

Setup of the Modeling Systems

88 The formulations of the four air quality modeling systems (the CMAQ modeling system in its

89 versions 4.6 and 4.7, TRACK-ADMS, and FRAME) as regards the treatment of atmospheric and

90 chemical processes are quite different, as are the requirements in terms of input datasets (e.g.

91 meteorology, emissions).

The CMAO modeling system is a state-of-the-science Eulerian air quality modeling system,¹¹ 93 94 which has been used extensively for a variety of applications (e.g. retrospective, forecasting, regulatory,^{12,13} process-level applications). It can simulate the dynamics and composition of the 95 atmosphere over a broad range of spatial and temporal scales in a consistent framework based on 96 97 first-principles solutions. The setup and operational evaluation of the CMAQ modeling system, version 4.6, at a horizontal resolution of 5 km for the UK is detailed by Chemel et al.¹⁰ The 98 99 calculations performed with the CMAQ modeling system, version 4.7, have been configured to 100 be as close as possible to those of version 4.6 (e.g. same grid coordinates, chemical schemes, 101 meteorological fields, similar treatment of chemical initial and boundary conditions for the outer domain). Foley et al.¹⁴ documented the major changes from version 4.6 to 4.7 and their impact 102 103 on model performance characteristics.

104

TRACK-ADMS is a modeling system used to produce annual high-resolution maps of air 105 concentration of SO₂, NO_x, and PM₁₀, and of deposition of non-sea-salt (nss) sulfur (SO_x) and 106 nitrogen across the UK. It combines the Trajectory Model with Atmospheric Chemical Kinetics 107 (TRACK),¹⁵ for long-range impacts and the Atmospheric Dispersion Modelling System 108 (ADMS),¹⁶ for short-range impacts. The setup of TRACK-ADMS is as that used for national 109 annual audits.⁵ The horizontal resolution is 20 km for the long-range Lagrangian chemistry-110 111 transport model, TRACK, at distances greater than 50 km from the source, and 1 km for the 112 short-range dispersion model, ADMS, at distances less than 50 km from the source. Model 113 outputs are adjusted by calibration factors used either to adjust modeled values based on measurements or to account for transport and sources not directly modeled.⁵ It is worth noting 114 that TRACK-ADMS does not discriminate between oxidized nitrogen (NO_v) and reduced 115 116 nitrogen (NH_x) deposition and provides only total deposition (i.e. sum of wet and dry 117 depositions) as a standard output. The modeled wet deposition of nss sulfur is calculated as the 118 wet deposition from long-range sources of sulfur alone. Short-range wet deposition of sulfur was assumed to be small compared with its short-range dry deposition,¹⁷ and was not modeled. 119 120 Basically, the travel time from the source is not long enough for significant oxidation of SO_2 to 121 take place, so that wet deposition is not an effective removal process. The modeled wet 122 deposition of nitrogen is calculated as the sum of the wet deposition from long-range sources of 123 nitrogen and the short-range wet deposition of ammonia (NH₂). The modeled dry deposition of

- 124 nitrogen is calculated in the same way but also includes the short-range dry deposition of NO_x.
- 125 Dry deposition is estimated from modeled ground-level concentration assuming a constant dry
- 126 deposition velocity, except for the short-range dry deposition of NH₃, which is derived on an
- 127 hour-by-hour basis throughout the year with a time-varying dry deposition velocity.
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129 FRAME is a Lagrangian chemistry-transport model used to simulate annual mean air concentrations of SO₂, NO_x, and NH₃, and depositions of nss SO_x, NO_y, and NH_x, along 130 131 straight-line trajectories at a horizontal resolution of 5 km in the UK. The setup of FRAME is as that used for national assessment of acid deposition.⁶ A detailed description of the original 132 version of FRAME and its development to improve the representation of sulfur and oxidized 133 nitrogen are given elsewhere.^{18,19} 134 135 136 All the model grids cover the UK (see Figure 1) and model results are presented for the 'UK 137 domain'. The horizontal resolution of the CMAQ modeling system and FRAME is 5 km. For 138 TRACK-ADMS, it is 20 km far from the source and 1 km close to the source. Outputs of the 139 modeling systems have been reprojected on a common grid with an effective horizontal 140 resolution of 5 km in order to accommodate the different grids and horizontal resolutions of the 141 models and to minimize the effects of interpolation due to the reprojection. Note that 142 interpolating outputs of TRACK-ADMS from a horizontal resolution of 1 km to 5 km results in a 143 smoothing effect that will not significantly change its overall performance (since it will perform

- 144 either slightly better or worse depending on location). The vertical resolution is different for each
- 145 modeling system. In the present study, we focus on ground surface air concentration and
- 146 deposition. The assessment of the impact of vertical resolution is being considered for future
- 147 work.
- 148
- 149 Figure 1 here
- 150

151 TRACK-ADMS and FRAME use annual mean observational data to derive the meteorological

152 fields (incl. precipitation map, and wind frequency and wind speed roses) for the chemistry-

- transport model,^{5,6} while the CMAQ modeling system uses outputs from a meteorological model,
- the Weather Research and Forecasting (WRF) model, version 3.0.1.1, nudged towards analyses

155 in the present study.¹⁰ A wind rose is used in FRAME to give the appropriate weighting to

156 directional air concentration and deposition for calculation of mean air concentration and total

157 deposition. It is not practical to harmonize input data for the meteorology in the present model

- 158 comparison exercise, so each modeling system has used its own input meteorological dataset.
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160 Chemical initial and boundary conditions were derived from calculations of larger-scale

161 chemistry-transport models for the CMAQ modeling system¹⁰ and FRAME calculations,⁷ while

- 162 TRACK-ADMS used data from remote sites to estimate the contributions of sources not directly
 163 modeled to air concentration and deposition.⁵
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165 For an effective model comparison, input emissions datasets for the modeling systems have been kept as consistent as possible. The four air quality modeling systems used the same annual 166 anthropogenic emissions data as that used by the CMAQ modeling system.¹⁰ The distribution of 167 168 the emissions in time was not prescribed. Since the focus of the present work is on regulated 169 industrial sources, a detailed emission inventory for point sources in the UK including stack parameters and emissions data by source sectors is required.²⁰ Such a detailed emission inventory 170 171 was specifically compiled for the purpose of the model comparison exercise. For the CMAQ 172 modeling system, emissions from point sources were mixed instantaneously in the entire grid cell 173 indentified at the level of sources plume rise. The Lagrangian plume-in-grid approach, which is 174 implemented in the CMAQ modeling system to resolve the spatial scale of large point sources plumes,²¹ was not used in the present study, as while this option is available in version 4.6, it is 175 176 not supported in version 4.7. Although large point source plumes cannot be represented 177 explicitly by Eulerian air quality modeling systems, their representation can be approximated by using fine grid spacings,²² as is the case in our work. TRACK-ADMS and FRAME are designed 178 179 to track plumes in a Lagrangian reference frame, so that there is no need to further resolve their 180 spatial scale.

181

182 Biogenic gas emissions were included in the CMAQ modeling system and TRACK but not in

183 ADMS and FRAME. They are important for studying regional ozone pollution, but this topic is

184 out of the scope of the present study. Sea-salt emissions contribute to PM₁₀ and sulfur deposition.

185 They were included in the CMAQ modeling system but considered only for PM₁₀ in TRACK-

ADMS and not considered at all in FRAME. In order to compare like to like, we did not considerthe sea-salt contribution to sulfur deposition in our work.

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Evaluation of the Modeling Systems

190 An evaluation of the model baseline calculations against UK monitoring network data for the 191 year 2003 is performed in order to gain insights into the performance characteristics of each 192 modeling system and to provide some guidance as regards the selection of a reference dataset for 193 the model footprint comparison. Modeled air concentrations of SO_2 , NO_x , and PM_{10} are compared with measurements from monitoring sites of the UK Automatic Urban and Rural 194 195 Network (AURN) and those run by the major power plant companies in the UK, information 196 from which is collated by JEP, comprising 82 and 34 sites, respectively. Airport, kerbside, 197 roadside, urban center, and urban industrial AURN monitoring sites were excluded as being non-198 representative of typical background concentrations, while all remote, rural, suburban, and urban 199 background sites were kept for the model evaluation. All the JEP monitoring sites are located in 200 the vicinity of power stations and can be classified as rural or urban background sites. Modeled 201 wet depositions of nss SO_x , NO_y , and NH_x are compared with observational data derived from the Secondary Acid Precipitation Monitoring Network (SAPMN), comprising 38 sites providing 202 203 collection of precipitation and measurements of ion concentrations. Precipitation was collected at 204 those sites using bulk precipitation samplers. The limitations of using this data for the evaluation 205 of wet deposition should be discussed. Previous experience has indicated that bulk collectors do 206 not measure precipitation very well because not all the rainwater is collected. Measurements 207 from such bulk samplers can be tainted by the dry deposition of gas and particles on the funnel surface, which are washed into the sample and thus included in it.^{23,24} Dry deposition was found 208 to contribute around 20% for sulfate (SO_4^{2-}) , 20-30% for nitrate (NO_3^{-}) , and 20-40% for 209 ammonium (NH_4^+) ion concentrations in the UK.²⁴ The dry contribution to wet deposition is not 210 211 quantified for each sample and is thus part of the observational error. In addition, wet depositions 212 derived from site-specific measurements may not be representative of grid cell averages, which 213 may be affected by the spatial variability of rainfall amounts and ion concentrations due to orographic enhancement effects.^{23,25,26} In order to examine the effects of spatial variability in 214 215 rainfall amounts on wet deposition data, we also consider using the UK Met Office precipitation 216 observations gridded at a 5-km horizontal resolution to be compared with that of the bulk

217 collectors. A quantification of the effects of spatial variability of ion concentrations on wet

218 deposition requires further research, which is kept in mind for future work. The spatial coverage

of the monitoring networks is displayed in Figure 1, along with the type (e.g. urban, rural) of theAURN sites.

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The fraction of the model predictions, within a factor of two of the observations (FO2), the correlation coefficient, and the normalized mean bias (NMB) are calculated considering all monitoring sites, for the annual mean air concentration and deposition of the measured species. A summary of the values of these statistical metrics is provided in Tables 1 to 3. Model budgets for nss sulfur and nitrogen deposition are given in Table 4. It should be noted that the year 2003 was very dry with the lowest annual precipitation of the last two decades. This resulted in lower than average wet deposition and higher dry deposition.

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Table 1 here

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- Table 2 here
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- Table 3 here
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- Table 4 here
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Model acceptance criteria for 'operational' evaluation have recently been defined in the UK.²⁷ It 238 239 is recommended that an air quality modeling system is considered acceptable if the FO2 values 240 are greater than 50% and if the NMB values lie within the range -20 - 20% for both air 241 concentration and deposition. Correlation coefficients are not recommended as evaluation 242 metrics because they can be strongly influenced by the presence of outliers when there are a 243 small number of pair values. However, they turn out to be informative in the present study to 244 investigate how sensitive model performance is to the derivation of wet deposition from the 245 measurements of precipitation and ion concentrations.

247 It is worth noting that all the models fulfill the first criterion (namely, FO2 > 50%) for SO_2 , NO_x , and PM_{10} air concentrations, and for nss SO_x , NO_y , and NH_x wet depositions (see Table 248 249 1). Using site-specific measurements of precipitation rather than the gridded UK Met Office 250 precipitation observations leads to larger FO2 values for all the species and modeling systems 251 considered, with the exception of NH, wet deposition for FRAME. Interestingly, the correlation 252 coefficients for wet deposition are increased significantly for all the modeling systems when 253 using the precipitation collected by the bulk collectors (see Table 2). The low correlation 254 coefficients obtained for SO_2 and PM_{10} do not indicate per se poor performance but are the result of a narrow range of concentrations and the presence of outliers for SO_2 . This result 255 256 suggests that the rainfall amounts collected by the bulk collectors provide a better spatial 257 representation of what was measured at the sites than the gridded UK Met Office precipitation 258 observations. In contrast to the first criterion, all models fail to fulfill the second criterion 259 (namely, NMB in the range -20 - 20%, see Table 3). Indeed, the NMB for one or more of the 260 species air concentration and/or wet deposition is outside the range for all the modeling systems. All modeling systems tend to under estimate the annual mean air concentrations of SO₂, NO_x, 261 and PM_{10} (as indicated by a negative NMB in Table 3). The NMB values for wet deposition 262 263 indicate that the precipitation collected by the bulk collectors is less than that derived using the 264 gridded UK Met Office precipitation observations. The NMB absolute values are smaller when 265 using the precipitation collected by the bulk collectors for the CMAQ modeling system, version 266 4.6. Conversely, these values are larger for FRAME. This result is to be expected since FRAME 267 is using an annual mean precipitation map derived from the gridded UK Met Office precipitation 268 observations. As for the CMAQ modeling system, version 4.7, no clear pattern is evident in the wet deposition results with the NMB absolute values increasing for nss SO_x and NH_x, and 269 270 decreasing for NO_v.

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The ranges of variation in the UK wet, dry, and total deposition budgets around the mean values calculated across the modeling systems are -33 - 19%, -40 - 21%, and -16 - 24%, respectively, for nss sulfur, and -26 - 24%, -19 - 13%, and -6 - 10%, respectively, for nitrogen (see Table 4). Overall, there appears more variability in the output of the modeling systems in terms of mass deposited in the UK for nss sulfur than for nitrogen. The annual total deposition of nss sulfur, as 277 calculated by each modeling system for the year 2003, is presented in Figure 2. In the CMAQ 278 modeling system precipitation is calculated explicitly by the WRF model while in the other 279 modeling systems it is derived from an annual mean precipitation map derived from the gridded 280 UK Met Office precipitation observations, and an enhanced washout rate is assumed over hilly areas due to the scavenging of cloud droplets by the seeder-feeder effect.²⁸ Interestingly, the 281 282 CMAO modeling system, version 4.7, produces 72% more nss sulfur wet deposition than version 283 4.6 (see Table 4), while the precipitation field is the same as in version 4.6. Foley et al.¹⁴ 284 incrementally evaluated the effect of the major changes from version 4.6 to 4.7 on model 285 performance characteristics. The changes most relevant to deposition, namely those changes to 286 the resolved cloud model and to the coarse particle treatment, were not found to have a 287 significant impact on sulfur and nitrogen deposition when averaged across monitoring stations. 288 We found that the difference between the two model calculations with the CMAQ modelling 289 system, in terms of nss sulfur wet deposition, is associated with more nss sulfate aerosols formed 290 in version 4.7 than in version 4.6, especially in Scotland. We keep this point in mind for future 291 work. Outside of Scotland, the spatial patterns of nss sulfur total deposition from the different 292 modeling systems are very similar with high deposition simulated over the North of England, the 293 Midlands, the hilly areas of Wales, and the Thames Estuary.

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No single modeling system among those considered in the model comparison exercise provides
the overall best performance but we would emphasize that the purpose of the comparison
exercise is not to identify and select the best performing modeling system.

298

299 FOOTPRINT CALCULATIONS

We have decided to select the CMAQ modeling system version 4.6 dataset as the reference
dataset for the model footprint comparison. The main reasons for the selection are summarized
below:

As opposed to TRACK-ADMS outputs, the CMAQ modeling system and FRAME
 outputs are not adjusted by calibration factors used either to adjust modeled values based
 on measurements or to account for transport and sources not directly modeled. Selecting
 the CMAQ modeling system or FRAME for the reference dataset is scientifically
 preferable because it does not involve any calibration of the outputs.

308 The CMAQ modeling system is the most sophisticated modeling system among those • 309 considered in the model comparison exercise. It can simulate complex physical processes 310 that transport and transform multiple pollutants in a physically realistic process-based 311 way in a dynamical environment. It has been applied to short-term episode modeling as 312 well as the production of annual statistics in a variety of places around the globe, 313 including the UK. Conversely, TRACK-ADMS and FRAME treat some of the chemical 314 processes in a more simplistic way, are limited in the species considered, have a simple 315 representation of meteorology, and have been applied essentially to produce annual 316 statistics in the UK.

- Selecting the CMAQ modeling system version 4.6 rather version 4.7 enables one to
 appreciate changes made to the CMAQ modeling system from one release version to the
 next release version.
- 320

The annual mean air concentration and total deposition footprints are calculated for a fossil-fuel power station located in the South-East of England (see Figure 1) for the year 2003. The choice of the power station is fairly arbitrary but it is not close to the coast, nor near other power plants so that its plume is isolated and directional analysis could be applied to monitoring sites around it. The method used to calculate the footprint of this source consists of calculating the difference between the baseline calculation and that with the source removed.

327

328 The power station emissions can account for a significant fraction of the local impacts for some 329 species for 2003 (see Tables 5 and 6), even though their contributions to the UK annual mean air concentrations of SO_2 , NO_x , and PM_{10} , and total deposition budgets are small (see Table 7). 330 The mean contributions calculated across the modeling systems are 2.45% for SO_2 , 0.60% for 331 332 NO_x , 0.30% for PM_{10} , 2.13% for nss sulfur deposition, and 0.22% for nitrogen deposition. These values are comparable to those reported for similar power stations elsewhere.²⁹ There are 333 334 rather large differences in the predicted maximum contributions of the power station to regional 335 air pollution and deposition (see Table 5). Overall, results from the footprint calculations suggest that the power station contributes, locally, more to SO_2 air concentration and nss sulfur 336 337 deposition than to NO_x and PM_{10} air concentrations, and nitrogen deposition. The maximum contributions for SO₂ air concentration and nss sulfur deposition are more than twice those of 338

339 the other species reported in Table 5, for all modeling systems except TRACK-ADMS, which 340 also predicts a relatively large maximum contribution for PM_{10} . This result reflects the large contribution of the power generation sector to SO_2 emissions in the UK (70% in 2003).¹ In 341 342 comparison, its contribution to NO_x and PM_{10} emissions in the UK in 2003 were 22% and 6%, respectively.¹ The contribution of the power station to regional total deposition of nss sulfur, as 343 344 calculated by each modeling system for the year 2003, is presented in Figure 3. The spatial 345 extent of the nss sulfur total deposition footprint is consistent across the modeling systems and is 346 limited to the South of England. The contribution of the power station to nss sulfur total 347 deposition is most significant close the source. While the footprints from the two calculations 348 with the CMAQ modeling system appear to be very similar, those from TRACK-ADMS and 349 FRAME show some differences. In contrast to the other modeling systems, TRACK-ADMS 350 does not predict the maximum contribution at the location of the source but at some distance 351 downwind of the source, and attaches more importance to the northeast sector. Also, the 352 calculations by TRACK and ADMS look to be loosely coupled. As for FRAME, it tends to give 353 more weight to the southeast direction than the CMAQ modeling system. These differences 354 indicate that the wind fields used by TRACK-ADMS and FRAME differ appreciably from those 355 used by the CMAQ modeling system, which are for the year 2003. 356 357 Table 5 here 358

359 Table 6 here

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The maximum distance from the power station at which its contribution is half of its maximum contribution depends strongly on the modeling system and species considered (see Table 6). This is partly explained by the shape of the distributions of the contributions of the power station to regional air pollution and deposition. Indeed, we found that those distributions for the CMAQ modeling system are more skewed (larger skewness), and more sharply peaked (larger kurtosis) than for TRACK-ADMS and FRAME. This indicates that the contributions are more localized in space for the CMAQ modeling system than for TRACK-ADMS and FRAME. This result can

³⁶¹ Table 7 here

also be inferred from Table 6. Further work is required to identify the reasons for the differencesbetween the shapes of the models distributions.

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373 We calculate the spatial correlation and the coefficient of variation of the root mean square error 374 (CVRMSE) between each model footprint and that calculated by the CMAQ modeling system 375 version 4.6, in the area indicated by a dashed polyline in Figure 3 (see Table 8). The correlation 376 coefficient quantifies model agreement in terms of spatial patterns, and the CVRMSE measures 377 the magnitude of the difference between model footprints. The CVRMSE is a dimensionless 378 measure that is extremely useful when comparing between datasets with different mean values. 379 A CVRMSE value of 10% for a modeling system would indicate that the mean variation in air 380 concentration (or deposition) between this modeling system and the reference modeling system 381 (the CMAQ modeling system version 4.6) is 10% of the mean value of the air concentration (or 382 deposition) calculated by the reference modeling system. Table 8 indicates that the two 383 calculations with the CMAQ modeling system are in good agreement with each other, both in 384 terms of spatial patterns and magnitude of the footprints. Larger differences are found between 385 the footprints produced by the CMAQ modeling system, TRACK-ADMS, and FRAME. At some 386 locations the magnitude of the footprints can differ by more than a factor of two.

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388 Table 8 here

389

390 SUMMARY AND CONCLUSIONS

391 Contributions of the emissions from a UK regulated fossil-fuel power station to regional air 392 pollution and deposition are estimated using four air quality modeling systems for the year 2003. 393 The modeling systems vary in complexity and emphasis in the way they treat atmospheric and 394 chemical processes, and include the CMAQ modeling system in its versions 4.6 and 4.7, 395 TRACK-ADMS, and FRAME. An evaluation of the baseline calculations against UK monitoring 396 network data has revealed that all modeling systems tend to under estimate the annual mean air 397 concentrations of SO_2 , NO_x , and PM_{10} , and that there is a high variability in the output of the 398 modeling systems for nss sulfur and nitrogen deposition. No individual modeling system was 399 found to provide the overall best performance. One needs caution in making regulatory or policy 400 decisions on the basis of one model. However, the agreement is good enough to make broad,

401 general decisions, but this will become more difficult as emissions reductions become harder to

402 implement. The CMAQ modeling system version 4.6 dataset was selected as the most

403 appropriate reference dataset for the model footprint comparison.

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405 The annual mean air concentration and total deposition increments due to the power station were 406 summarized for each modeling system and compared using a range of diagnostic metrics. Differences between model results depend, inter alia, on the treatment of plume chemistry.³⁰ and 407 408 emissions data processing. For instance, for the CMAQ modeling system, emissions from point 409 sources were mixed instantaneously into the entire grid cell indentified at the level of sources 410 plume rise, while for TRACK-ADMS and FRAME point sources plumes are tracked in a Lagrangian reference frame. In addition, the current theoretical understanding of the processes 411 leading to acid deposition is limited.³¹ Detailed process-level studies are needed to pinpoint 412 413 deficiencies in acid deposition modeling. This wide area of research is kept for future work. 414

415 There are large uncertainties in the assessment of contributions of industrial sources to regional 416 air pollution and deposition. A critical question that remains to be examined is whether 417 uncertainties such as those reported in the present work still render such model footprints 418 meaningful for policy applications. Quantifying the uncertainty associated with a single modeling system is extremely difficult given the range of inputs and process calculations.^{32,33} 419 420 Hence, an ensemble average of model calculations could be used to provide an estimate of the 421 uncertainty associated with an industrial source footprint. It has to be recognized that air quality 422 modeling systems such as TRACK-ADMS and FRAME still have run times much faster than 423 those of advanced systems such the CMAQ modeling system. For this reason, such modeling 424 systems are attractive for source-receptor calculations involving a large number of model 425 calculations.

426

Other modeling systems have been used extensively to map sulfur and nitrogen deposition in the
UK, namely the Concentration Based Estimated Deposition (CBED)³⁴ modeling system and the
Hull Acid Rain Model (HARM).³⁵ CBED is the operational observation-based modeling system
used to inform policy makers about current levels of sulfur and nitrogen deposition in the UK.
HARM has been used to support the development of emissions abatement strategies for reducing

432 acid deposition in the UK.³⁶ A comparison of the model deposition budget predictions reported
433 in our work with those of these modeling systems will be undertaken in a future study.

434

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537 **TABLES**

538 Table 1. Percentage fraction of predictions, within a factor of two of observations (FO2), 539 considering all monitoring sites within the Automatic Urban and Rural Network (AURN) and the 540 Joint Environmental Programme (JEP) monitoring sites, for the annual mean air concentrations of SO₂, NO_x, and PM₁₀, and within the Secondary Acid Precipitation Monitoring Network 541 542 (SAPMN) for non-sea-salt (nss) SO_x , NO_y , and NH_x wet depositions, for each modeling 543 system for the year 2003. The figures for wet deposition that are given in brackets correspond to 544 observational data derived using the gridded UK Met Office precipitation observations (see text 545 for details).

| | CMAQ V4.6 | CMAQ V4.7 | TRACK-ADMS | FRAME |
|--------------------------------|--------------|--------------|------------|-------------|
| SO ₂ | 87.7 | 87.7 | 69.2 | 78.5 |
| NO _x | 72.6 | 58.9 | 91.8 | 84.9 |
| PM ₁₀ | 88.2 | 100.0 | 100.0 | NA |
| Nss SO_x wet deposition | 100.0 (86.5) | 83.8 (83.8) | NA | 81.1 (81.1) |
| NO_y wet deposition | 97.3 (86.5) | 100.0 (89.2) | NA | 91.9 (83.8) |
| NH _x wet deposition | 97.3 (75.7) | 86.5 (81.1) | NA | 62.2 (70.3) |

| | CMAQ V4.6 | CMAQ V4.7 | TRACK-ADMS | FRAME |
|--------------------------------|-------------|-------------|------------|-------------|
| SO ₂ | 0.27 | 0.29 | 0.40 | 0.28 |
| NO _x | 0.76 | 0.77 | 0.76 | 0.76 |
| PM ₁₀ | 0.09 | 0.00 | 0.45 | NA |
| Nss SO_x wet deposition | 0.82 (0.43) | 0.75 (0.41) | NA | 0.83 (0.44) |
| NO_y wet deposition | 0.85 (0.51) | 0.86 (0.54) | NA | 0.77 (0.27) |
| NH _x wet deposition | 0.78 (0.34) | 0.67 (0.31) | NA | 0.65 (0.19) |

Table 2. Same caption as Table 1 for the correlation coefficient.

Table 3. Same caption as Table 1 for the normalized mean bias (NMB), as a percentage.

| | CMAQ V4.6 | CMAQ V4.7 | TRACK-ADMS | FRAME |
|---------------------------|---------------|--------------|------------|-------------|
| SO ₂ | -6.7 | -8.6 | -39.5 | -11.7 |
| NO _x | -41.5 | -47.2 | -15.5 | -6.4 |
| PM_{10} | -32.7 | -8.9 | -20.2 | NA |
| Nss SO_x wet deposition | -2.7 (-12.6) | 50.9 (35.5) | NA | 70.8 (53.4) |
| NO_y wet deposition | -12.0 (-22.6) | -9.0 (-20.0) | NA | 39.9 (23.0) |
| NH_x wet deposition | -13.3 (-22.8) | 32.7 (18.2) | NA | 67.8 (49.4) |

551 Table 4. UK deposition budgets for non-sea-salt (nss) sulfur (in Gg S) and nitrogen deposition

| 552 | (in Gg N), as calculated by each modeling system for the year 2003. |
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| | CMAQ V4.6 | CMAQ V4.7 | TRACK-ADMS | FRAME | |
|--------------------------------|-----------|-----------|------------|-------|--|
| Nss SO_x wet deposition | 57 | 98 | NA | 102 | |
| Nss SO_x dry deposition | 130 | 131 | NA | 65 | |
| Nss S total deposition | 187 | 229 | 154 | 167 | |
| NO_y wet deposition | 46 | 50 | NA | 67 | |
| NO _y dry deposition | 75 | 79 | NA | 61 | |
| NH_x wet deposition | 48 | 79 | NA | 90 | |
| NH _x dry deposition | 97 | 103 | NA | 69 | |
| N total deposition | 266 | 311 | 266 | 287 | |

Table 5. Maximum percentage contribution of the power station to regional air concentration for SO₂, NO_x, and PM₁₀, and non-sea-salt (nss) sulfur and nitrogen total deposition, for each modeling system for the year 2003.

| | CMAQ V4.6 | CMAQ V4.7 | TRACK-ADMS | FRAME |
|------------------------|-----------|-----------|------------|-------|
| SO ₂ | 70.2 | 68.1 | 22.7 | 38.7 |
| NO _x | 22.5 | 19.5 | 2.9 | 7.7 |
| PM_{10} | 6.0 | 3.0 | 10.1 | NA |
| Nss S total deposition | 67.1 | 60.2 | 15.6 | 32.0 |
| N total deposition | 7.3 | 6.0 | 1.1 | 2.7 |

Table 6. Maximum distance (in km) from the power station at which its contribution to regional

560 air concentration for SO_2 , NO_x , and PM_{10} , and non-sea-salt (nss) sulfur and nitrogen total

| 561 c | deposition, | is half | of its ma | ximum | contribution, | for each | modeling | system | for the y | ear 2003. |
|-------|-------------|---------|-----------|-------|---------------|----------|----------|--------|-----------|-----------|
|-------|-------------|---------|-----------|-------|---------------|----------|----------|--------|-----------|-----------|

| | CMAQ V4.6 | CMAQ V4.7 | TRACK-ADMS | FRAME | |
|------------------------|-----------|-----------|------------|-------|--|
| SO ₂ | 10 | 15 | 115 | 60 | |
| NO _x | 5 | 5 | 140 | 70 | |
| PM ₁₀ | 5 | 35 | 10 | NA | |
| Nss S total deposition | 15 | 20 | 115 | 35 | |
| N total deposition | 5 | 5 | 115 | 55 | |

Table 7. Percentage contribution of the power station to the UK annual mean air concentrations 564 of SO_2 , NO_x , and PM_{10} , and non-sea-salt (nss) sulfur and nitrogen total deposition budgets, for

565 each modeling system for the year 2003.

| | CMAQ V4.6 | CMAQ V4.7 | TRACK-ADMS | FRAME | |
|------------------------|-----------|-----------|------------|-------|--|
| SO ₂ | 2.19 | 2.17 | 2.58 | 2.85 | |
| NO _x | 0.67 | 0.63 | 0.47 | 0.62 | |
| PM_{10} | 0.34 | 0.28 | 0.28 | NA | |
| Nss S total deposition | 2.24 | 1.85 | 1.87 | 2.55 | |
| N total deposition | 0.19 | 0.16 | 0.13 | 0.39 | |

Table 8. Spatial correlation coefficient and coefficient of variation of the root mean square error (CVRMSE, in percents), reflecting similarities between the footprints of air concentrations of SO₂, NO_x, and PM₁₀, and total depositions of non-sea-salt (nss) sulfur and nitrogen, with respect to the reference modeling system (the CMAQ modeling system, version 4.6), for each modeling system in the area indicated by a dashed polyline in Figure 3, for the year 2003.

| | Spati | al correlation coeffi | cient | CVRMSE | | | |
|------------------------|-----------|-----------------------|-------|-----------|------------|-------|--|
| | CMAQ V4.7 | TRACK-ADMS | FRAME | CMAQ V4.7 | TRACK-ADMS | FRAME | |
| SO ₂ | 0.98 | 0.36 | 0.61 | 16.7 | 116.4 | 98.4 | |
| NO _x | 0.98 | 0.34 | 0.62 | 24.9 | 101.4 | 96.0 | |
| PM ₁₀ | 0.82 | 0.61 | NA | 39.8 | 105.8 | NA | |
| Nss S total deposition | n 0.98 | 0.46 | 0.78 | 17.8 | 108.0 | 71.8 | |
| N total depostion | 0.90 | 0.42 | 0.43 | 46.3 | 101.1 | 195.5 | |

574

575 **LIST OF FIGURE CAPTIONS**

- Location and type (remote, rural, suburban, urban background) of monitoring sites in the UK Automatic Urban and Rural Network (AURN) and the Joint Environmental Programme (JEP) monitoring sites, as filled circles, and the Secondary Acid Precipitation Monitoring Network (SAPMN), as open circles, used for the evaluation of the model baseline calculations. The grey-filled area corresponds to the 'UK domain' used for the model comparison exercise. The location of the fossil-fuel power station considered for the footprint calculations is marked by a cross symbol.
- 2. Annual total deposition of non-sea-salt (nss) sulfur, as calculated by each modeling
 system in the 'UK domain' for the year 2003: (a) CMAQ version 4.6, (b) CMAQ version
 4.7, (c) TRACK-ADMS, and (d) FRAME.
- 586 3. Percentage contribution of the power station to regional non-sea-salt (nss) sulfur total
- 587 deposition, as calculated by each modeling system in the 'UK domain' for the year 2003:
- 588 (a) CMAQ version 4.6, (b) CMAQ version 4.7, (c) TRACK-ADMS, and (d) FRAME.
- 589 Note that the color scale is not linear. The dashed polyline represents the area over which
- 590 the statistics reported in Table 8 are calculated.